OXFORD IB STUDY GUIDES

Geoffrey Neuss

Chemistry

FOR THE IB DIPLOMA

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OXFORD

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Introduction and acknowledgements

This book replaces the very success ul rst and second editions. Like earlier editions it is written speci cally or students studying Chemistry or the International Baccalaureate Diploma although many students ollowing their own national systems will also nd it help ul. It comprehensively covers the new programme that will be examined rom 2016 onwards. All the in ormation required or each topic is set out in separate boxes with clear titles that ollow aith ully the layout o the syllabus. The rst 11 topics cover both the Core content needed by all students and the Additional Higher Level material under the main topic headings. The di erence between the two levels is clearly distinguished. Both Higher Level and Standard Level students must study one o the our options and each option stands in its own right even i this has meant repeating small amounts o some o the material. Worked examples are included where they are appropriate and at the end o each main topic there are both multiple choice and short-answer practice questions. Similarly at the end o each option I have included shortanswer questions. Many o these questions are taken rom past IB examination papers and I would like to thank the International Baccalaureate Organization or giving me permission to use them. The remaining questions are written to the same IB standard speci cally or this book. Worked answers to the questions are provided. These answers are not necessarily ull 'model' answers but they do contain all the in ormation needed to score each possible mark. I have included one chapter on the underlying philosophy o the course to explain and give examples o the Essential Ideas, the Nature o Science, the International Dimension, and Utilization. To help you, the student, gain the highest grade possible the nal chapter is devoted to giving you advice on how to study and prepare or the nal examination. It also advises you on how to excel at the internally assessed practical component o the course. For those who opt or Chemistry as the subject or their Extended Essay it gives advice and guidance on how to choose the topic and write your Essay. A comprehensive Extended Essay checklist is included to help you gain bonus points towards your IB Diploma.

I have stuck rigorously to the syllabus to produce this study guide which contains all the necessary subject content required or the examination in one easily accessible and compact ormat. IB Chemistry is, o course, about much more than the nal examination and you are also encouraged to read widely around the subject to urther your knowledge and understanding as well as your enjoyment o chemistry. Many more multiple choice tests and short answer questions, as well as a blog and much more background in ormation (e.g. videos) about each topic and option can be ound on my website or IB Diploma Chemistry teachers at www.thinkib.net/chemistry.

I have been ortunate at Atlantic College to teach many highly motivated and gi ted students who have o ten challenged me with searching questions. During my association with the International Baccalaureate, the European Baccalaureate and the United World Colleges I have been privileged to meet, work alongside and exchange ideas with many excellent Chemistry teachers who exude a real enthusiasm or their subject. Many o these have influenced me greatly in particular, John Devonshire, a ellow teacher at Atlantic College and Jacques Furnemont, an Inspector o Chemistry in Belgium. I value greatly their advice, opinions and knowledge. I would also like to pay tribute to two ormer Chie Examiners or the IB, Ron Ragsdale and Arden Zipp. The high regard with which IB Chemistry is held today owes much to both o them.

Finally I should like to thank my wi e Chris and my riend and colleague John or their patience and unstinting support throughout.

Dr Geo rey Neuss

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1 STOICHIOMETRIC RELATIONSHIPS

Particulate nature of matter

ELEMENTS

All substances are made up o one or more elements. An element cannot be broken down by any chemical process into simpler substances. There are just over 100 known elements. The smallest part o an element is called an atom.

Names o the first 20 elements

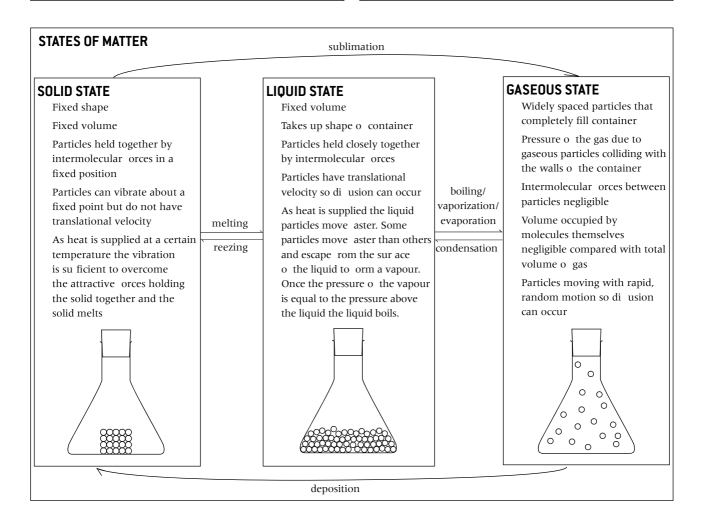
Atomic Number	Name	Symbol	Relative atomic mass
1	hydrogen	Н	1.01
2	helium	He	4.00
3	lithium	Li	6.94
4	beryllium	Be	9.01
5	boron	В	10.81
6	carbon	С	12.01
7	nitrogen	Ν	14.01
8	oxygen	0	16.00
9	uorine	F	19.00
10	neon	Ne	20.18
11	sodium	Na	22.99
12	magnesium	Mg	24.31
13	aluminium	Al	26.98
14	silicon	Si	28.09
15	phosphorus	Р	30.97
16	sul ur	S	32.07
17	chlorine	Cl	35.45
18	argon	Ar	39.95
19	potassium	K	39.10
20	calcium	Ca	40.08

COMPOUNDS

Some substances are made up o a single element, although there may be more than one atom o the element in a particle o the substance. For example, oxygen is diatomic, that is, a molecule o oxygen contains two oxygen atoms and has the ormula O_2 . A compound contains more than one element combined chemically in a fixed ratio. For example, a molecule o water contains two hydrogen atoms and one oxygen atom. It has the ormula H_2O . Water is a compound not an element because it can be broken down chemically into its constituent elements: hydrogen and oxygen. Compounds have di erent chemical and physical properties rom their component elements.

MIXTURES

The components o a mixture may be elements or compounds. These components are not chemically bonded together. Because they are not chemically combined, the components o a mixture retain their individual properties. All the components o a mixture may be in the same phase, in which case the mixture is said to be **homogeneous**. Air is an example o a gaseous homogenous mixture. I the components o a mixture are in di erent phases the mixture is said to be **heterogeneous**. There is a physical boundary between two phases. A solid and a liquid is an example o a two-phase system. It is possible to have a single state but two phases. For example, two immiscible liquids such as oil and water orm a heterogeneous mixture.



The mole concept and chemical formulas

MOLE CONCEPT AND AVOGADRO S CONSTANT

A single atom of an element has an extremely small mass. For example, an atom of carbon-12 has a mass of 1.993 10^{-23} g. This is far too small to weigh. A more convenient amount to weigh is 12.00 g. 12.00 g of carbon-12 contains 6.02 10^{23} atoms of carbon-12. This number is known as Avogadro s constant (N_{4} or L).

Chemists measure amounts of substances in moles. A mole is the amount of substance that contains *L* particles of that substance. The mass of one mole of **any** substance is known as the **molar mass** and has the symbol *M*. For example, hydrogen atoms have $\frac{1}{12}$ of the mass of carbon-12 atoms so a mole of hydrogen atoms contains 6.02 10²³ hydrogen atoms and has a mass of 1.01 g. In reality elements are made up of a mixture of isotopes.

The **relative atomic mass** of an element A_i is the weighted mean of all the naturally occurring isotopes of the element relative to carbon-12. This explains why the relative atomic masses given for the elements on page 1 are not whole numbers. The units of molar mass are g mol⁻¹ but relative molar masses M_i have no units. For molecules **relative molecular mass** is used. For example, the M_i of glucose, $C_6H_{12}O_6 = (6 \quad 12.01) + (12 \quad 1.01) + (6 \quad 16.00) = 180.18$. For ionic compounds the term **relative formula mass** is used. Be careful to distinguish between the words **mole** and **molecule**. A molecule of hydrogen gas contains two atoms of hydrogen atoms.

FORMULAS OF COMPOUNDS

Compounds can be described by different chemical formulas.

Empirical formula (literally the formula obtained by experiment)

This shows the simplest whole number ratio of atoms of each element in a particle of the substance. It can be obtained by either knowing the mass of each element in the compound or from the percentage composition by mass of the compound. The percentage composition can be converted directly into mass by assuming 100 g of the compound are taken.

Example: A compound contains 40.00% carbon, 6.73% hydrogen and 53.27% oxygen by mass, determine the empirical formula.

 Amount / mol
 Ratio

 C
 40.00/12.01 = 3.33 1

 H
 6.73/1.01 = 6.66 2

 O
 53.27/16.00 = 3.33 1

 Molecular formula

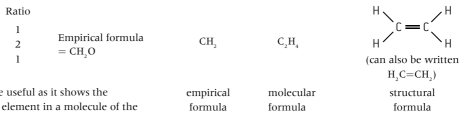
substance. It can be obtained from the empirical formula if the molar mass of the compound is also known.

Methanal CH₂O ($M_r = 30$), ethanoic acid C₂H₄O₂ ($M_r = 60$) and glucose C₆H₁₂O₆ ($M_r = 180$) are different substances with different molecular formulas but all with the same empirical formula CH₂O. Note that subscripts are used to show the number of atoms of each element in the compound.

Structural formula

This shows the arrangement of atoms and bonds within a molecule and is particularly useful in organic chemistry.

The three different formulas can be illustrated using ethene:



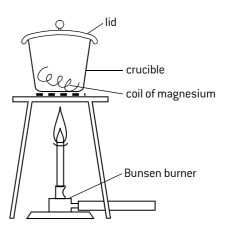
For molecules this is much more useful as it shows the actual number of atoms of each element in a molecule of the

EXPERIMENTAL DETERMINATION OF AN EMPIRICAL FORMULA

The empirical formula of magnesium oxide can be determined simply in the laboratory. A coil of magnesium ribbon about 10 cm long is placed in a pre-weighed crucible and its mass recorded. The crucible is placed on a clay triangle and heated strongly. When the magnesium ribbon starts to burn the lid is lifted slightly to allow more air to enter and the heating is continued until all the magnesium has burned. After cooling the crucible, its lid and its contents are reweighed.

Table of typical raw quantitative data:

	Mass / g (0.001 g)	
Mass of crucible + lid	30.911	
Mass of crucible + lid + magnesium	31.037	
Mass of crucible + lid + magnesium oxide	31.106	
Mass of magnesium = $31.037 - 30.911 = 0.126$ g Mass of magnesium oxide = $31.106 - 30.911 = 0.195$ g Mass of oxygen combining with magnesium = $0.195 - 0.126 = 0.069$ g Amount of magnesium = $\frac{0.126}{24.31} = 5.2$ 10 ³ mol		
Amount of oxygen $=$ $\frac{0.069}{16.00} = 4.3$ 10 ³ mol Ratio of Mg to O $=$ $\frac{5.2}{4.3}$ $\frac{10}{10^3} = 1.2$ to 1		
Convert to whole number ratio = $6:5$ Empirical formula of magnesium oxide as determined by this experiment is Mg _e O _e .		



2

Chemical reactions and e uations

PROPERTIES OF CHEMICAL REACTIONS

In all chemical reactions:

new substances are ormed

bonds in the reactants are broken and bonds in the products are ormed resulting in an energy change between the reacting system and its surroundings

there is a fixed relationship between the number o particles o reactants and products resulting in no overall change in mass this is known as the stoichiometry o the reaction.

CHEMICAL EQUATIONS

Chemical reactions can be represented by chemical equations. Reactants are written on the le t-hand side and products on the right-hand side. The number o moles o each element must be the same on both sides in a balanced chemical equation, e.g. the reaction o nitric acid (one o the acids present in acid rain) with calcium carbonate (the main constituent o marble statues).

$CaCO_{3}(s)$	+	2HNO ₃ (aq)	$Ca(NO_3)_2(aq)$	+ $CO_{2}(g)$ +	$H_{2}O(l)$
calcium carbonate		nitric acid	calcium nitrate	carbon dioxide	water
RE	ACTAN	ГS		PRODUCTS	

STATE SYMBOLS

Because the physical state that the reactants and products are in can a ect both the rate o the reaction and the overall energy change it is good practice to include the state symbols in the equation.

> (s) - solid (l) - liquid (g) - gas(aq) - in aqueous solution

OR

e.g.

A single arrow is used i the reaction goes to completion. Sometimes the reaction conditions are written on the arrow:

Ni catalyst, 180 C $C_2H_4(g) + H_2(g)$

 $C_{2}H_{2}(g)$

Reversible arrows are used or reactions where both the reactants and products are present in the equilibrium mixture:

e.g.
$$3H_2(g) + N_2(g) \xleftarrow{Fe(s), 550 C}{250 \text{ Atm}} 2NH_3(g)$$

COEFFICIENTS AND MOLAR RATIO

The coe ficient re ers to the number in ront o each reactant and product in the equation. The coe ficients give in ormation on the molar ratio. In the first example above, two moles o nitric acid react with one mole o calcium carbonate to produce one mole o calcium nitrate, one mole o carbon dioxide and one mole o water. In the reaction between hydrogen and nitrogen above, three moles o hydrogen gas react with one mole o nitrogen gas to produce two moles o ammonia gas.

IONIC EQUATIONS

Because ionic compounds are completely dissociated in solution it is sometimes better to use ionic equations. For example, when silver nitrate solution is added to sodium chloride solution a precipitate o silver chloride is ormed.

 $Ag^{+}(aq) + NO_{3}^{-}(aq) + Na^{+}(aq) + Cl^{-}(aq)$ $AgCl(s) + Na^{+}(aq) + NO_{3}^{-}(aq)$

 $Na^+(aq)$ and $NO_3^-(aq)$ are spectator ions and do not take part in the reaction. So the ionic equation becomes:

 $Ag^{+}(aq) + Cl^{-}(aq)$ AgCl(s)

From this we can deduce that any soluble silver salt will react with any soluble chloride to orm a precipitate o silver chloride.

Mass and gaseous volume relationships

SOLIDS

Normally measured by weighing to obtain the mass.

1.000 kg = 1000 g

When weighing a substance the mass should be recorded to show the accuracy of the balance. For example, exactly 16 g of a substance would be recorded as 16.00 g on a balance weighing to + or

- 0.01 g but as 16.000 g on a balance weighing to + or - 0.001 g.

MEASUREMENT OF MOLAR QUANTITIES

In the laboratory moles can conveniently be measured using either mass or volume depending on the substances involved.

LIQUIDS

Pure liquids may be weighed or the volume recorded.

The density of the liquid $= \frac{\text{mass}}{\text{volume}}$ and is

usually expressed in g cm⁻³.

GASES Mass or volume may be used for gases.

SOLUTIONS

Volume is usually used for solutions.

 $1.000 \text{ litre} = 1.000 \text{ dm}^3 = 1000 \text{ cm}^3$

Concentration is the amount of solute (dissolved substance) in a known volume of solution (solute plus solvent). It is expressed either in g dm⁻³, or, more usually in mol dm⁻³. For very dilute solutions it is also sometimes expressed in parts per million, ppm. A solution of known concentration is known as a standard solution.

To prepare a 1.00 mol dm⁻³ solution of sodium hydroxide dissolve 40.00 g of solid sodium hydroxide in distilled water and then make the total volume up to 1.00 dm³. Concentration is often represented by

square brackets, e.g.

 $[NaOH(aq)] = 1.00 \text{ mol } dm^{-3}$

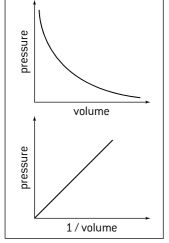
A 25.0 cm³ sample of this solution contains

 $1.00 \quad \frac{25.0}{1000} = 2.50 \quad 10^{2} \text{ mol of NaOH}$

CHANGING THE VARIABLES FOR A FIXED MASS OF GAS

$P = \frac{1}{V}$ (or PV = constant)

At constant temperature: as the volume decreases the concentration of the particles increases, resulting in more collisions with the container walls. This increase in pressure is inversely proportional to the volume, i.e. doubling the pressure halves the volume.



 $P \quad T \text{ (or } \frac{P}{T} = \text{constant})$

At constant volume: increasing the temperature increases the average kinetic energy so the force with which the particles collide with the container walls increases. Hence pressure increases and is directly proportional to the absolute temperature, i.e. doubling the absolute temperature doubles the pressure.

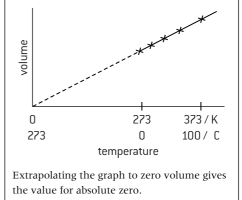
IDEAL GAS EQUATION The different variables for a gas are all related by the ideal gas equation. PV = nRT $P = \text{pressure in Pa} (\text{N m}^{-2})$

- $(1 \text{ atm} = 1.013 \quad 10^5 \text{ Pa})$
- T = absolute temperature in K V = volume in m³
 - $(1 \text{ cm}^3 = 1 \quad 10^6 \text{ m}^3)$
- n = number of moles

$R = \text{gas constant} = 8.314 \text{ J K}^{-1} \text{ mol}^{-1}$

V T (or $\frac{V}{T}$ = constant)

At constant pressure: at higher temperatures the particles have a greater average velocity so individual particles will collide with the container walls with greater force. To keep the pressure constant there must be fewer collisions per unit area so the volume of the gas must increase. The increase in volume is directly proportional to the absolute temperature, i.e. doubling the absolute temperature doubles the volume.



UNITS

The gas constant can be expressed in different units but it is easier to use SI units.

$$R = \frac{PV}{nT} = \frac{N \text{ m}^{-2} \text{ m}^3}{\text{mol} \text{ K}} = N \text{ m mol}^{-1} \text{ K}^{-1}$$
$$= J \text{ K}^{-1} \text{ mol}^{-1}$$

REAL GASES

An ideal gas exactly obeys the gas laws. Real gases do have some attractive forces between the particles and the particles themselves do occupy some space so they do not exactly obey the laws. If they did they could never condense into liquids. A gas behaves most like an ideal gas at high temperatures and low pressures.

Molar volume of a gas and calculations

MOLAR VOLUME OF A GAS

The ideal gas equation depends on the amount o gas (number o moles o gas) but not on the nature o the gas. Avogadro s Law states that equal volumes o di erent gases at the same temperature and pressure contain the same number o moles. From this it ollows that one mole o any gas will occupy the same volume at the same temperature and pressure. This is known as the molar volume o a gas. At 273K and 1.00 10^5 Pa pressure this volume is 22.7 10^2 m³ (22.7 dm³ or 22 700 cm³).

When the mass o a particular gas is fixed (nR is constant) a use ul expression to convert the pressure, temperature and volume under one set o conditions (1) to another set o conditions (2) is:

$$\frac{P_{1}V_{1}}{T_{1}} = \frac{P_{2}V_{2}}{T_{2}}$$

In this expression there is no need to convert to SI units as long as the same units or pressure and volume are used on both sides o the equation. However do not orget that T re ers to the absolute temperature and must be in kelvin.

CALCULATIONS FROM EQUATIONS

Work methodically.

Step 1. Write down the correct ormulas or all the reactants and products.

Step 2. Balance the equation to obtain the correct stoichiometry o the reaction.

Step 3. I the amounts o all reactants are known work out which are in **excess** and which one is the limiting reagent. By knowing the **limiting reagent** the maximum **yield** o any o the products can be determined.

Step 4. Work out the amount (in mol) o the substance required.

Step 5. Convert the amount (in mol) into the mass or volume.

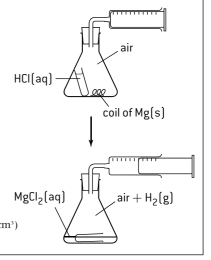
Step 6. Express the answer to the correct number o significant figures and include the appropriate units.

WORKED EXAMPLES

(a) Calculate the volume o hydrogen gas evolved at 273 K and 1.00 10⁵ Pa when 0.623 g o magnesium reacts with 27.3 cm³ o 1.25 mol dm⁻³ hydrochloric acid.

Equation:

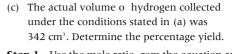
 $Mg(s) + 2HCl(aq) \qquad H_2(g) + MgCl_2(aq) \qquad H_2(g) + MgCl_2(g) + MgCl_2(g) \qquad H_2(g) + MgCl_2(g) \qquad H_2(g) + MgCl_2(g) \qquad H_2(g) + MgCl_$



(b) Calculate the volume occupied by the hydrogen evolved. In the example
above i it had been collected at 22 C and at a pressure o 1.12 10⁵ Pa
Step 1. Express the temperature as an absolute temperature
22 C = 295 K
Step 2. Apply the ideal gas equation
$$pV = nRT$$

1.12 10⁵ $V = 1.705$ 10⁻² 8.314 295
 $V = \frac{1.705 \ 10^{-2} \ 8.314 \ 295}{1.12 \ 10^{5}} = 3.73 \ 10^{-4} \text{ m}^{3} (373 \text{ cm}^{3})$
This could also be solved using $\frac{P_{1}V_{1}}{T_{1}} = \frac{P_{2}V_{2}}{T_{2}}$
 $V_{2} = V_{1} \quad \frac{P_{1}}{P_{2}} \quad \frac{T_{2}}{T_{1}} = 0.387 \quad \frac{1.00 \ 10^{-5}}{1.12 \ 10^{-5}} \quad \frac{295}{273}$
 $= 0.373 \text{ cm}^{3} (373 \text{ cm}^{3})$

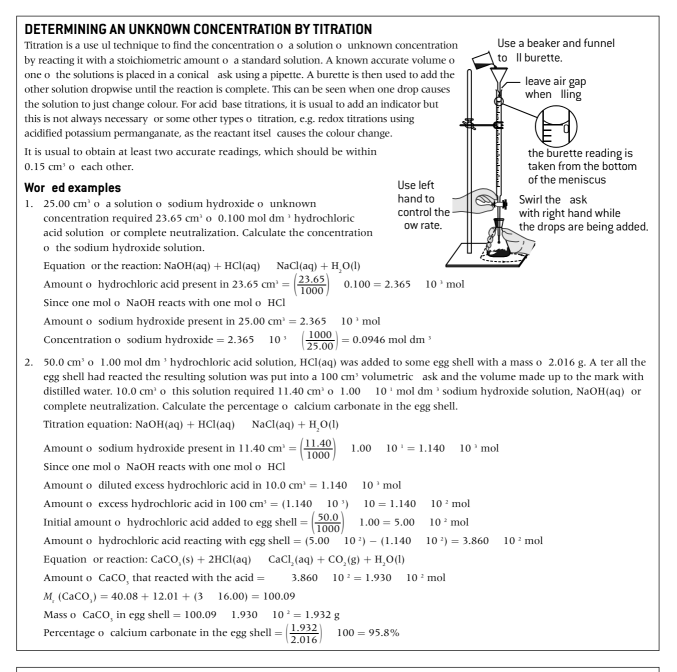
Ť



Step 1. Use the mole ratio rom the equation and the amounts o reactants to determine the limiting reagent and hence the theoretical maximum yield. From part (a) theoretical yield = 387 cm³
Step 2. Apply the relationship:

Percentage yield = $\frac{\text{Experimental yield}}{\text{Theoretical yield}}$ 100 Percentage yield = $\frac{342}{387}$ 100 = 88.4%

Titration and atom econom



ATOM ECONOMY (AN EXAMPLE OF UTILIZATION)

As well as trying to achieve high yields in industrial processes, chemists try to increase the conversion e ficiency o a chemical process. This is known as atom economy. Ideally in a chemical process no atom is wasted. The atom economy is a measure o the amount o starting materials that become use ul products. A high atom economy means that ewer natural resources are used and less waste is created. The atom economy can be calculated by using the ollowing steps:

- 1. Write the balanced equation or the reaction taking place.
- 2. Calculate the relative molecular mass o each product and then the total mass o each product ormed assuming molar quantities. Note that this is the same as the total mass o the reactants.
- 3. Calculate the relative molecular mass o each desired product and then the total mass o each desired product ormed assuming molar quantities.
- 4. Atom economy = $\frac{\text{total mass o desired product(s)}}{\text{total mass o all products}}$ 100

For example, consider the production o iron by the reduction o iron(III) oxide using the thermite reaction.

 $2Al(s) + Fe_2O_3(s) = 2Fe(s) + Al_2O_3(s)$

The total mass o products ormed = 2 - 55.85 + [(2 - 26.98) + (3 - 16.00)] = 213.66 g

The total amount o iron (the desired product) ormed = 2 - 55.85 = 111.70 g

The atom economy or this reaction is $\frac{111.70}{213.66}$ 100 = 52.3%

Obviously i a use can also be ound or all the aluminium oxide produced then the atom economy or this reaction will increase to 100%.

6 STOICHIOMETRIC RELATIONSHIPS

MULTIPLE CHOICE QUESTIONS STOICHIOMETRIC RELATIONSHIPS

1. How many oxygen **atoms** are in 0.100 mol o $CuSO_4.5H_2O$? 1

A. 5.42	1022	C. 2.41	1023
B. 6.02	1022	D. 5.42	1023

2. Which is not a true statement?

A. One mole o methane contains our moles o hydrogen atoms

- B. One mole o ¹²C has a mass o 12.00 g
- C. One mole o hydrogen gas contains 6.02 10²³ atoms o hydrogen
- D. One mole o methane contains 75% o carbon by mass
- 3. A pure compound contains 24 g o carbon, 4 g o hydrogen and 32 g o oxygen. No other elements are present. What is the empirical ormula o the compound?

A.	$C_2H_4O_2$	C.	$\mathrm{CH}_4\mathrm{O}$
В.	CH_O	D.	СНО

4. What is the mass in grams o one molecule o ethanoic acid CH,COOH?

22

A. 0.1		С.	1	10
B. 3.6	1025	D.	60	

5.	What is the relative molecular mass, A	М _. , о	carbon	dioxide,
	CO ₂ ?			

A. 44.01 g mol ¹	C. 44.01 kg mol ¹
B. 44.01 mol g ⁻¹	D. 44.01

6. Which o the ollowing changes o state is an exothermic process?

A. melting	C. vaporizing
B. condensing	D. boiling

7. What is the empirical ormula or the compound $C_6H_5(OH)_2$?

A. C_6H_6O	C. C ₆ H ₇ O
B. $C_6H_5O_2H_2$	D. $C_{6}H_{7}O_{2}$

8. Phosphorus burns in oxygen to produce phosphorus pentoxide P₄O₁₀. What is the sum o the coe ficients in the balanced equation?

$_{-}P_{_{4}}(s) + _{-}O_{_{2}}(g)$	$-P_4O_{10}(s)$		
A. 3		C.	6
B. 5		D.	7

9. Magnesium reacts with hydrochloric acid according to the ollowing equation:

 $Mg(s) + 2HCl(aq) MgCl_2(aq) + H_2(g)$

What mass o hydrogen will be obtained i 100 cm³ o 2.00 mol dm ³ HCl are added to 4.86 g o magnesium?

A. 0.2 g	C. 0.8 g
B. 0.4 g	D. 2.0 g

10. Butane burns in oxygen according to the equation below.

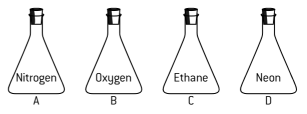
 $2C_4H_{10}(g) + 13O_2(g) = 8CO_2(g) + 10H_2O(l)$

I 11.6 g o butane is burned in 11.6 g o oxygen which is the limiting reagent?

A. Butane	C. Neither

B. Oxygen D. Oxygen and butane

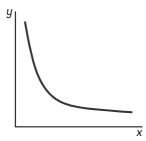
11. Four identical containers under the same conditions are filled with gases as shown below. Which container and contents will have the highest mass?



12. What is the amount, in moles, o sul ate ions in 100 cm³ o $0.020 \text{ mol } dm^{-3} \text{ FeSO}_4(aq)$?

A. 2.0	10-3	C.	2.0	10-
B. 2.0	10-2	D.	2.0	

- 13. 300 cm³ o water is added to a solution o 200 cm³ o
 0.5 mol dm⁻³ sodium chloride. What is the concentration o sodium chloride in the new solution?
 - A. 0.05 mol dm⁻³ C. 0.2 mol dm⁻³
 - B. 0.1 mol dm⁻³ D. 0.3 mol dm⁻³
- 14. Separate samples o two gases, each containing a pure substance, are ound to have the same density under the same conditions o temperature and pressure. Which statement about these two samples must be correct?
 - A. They have the same volume
 - B. They have the same relative molecular mass
 - C. There are equal numbers o moles o gas in the two samples
 - D. They condense at the same temperature
- 15. The graph below represents the relationship between two variables in a fixed amount o gas.



Which variables could be represented by each axis?

	x-axis	y-axis
A.	pressure	temperature
В.	volume	temperature
C.	pressure	volume
D.	temperature	volume

16. Sul uric acid and sodium hydroxide react together according to the equation:

$$\begin{split} H_2SO_4(aq) &+ 2NaOH(aq) & Na_2SO_4(aq) + 2H_2O(l) \\ What volume o & 0.250 mol dm ³ NaOH is required to neutralize exactly 25.0 cm³ o & 0.125 mol dm ³ H_2SO_4? \\ A. 25.0 cm³ & C. 50 cm³ \end{split}$$

B. 12.5 cm³ D. 6.25 cm³

SHORT ANSWER QUESTIONS STOICHIOMETRIC RELATIONSHIPS

	$C_4H_6O_3$, $C_9H_8O_4$, is made by reacting ethanoic anhydride, ($M_r = 102.1$), with 2-hydroxybenzoic acid 38.1), according to the equation:			b) The combustion o unleaded gasoline still produces pollution with both local and global consequences.Identi y one exhaust gas that causes local pollution	
	2C ₇ 1	$H_6O_3 + C_4H_6O_3 = 2C_9H_8O_4 + H_2O_3$			and one exhaust gas that causes global pollution.	[2]
	etha	5.0 g 2-hydroxybenzoic acid is reacted with 15.0 g noic anhydride, determine the limiting reagent in reaction.	[3]	5.	An experiment was per ormed to determine the percentage or iron present in a sample or iron ore. 3.682 10 ⁻¹ g or the sam was dissolved in acid and all or the iron was converted to Fe ²	nple
	be o	ulate the maximum mass o aspirin that could btained in this reaction.	[2]		The resulting solution was titrated with a standard solution o potassium manganate(VII), $KMnO_4$ with a concentration o 2.152 10 ² mol dm ³ . The end point was indicated with	n
	calcu	e mass obtained in this experiment was 13.7 g, late the percentage yield o aspirin. o a metal sul ate with the ormula M ₂ SO ₄ was	[1]		one drop caused a slight pink colour to remain. It was out that 22.50 cm ³ o the potassium manganate(VII) solution v	nd
	dissolve added in	d in water. Excess barium nitrate solution was n order to precipitate all the sul ate ions in the			required to reach the end point. In acidic solution, MnO_4 reacts with Fe^{2+} ions to orm Mn^{2+} and Fe^{3+} ions according to the ollowing equation:	
	a) Calc	barium sul ate. 9.336 g o precipitate was obtained. ulate the amount o barium sul ate BaSO ₄	[2]		MnO_4^- (aq) + 5Fe ²⁺ (aq) + 8H ⁺ (aq) $Mn^{2+}(aq)$ + 5Fe ³⁺ + 4H ₂ O(l)	(aq)
	b) Calc	ipitated. ulate the amount o sul ate ions present in 14.48 g o M ₂ SO ₄ .	[2] [1]		 a) Calculate the amount (in mol) o MnO₄⁻ used in the titration. 	[2]
		uce the relative molar mass o M_2SO_4 .	[1]		b) Calculate the amount (in mol) o Fe present in	
	d) Calc	ulate the relative atomic mass o M and hence ti y the metal.	[2]		 the 3.682 10⁻¹ g sample o iron ore. c) Determine the percentage by mass o Fe present 	[2]
3.	A stude 15.0 cm	nt added 7.40 10 ² g o magnesium ribbon to ³ o 2.00 mol dm ³ hydrochloric acid. The hydrogen		6.	in the 3.682 10 ⁻¹ g sample o iron ore. Copper metal may be produced by the reaction o copper(oxide and copper(I) sulfide according to the equation.	[2] I)
	gas prod and 1.0	duced was collected using a gas syringe at 20.0 C $0 = 10^5$ Pa.			$2Cu_{,}O(s) + Cu_{,}S(s) = 6Cu(s) + SO_{,}(g)$	
		e the equation or the reaction between			A mixture o 10.0 kg o copper(I) oxide and 5.00 kg o	
		nesium and hydrochloric acid.	[1]		copper(I) sulfide was heated until no urther reaction	
	b) Dete	ermine the limiting reactant.	[3]		occurred.	[2]
	c) Calc	ulate the theoretical yield o hydrogen gas:			a) Determine the limiting reagent in this reaction.b) Calculate the maximum mass o copper that could	[3]
	()	in mol	[1]		be obtained rom these masses o reactants.	[2]
	. ,	in cm ³ , under the stated conditions o temperature and pressure.	[2]		c) Assuming the reaction to produce copper goes to completion according to the equation deduce the	
		actual volume o hydrogen measured was lower than calculated theoretical volume. Suggest two reasons			atom economy or this reaction.	[3]
		the volume o hydrogen gas obtained was less.	[2]	7.	The empirical ormula o magnesium oxide is MgO. Suggest our assumptions that were made in the	
		Thomas Midgley discovered that the addition downward could improve the combustion o			experiment detailed on page 2 that may not be true and which might account or the wrong result being obtained.	[4]
	hydroca	arbons in automobile (car) engines. This was the		8.	The percentage composition by mass o a hydrocarbon	[4]
		ng o the use o leaded gasoline (petrol). centage composition, by mass, o the lead compou	nd		is C: 85.6% and H: 14.4%.	
	-	Midgley is Pb: 64.052%, C: 29.703% and H: 6.245			a) Calculate the empirical ormula o the hydrocarbon.	[2]
	a) (i)	Determine the empirical ormula o the lead compound.	[3]		 b) A 1.00 g sample o the hydrocarbon at a temperature o 273 K and a pressure o 1.00 10⁵ Pa has a volume o 0.405 dm³. 	
	(ii)	Leaded gasoline has been phased out because the			(i) Calculate the molar mass of the hydrocarbon.	[2]
		lead(IV) oxide, PbO ₂ , produced as a side product in the combustion reaction may cause brain			(ii) Deduce the molecular ormula o the hydrocarbon.	[2]
		damage in children. 0.01 mol o Midgley s lead compound produces			c) Explain why the incomplete combustion o	
		0.01 mol o lead(IV) oxide. Deduce the molecular ormula o Midgley s compound.	[1]		hydrocarbons is harm ul to humans.	[2]
	(iii)	Determine the equation or the complete combustion o Midgley s compound.	[2]			

8

ATOMIC STRUCTURE

T e uc ear atom

COMPOSITION OF ATOMS

The smallest part o an element is an atom. It used to be thought that atoms are indivisible but they can be broken down into many di erent sub-atomic particles. All atoms, with the exception o hydrogen, are made up o three undamental sub-atomic particles protons, neutrons and electrons.

The hydrogen atom, the simplest atom o all, contains just one proton and one electron. The actual mass o a proton is $1.673 10^{-24}$ g but it is assigned a relative value o 1. The mass o a neutron is virtually identical and also has a relative mass o 1. Compared with a proton and a neutron an electron has negligible mass with a relative mass o only $\frac{1}{2000}$. Neutrons are neutral particles. An electron has a charge o $1.602 10^{-19}$ coulombs which is assigned a relative value o -1. A proton carries the same charge as an electron but o an opposite sign so has a relative value o +1. All atoms are neutral so must contain equal numbers o protons and electrons.

SUMMARY OF RELATIVE MASS AND CHARGE

Particle	Relative mass	Relative charge
proton	1	+1
neutron	1	0
electron	5 10 ⁻⁴	-1

SIZE AND STRUCTURE OF ATOMS

Atoms have a radius in the order o 10^{-10} m. Almost all o the mass o an atom is concentrated in the nucleus which has a very small radius in the order o 10^{-14} m. All the protons and neutrons (collectively called nucleons) are located in the nucleus. The electrons are to be ound in energy levels or shells surrounding the nucleus. Much o the atom is empty space.

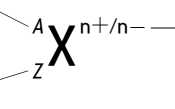
MASS NUMBER A

Equal to the number o protons and neutrons in the nucleus.

ATOMIC NUMBER Z

Equal to the number o protons in the nucleus and to the number o electrons in the atom. The atomic number defines which element the atom belongs to and consequently its position in the periodic table.

SHORTHAND NOTATION For an atom or ion



CHARGE Atoms have no charge so n = 0 and this is le t blank. However by losing one or more electrons atoms become positive ions, or by gaining one or more electrons atoms orm negative ions.

EXAMPLE	S				
Symbol	Atomic number	Mass number	Number of protons	Number of neutrons	Number of electrons
⁹ ₄ Be	4	9	4	5	4
${}^{40}_{20}Ca^{2+}$	20	40	20	20	18
$^{37}_{17}{\rm Cl}^{-}$	17	37	17	20	18

ISOTOPES

All atoms o the same element must contain the same number o protons, however they may contain a di erent number o neutrons. Such atoms are known as isotopes. Chemical properties are related to the number o electrons so isotopes o the same element have identical chemical properties. Since their mass is di erent their physical properties such as density and boiling point are di erent.

Examples o	isotopes:	$^{1}_{1}$ H	${}^{2}_{1}H {}^{3}_{1}$	H $^{12}_{6}C$	$^{14}_{6}C$	³⁵ ₁₇ Cl	³⁷ ₁₇ Cl
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RELATIVE ATOMIC MASS

The two isotopes o chlorine occur in the ratio o 3:1. That is, naturally occurring chlorine contains 75% $^{35}_{17}$ Cl and 25% $^{37}_{17}$ Cl. The weighted mean molar mass is thus:

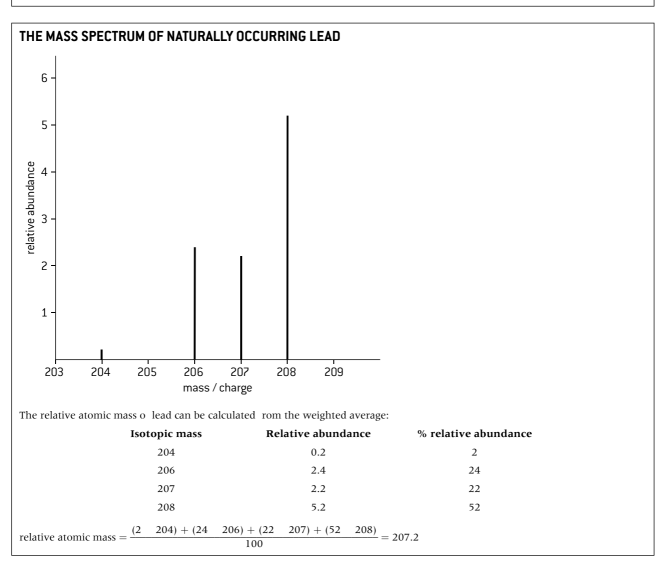
$$\frac{(75 \quad 35) + (25 \quad 37)}{100} = 35.5 \text{ g mol}^{-1}$$

and the relative atomic mass is 35.5. Accurate values to 2 d.p. or all the relative atomic masses o the elements are given in Section 6 o the IB data booklet. These are the values that must be used when per orming calculations in the examinations.

MASS SPECTROMETER

Relative atomic masses can be determined using a mass spectrometer. A vaporized sample is injected into the instrument. Atoms o the element are ionized by being bombarded with a stream o high energy electrons in the ionization chamber. In practice the instrument is set so that only ions with a single positive charge are ormed. The resulting unipositive ions pass through holes in parallel plates under the influence o an electric eld where they are accelerated. The ions are then deflected by an external magnetic eld.

The amount o deflection depends both on the mass o the ion and its charge. The smaller the mass and the higher the charge the greater the deflection. Ions with a particular mass/charge ratio are then recorded on a detector which measures both the mass and the relative amounts o all the ions present.



USES OF RADIOACTIVE ISOTOPES

Isotopes have many uses in chemistry and beyond. Many, but by no means all, isotopes o elements are radioactive as the nuclei o these atoms break down spontaneously. When they break down these radioisotopes emit radiation which is dangerous to living things. There are three di erent orms o radiation. Gamma () radiation is highly penetrating whereas alpha () radiation can be stopped by a ew centimetres o air and beta () radiation by a thin sheet o aluminium. Radioisotopes can occur naturally or be created arti cially. Their uses include nuclear power generation, the sterilization o surgical instruments in hospitals, crime detection, nding cracks and stresses in metals and the preservation o ood. ${}^{14}_{6}$ C is used or carbon dating, ${}^{60}_{27}$ Co is used in radiotherapy and ${}^{131}_{53}$ I and ${}^{125}_{53}$ I are used as tracers in medicine or treating and diagnosing illness.

Emissio s ectra

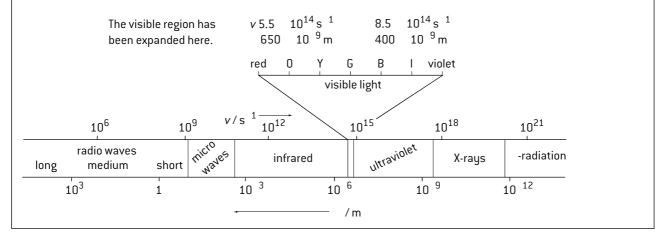
THE ELECTROMAGNETIC SPECTRUM

Electromagnetic waves can travel through space and, depending on the wavelength, also through matter. The velocity o travel *c* is related to its wavelength and its requency *v*. Velocity is measured in m s⁻¹, wavelength in m and requency in s⁻¹ so it is easy to remember the relationship between them:

c = v

 $(m \ s^{-1})$ (m) (s^{-1})

Electromagnetic radiation is a orm o energy. The smaller the wavelength and thus the higher the requency the more energy the wave possesses. Electromagnetic waves have a wide range o wavelengths ranging rom low energy radio waves to high energy -radiation. Visible light occupies a very narrow part o the spectrum.

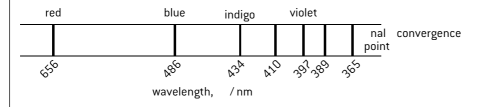


ATOMIC EMISSION SPECTRA

White light is made up o all the colours o the spectrum. When it is passed through a prism a **continuous spectrum** o all the colours can be obtained.

When energy is supplied to individual elements they emit a spectrum which only contains emissions at particular wavelengths. Each element has its own characteristic spectrum known as a **line spectrum** as it is not continuous.

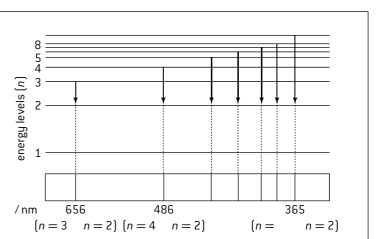
The visible hydrogen spectrum



Note that the spectrum consists o discrete lines and that the lines converge towards the high energy (violet) end o the spectrum. A similar series o lines at even higher energy also occurs in the ultraviolet region o the spectrum and several other series o lines at lower energy can be ound in the in rared region o the spectrum.

EXPLANATION OF EMISSION SPECTRA

When energy is supplied to an atom electrons are excited (gain energy) rom their lowest (ground) state to an excited state. Electrons can only exist in certain fixed energy levels. When electrons drop rom a higher level to a lower level they emit energy. This energy corresponds to a particular wavelength and shows up as a line in the spectrum. When electrons return to the first level (n = 1) the series o lines occurs in the ultraviolet region as this involves the largest energy change. The visible region spectrum is ormed by electrons dropping back to the n = 2 level and the first series in the in rared is due to electrons alling to the n = 3 level. The lines in the spectrum converge because the energy levels themselves converge.



E ectro ic co fi uratio

TYPES 0 ORBITAL

Electrons are ound in orbitals. Each orbital can contain a maximum o two electrons each with opposite spins. The first level (n = 1) contains just one orbital, called an s orbital. The second level (n = 2) contains one s orbital and three p orbitals. The 2p orbitals are all o equal energy but the sub-level made up o these three 2p orbitals is slightly higher in energy than the 2s orbital.

Principal level (shell)	Number	Number of each type of orbital		pe of orbital	Maximum number of		Relative energies of
п	S	р	d		electrons in level ($=2n^2$)		sub-levels within an atom
1	1				2	1	
2	1	3			8		$\frac{4s}{3s}$ $\frac{4p}{3p}$ $\frac{3d}{3d}$
3	1	3	5		18	Ŋ	
4	1	3	5	7	32	energy	2s ² p
The relative position o all the sub-levels or the first our main energy levels is shown.							
Note that the 4s sub-level is below the 3d sub-level. This explains why the third level is sometimes stated to hold 8 or 18 electrons.							

ELECTRONIC CON IGURATION AND AU BAU PRINCIPLE

The electronic configuration can be determined by ollowing the au bau (building up) principle. The orbitals with the lowest energy are filled first. Each orbital can contain a maximum o two electrons. Orbitals within the same sub-shell are filled singly first this is known as Hund s rule,

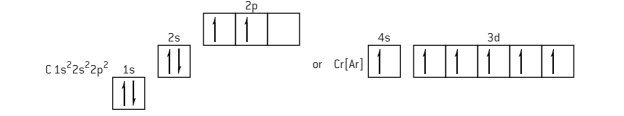
e.g.	H $1s^{1}$	Li $1s^{2}2s^{1}$	Na $1s^22s^22p^63s^1$	K 1s ² 2s ² 2p ⁶ 3s ² 3p ⁶ 4s ¹
c. <u>.</u> .	11 15	EI 15 25	nu 15 25 2p 55	R 15 25 2p 55 5p 15
	He 1s ²	Be $1s^22s^2$	$Mg \ 1s^2 2s^2 2p^6 3s^2$	Ca 1s ² 2s ² 2p ⁶ 3s ² 3p ⁶ 4s ²
		B $1s^22s^22p^1$	Al 1s ² 2s ² 2p ⁶ 3s ² 3p ¹	Sc 1s ² 2s ² 2p ⁶ 3s ² 3p ⁶ 4s ² 3d ¹
		C $1s^22s^22p^2$	Si 1s ² 2s ² 2p ⁶ 3s ² 3p ²	Ti 1s ² 2s ² 2p ⁶ 3s ² 3p ⁶ 4s ² 3d ²
		$N 1s^2 2s^2 2p^3$	P 1s ² 2s ² 2p ⁶ 3s ² 3p ³	V 1s ² 2s ² 2p ⁶ 3s ² 3p ⁶ 4s ² 3d ³
		O 1s ² 2s ² 2p ⁴	$S 1s^2 2s^2 2p^6 3s^2 3p^4$	
		F $1s^22s^22p^5$	Cl 1s ² 2s ² 2p ⁶ 3s ² 3p ⁵	
		Ne $1s^22s^22p^6$	Ar 1s ² 2s ² 2p ⁶ 3s ² 3p ⁶	

To save writing out all the lower levels the configuration may be shortened by building on the last noble gas configuration. For example, continuing on rom titanium, vanadium can be written [Ar]4s²3d³, then Cr [Ar]4s¹3d⁵, Mn [Ar]4s²3d⁵, Fe [Ar]4s²3d⁶ etc.

Note that the electron configurations o the transition metals show two irregularities. When there is the possibility o the d sub-level becoming hal - ull or completely ull it takes precedence over completely filling the 4s level first so chromium has the configuration $[Ar]4s^{1}3d^{5}$ (rather than $[Ar]4s^{2}3d^{4}$) and copper has the configuration $[Ar]4s^{1}3d^{10}$ (rather than $[Ar]4s^{2}3d^{9}$). When transition metals orm ions the 4s electrons are removed first so Fe²⁺ has the configuration $[Ar]3d^{6}$. The IB requires that you can write the configuration or any element or ion up to krypton (Z = 36). The ull electronic configuration or krypton is $1s^{2}2s^{2}2p^{6}3s^{2}3p^{6}4s^{2}3d^{10}4p^{6}$ or it can be shortened to $[Ar]4s^{2}3d^{10}4p^{6}$.

(When writing electronic configurations check that or a neutral atom the sum o the superscripts adds up to the atomic number o the element.)

Sometimes boxes are used to represent orbitals so the number o unpaired electrons can easily be seen, e.g.



SHAPES O ORBITALS

An electron has the properties o both a particle and a wave. Heisenberg s uncertainty principle states that it is impossible to know the exact position o an electron at a precise moment in time. An orbital describes the three-dimensional shape where there is a high probability that the electron will be located.

dimensional shape where there is a high probability that p_x s p_x s orbitals are spherical and the three p orbitals are orthogonal (at right angles) to each other.

🚥 Evidence from ionization energies

EVIDENCE FROM IONIZATION ENERGIES

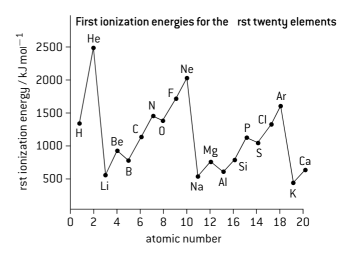
The first ionization energy o an element is defined as the energy required to remove one electron rom an atom in its gaseous state. It is measured in kJ mol⁻¹.

$$X(g) = X^+(g) + e^{-2g}$$

A graph o first ionization energies plotted against atomic number shows a repeating pattern.

It can be seen that the highest value is or helium, an atom that contains two protons and two electrons. The two electrons are in the lowest level and are held tightly by the two protons. For lithium it is relatively easy to remove an electron, which suggests that the third electron in lithium is in a higher energy level than the first two. The value then generally increases until element 10, neon, is reached be ore it drops sharply or sodium. This graph provides evidence that the levels can contain di erent numbers o electrons be ore they become ull.

Electrons with opposite spins tend to repel each other. When orbitals o the same energy (degenerate) are filled



the electrons will go singly into each orbital first be ore they pair up to minimize repulsion. This explains why there is a regular increase in the first ionization energies going rom B to N as the three 2p orbitals each gain one electron. Then there is a slight decrease between N and O as one o the 2p orbitals gains a second electron be ore a regular increase again.

EVIDENCE FOR SUB-LEVELS

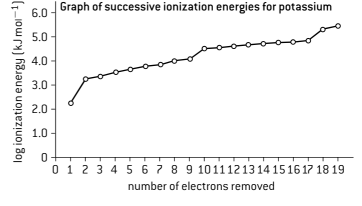
The graph already shown above was or the first ionization energy or the first 20 elements. Successive ionization energies or the same element can also be measured, e.g. the second ionization energy is given by:

$$X^{+}(g) = X^{2+}(g) + e^{-g}$$

As more electrons are removed the pull o the protons holds the remaining electrons more tightly so increasingly more energy is required to remove them, hence a logarithmic scale is usually used. A graph o the successive ionization energies or potassium also provides evidence o the number o electrons in each main level.

By looking to see where the first large jump occurs in successive ionization energies one can determine the number o valence electrons (and hence the group

in the periodic table to which the element belongs).



I the graph or first ionization energies is examined more closely then it can be seen that the graph does not increase regularly. This provides evidence that the main levels are split into sub-levels.

IONIZATION ENERGIES FROM EMISSION SPECTRA

It can be seen rom the emission spectrum o hydrogen that the energy levels converge. Hydrogen contains just one electron, which will be in the lowest energy level in its ground state. I su ficient energy is supplied it can be promoted to the infinite level that is it has been removed rom the atom and the atom has become ionized to orm the H⁺ ion. This amount o energy corresponds to the energy it would emit i it ell back rom n = to n = 1 which produces a line in the ultraviolet region o the spectrum at a wavelength o 91.2 nm. We can use this value to calculate the energy involved. Wavelength and requency are related by the expression c = v where *c* is the velocity o light. Energy and requency are related by the expression E = hv where *h* is Planck s constant and has the value 6.63 10^{-34} J s.

The energy to remove one electron = $hv = \frac{hc}{m} = \frac{6.63 \ 10^{-34} \text{ J s}}{91.2 \ 10^{-9} \text{ m}} = 2.18 \ 10^{-18} \text{ J}$

For one mole o electrons we need to multiply by Avogadro's constant (6.02 10^{23}) to give 1.312 10^{6} J mol⁻¹ or 1312 kJ mol⁻¹. This value is exactly the same as the experimentally determined value or the first ionization energy o hydrogen.

MULTIPLE CHOICE QUESTIONS ATOMIC STRUCTURE

1. Which o the ollowing particles contain more neutrons than electrons?

I. ${}^{1}_{1}H^{+}$	II. ⁷⁹ ₃₅ Br	III. ${}^{23}_{11}Na^+$
A. I and II on	ly	C. II and III only
B. I and III of	nly	D. I, II and III

- 2. Which one o the ollowing sets represents a pair o isotopes?
 - A. ${}^{31}_{15}P$ and ${}^{32}_{15}P$ C. Diamond and C_{40}
 - B. ${}^{24}_{12}Mg$ and ${}^{24}_{12}Mg^{2+}$ D. ${}^{40}_{18}Ar$ and ${}^{40}_{20}Ca$
- 3. Which species contains 16 protons, 17 neutrons and 18 electrons?

A. ³² S	C. ³⁴ S
B. ³³ S ²	D. ³⁵ S ²

- 4. Which quantities are the same or all atoms o chlorine?
 - I. Number o protons
 - II. Number o electrons
 - III. Number o neutrons
 - A. I and II only C. II and III only
 - B. I and III only D. I, II and III
- 5. A sample o zinc has the ollowing composition:

Isotope	% abundance
⁶⁴ Zn	55
66Zn	40
⁶⁸ Zn	5

What is the relative atomic mass o the zinc in this sample?

A. 64.5	C. 65.9
B. 65.0	D. 66.4

- 11. Which is the correct definition or the second ionization energy o carbon?
 - A. C(s)C(s) + 2eC. $C^+(s)$ $C^{2+}(s) + 2e$ B. C(g) $C(g) + 2e^-$ D. $C^+(g)$ $C^{2+}(g) + 2e$
- 12. The first five ionization energies, in kJ mol⁻¹, or a certain element are 577, 1980, 2960, 6190 and 8700 respectively. Which group in the periodic table does this element belong to?

A. 1	C. 3
B. 2	D. 4

13. Which transition in the hydrogen emission spectrum corresponds to the first ionization energy o hydrogen?

A. <i>n</i> =	n = 1	C. <i>n</i> =	<i>n</i> = 2
B. <i>n</i> = 2	n = 1	D. <i>n</i> = 4	<i>n</i> = 2

- 14. Which ionization requires the most energy?

6. In the electromagnetic spectrum, which will have the shortest wavelength **and** the greatest energy?

	Shortest wavelength	Greatest energy
А.	ultraviolet	ultraviolet
в.	in rared	in rared
C.	ultraviolet	in rared
D.	in rared	ultraviolet

7. Which electronic transition in a hydrogen atom releases the most energy?

A.
$$n = 1$$
 $n = 2$
B. $n = 7$ $n = 6$
C. $n = 6$ $n = 7$
D. $n = 2$ $n = 1$

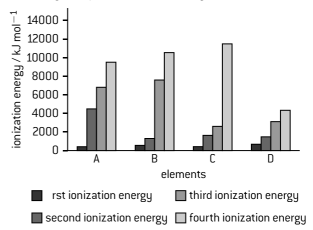
- 8. Which shows the sub-levels in order o **increasing** energy in the ourth energy level o an atom?
 - $\begin{array}{ll} A. & < d < p < s \\ B. \ p < d < & < s \\ C. \ d < & < p < s \\ D. \ s < p < d < \end{array}$
- 9. What is the electron configuration o copper?

A. $[Ar]4s^23d^9$ C. $1s^22s^22p^63s^23p^64s^13d^9$	d^{10}
---	----------

- B. $1s^22s^22p^63s^23p^63d^{10}$ D. [Ar] $3d^9$
- 10. How many unpaired electrons are present in an atom o sul ur in its ground state?

A. 1	С.	4
B. 2	D.	6

- 15. All o the ollowing actors a ect the value o the ionization energy o an atom **except** the:
 - A. mass o the atom
 - B. charge on the nucleus
 - C. size o the atom
 - D. main energy level rom which the electron is removed
- 16. The graph below shows the first our ionization energies o our elements A, B, C and D (the letters are not their chemical symbols). Which element is magnesium?



SHORT ANSWER QUESTIONS ATOMIC STRUCTURE

[1]

[2]

[1]

[1]

[2]

[1]

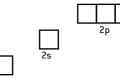
[2]

[2]

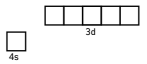
- a) Define the term relative atomic mass, A_r.
 b) The relative atomic mass o naturally occurring chlorine is 35.45. Calculate the abundances o ³⁵Cl and ³⁷Cl in naturally occurring chlorine.
 - c) (i) State the electron configuration o chlorine.(ii) State the electron configuration or a chloride ion. Cl.
 - d) Explain how ³⁵Cl and ³⁷Cl di er in their chemical properties.
- a) Explain why the relative atomic mass o cobalt is greater than the relative atomic mass o nickel, even though the atomic number o nickel is greater than the atomic number o cobalt.
 - b) Deduce the numbers o protons and electrons in the Co²⁺ ion. [1]
 - c) (i) Deduce the electron configuration o the Co atom. [1]
 (ii) Deduce the electron configuration o the Co²⁺ ion. [1]
- 3. $\frac{99}{43}$ Tc is a radioactive isotope o technetium.
 - a) (i) Define the term isotope. [1]
 - (ii) Determine the number o neutrons in one atom o technetium-99. [1]
 - (iii) Technetium-99 is used as a tracer in medicine.Suggest a reason why it is potentially dangerous. [2]
 - b) Carbon in living organisms consists o two isotopes,
 ¹²C and ¹⁴C, in a fixed ratio. This ratio remains constant in a living organism as the carbon is constantly being replaced through photosynthesis. Once an organism dies the ¹⁴C slowly decays to ¹⁴N with a hal -li e o 5300 years.
 - (i) Identi y the number o protons, neutrons and electrons in carbon-12 and in carbon-14. [2]

 (ii) Suggest the identity o the particle that is emitted when an atom o ¹⁴C is converted into an atom o ¹⁴N. [1]

- (iii) Discuss how the decay o carbon-14 can be used in carbon dating. [2]
- 4. Annotate the 2s and 2p boxes, using or to represent a spinning electron, to complete the electron configuration or an oxygen atom. [1]



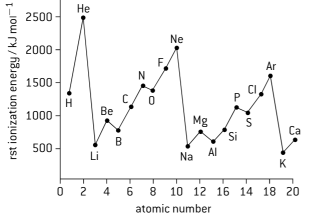
- Draw and label an energy level diagram or the hydrogen atom. In your diagram show how the series o lines in the ultraviolet and visible regions o its emission spectrum are produced, clearly labelling each series. [4]
- The electron configuration o chromium can be expressed as [Ar]4s^x3d^y.
 - (i) Explain what the square brackets around argon,[Ar], represent. [1]
 - (ii) State the values o x and y. [1]
 - (iii) Annotate the diagram below showing the 4s and 3d orbitals or a chromium atom using an arrow, or , to represent a spinning electron. [1]



H

7. The graph below shows the first ionization energy plotted against atomic number or the first 20 elements.

First ionization energies for the rst twenty elements



a) Define the term *first ionization energy*.

- b) Explain the ollowing:
 - (i) why there is a general increase in the value or the first ionization energy across period 2 rom Li to Ne

- (ii) why the first ionization energy o neon is higher than that o sodium [2]
- (iii) why the first ionization energy o beryllium ishigher than that o boron [2]
- (iv) why the first ionization energy o sul ur is lower than that o phosphorus. [2]
- c) Predict how the graph or the second ionization energy plotted against atomic number or the first 20 elements di ers rom the graph shown above. [3]
- Electrons are much too small to ever be seen . Discuss the evidence that electrons exist in fixed energy levels and that these levels can be split into sub-levels. [5]
- 9. The first ionization energy o hydrogen is 1312 kJ mol⁻¹. Determine the requency and wavelength o the convergence line in the ultraviolet emission spectrum o hydrogen. (Use in ormation given in Sections 1 and 2 o the IB data booklet.) [3]

3 PERIODICITY

T t

2 	3	4				ar			THE PERIODIC TABLE											
	3	4				g	oup nu 人	mber												
 ock		•	5	6	7	8	9	10	11	12	13	14	15	16	17	18				
													n-h	lock						
				_			_						P 8	lock		² He				
4 Be							r				5 B	е С	7 N	8 0	9 F	¹⁰ Ne				
12 Mg					d-b	lock					13 Al	¹⁴ Si	15 P	16 S	17 CI	18 Ar				
20 Ca	²¹ Sc	22 Ti	23 V	24 Cr	²⁵ Mn	26 Fe	27 Co	28 Ni	29 Cu	³⁰ Zn	31 Ga	32 Ge	33 As	³⁴ Se	35 Br	36 Kr				
38 Sr	³⁹ Ү	40 Zr	41 N b	42 Mo	43 Tc	44 Ru	45 Rh	46 Pd	47 Ag	48 C d	49 In	50 Sn	51 Sb	52 Te	53 	⁵⁴ Xe				
56 Ba	⁵⁷ * La	72 H f	73 Ta	74 W	75 Re	76 Os	77 Ir	⁷⁸ Pt	79 Au	⁸⁰ Hg	81 Ti	⁸² Pb	83 Bi	⁸⁴ Po	⁸⁵ At	86 Rn				
⁸⁸ Ra	⁸⁹ * Ac *	104 Rf	105 Db	106 Sg	107 Bh	¹⁰⁸ Hs	109 Mt	110 Ds	111 Rg	112 Cn	¹¹³ Uut	¹¹⁴ Uuq	115 Uup	¹¹⁶ Uuh	¹¹⁷ Uus	¹¹⁸ Uuo				
f-block																				
	*	58 Ce	⁵⁹ Pr	60 Nd	⁶¹ Pm	62 Sm	63 Eu	64 G d	65 Tb	⁶⁶ Dу	67 Ho	68 Er	69 Tm	70 Yb	71 Lu					
	* *	90 Th	91 Pa	92 U	93 Np	94 Pu	95 Am	96 Cm	97 Bk	98 Cf	99 Es	100 Fm	¹⁰¹ M d	102 No	103 Lr					
	Be 12 Mg 20 Ca 38 Sr 56 Ba 88	Be 12 Mg 20 21 Ca Sc Sc 38 39 Sr Y 56 57 Ba La 88 89 88 89 Ra Ac *	Be ¹² Mg ²⁰ ²¹ ²⁰ ²¹ ²² ²⁰ ²¹ ²¹ ²² ²¹ ²² ²¹ ²² ²¹ ²² ²¹ ²² ²¹ ²² ²¹ ²² ²¹ ²² ²¹ ²² ²¹ ²² ²¹ ²² ²¹ ²² ²¹ ²² ²¹ ²² ²¹ ²² ²¹ ²² ²¹ ²² ²¹ ³⁸ ³⁹ ³⁹ ⁴⁰ ^{2r} ²⁷ ²⁷ ⁵⁶ ⁵⁷ ⁸⁸ ⁸⁹ ⁸⁹ ⁸⁹ ⁸¹ ⁴⁰ ^{2r} ⁵⁶ ⁵⁷ ⁸⁸ ⁸⁹ ⁸⁹ ⁸⁹ ⁸¹ ¹⁰⁴ ⁸⁸ ⁸⁸ ⁸⁸ ⁸⁹ ⁸ ⁸ ⁸ ⁵⁷ ⁷² ⁵⁸ ⁵⁷ ⁸⁷ ⁵⁸ ⁵⁸ ⁵⁷ ⁸⁷ ⁵⁸	Be I2 I2 <thi2< th=""> I2 I2 I2<!--</td--><td>Be 12 Mg 20 21 22 23 24 Ca Sc Ti V Cr 38 39 40 41 42 Sr Y Zr Nb Mo 56 57 * 72 73 74 Ba La Hf Ta W 88 89 * 104 105 106 Ra Ac * Rf Db Sg * 90 91 92 91 * Tb Pc 11</td><td>Be eler 12 Mg </td><td>Be element 12 Mg </td><td>Be element 12 Mg d-block 20 21 22 23 24 25 26 27 Ca Sc Ti V Cr Mn Fe Co 38 39 40 41 42 43 44 45 Sr Y Zr Nb Mo Tc Ru Rh 56 57 * 72 73 74 75 76 77 Ba La Hf Ta W Re Os Ir 88 89 * 104 105 106 107 108 109 Ra Ac * Rf Db Sg Bh Hs Mt * Tb P2 93 94 95</td><td>Be element 12 Mg 4<td>Be element 12 Mg </td><td>Be element 12 Mg </td><td>Be element B 12 Mg element 13 Al 20 21 22 23 24 25 26 27 28 29 30 31 Al 20 21 22 23 24 25 26 27 28 29 30 31 Al 20 21 22 23 24 25 26 27 28 29 30 31 Ga 38 39 40 41 42 43 44 45 46 47 48 49 Sr Y Zr Nb Mo Tc Ru Rh Pd Ag Cd In 56 57 * 72 73 74 75 76 77 78 79 80 81 Ba La Hf Ta W Re Os Ir Pt Au Hg Ti 88 89 * 104 105 Sg Bh Hs Mt Ds<!--</td--><td>Be element B C 12 Mg </td><td>Be element B C N 12 Mg </td><td>Be element B C N O 12 Mg </td><td>Be Image Im</td></td></td></thi2<>	Be 12 Mg 20 21 22 23 24 Ca Sc Ti V Cr 38 39 40 41 42 Sr Y Zr Nb Mo 56 57 * 72 73 74 Ba La Hf Ta W 88 89 * 104 105 106 Ra Ac * Rf Db Sg * 90 91 92 91 * Tb Pc 11	Be eler 12 Mg	Be element 12 Mg	Be element 12 Mg d-block 20 21 22 23 24 25 26 27 Ca Sc Ti V Cr Mn Fe Co 38 39 40 41 42 43 44 45 Sr Y Zr Nb Mo Tc Ru Rh 56 57 * 72 73 74 75 76 77 Ba La Hf Ta W Re Os Ir 88 89 * 104 105 106 107 108 109 Ra Ac * Rf Db Sg Bh Hs Mt * Tb P2 93 94 95	Be element 12 Mg 4 <td>Be element 12 Mg </td> <td>Be element 12 Mg </td> <td>Be element B 12 Mg element 13 Al 20 21 22 23 24 25 26 27 28 29 30 31 Al 20 21 22 23 24 25 26 27 28 29 30 31 Al 20 21 22 23 24 25 26 27 28 29 30 31 Ga 38 39 40 41 42 43 44 45 46 47 48 49 Sr Y Zr Nb Mo Tc Ru Rh Pd Ag Cd In 56 57 * 72 73 74 75 76 77 78 79 80 81 Ba La Hf Ta W Re Os Ir Pt Au Hg Ti 88 89 * 104 105 Sg Bh Hs Mt Ds<!--</td--><td>Be element B C 12 Mg </td><td>Be element B C N 12 Mg </td><td>Be element B C N O 12 Mg </td><td>Be Image Im</td></td>	Be element 12 Mg	Be element 12 Mg	Be element B 12 Mg element 13 Al 20 21 22 23 24 25 26 27 28 29 30 31 Al 20 21 22 23 24 25 26 27 28 29 30 31 Al 20 21 22 23 24 25 26 27 28 29 30 31 Ga 38 39 40 41 42 43 44 45 46 47 48 49 Sr Y Zr Nb Mo Tc Ru Rh Pd Ag Cd In 56 57 * 72 73 74 75 76 77 78 79 80 81 Ba La Hf Ta W Re Os Ir Pt Au Hg Ti 88 89 * 104 105 Sg Bh Hs Mt Ds </td <td>Be element B C 12 Mg </td> <td>Be element B C N 12 Mg </td> <td>Be element B C N O 12 Mg </td> <td>Be Image Im</td>	Be element B C 12 Mg	Be element B C N 12 Mg	Be element B C N O 12 Mg	Be Image Im				

FEATURES OF THE PERIODIC TABLE

In the periodic table elements are placed in order of increasing atomic number.

Elements are arranged into four blocks associated with the four sub-levels s, p, d and f.

The period number (*n*) is the outer energy level that is occupied by electrons.

Elements in the same vertical group contain the same number of electrons in the outer energy level.

The number of the principal energy level and the number of the valence electrons in an atom can be deduced from its position on the periodic table.

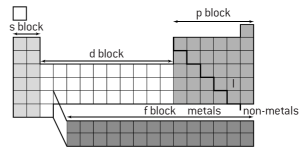
The periodic table shows the positions of metals, nonmetals and metalloids.

Certain groups have their own name. For example, group 1 is known as the alkali metals, group 17 as the halogens and group 18 as the noble gases.

The d-block elements (groups 3 to 12) are known as the transition metals and the f-block elements form two distinct groups the lanthanoids (from La to Lu) and the actinoids (from Ac to Lr).

Metals are on the left and in the centre of the table and non-metals are on the right (distinguished by the thick line). Metalloids, such as boron, silicon and germanium have properties intermediate between those of a metal and a non-metal.

ELECTRON CONFIGURATION AND THE PERIODIC TABLE



An element s position in the periodic table is related to its valence electrons so the electronic con guration of any element can be deduced from the table, e.g. iodine (Z = 53) is a p block element. It is in group 17 so its con guration will contain ns^2np^5 . If one takes H and He as being the rst period then iodine is in the fth period so n = 5. The full con guration for iodine will therefore be:

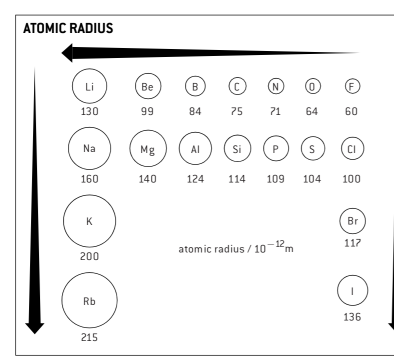
 $1s^22s^22p^63s^23p^64s^23d^{10}4p^65s^24d^{10}5p^5 \ or \ [Kr] \ 5s^24d^{10}5p^5$

P t n s (1)

PERIODICITY

Elements in the same group tend to have similar chemical and physical properties. There is a change in chemical and physical properties across a period. The repeating pattern of physical and chemical properties shown by the different periods is known as **periodicity**.

These periodic trends can clearly be seen in atomic radii, ionic radii, ionization energies, electronegativities, electron af nities and melting points.



The atomic radius is the distance from the nucleus to the outermost electron. Since the position of the outermost electron can never be known precisely, the atomic radius is usually de ned as half the distance between the nuclei of two bonded atoms of the same element.

As a group is descended, the outermost electron is in a higher energy level, which is further from the nucleus, so the radius increases.

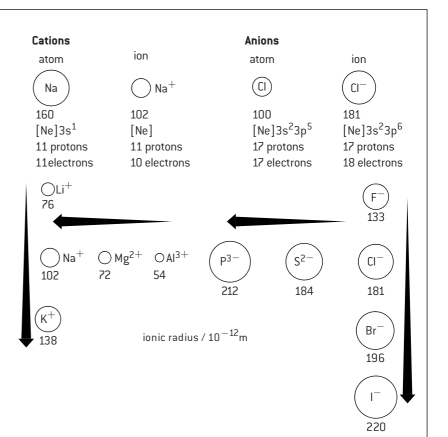
Across a period electrons are being added to the same energy level, but the number of protons in the nucleus increases. This attracts the energy level closer to the nucleus and the atomic radius decreases across a period.

IONIC RADIUS

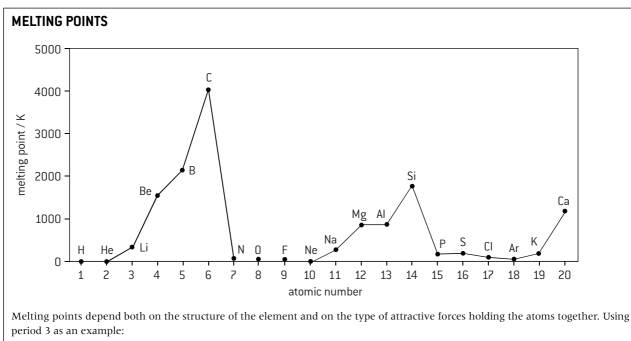
It is important to distinguish between positive ions (**cations**) and negative ions (**anions**). Both cations and anions increase in size down a group as the outer level gets further from the nucleus.

Cations contain fewer electrons than protons so the electrostatic attraction between the nucleus and the outermost electron is greater and the ion is smaller than the parent atom. It is also smaller because the number of electron shells has decreased by one. Across the period the ions contain the same number of electrons (**isoelectronic**), but an increasing number of protons, so the ionic radius decreases.

Anions contain more electrons than protons so are larger than the parent atom. Across a period the size decreases because the number of electrons remains the same but the number of protons increases.



Pdtnds(2)



At the left of the period elements exhibit metallic bonding (Na, Mg, Al), which increases in strength as the number of valence electrons increases.

Silicon, the metalloid, in the middle of the period has a macromolecular covalent structure with very strong bonds resulting in a very high melting point.

Elements in groups 15, 16, and 17 (P_4 , S_8 , and Cl_2) show simple molecular structures with weak intermolecular forces of attraction between the molecules.

The noble gases (Ar) exist as **monatomic molecules** (single atoms) with extremely weak forces of attraction between the atoms.

Within groups there are also clear trends:

In group 1 the melting point decreases down the group as the atoms become larger and the strength of the metallic bond decreases. Li Na K Rb Cs

M. pt / K	454	371	336	312	302
In group 17 the intermolecular a	ttractive force	s between	the diatom	nic molecul	les increase down the group so the melting points increase.
	F ₂	Cl ₂	Br ₂	I_{2}	
M. pt / K	53	172	266	387	

FIRST IONIZATION ENERGY

The de nition of rst ionization energy and a graph showing the values for the rst 20 elements has already been given on page 13. The values decrease down each group as the outer electron is further from the nucleus and therefore less energy is required to remove it, e.g. for the group 1 elements, Li, Na and K.

Element:	Li	Na	Κ
Electron con guration	$1s^{2}2s^{1}$	$[Ne]3s^1$	[Ar]4s ¹
First ionization energy (kJ mol ⁻¹)	520	496	419

Generally the values increase across a period. This is because the extra electrons are lling the same energy level and the extra protons in the nucleus attract this energy level closer making it harder to remove an electron.

The values do not increase regularly across a period because new sub-levels are being lled. The p sub-level is higher in energy than the s sub-level. This explains why the value for B $(1s^22s^22p^1)$ is slightly lower than the value for Be (Is^22s^2) and the value for Al $([Ne]3s^23p^1)$ is slightly lower than Mg $([Ne]3s^2)$. There is also a drop in value between N $(1s^22s^22p^3)$ and O $(1s^22s^22p^4)$ and between P $([Ne]3s^23p^3)$ and S $([Ne]3s^23p^4)$. This is because when electrons pair up in an orbital there is increased repulsion so the paired electron is easier to remove compared with when the three electrons are all unpaired, one each in the three separate p orbitals.

ELECTRONEGATIVITY

Electronegativity is a relative measure of the attraction that an atom has for a shared pair of electrons when it is covalently bonded to another atom. As the size of the atom decreases the electronegativity increases, so the value increases across a period and decreases down a group. The three most electronegative elements are F, N, and O.

ELECTRON AFFINITY

The electron af nity is the energy change when an electron is added to an isolated atom in the gaseous state, i.e.

$$X(g) + e^- = X^-(g)$$

Atoms want an extra electron so electron af nity values are negative for the addition of the rst electron. However, when oxygen forms the O^{2-} ion the overall process is endothermic.

	$O(g) + e^{-}$	O- (g)	$H^{\ominus} = -141 \text{ kJ mol}^{-1}$
	$O^-(g) + e^-$	$O^{2-}(g)$	$H^{\ominus} = +753 \text{ kJ mol}^{-1}$
overall	$O(g) + 2e^{-}$	$O^{2-}(g)$	$H^{\ominus} = +612 \text{ kJ mol}^{-1}$

CHEMICAL PROPERTIES OF ELEMENTS IN THE SAME GROUP

Group 1 the alkali metals



Ρ

Lithium, sodium, and potassium all contain one electron in their outer shell. They are all reactive metals and are stored under liquid paraf n to prevent them reacting with air. They react by losing their outer electron to form the metal ion. Because they can readily lose an electron they are good reducing agents. The reactivity increases down

reactivity the group as the outer electron is in successively higher energy levels and less energy is required to remove it.

They are called alkali metals because they all react with water to form an alkali solution of the metal hydroxide and hydrogen gas. Lithium oats and reacts quietly, sodium melts into a ball which darts around on the surface, and the heat generated from the reaction with potassium ignites the hydrogen.

 $2Li(s) + 2H_2O(l) = 2Li^+(aq) + 2OH^-(aq) + H_2(g)$

 $2Na(s) + 2H_2O(l) = 2Na^+(aq) + 2OH^-(aq) + H_2(g)$

 $2K(s) + 2H_2O(l) = 2K^+(aq) + 2OH^-(aq) + H_2(g)$

They all also react readily with chlorine, bromine and iodine to form ionic salts, e.g.

 $2Na(s) + Cl_{2}(g) = 2Na^{+}Cl^{-}(s)$

 $2K(s) + Br_2(l) = 2K^+Br^-(s)$

 $2Li(s) + I_2(g) = 2Li^+I^-(s)$

Group 17 the halogens

reactivity



The halogens react by gaining one more electron to form halide ions. They are good oxidizing agents. The reactivity decreases down the group as the outer shell is increasingly at higher energy levels and further from the nucleus. This, together with the fact that there are more electrons between the nucleus and the outer shell, decreases the attraction for an extra electron.

Chlorine is a stronger oxidizing agent than bromine, so can remove the electron from bromide ions in solution to form chloride ions and bromine. Similarly both chlorine and bromine can oxidize iodide ions to form iodine.

 $Cl_2(aq) + 2Br^-(aq) = 2Cl^-(aq) + Br_2(aq)$ $Cl_{2}(aq) + 2I^{-}(aq) = 2Cl^{-}(aq) + I_{2}(aq)$ $Br_2(aq) + 2I^-(aq) = 2Br^-(aq) + I_2(aq)$

Test for halide ions

The presence of halide ions in solution can be detected by adding silver nitrate solution. The silver ions react with the halide ions to form a precipitate of the silver halide. The silver halides can be distinguished by their colour. These silver halides react with light to form silver metal. This is the basis of old-fashioned lm photography.

> where X = Cl, Br, or I $Ag^+(aq) + X^-(aq)$ AgX(s) AgCl white light AgBr cream $Ag(s) + X_2$ AgI yellow

CHANGE FROM METALLIC TO NON-METALLIC NATURE OF THE ELEMENTS ACROSS PERIOD 3

Metals tend to be shiny and are good conductors of heat and electricity. Sodium, magnesium and aluminium all conduct electricity well. Silicon is a semiconductor and is called a metalloid as it possesses some of the properties of a metal and some of a non-metal. Phosphorus, sulfur, chlorine and argon are non-metals and do not conduct electricity. Metals can also be distinguished from non-metals by their chemical properties. Metal oxides tend to be basic, whereas non-metal oxides tend to be acidic.

Sodium oxide and magnesium oxide are both basic and react with water to form hydroxides,

 $Na_2O(s) + H_2O(l)$ 2NaOH(aq) $MgO(s) + H_2O(l)$ Mg(OH), e.g.

Aluminium is a metal but its oxide is amphoteric, that is, it can be either basic or acidic depending on whether it is reacting with an acid or a base.

The remaining elements in period 3 have acidic oxides. For example, sulfur trioxide reacts with water to form sulfuric acid, phosphorus pentoxide reacts with water to form phosphoric(V) acid and nitrogen(IV)oxide reacts with water to form nitric acid.

 $P_4O_{10}(s) + 6H_2O(l) - 4H_3PO_4(aq)$ $3NO_2(g) + H_2O(l) = 2HNO_3(aq) + NO(g)$ $SO_3(g) + H_2O(l) = H_2SO_4(aq)$

ம The transition metals

THE FIRST ROW TRANSITION ELEMENTS							i I		
Element (Sc) Ti V	Cr	Mn	Fe	Со	Ni	Cu	(Zn)		
Electron con guration [Ar] 4s ² 3d ¹ 4s ² 3d ² 4s ² 3d ³	$4s^13d^5$	4s²3d⁵	4s²3d6	4s²3d7	$4s^23d^8$	4s13d10	4s ² 3d ¹⁰		
A transition element is de ned as an element that possesses an incomplete d sub-level in one or more of its oxidation states. Scandium is not a typical transition metal as its common ion Sc ³⁺ has no d electrons. Zinc is not a transition metal as it contains a full d sub-level in all its oxidation states. (Note: for Cr and Cu it is more energetically favourable to half- ll and completely ll the d sub-level respectively so they contain only one 4s electron). Transition elements have variable oxidation states, form complex ions with ligands, have coloured compounds, and display catalytic and magnetic properties.									
VARIABLE OXIDATION STATES The 3d and 4s sub-levels are very similar in energy. W transition metals lose electrons they lose the 4s electron	ons rst.	Ma	CATALYTIC BEHAVIOUR Many transition elements and their compounds are very ef cient catalysts, that is, they increase the rate of chemical						
All transition metals can show an oxidation state of + of the transition metals can form the +3 or +4 ion (e Mn ⁴⁺) as the ionization energies are such that up to t electrons can also be lost. The M ⁴⁺ ion is rare and in t	the and	reactions. This helps to make industrial processes, such as the production of ammonia and sulfuric acid, more ef cient and economic. Platinum and palladium are used in catalytic converters tted to cars. In the body, iron is found in haem							
oxidation states the element is usually found not as the metal ion but either covalently bonded or as the oxyat such as MnO ₄ ⁻ . Some common examples of variable of	ne free mion,				itamin B ₁₂	. Other con	mmon examples		
states in addition to $+2$ are:		На	ber proces	s	$3H_{_2}(g) +$	$N_2 \stackrel{Fe(s)}{=} 2$	NH ₃ (g)		
Cr(+3) CrCl ₃ chromium(III) chlor	ide		nadium(V))		V ₂ O ₅	(s) $= 2SO_3(g)$		
$Cr(+6)$ $Cr_2O_7^{2-}$ dichromate(VI) ion			ide in the ntact proce		$O_{2}(g) + O$	$_{2}(g) =$	= 2SO ₃ (g)		
$Mn(+4)$ MnO_2 manganese(IV) oxid	e	: :	rhact proce	1.33					
$Mn(+7)$ MnO_4^- manganate(VII) ion			drogenatio	on C.H	$(g) + H_{2}(g)$	g) <u>Ni(s</u>	$) \longrightarrow C_2 H_6(g)$		
$Fe(+3)$ Fe_2O_3 iron(III) oxide		*	ctions		4(8)2(6/	1		
$Cu(+1)$ Cu_2O copper(I) oxideA full list of the common oxidation states can be fourSection 14 of the IB data booklet.	l l ox	anganese(I ide with drogen per	2H	$_{2}O_{2}(aq) - \frac{N}{2}$	$\xrightarrow{\text{AnO}_2(s)} 2$	$H_2O(l) + O_2(g)$			

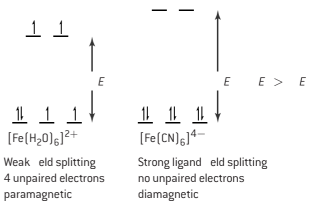
MAGNETIC PROPERTIES

Transition metals and their complexes that contain unpaired electrons can exhibit magnetism. Iron metal and some other metals (e.g. nickel and cobalt) show ferromagnetism. This is a permanent type of magnetism. In this type of magnetism the unpaired electrons align parallel to each other in domains irrespective of whether an external magnetic or electric eld is present. This property of iron has been utilized for centuries to make compasses, which align with the Earth's magnetic eld to point north.

Many complexes of transition metals contain unpaired electrons. Unlike paired electrons, where the spins cancel each other out, the spinning unpaired electrons create a small magnetic eld and will line up in an applied electric or magnetic eld to make the transition metal complex weakly magnetic when the eld is applied, i.e. they reinforce the external magnetic eld. This type of magnetism is known as paramagnetism. The more unpaired electrons there are in the complex the more paramagnetic the complex will be.

When all the electrons in a transition metal complex are paired up the complex is said to be diamagnetic.

Iron(II) complex ions ([Ar]3d⁶) can be paramagnetic or diamagnetic. The ve d orbitals are split by the ligands according to the spectrochemical series (see page 22). If the ligands are low in the spectrochemical series (e.g. H_2O) and split the d orbitals by only a small amount, the electrons can occupy all the d orbitals giving four units of paramagnetism as there will be four unpaired electrons. This is known as weak ligand eld splitting. Ligands high in the spectrochemical series (e.g. CN^-) will cause a larger splitting. Only the lower d orbitals will be occupied and the complex will be diamagnetic as there are no unpaired electrons.



_ _ _ _ _ _ _ _ _ _ _ _ _ _ _ _ _

🕕 Transition metal complex ions

FORMATION OF COMPLEX IONS

Because of their small size d-block ions act as Lewis acids and attract species that are rich in electrons. Such species are known as **ligands**. Ligands are neutral molecules or anions which contain a non-bonding pair of electrons. These electron pairs can form co-ordinate covalent bonds with the metal ion to form **complex ions**.

A common ligand is water and most (but not all) transition metal ions exist as hexahydrated complex ions in aqueous solution, e.g. $[Fe(H_2O)_6]^{3+}$. Ligands can be replaced by other ligands. A typical example is the addition of ammonia to an aqueous solution of copper(II) sulfate to give the deep blue colour of the tetraaminecopper(II) ion. Similarly if concentrated hydrochloric acid is added to a solution of Cu^{2+} (aq) the yellow tetrachlorocopper(II) anion is formed.

Note: in this ion the overall charge on the ion is -2 as the four ligands each have a charge of -1.

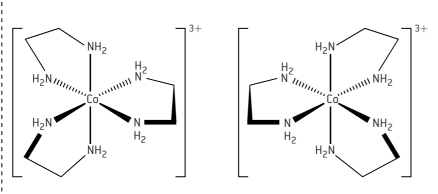
$$[\operatorname{CuCl}_4]^{2-} \xrightarrow[\text{H}_2O]{} [\operatorname{Cu}(\operatorname{H}_2O)_4]^{2+} \xrightarrow[\text{H}_2O]{} [\operatorname{Cu}(\operatorname{NH}_3)_4]$$

The number of lone pairs bonded to the metal ion is known as the **coordination number**. Compounds with a coordination number of six are octahedral in shape, those with a coordination number of four are tetrahedral or square planar, whereas those with a coordination number of two are usually linear.

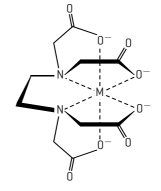
 $\begin{array}{c|c} \mbox{Coordination number } 6 & 4 & 2 \\ \mbox{Examples} & [Fe(CN)_6]^{3-} & [CuCl_4]^{2-} & [Ag(NH_3)_2]^+ \\ & [Fe(OH)_3(H_2O)_3][Cu(NH_3)_4]^{2+} \end{array}$

POLYDENTATE LIGANDS

Ligands such as water and cyanide ions are known as monodentate ligands as they utilize just one non-bonding pair to form a coordinate covalent bond to the metal ion. Some ligands contain more than one non-bonding pair and can form two or more coordinate bonds to the metal ion. Three common examples are ethylenediamine, $H_2NCH_2CH_2NH_2$, oxalate ions $(COO^-)_2$, both of which can use two non-bonding pairs to form bidentate ligands, and EDTA (EthyleneDiamineTetraAcetic acid) or its ion, EDTA⁴⁻, which can act as hexadentate ligands.



The Co³⁺ ion can form two different stereoisomers (mirror images) with the three separate ethylenediamine molecules acting as bidentate ligands. The coordination number is six as although each metal ion is surrounded by only three ligands the metal ion forms a total of six coordinate bonds with the ligands.



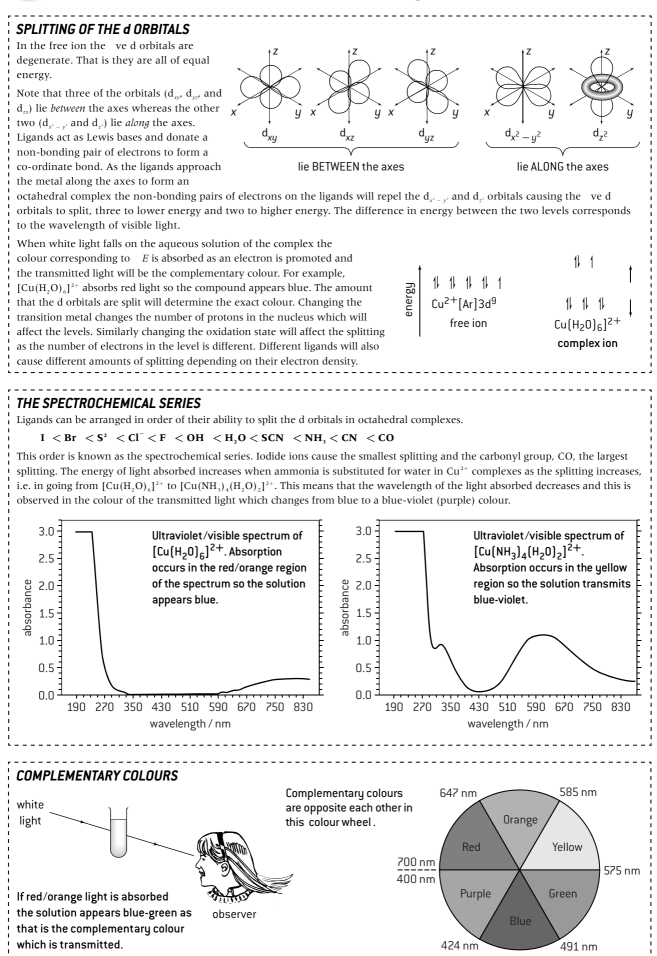
EDTA⁴⁻ acting as a hexadentate ligand with a transition metal, M to form a complex such as [Cu(EDTA)]²⁻. Note that the coordination number is still six even though only one ligand surrounds the metal ion.

FACTORS AFFECTING THE COLOUR OF TRANSITION METAL COMPLEXES

Transition metals are de ned as elements having an incomplete d sub-level in one or more of their oxidation states. Compounds of Sc³⁺ which have no d electrons and of Cu⁺ and Zn²⁺ which both have complete d sub-shells are colourless. This strongly suggests that the colour of transition metal complexes is related to an incomplete d level. The actual colour is determined by four different factors.

- 1. The nature of the transition element. For example $Mn^{2+}(aq)$ and $Fe^{3+}(aq)$ both have the conguration [Ar]3d⁵. $Mn^{2+}(aq)$ is pink whereas $Fe^{3+}(aq)$ is yellow.
- The oxidation state. Fe²⁺(aq) is green whereas Fe³⁺(aq) is yellow.
- The identity of the ligand. [Cu(H₂O)₆]²⁺ (sometimes shown as [Cu(H₂O)₄]²⁺), is blue, [Cu(NH₃)₄(H₂O)₂]²⁺ (sometimes shown as [Cu(NH₃)₄]²⁺), is blue/violet whereas [CuCl₄]²⁻ is yellow (green in aqueous solution).
- 4. The stereochemistry of the complex. The colour is also affected by the shape of the molecule or ion. In the above example [Cu(H₂O)₆]²⁺ is octahedral whereas [CuCl₄]²⁻ is tetrahedral. However for the IB only octahedral complexes in aqueous solution will be considered.

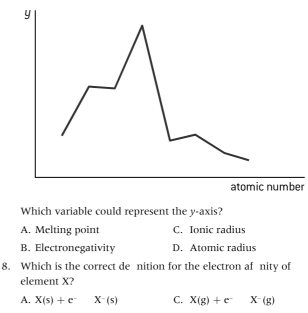
Colour of transition metal omplexes



MULTIPLE CHOICE QUESTIONS PERIODICITY

- 1. Where in the periodic table would the element with the electronic structure $1s^22s^22p^63s^23p^64s^23d^{10}4p^2$ be located?
 - A. s block C. d block
 - B. p block D. lanthanoids
- 2. Which property decreases with increasing atomic number in group 17 (the halogens)?
 - A. Melting point
 - B. Electronegativity
 - C. Atomic radius
 - D. Ionic radius of the negative ion
- 3. Which property increases with increasing atomic number for both the halogens and the alkali metals?
 - A. Reactivity with water C. Electron af nity
 - B. Electronegativity D. Atomic radius
- 4. Which one of the following series is arranged in order of increasing size?
 - A. P³⁻, S²⁻, Cl⁻ B. Cl, Ar, K D. H⁺, H, H⁻
- 5. Which element has the lowest electronegativity value?
 - A. potassium C. iodine
 - B. uorine D. hydrogen
- 6. Which statements about the periodic table are correct?
 - I. The elements Mg, Ca and Sr have similar chemical properties.
 - II. Elements in the same period have the same number of main energy levels.
 - III. The oxides of Na, Mg and P are basic.
 - A. I and II only C. II and III only
 - B. I and III only D. I, II and III

7. The *x*-axis of the graph below represents the atomic number of the elements in period 3.



- $B. \ X(s) \qquad X^{\scriptscriptstyle +}(s) + e^{\scriptscriptstyle -} \qquad \qquad D. \ X(g) \qquad X^{\scriptscriptstyle +}(g) + e^{\scriptscriptstyle -}$
- 9. Which oxide will react with water to give a solution with a pH greater than 7?
 - A. NO₂ C. MgO
- B. P_4O_{10} D. SiO_2
- 10. Which reaction does **not** occur readily?
 - A. $Cl_2(aq) + 2Br^-(aq) = Br_2(aq) + 2Cl^-(aq)$
 - B. $2Na(s) + H_2O(l) = 2NaOH(aq) + H_2(g)$
 - C. $Na_2O(s) + H_2O(l) = 2NaOH(aq)$
 - D. $I_2(aq) + 2Br^-(aq) = Br_2(aq) + 2l^-(aq)$

HL

- 11. Which complex ion is colourless in aqueous solution?
 - A. $[Mn(H_2O)_6]^{2+}$ C. $[Cu(NH_3)_4]^{2+}$
 - B. $[Zn(H_2O)_4]^{2+}$ D. $[CuCl_4]^{2-}$
- 12. A certain element has the electron conguration 1s²2s²2p⁶3s²3p⁶4s²3d³. What oxidation state(s) would this element most likely show?

A. $+2$ only	C. $+2$ and $+5$ only
B. $+3$ only	D. +2, +3, +4 and +5

13. What is the overall charge on the complex ion formed by Fe(III) and six CN⁻ ligands?

A. +3 C. -3 B. -6 D. +6

- 14. Which statements about transition metal complexes are correct?
 - I. The colour of the complex is due to light being emitted when an electron falls from a higher split d sub-level to a lower split d sub-level.
 - II. The difference in energy between the split d sub-levels depends on the nature of the surrounding ligands.
 - III. The colour of the complex is in uenced by the oxidation state of the metal.
 - A. I and II only C. II and III only
 - B. I and III only D. I, II and III

- 15. What is the electron con guration of the Mn^{2+} ion?
 - A. 1s²2s²2p⁶3s²3p⁶3d⁵
 - B. 1s²2s²2p⁶3s²3p⁶4s²3d⁵
 - C. 1s²2s²2p⁶3s²3p⁶4s²3d³
 - D. 1s²2s²2p⁶3s²3p⁶4s¹3d⁴
- 16. Which statement is true for all polydentate ligands?
 - A. They contain more than one pair of non-bonding electrons.
 - B. They can only form one coordinate bond with a transition metal ion.
 - C. They do not affect the size of the splitting of the d sub-level.
 - D. They can only form complexes with transition metal ions.

SHORT ANSWER QUESTIONS PERIODICITY

1.	Carbon, silicon and tin belong to the same group in the periodic table.	
	a) Distinguish between the terms <i>group</i> and <i>period</i> in terms of electron con guration.	[2]
	b) State in which block of the periodic table (s, p, d or f) these three elements are located.	[1]
	c) Explain why the rst ionization energy of silicon is lower than that of carbon.	[2]
	 d) Describe a simple experiment to show the change in non-metallic to metallic properties as the atomic number of the elements in the group increases. 	[3]
2.	a) The maximum number of electrons in each energy level is determined by the expression 2n ² where n is the number of the level. Explain why the third level is sometimes said to contain eight electrons and sometimes said to contain eighteen electrons.	[2]
	b) In terms of electron con guration describe the essential difference between the lanthanoids and the actinoids.	[2]
3.	a) Describe how the acid base nature of the oxides changes across period 3 (Na Cl).b) State the equation for the reaction of water with	[3]
	/ *	[1]
	(ii) phosphorus(V) oxide, P_4O_{10}	[1]
	(iii) sulfur(VI) oxide, SO ₃	[1]
	c) Explain why oxides of nitrogen and sulfur cause damage to many types of buildings. Include relevant	
4	* * *	[4]
4.	a) State the electron conguration for the P^{3-} ion.	[1]
	 b) Explain why the ionic radius of the S²⁻ ion is smaller than that of the P³⁻ ion. 	[2]

	c) Potassium and bromine are in the same period. Explain why the atomic radius of bromine is considerably smaller than the atomic radius of potassium.	[2]	
	d) Suggest a reason why no value is given for the atomi radius of neon in the IB data booklet.	c [1]	
5.	The periodic table shows the relationship between electron con guration and the properties of elements and is a valuabl tool for making predictions in chemistry.		
	a) (i) Identify the property used to arrange elements in the periodic table.	[1]	
	 (ii) Outline two reasons why electronegativity increases across period 3 in the periodic table and one reason why noble gases are not assigne electronegativity values. 	d [3]	
	b) (i) De ne the term rst ionization energy of an atom	n. [2]	
	(ii) Explain the general increasing trend in the rst ionization energies of the period 2 elements,		
	Li to Ne.	[2]	
	(iii) Explain why the rst ionization energy of borom is lower than that of beryllium.	[2]	
	 c) Explain why sodium conducts electricity but phosphorus does not. 	[2]	
6	In many cities around the world, public transport	[4]	

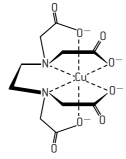
6. In many cities around the world, public transport vehicles use diesel, a liquid hydrocarbon fuel, which often contains sulfur impurities and undergoes incomplete combustion. All public transport vehicles in New Delhi, India, have been converted to use compressed natural gas (CNG) as fuel. Suggest two ways in which this improves air quality, giving a reason for your answer. [3]



- Transition elements, such as iron, have many characteristic properties and uses.
 - a) State and explain in terms of **acid base properties** the type of reaction that occurs between Fe^{2+} ions and water to form $[Fe(H_2O)_6]^{2+}$. [2]

[2]

- b) Explain why the colour of $[Fe(H_2O)_6]^{2+}$ is different to that of $[Fe(H_2O)_6]^{3+}$.
- c) Explain why iron metal is magnetic and why some of its complexes such as [Fe(H₂O)₆]²⁺ are paramagnetic whereas others such as [Fe(CN)₆]⁴⁻ are diamagnetic. [4]
- d) The Haber process to form ammonia is exothermic. Explore the economic signi cance of using an iron catalyst. [3]
- 8. The diagram shows the $[Cu(EDTA)]^{4-}$ ion.



a) Deduce the formula mass of this complex ion.	[2]
b) State the coordination number and the shape of the ion.	[2]
c) Explain how the EDTA ion is able to bond to the C_{12}^{12} i.e.	[2]
Cu^{2+} ion.	[3]

9. Ligands can be listed in terms of their ability to split the d orbitals. This is known as the spectrochemical series.

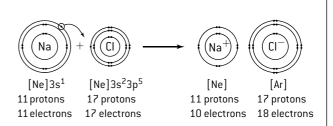
- a) Suggest a reason why different ligands cause the d orbitals to be split by different amounts. [2]
- b) Ammonia molecules are higher in the spectrochemical series (cause greater splitting) than water molecules. Explain why the light blue colour of $[Cu(H_2O)_6]^{2+}$ changes to a more purple colour when excess ammonia is added to copper(II) sulfate solution to form the $[Cu(NH_3)_4(H_2O)_2]^{2+}$ ion. [3]
- c) Explain why aqueous solutions of scandium(III) compounds are not coloured. [2]

4 CHEMICAL BONDING AND STRUCTURE

lonic bonding

IONIC BOND

When atoms combine they do so by trying to achieve a noble gas configuration. Ionic compounds are ormed when electrons are trans erred rom one atom to another to orm ions with complete outer shells o electrons. In an ionic compound the positive and negative ions are attracted to each other by strong electrostatic orces, and build up into a strong lattice. Ionic compounds have high melting points as considerable energy is required to overcome these orces o attraction.



The classic example o an ionic compound is sodium chloride Na^+Cl^- , ormed when sodium metal burns in chlorine. Chlorine is a covalent molecule, so each atom already has a noble gas configuration. However, the energy given out when the ionic lattice is ormed is su ficient to break the bond in the chlorine molecule to give atoms o chlorine. Each sodium atom then trans ers one electron to a chlorine atom to orm the ions.

The charge carried by an ion depends on the number o electrons the atom needed to lose or gain to achieve a ull outer shell.

	Cations			Anions	
Group 1	Group 2	Group 3	Group 15	Group 16	Group 17
+1	+2	+3	-3	-2	-1
Li^+ Na^+ K^+	$Mg^{_{2+}} Ca^{_{2+}}$	$\mathrm{Al}^{_{3+}}$	N ³⁻ P ³⁻	$O^{2-} S^{2-}$	$F^ Cl^ Br^-$

Thus, in magnesium chloride, two chlorine atoms each gain one electron rom a magnesium atom to orm $Mg^{2+}Cl_2^-$. In magnesium oxide two electrons are trans erred rom magnesium to oxygen to give $Mg^{2+}O^{2-}$. Transition metals can orm more than one ion. For example, iron can orm Fe^{2+} and Fe^{3+} and copper can orm Cu^+ and Cu^{2+} .

FORMULAS OF IONIC COMPOUNDS

It is easy to obtain the correct ormula as the overall charge o the compound must be zero.

lithium uoride $Li^+F^$ sodium oxide Na^+, O^{2-}

calcium sulfide Ca²⁺S²⁻

magnesium chloride Mg²⁺Cl⁻

aluminium bromide Al³⁺Br⁻, iron(III) oxide Fe³⁺₂O²⁻, iron(II) oxide Fe²⁺O²⁻

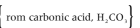
potassium nitride $K_{3}^{+}N^{3-}$ calcium phosphide $Ca_{3}^{2}P_{2}^{3-}$ iron(II) oxide $Fe^{2+}O^{2-}$ Note: the ormulas above have been written to show the charges carried by the ions. Unless asked specifically to do this it is common practice to omit the charges and simply write LiF, MgCl₂, etc.

IONS CONTAINING MORE THAN ONE ELEMENT (POLYATOMIC IONS)

In ions ormed rom more than one element the charge is o ten spread (delocalized) over the whole ion. An example o a positive ion is the ammonium ion NH_4^+ , in which all our N H bonds are identical. Negative ions are sometimes known as acid radicals as they are ormed when an acid loses one or more H^+ ions.

hydroxide OH⁻ nitrate NO⁻₃ sul ate SO₄²⁻

(rom nitric acid, HNO₃) GO^{-} {rom sul uric acid, H₂SO₄} carbonate CO₃²⁻ hydrogencarbonate HCO₃⁻ ethanoate CH₃COO⁻ phosphate PO ³⁻



(rom ethanoic acid, CH₃COOH)

hydrogensul ate $HSO_4^ HSO_4^ HSO_4^-$

IONIC BOND AND PROPERTIES OF IONIC COMPOUNDS

Ionic compounds are ormed between metals on the let o the periodic table and non-metals on the right o the periodic table; that is, between elements in groups 1, 2, and 3 with a low electronegativity (electropositive elements) and elements with a high electronegativity in groups 15, 16, and 17. Generally the di erence between the electronegativity values needs to be greater than about 1.8 or ionic bonding to occur.

Ions in solid ionic compounds are held in a crystal lattice. The ionic bond is the sum o all the electrostatic attractions (and repulsions) within the lattice. A large amount o energy is required to break the lattice so ionic compounds tend to have high melting points. Many are soluble in water as the hydration energy o the ions provides the energy to overcome the lattice enthalpy. Solid ionic compounds cannot conduct electricity as the ions are held in fixed positions. When molten the ions are ree to move and conduct electricity as they are chemically decomposed at the respective electrodes.



Sodium chloride (melting point 801 C) lons held strongly in ionic lattice

Co alent bonding

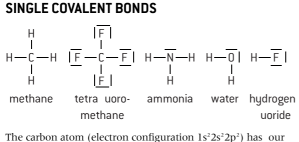
SINGLE COVALENT BONDS

Covalent bonding involves the sharing o one or more pairs o electrons so that each atom in the molecule achieves a noble gas configuration. The simplest covalent molecule is hydrogen. Each hydrogen atom has one electron in its outer shell. The two electrons are shared and attracted electrostatically by both positive nuclei resulting in a directional bond between the two atoms to orm a molecule. When one pair o electrons is shared the resulting bond is known as a single covalent bond. Another example o a diatomic molecule with a single covalent bond is chlorine, Cl₂.

LEWIS STRUCTURES

In the Lewis structure (also known as electron dot structure) all the valence electrons are shown. There are various di erent methods o depicting the electrons. The simplest method involves using a line to represent one pair o electrons. It is also acceptable to represent single electrons by dots, crosses or a combination o the two. The our methods below are all correct ways o showing the Lewis structure o uorine.

Sometimes just the shared pairs o electrons are shown, e.g. F F. This gives in ormation about the bonding in the molecule, but it is not the Lewis structure as it does not show all the valence electrons.



electrons in its outer shell and requires a share in our more electrons. It orms our single bonds with elements that only require a share in one more electron, such as hydrogen or chlorine. Nitrogen $(1s^22s^22p^3)$ orms three single bonds with hydrogen in ammonia leaving one non-bonded pair o electrons (also known as a lone pair). In water there are two non-bonded pairs and in hydrogen uoride three non-bonded pairs.

BOND LENGTH AND BOND STRENGTH

The strength o attraction that the two nuclei have or the shared electrons a ects both the length and strength o the bond. Although there is considerable variation in the bond lengths and strengths o single bonds in di erent compounds, double bonds are generally much stronger and shorter than single bonds. The strongest covalent bonds are shown by triple bonds.

		Length / nm	Strength / kJ mol ¹		
Single bonds	Cl Cl	0.199	242		
	C C	0.154	346		
Double bonds	C = C	0.134	614		
	0=0	0.121	498		
Triple bonds	C C	0.120	839		
	N N	0.110	945		
e.g. ethanoic acid: 0.124 nm 0 II C 0.143 nm					
	H-1	OH H double bond between C and O shorter and stronger than single bond			

MULTIPLE COVALENT BONDS

In some compounds atoms can share more than one pair o electrons to achieve an noble gas configuration.

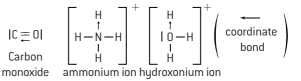
$$(0=0)$$
 |N=N| $(0=C=0)$ $\underset{H}{\overset{H}{\sim}}C=C\underset{H}{\overset{H}{\sim}}H-C=C-H$
oxygen nitrogen carbon dioxide ethene ethyne

CI

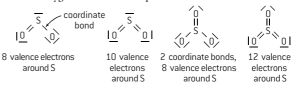
C

COORDINATE (DATIVE) BONDS

The electrons in the shared pair may originate rom the same atom. This is known as a coordinate covalent bond.

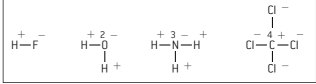


Sul ur dioxide and sul ur trioxide are both sometimes shown as having a coordinate bond between sul ur and oxygen or they are shown as having double bonds between the sul ur and the oxygen. Both are acceptable.



BOND POLARITY

In diatomic molecules containing the same element (e.g. H_2 or Cl_2) the electron pair will be shared equally, as both atoms exert an identical attraction. However, when the atoms are di erent the more electronegative atom exerts a greater attraction or the electron pair. One end o the molecule will thus be more electron rich than the other end, resulting in a polar bond. This relatively small di erence in charge is represented by + and -. The bigger the di erence in electronegativities the more polar the bond.

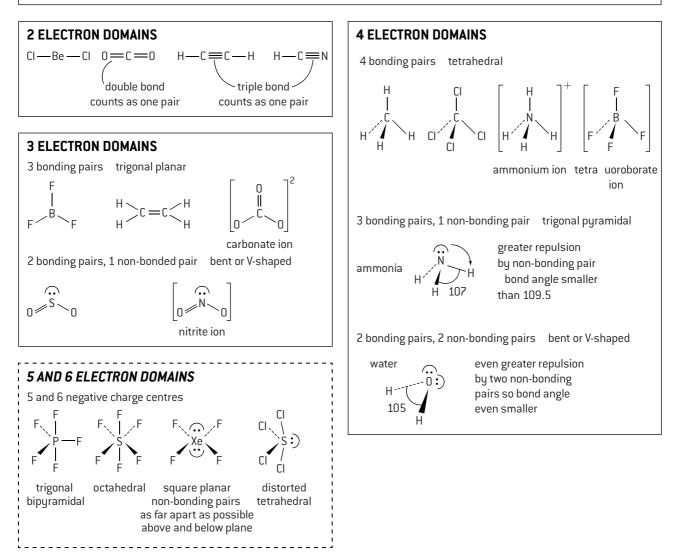


Shapes of simple molecules and ions

VSEPR THEORY The shapes o simple molecules and ions can be	No. of electron domains	Shape	Name of shape	Bond angle(s)
determined by using the valence shell electron	2	0-0-0	linear	180
pair repulsion (VSEPR) theory. This states that pairs o electrons arrange themselves around the central atom so that they are as ar apart rom each other as possible. There will be greater	3		trigonal planar	120
repulsion between non-bonded pairs o electrons than between bonded pairs. Since all the electrons in a multiple bond must lie in the same direction, double and triple bonds count as one pair o electrons. Strictly speaking the theory re ers to	4		tetrahedral	109.5
electron domains, but or most molecules this equates to pairs o electrons. This results in five basic shapes depending on the number o pairs. For 5 and 6 electron domains	5		trigonal bipyramidal	90 , 120 , 180
the octet needs to be expanded and this can only happen i there are readily available d orbitals present that can also be utilized. For Standard Level only 2, 3 and 4 electron domains need to be considered.	6		octahedral	90 , 180

WORKING OUT THE ACTUAL SHAPE

To work out the actual shape o a molecule calculate the number o pairs o electrons around the *central* atom, then work out how many are bonding pairs and how many are non-bonding pairs. (For ions, the number o electrons that equate to the charge on the ion must also be included when calculating the total number o electrons.)



Resonance hybrids and allotropes of carbon

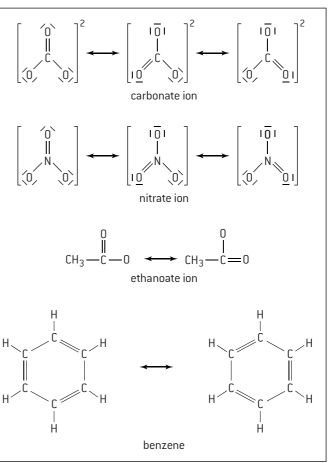
diamond

RESONANCE STRUCTURES

When writing the Lewis structures for some molecules it is possible to write more than one correct structure. For example, ozone can be written:

$$(\underline{0}^{\overline{0}},\underline{0}) \longleftrightarrow (\underline{0}^{\overline{0}},\underline{0})$$

These two structures are known as resonance hybrids. They are extreme forms of the true structure, which lies somewhere between the two. Evidence that this is true comes from bond lengths, as the bond lengths between the oxygen atoms in ozone are both the same and are intermediate between an O=O double bond and an O O single bond. Resonance structures are usually shown with a double headed arrow between them. Other common compounds which can be written using resonance structures are shown here.



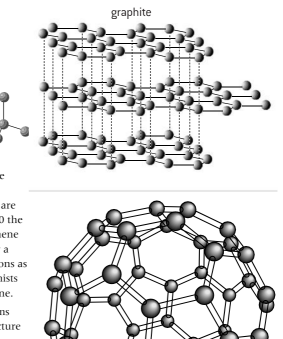
ALLOTROPES OF CARBON

Allotropes occur when an element can exist in different crystalline forms. In diamond each carbon atom is covalently bonded to four other carbon atoms to form a giant covalent structure. All the bonds are equally strong and there is no plane of weakness in the molecule so diamond is exceptionally hard, and because all the electrons are localized it does not conduct electricity. Both silicon and silicon dioxide, SiO₂, form similar giant tetrahedral structures.

In graphite each carbon atom has very strong

bonds to three other carbon atoms to give layers of hexagonal rings. There are only very weak bonds between the layers. The layers can slide over each other so graphite is an excellent lubricant and because the electrons are delocalized between the layers it is a good conductor of electricity. In 2010 the Nobel prize for Physics was awarded for the discovery of graphene. Graphene is a single layer of hexagonally arranged carbon atoms, i.e. it is essentially a form of graphite which is just one atom thick. It is extremely light, functions as a semiconductor and is 200 times stronger than steel. More recently chemists have developed a new magnetic form of graphene which is called graphone.

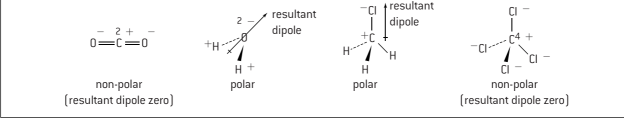
A third allotrope of carbon is buckminsterfullerene, $C_{_{\delta 0}}$. Sixty carbon atoms are arranged in hexagons and pentagons to give a geodesic spherical structure similar to a football. Following the initial discovery of buckminsterfullerene many other similar carbon molecules have been isolated. This has led to a new branch of science called nanotechnology.



Intermolecular forces

MOLECULAR POLARITY

Whether a molecule is polar, or not, depends both on the relative electronegativities o the atoms in the molecule and on its shape. I the individual bonds are polar then it does not necessarily ollow that the molecule will be polar as the resultant dipole may cancel out all the individual dipoles.



LONDON DISPERSION FORCES

Even in non-polar molecules the electrons can at any one moment be unevenly spread. This produces temporary instantaneous dipoles. An instantaneous dipole can induce another dipole in a neighbouring particle resulting in a weak attraction between the two particles. London dispersion orces increase with increasing mass.

Cl2

70.9

C₂H₆

30.0

886

34.0

increasing London dispersion forces

increasing London dispersion forces

Br₂ 160

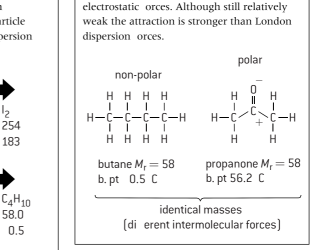
58.0

 C_3H_8

44.0

42 2

DIPOLE DIPOLE FORCES Polar molecules are attracted to each other by



INTERMOLECULAR FORCES

 F_2

Μ,

M,

b.pt/C

b.pt/C

38.0

 CH_4

16.0

162

188

The covalent bonds between the atoms within a molecule are very strong. The orces o attraction between the molecules are much weaker. These intermolecular orces depend on the polarity o the molecules. Be care ul with the terminology. The weakest intermolecular orces are called London dispersion orces. This description re ers to instantaneous dipole-induced dipole orces that exist between any atoms or groups o atoms and should be used or non-polar entities. A more general inclusive term is van der Waals orces which includes dipole dipole and dipole-induced dipole as well as London dispersion orces.

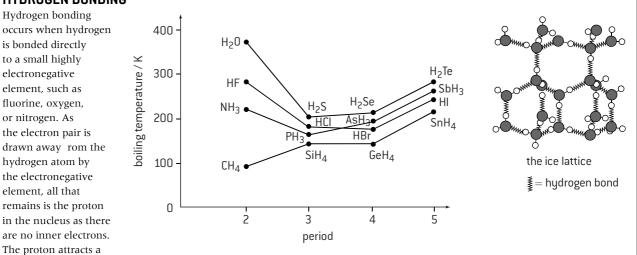
254

183

58.0

0.5

HYDROGEN BONDING



non-bonding pair o electrons rom the F, N, or O resulting in a much stronger dipole dipole attraction. Water has a much higher boiling point than the other group 16 hydrides as the hydrogen bonding between water molecules is much stronger than the dipole dipole bonding in the remaining hydrides. A similar trend is seen in the hydrides o group 15 and group 17. Hydrogen bonds between the molecules in ice result in a very open structure. When ice melts the molecules can move closer to each other so that water has its maximum density at 4 C.

Physical properties related to bonding type

MELTING AND BOILING POINTS

When a liquid turns into a gas the attractive orces between the particles are completely broken so boiling point is a good indication o the strength o intermolecular orces. When solids melt, the crystal structure is broken down, but there are still some attractive orces between the particles. Melting points do give an indication o the strength o intermolecular orces but they are also determined by the way the particles pack in the crystal state. They are also a ected by impurities. Impurities weaken the structure and result in lower melting points.

Covalent bonds are very strong so macromolecular covalent structures have extremely high melting and boiling points. For example, diamond, which has a giant tetrahedral structure, melts in the region o 4000 C and silicon dioxide, SiO₂, which has a similar structure, melts at over 1600 C. Graphite has very strong bonds between the carbon atoms in its hexagonal layers and has a similar melting point to diamond. Metals (see next page) and ionic compounds also tend to have relatively high melting and boiling points due to ionic attractions. Although it might be expected that ionic compounds with smaller more highly charged ions have higher melting points and boiling points the acts do not support this.

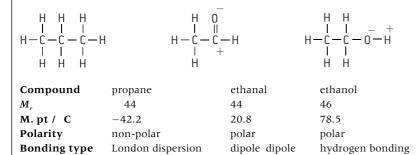
Ionic compound	Melting point / C	Boiling point / C	
LiCl	605	1382	
NaCl	801	1413	
KCl	770	1420	
MgCl ₂	714	1412	

The melting and boiling points o simple covalent molecules depend on the type o orces o attraction between the molecules. These ollow the order:

hydrogen bonding > dipole dipole > London dispersion orces

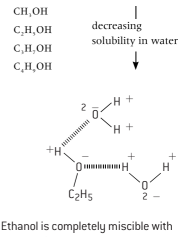
The weaker the attractive orces the more volatile the substance.

For example, propane, ethanal and ethanol have similar molar masses but there is a considerable di erence in their melting points.



SOLUBILITY

Like tends to dissolve like . Polar substances tend to dissolve in polar solvents, such as water, whereas nonpolar substances tend to dissolve in non-polar solvents, such as heptane or tetrachloromethane. Organic molecules o ten contain a polar head and a nonpolar carbon chain tail. As the nonpolar carbon chain length increases in an homologous series the molecules become less soluble in water. Ethanol itsel is a good solvent or other substances as it contains both polar and non-polar ends.



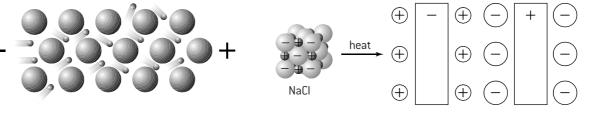
Ethanol is completely miscible with water as it can hydrogen-bond to water molecules.

CONDUCTIVITY

For conductivity to occur the substance must possess electrons or ions that are ree to move. Metals (and graphite) contain delocalized electrons and are excellent conductors. Molten ionic salts also conduct electricity, but are chemically decomposed in the process. Where all the electrons are held in fixed positions, such as diamond or in simple molecules, no electrical conductivity occurs.

When a potential gradient is applied to the metal,When an ithe delocalized electrons can move towards theoppositelypositive end of the gradient carrying charge.it is the io

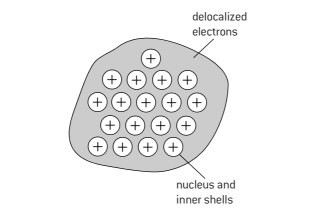
When an ionic compound melts, the ions are free to move to oppositely charged electrodes. Note: in molten ionic compounds it is the ions that carry the charge, not free electrons.



Metals and alloys

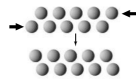
METALLIC BONDING

The valence electrons in metals become detached from the individual atoms so that metals consist of a close packed lattice of positive ions in a sea of delocalized electrons. A metallic bond is the attraction that two neighbouring positive ions have for the delocalized electrons between them.



Generally the strength of a metallic bond depends on the charge of the ions and the radius of the metal ion.

Metals are malleable, that is, they can be bent and reshaped under pressure. They are also ductile, which means they can be drawn out into a wire.



Metals are malleable and ductile because the close-packed layers of positive ions can slide over each other without breaking more bonds than are made.

MELTING POINTS OF METALS

Although most metals tend to have quite high melting points, mercury is a liquid at room temperature and the group 1 elements (alkali metals) all melt below 181 C. The trend in group 1 clearly follows the pattern that the smaller the metal ion formed when the valence electrons delocalize the stronger the metallic bond and the higher the melting point.

		Li	Na	K	Rb	Cs
M. pt /	С	181	97.8	63.5	39.3	28.5

However this logic only just holds true across period 3 (Na to Al) even though the charge on the ion is also increasing at the same time as the size of the ion is decreasing. It breaks down in group 14 as tin, which has a smaller ionic radius than lead, has a lower boiling point.

		Na	Mg	Al	Sn	Pb
M. pt /	С	97.8	650	660	232	328

This is because the melting point does not only depend upon the size and charge of the ion formed when the valence electrons are delocalized but also on the way in which the atoms are arranged in the solid metal.

ALLOYS

Alloys are sometimes termed a metallic solid solution. They are usually made up of more than one metal although steel is an alloy of iron and carbon. Some common alloys are brass, bronze, solder, pewter and amalgams.

Alloy	Principal metal	Added metal(s)
Brass	copper	zinc
Bronze	copper	tin
Solder	lead	tin (some may have more tin than lead)
Pewter	tin	copper, antimony, bismuth or lead
Amalgams	mercury	e.g. tin, silver, gold or sodium

The addition of another metal to a metallic element alters its properties. The added metals are likely to have a different radius and may have a different charge and so distort the structure of the original metal as the bonding is less directional. One obvious example of this is that alloys may have lower melting points than their component metals. For example, before copper and plastic were used for water piping lead tended to be used (the origin of the word *plumber*). Lead melts at 328 C and when pipes were being joined or repaired there was a danger of melting the actual pipe if too much heat was employed. Solder has a much lower melting point (typically 180–190–C) and can be used to weld lead pipes together or to secure wires to terminals in an electrical circuit.

Generally alloys are less ductile and less malleable than pure metals as the added impurities disturb the lattice. This also tends to make alloys harder than the pure metals they are derived from. For example, aluminium is a soft, ductile and malleable metal. When it is alloyed with another soft metal such as copper the resulting aluminium alloy is much harder and stronger and yet still retains much of its low density. Small amounts of carbon added to iron produce steel with a high tensile strength. If chromium is also added it produces stainless steel, an alloy of steel with a much increased resistance to corrosion.

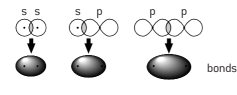


COMBINATION OF ATOMIC ORBITALS TO FORM MOLECULAR ORBITALS

Although the Lewis representation is a use ul model to represent covalent bonds it does make the alse assumption that all the valence electrons are the same. A more advanced model o bonding considers the combination o atomic orbitals to orm molecular orbitals.

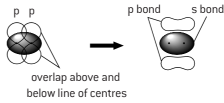
bonds

A (sigma) bond is ormed when two atomic orbitals on di erent atoms overlap along a line drawn through the two nuclei. This occurs when two s orbitals overlap, an s orbital overlaps with a p orbital, or when two p orbitals overlap head on .



bonds

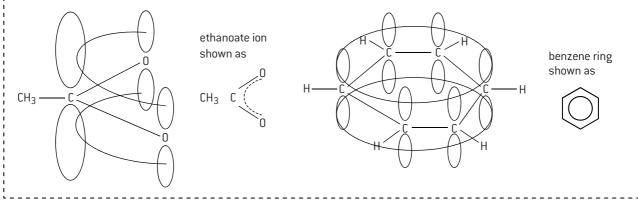
A (pi) bond is ormed when two p orbitals overlap sideways on . The overlap now occurs above and below the line drawn through the two nuclei. A bond is made up o two regions o electron density.



DELOCALIZATION OF ELECTRONS

Resonance involves using two or more Lewis structures to represent a particular molecule or ion where the structure cannot be described by using a single Lewis structure. They can also be explained by the delocalization o electrons. For example, in the ethanoate ion the carbon atom and the two oxygen atoms each have a p orbital containing one electron a ter the bonds have been ormed. Instead o orming just one double bond between the carbon atom and one o the oxygen atoms the electrons can delocalize over all three atoms. This is energetically more avourable than orming just one double bond.

Delocalization can occur whenever alternate double and single bonds occur between carbon atoms. The delocalization energy in benzene is about 150 kJ mol⁻¹, which explains why the benzene ring is so resistant to addition reactions.



FORMAL CHARGE

Formal charges

Formal charge is a technique used in chemistry that is based on a alse assumption but which can be use ul or determining which o several potential Lewis structures is pre erred when two or more are possible. It assumes that all atoms in a molecule or ion have the same electronegativity (the alse assumption) and is equal to the (Number o Valence electrons) – (Number o Non-bonding electrons) – (Number o Bonding electrons). This can be described by the ormula FC = V - N - B although note that this ormula is not given in the IB data booklet. The pre erred structure is the one where the individual atoms have the lowest possible ormal charge.

For example, consider two possible structures or carbon dioxide, both o which obey the octet rule.

(0) = 0 = 0 (0) (0) (0)		—
C = 4 - 0 - (8 $) = 0$	C = 4 - 0 - ((8) = 0
O = 6 - 4 - (4) = 0	O = 6 - 2 - (6) = +1 (O with triple bond)
	O = 6 - 6 - (2) = -1 (O with single bond)
l charge o zero but the pre-erred structure	e is the first one wit	th the two double bonds as the individ

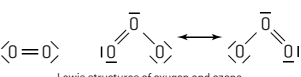
Both give a total ormal charge o zero but the pre erred structure is the first one with the two double bonds as the individual atoms have the lowest ormal charges.



IMPORTANCE OF THE OZONE LAYER

From the Lewis structures o both oxygen and ozone it can be seen that the double bond in oxygen is stronger than the one and a hal bond between the oxygen atoms in ozone.

This di erence in bond enthalpies helps to protect us rom the Sun s harm ul ultraviolet radiation. The ozone layer occurs in the stratosphere between about 12 km and 50 km above the sur ace o the Earth. Stratospheric ozone is in dynamic



Lewis structures of oxygen and ozone

equilibrium with oxygen and is continually being ormed and decomposed. The strong double bond in oxygen is broken by high energy ultraviolet light rom the Sun to orm atoms. These oxygen atoms are called radicals as they possess an unpaired electron and are very reactive. One oxygen radical can then react with an oxygen molecule to orm ozone.

$$\begin{array}{c} O=O (g) \xrightarrow{\text{UV (high energy)}} 2O (g) \\ O (g) + O_2(g) & O_2(g) \end{array}$$

The weaker bonds in ozone require ultraviolet light o less energy to break them. When they are broken the reverse process happens and the ozone breaks down back to an oxygen molecule and an oxygen radical. The radical can then react with another ozone molecule to orm two oxygen molecules.

$$O_3(g) \xrightarrow{\text{UV (lower energy)}} O_2(g) + O(g)$$

 $O_3(g) + O(g) = 2O_2(g)$

Overall the rate o production o ozone is equal to the rate o ozone destruction this process, during which a wide range o ultraviolet light is absorbed, is known as a steady state. Human-made pollutants such as CFCs and oxides o nitrogen can disrupt this process and in recent years large holes in the ozone layer have appeared particularly in winter and early spring over the South and North Poles.

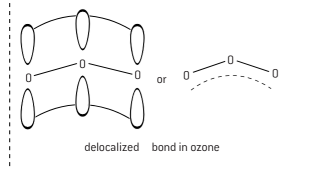
WAVELENGTH OF UV LIGHT NECESSARY FOR O₂ AND O₂ DISSOCIATION

The bond enthalpy or the O=O double bond is given as 498 kJ mol⁻¹ in the IB data booklet. For just one double bond this equates to 8.27 10^{-19} J. The wavelength o light that corresponds to this enthalpy value (*E*) can be calculated by

combining the expressions E = h and c = to give $= \frac{hc}{E}$ where *h* is Planck s constant and *c* is the velocity o light.

$$= \frac{6.63 \quad 10^{-34} \text{ (J s)} \quad 3.00 \quad 10^8 \text{ (m s}^{-1})}{8.27 \quad 10^{-19} \text{ (J)}} = 2.41 \text{ nm}$$

This is in the high energy region o the ultraviolet spectrum. Ozone is described above as two resonance hybrids. An alternative bonding model is to consider the electrons to be delocalized over all three oxygen atoms. In both models the bond order is 1.5, i.e. with an enthalpy between an O O single bond (144 kJ mol⁻¹) and an O=O double bond (498 kJ mol⁻¹) so ultraviolet light with a longer wavelength (lower energy) is absorbed in breaking the ozone bonds. The actual wavelength required is 330 nm. Working backwards this gives the strength o the O–O bond in ozone as 362 kJ mol⁻¹.



CATALYSIS AND OZONE DESTRUCTION BY CFC AND NO_v

CFCs catalyse the destruction o ozone because the high energy ultraviolet light in the stratosphere causes the homolytic fission o the C Cl bond to produce chlorine radicals. Note that it is the C Cl bond that breaks, not the C F bond, as the C Cl bond strength is weaker. These radicals then break down ozone molecules and regenerate more radicals so that the process continues until the radicals eventually escape or terminate. It has been estimated that one molecule o a CFC can catalyse the breakdown o up to 100 000 molecules o ozone.

$$CCl_2F_2(g) \longrightarrow CClF_2(g) + Cl (g)$$
(radical initiation)

$$Cl (g) + O_3(g) \longrightarrow ClO (g) + O_2(g)$$
(propagation

$$Cl (g) + O (g) \longrightarrow Cl (g) + O_2(g)$$
o radicals)

Evidence to support this mechanism is that the increase in the concentration o chlorine monoxide in the stratosphere over the Antarctic has been shown to mirror the decrease in the ozone concentration.

Nitrogen oxides also catalytically decompose ozone by a radical mechanism. The overall mechanism is complex. Essentially oxygen radicals are generated by the breakdown o NO_2 in ultraviolet light.

 $NO_2(g) \longrightarrow NO(g) + O(g)$

The oxygen radicals then react with ozone

 $O(g) + O_3(g) \longrightarrow 2O_2(g)$

The nitrogen oxide can also react with ozone to regenerate the catalyst

 $NO(g) + O_3(g) \longrightarrow NO_2(g) + O_2(g)$

The overall reaction can be simplified as:

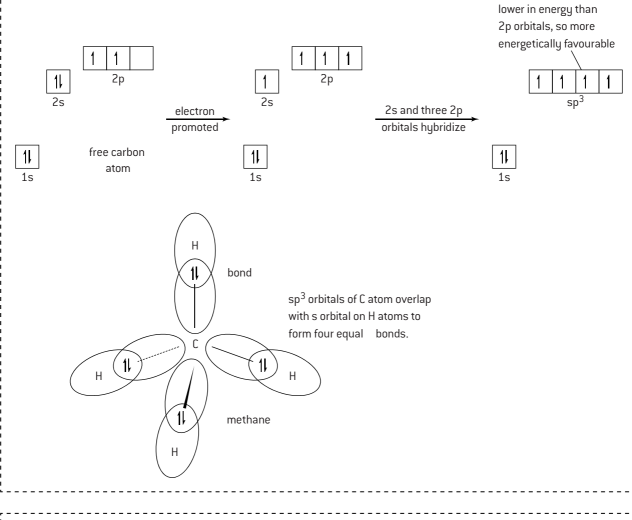
 $2O_3(g) \longrightarrow 3O_2(g)$

🕕 Hybridization (1)

sp³ HYBRIDIZATION

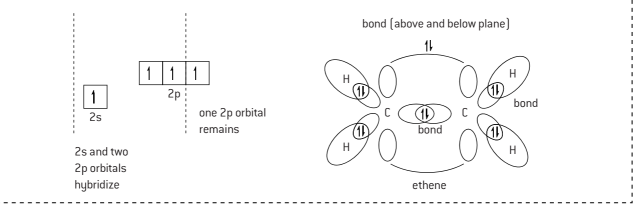
Methane provides a good example o sp³ hybridization. Methane contains our equal C H bonds pointing towards the corners o a tetrahedron with bond angles o 109.5 . A ree carbon atom has the configuration $1s^22s^22p^2$. It cannot retain this configuration in methane. Not only are there only two unpaired electrons, but the p orbitals are at 90 to each other and will not give bond angles o 109.5 when they overlap with the s orbitals on the hydrogen atoms.

When the carbon bonds in methane one o its 2s electrons is promoted to a 2p orbital and then the 2s and three 2p orbitals *hybridize* to orm our new hybrid orbitals. These our new orbitals arrange themselves to be as mutually repulsive as possible, i.e. tetrahedrally. Four equal bonds can then be ormed with the hydrogen atoms.



sp² HYBRIDIZATION

sp² hybridization occurs in ethene. A ter a 2s electron on the carbon atom is promoted the 2s orbital hybridizes with two o the 2p orbitals to orm three new planar hybrid orbitals with a bond angle o 120 between them. These can orm bonds with the hydrogen atoms and also a bond between the two carbon atoms. Each carbon atom now has one electron remaining in a 2p orbital. These can overlap to orm a bond. Ethene is thus a planar molecule with a region o electron density above and below the plane.



🕕 Hybridization (2)

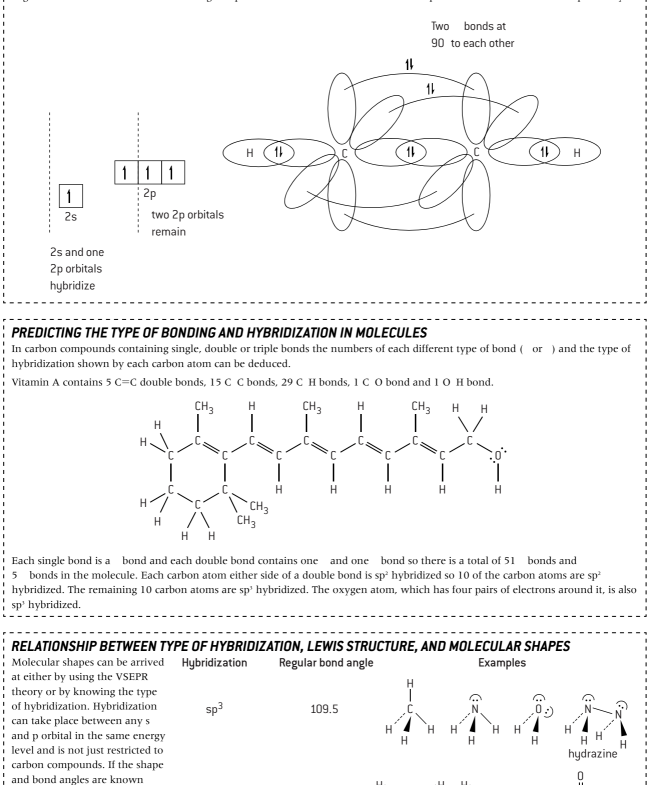
p HYBRIDIZATION

from using Lewis structures then

the type of hybridization can be deduced. Similarly if the type of hybridization is known the shape and bond angles can be deduced. sp²

sp

sp hybridization occurs when the 2s orbital hybridizes with just one of the 2p orbitals to form two new linear sp hybrid orbitals with an angle of 180 between them. The remaining two p orbitals on each carbon atom then overlap to form two bonds. An example is ethyne.



120

180

 $H - C \equiv C - H$

 $(: N \equiv N:)$

MULTIPLE CHOICE QUESTIONS CHEMICAL BONDING AND STRUCTURE

	Nitrate	rate Phosphate Carbonate		Ammonium
Α.	NO_3^-	PO_{4}^{3-}	CO ₃ -	NH_{3}^{+}
В.	NO ₃ ²⁻	PO_{3}^{2-}	CO ₃ ²⁻	$\mathrm{NH_3^+}$
C.	NO_3^{-}	PO_{4}^{3-}	CO3 ²⁻	$\mathrm{NH_4^+}$
D.	NO ₃ ²⁻	PO ₃ ²⁻	CO ₃ ²⁻	NH_4^+

1. What are the correct ormulas o the ollowing ions?

2. Which is the correct Lewis structure or ethene?

A. H. * . H	C. H H
*C * C *	:*C*C:*
H * * * H	H H
B. H H H C C C H H C C C H H H	D. H × H ×C×C× .H × H.

Given the ollowing electronegativities, H: 2.2 N: 3.0
 O: 3.5 F: 4.0, which bond would be the most polar?

A. O H in H_2O	C. N O in NO_2
B. N F in NF ₂	D. N H in NH ₃

4. Which substance is made up o a lattice o positive ions and ree moving electrons?

A. Graphite	C. Sul ur
B. Sodium chloride	D. Sodium

5. When CH₄, NH₃, H₂O, are arranged in order o **increasing** bond angle, what is the correct order?

A. CH_4 , NH_3 , H_2O	C. $NH_{3'}$ $CH_{4'}$ $H_{2}O$
B. NH ₃ , H ₂ O, CH ₄	D. H ₂ O, NH ₃ , CH ₄

6. Which order is correct when the ollowing compounds are arranged in order o **increasing** melting point?

A. $CH_4 < H_2S < H_2O$	$\text{C. } \text{CH}_{_4} < \text{H}_{_2}\text{O} < \text{H}_{_2}\text{S}$
B. $H_2S < H_2O < CH_4$	D. $H_{3}S < CH_{4} < H_{2}O$

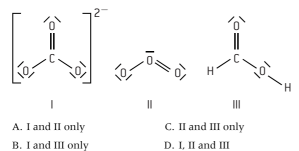
16		
1.5	п	
1.1	-	

- 11. Which o the ollowing species contains at least one atom that is sp² hybridized?
 - A. hydrogen cyanide, HCN
 - B. 2-methylpropane, CH₃CH(CH₃)CH₃
 - C. propanone, CH₃COCH₃
 - D. ethanol, CH₃CH₂OH
- 12. How many and bonds are present in propanal, CH₃CHCHO?

A.	8	and	2	C.	5	and	1

B. 8 and 1	D.	9	and	1
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13. Which species have delocalized electrons?



- 7. Which species contain a coordinate covalent bond?
 - I. HCHO

II. CO	
III. $H_{3}O^{+}$	
A. I and II only	C. II and III only
B. I and III only	D. I, II and III

8. Which is the correct order or **decreasing** H N H bond angles in the species NH₂⁻, NH₃ and NH₄⁺ (largest bond angle first)?

A. NH_3 , NH_2^- , NH_4^+	C. NH_{2}^{-} , NH_{4}^{+} , NH_{3}^{-}
B. NH_4^+ , NH_3 , NH_2^-	D. NH_{2}^{-} , NH_{3} , NH_{4}^{+}

- 9. Which is the correct order or **increasing** intermolecular orces o attraction (smallest orce first)?
 - A. Covalent bonds, hydrogen bonds, dipole dipole, London dispersion orces
 - B. London dispersion orces, dipole dipole, hydrogen bonds, covalent bonds
 - C. London dispersion orces, hydrogen bonds, dipole dipole, covalent bonds
 - D. Covalent bonds, dipole dipole, hydrogen bonds, London dispersion orces
- 10. Which statement best explains why alloys tend to be less malleable than pure metals?
 - A. The added metal has more valence electrons so increases the amount o delocalization.
 - B. The added metal prevents the layers rom being drawn out into a wire.
 - C. The added metal disturbs the lattice so the layers are less able to slide over each other.
 - D. The added metal acts as an impurity and so lowers the melting point.
- 14. What is the ormal charge on the oxygen atom in the hydronium ion H_3O^+ ?

A2	C. 0
B1	D. +1

- 15. Which describes the shape o the SF_4 molecule?
 - A. Tetrahedral C. Square planar
 - B. Distorted tetrahedral D. Trigonal bipyramidal
- 16. Which statements about graphene are correct?
 - I. It can be considered as a single layer o graphite
 - II. The hybridization o the carbon atoms is $\ensuremath{\text{spin}}^2$
 - III.It is an allotrope o carbon
 - A. I and II only
 - B. I and III only
 - C. II and III only
 - D. I, II and III

SHORT ANSWER QUESTIONS CHEMICAL BONDING AND STRUCTURE

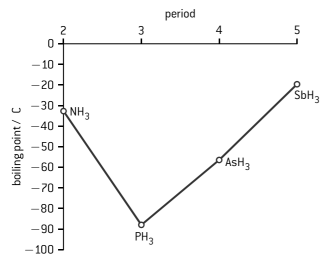
[8]

[5]

[2]

[4]

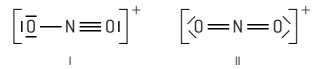
- PF₃, SF₂ and SiF₄ have di erent shapes. Draw their Lewis structures and use the VSEPR theory to predict the name o the electron domain geometry and the molecular shape o each molecule.
- 2. a) (i) Draw the Lewis structure o NH₃, state its molecular shape and deduce and explain the H–N–H bond angle in NH₃. [4]
 - (ii) The graph below shows the boiling points o the hydrides o group 15. Discuss the variation in the boiling points. [4]



b) Explain, using diagrams, why CO and NO₂ are polar molecules but CO₂ is a non-polar molecule.



- 6. a) Ozone and sul ur hexa uoride are greenhouse gases.
 - (i) Draw the Lewis structure o sul ur hexa uoride. [1](ii) Explain why sul ur can expand its octet whereas
 - oxygen cannot. [1] (iii) Deduce the electron domain geometry or both ozone and sul ur hexa uoride and deduce their molecular shape. [2]
 - (iv) Deduce the bond angles in ozone and sul ur hexa uoride
 - b) Another greenhouse gas is dichlorodi uoromethane, CCl₂F₂. This gas can also cause destruction o the ozone layer.
 - (i) Determine the wavelength o light required to break the C F and the C Cl bonds. [4]
 - (ii) Suggest why dichlorodi uoromethane is unreactive in the atmosphere near the sur ace o the Earth but reactive in the ozone layer. [2]
- Two Lewis structures that obey the octet rule can be proposed or the nitronium ion, NO⁺₂.



Deduce the ormal charge or each atom in both o the two proposed structures and determine which structure is the most likely.

- 3. Ethane, C₂H₂, and disilane, Si₂H₂, are both hydrides o group 14 elements with similar structures but with di erent chemical properties. a) Deduce the Lewis (electron dot) structure or Si₂H₂ showing all valence electrons. [1] b) State and explain the H Si H bond angle in Si₂H₂. [2] c) State which o the bonds, Si H or C H, is more polar. Explain your choice. [2] d) Predict, with an explanation, the polarity o the two molecules. [2] e) Explain why disilane has a higher boiling point than ethane. [2] a) State and explain which o propan-1-ol, CH₂CH₂CH₂OH, 4. and methoxyethane, CH, OCH, CH,, is more volatile. [3] b) Propan-1-ol, CH, CH, CH, OH, and hexan-1-ol, CH₃(CH₂)4CH₂OH, are both alcohols. State and explain which compound is more soluble in water. [2] c) Graphite is used as a lubricant and is an electrical conductor. Diamond is hard and does not conduct electricity. Explain these statements in terms o the structure and bonding o these allotropes o carbon. [6] 5. a) State the **full** and the **condensed** electron configuration or chlorine. [2] b) Deduce the orbital diagram or silicon using a box to represent an orbital and and to represent
 - electrons with opposite spins. [2] c) Explain why chlorine orms an ionic compound with sodium but a covalent compound with silicon. [2]
- 8. a) Describe the bonding within the carbon monoxide molecule.
 - b) Describe the delocalization o (pi) electrons and explain how this can account or the structure and stability o the carbonate ion, CO₃²⁻. [3]

[2]

- c) Explain the meaning o the term hybridization.State the type o hybridization shown by the carbon atoms in carbon dioxide, diamond, graphite and the carbonate ion. [5]
- d) (i) Explain the electrical conductivity o molten sodium oxide and liquid sul ur trioxide. [2]
 - (ii) Samples o sodium oxide and solid sul ur trioxide are added to separate beakers o water.
 Deduce the equation or each reaction and predict the electrical conductivity o each o the solutions ormed. [3]

5 ENERGETICS / THERMOCHEMISTRY

Me s ring enth y ch nges

EXOTHERMIC AND ENDOTHERMIC REACTIONS

Energy is defined as the ability to do work, that is, move a orce through a distance. It is measured in joules.

Energy = orce distance

(J) (N m)

In a chemical reaction energy is required to break the bonds in the reactants, and energy is given out when new bonds are ormed in the products. The most important type o energy in chemistry is heat. I the bonds in the products are stronger than the bonds in the reactants then the reaction is said to be **exothermic**, as heat is given out to the surroundings. Examples o exothermic

processes include combustion and neutralization. In **endothermic** reactions heat is absorbed rom the surroundings because the bonds in the reactants are stronger than the bonds in the products.

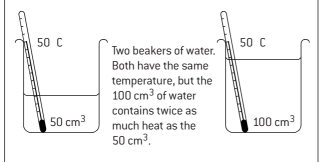
The internal energy stored in the reactants is known as its **enthalpy**, *H*. The absolute value o the enthalpy o the reactants cannot be known, nor can the enthalpy o the products, but what can be measured is the di erence between them, *H*. By convention *H* has a negative value or exothermic reactions and a positive value or endothermic reactions. It is normally measured under standard conditions o 100 kPa pressure at a temperature o 298 K. The **standard enthalpy change of a reaction** is denoted by H^{Θ} .

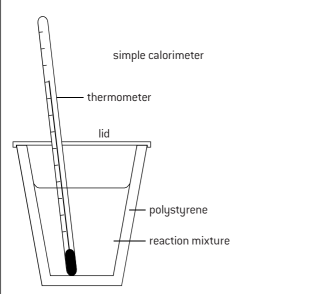
TEMPERATURE AND HEAT

It is important to be able to distinguish between heat and temperature as the terms are o ten used loosely.

Heat is a measure o the total energy in a given amount o substance and there ore depends on the amount o substance present.

Temperature is a measure o the hotness o a substance. It represents the average kinetic energy o the substance, but is independent o the amount o substance present.





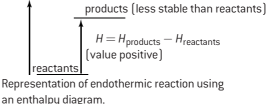
enthalpy, H

reactants

 $H = H_{\text{products}} - H_{\text{reactants}}$ [value negative]
products [more stable than reactants]

Representation of exothermic reaction using an enthalpy diagram.

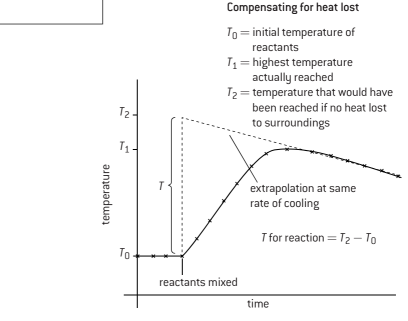




CALORIMETRY

The enthalpy change or a reaction can be measured experimentally by using a calorimeter. In a simple calorimeter all the heat evolved in an exothermic reaction is used to raise the temperature o a known mass o water. For endothermic reactions the heat trans erred rom the water to the reaction can be calculated by measuring the lowering o temperature o a known mass o water.

To compensate or heat lost by the water in exothermic reactions to the surroundings as the reaction proceeds a plot o temperature against time can be drawn. By extrapolating the graph, the temperature rise that would have taken place had the reaction been instantaneous can be calculated.



Hcc tions

CALCULATION OF ENTHALPY CHANGES

The heat involved in changing the temperature o any substance can be calculated rom the equation:

Heat energy = mass (m) specific heat capacity (c) temperature change (T)

The specific heat capacity o water is 4.18 kJ kg⁻¹ K⁻¹. That is, it requires 4.18 kilojoules o energy to raise the temperature o one kilogram o water by one kelvin.

Enthalpy changes are normally quoted in kJ mol⁻¹, or either a reactant or a product, so it is also necessary to work out the number o moles involved in the reaction which produces the heat change in the water.

WORKED EXAMPLE 1

50.0 cm³ o 1.00 mol dm⁻³ hydrochloric acid solution was added to 50.0 cm³ o 1.00 mol dm⁻³ sodium hydroxide solution in a polystyrene beaker. The initial temperature o both solutions was 16.7 C. A ter stirring and accounting or heat loss the highest temperature reached was 23.5 C. Calculate the enthalpy change or this reaction.

Step 1. Write equation or reaction

 $HCl(aq) + NaOH(aq) = NaCl(aq) + H_2O(l)$

Step 2. Calculate molar quantities

Amount o $HCl = \frac{50.0}{1000}$ 1.00 = 5.00 10^{-2} mol

Amount o $NaOH = \frac{50.0}{1000}$ 1.00 = 5.00 10^{-2} mol

There ore the heat evolved will be or $5.00 \quad 10^{-2}$ mol

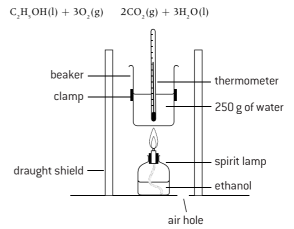
Step 3. Calculate heat evolved

Total volume o solution = $50.0 + 50.0 = 100 \text{ cm}^3$ Assume the solution has the same density and specific heat capacity as water then mass o water = 100 g = 0.100 kgTemperature change = 23.5 - 16.7 = 6.8 C = 6.8 KHeat evolved in reaction = $0.100 \quad 4.18 \quad 6.8 = 2.84 \text{ kJ}$ $= 2.84 \text{ kJ} \text{ (or } 5.00 \quad 10^{-2} \text{ mol})$ $H \text{ or reaction} = -2.84 \quad \frac{1}{5.00 \quad 10^{-2}} = -56.8 \text{ kJ mol}^{-1}$

(negative value as the reaction is exothermic)

WORKED EXAMPLE 2

A student used a simple calorimeter to determine the enthalpy change or the combustion o ethanol.



When 0.690g (0.015 mol) o ethanol was burned it produced a temperature rise o 13.2 K in 250 g o water. Calculate H or the reaction.

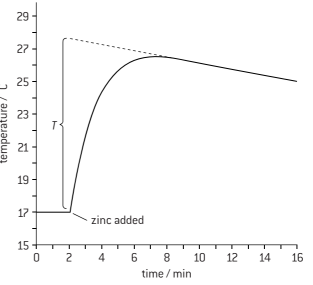
Heat evolved by 0.015 mol = $\frac{250}{1000}$ 4.18 13.2 = 13.79 kJ H = -13.79 $\frac{1}{0.015}$ = -920 kJ mol⁻¹

Note: the IB data booklet value is -1371 kJ mol⁻¹. Reasons or the discrepancy include the act that not all the heat produced is trans erred to the water, the water loses some heat to the surroundings, and there is incomplete combustion o the ethanol.

WORKED EXAMPLE 3

 $50.0 \text{ cm}^3 \text{ o } 0.200 \text{ mol } \text{dm}^{-3} \text{ copper(II)}$ sul ate solution was placed in a polystyrene cup. A ter two minutes 1.20 g o powdered zinc was added. The temperature was taken every 30 seconds and the ollowing graph obtained. Calculate the enthalpy change or the reaction taking place.

Step 1. Write the equation or the reaction
 $Cu^{2+}(aq) + Zn(s)$ $Cu(s) + Zn^{2+}(aq)$ Step 2. Determine the limiting reagent
Amount o $Cu^{+2}(aq) = \frac{50.0}{1000}$ 0.200 = 0.0100 mol
Amount o $Zn(s) = \frac{1.20}{65.37} = 0.0184$ mol
 $Cu^{2+}(aq)$ is the limiting reagentStep 3. Extrapolate the graph (*already done*) to compensate or
heat loss and determine T
T = 10.4 CStep 4. Calculate the heat evolved in the experiment or
0.0100 mol o reactants
Heat evolved $= \frac{50.0}{1000}$ 4.18 10.4 C = 2.17 kJStep 5. Express this as the enthalpy change or the reaction
H = -2.17 $\frac{1}{0.0100} = -217$ kJ mol⁻¹



Hess L w n st n r enth y ch nges

HESS LAW

Hess law states that the enthalpy change or a reaction depends only on the di erence between the enthalpy o the products and the enthalpy o the reactants. It is independent o the

reaction pathway.

goes via an intermediate.

The enthalpy change going rom A to

B is the same whether the reaction proceeds directly to A or whether it

 $H_1 \downarrow H_3 \qquad H_1 = H_2 +$

This law is a statement o the law o conservation o energy. It can be used to determine enthalpy changes, which cannot be measured directly. For example, the enthalpy o combustion o both carbon and carbon monoxide to orm carbon dioxide can easily be measured directly, but the combustion o carbon to carbon monoxide cannot. This can be represented by an energy cycle.

$$\begin{array}{c} C(S) + & 0_2(g) & \xrightarrow{H_{\chi}} CO(g) \\ -394 \\ \text{kJ mol} & 1 \\ 0_2(g) & 0_2(g) \\ CO_2(g) & -283 \text{ kJ mol} & 1 \end{array}$$

Hess law problems can also be solved by using simultaneous equations. Write the equations that are known and then manipulate them to arrive at the required equation. For example,

I C(s)	$+ O_{2}(g)$	$CO_2(g)$	$H = -394 \text{ kJ mol}^{-1}$
II CO(g	$() + O_2()$	(g) $CO_2(g)$	$H = -283 \text{ kJ mol}^{-1}$
Subtract II	rom I		
C(s) +	$O_{_{2}}(g) -$	CO(g) = 0	$H = -394 - (-283) \text{ kJ mol}^{-1}$
Rearrange e	equation		
C(s) +	$O_{2}(g)$	CO(g)	$H = -111 \text{ kJ mol}^{-1}$

WORKED EXAMPLE

Calculate the standard enthalpy change when one mole o methane is ormed rom its elements in their standard states. The standard enthalpies o combustion H_c^{\ominus} o carbon, hydrogen, and methane are -394, -286, and -890 kJ mol⁻¹ respectively.

Step 1. Write the equation or the enthalpy change with the unknown H^{Θ} value. Call this value H_{x}^{Θ}

$$C(s) + 2H_2(g) \xrightarrow{H_x^{\ominus}} CH_4(g)$$

Step 2. Construct an energy cycle showing the di erent routes to the products (in this case the products o combustion)

$$\begin{array}{c} \mathsf{C}(\mathsf{S}) + 2\mathsf{H}_2(\mathsf{g}) & \xrightarrow{\mathsf{H}_{\chi}} \mathsf{C}\mathsf{H}_4(\mathsf{g}) \\ \downarrow \mathsf{O}_2(\mathsf{g}) & \downarrow \mathsf{O}_2(\mathsf{g}) \\ \mathsf{C}\mathsf{O}_2(\mathsf{g}) + 2\mathsf{H}_2\mathsf{O}(\mathsf{I}) & \xrightarrow{\mathsf{H}_{\chi}} \mathsf{C}\mathsf{O}_2(\mathsf{g}) \end{array}$$

Step 3. Use Hess law to equate the energy changes or the two di erent routes $\underbrace{H_c^{\ominus}(C) + 2 \quad H_c^{\ominus}(H_2)}_{\text{direct route}} = \underbrace{H_x^{\ominus} + \quad H_c^{\ominus}(CH_4)}_{\text{route via methane}}$

Step 4. Rearrange the equation and substitute the values to give the answer $H_x^{\ominus} = H_c^{\ominus}(C) + 2 H_c^{\ominus}(H_2) - H_c^{\ominus}(CH_4)$ $= -394 + (2 - 286) - (-890) \text{ kJ mol}^{-1}$ $= -76 \text{ kJ mol}^{-1}$

STANDARD ENTHALPY CHANGES OF FORMATION H° and combustion $H^{\circ}_{.}$

The standard enthalpy change o ormation o a compound is the enthalpy change when one mole o the compound is ormed rom its elements in their standard states at 298 K and 100 kPa pressure. From this it ollows that H^{\ominus} or an element in its standard state will be zero. The standard enthalpy change o combustion, H_c^{\ominus} , is the enthalpy change when one mole o a substance is completely combusted in oxygen under standard conditions (298 K and 100 kPa pressure).

APPLICATION OF HESS LAW

The standard enthalpy change o ormation o ethanol, $C_2H_5OH(l)$ cannot be determined directly but an accurate value can be obtained indirectly by using the experimental values (in kJ mol⁻¹) or the standard enthalpy changes o combustion o carbon (-393.5), hydrogen (-285.8) and ethanol (-1371). Although this could be solved by just using simultaneous equations it is neater to show the energy cycle. Hess law is then applied by equating the energy changes involved in combusting carbon and hydrogen directly with the energy changes involved when they are first combined to orm ethanol and then combusting the ethanol.

$$2C(s) + 3H_{2}(g) + 0_{2}(g) +$$

Bon enth ies

BOND ENTHALPIES

Enthalpy changes can also be calculated directly rom bond enthalpies. The bond enthalpy is defined as the enthalpy change or the process

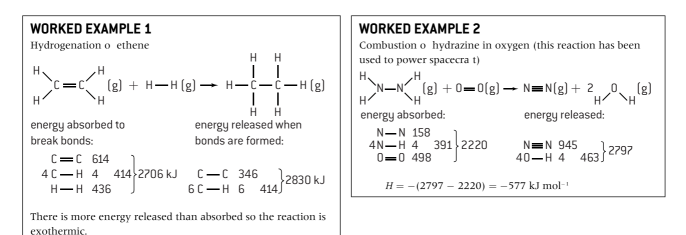
X-Y(g) = X(g) + Y(g) Note the gaseous state.

For bond ormation the value is negative as energy is evolved and or bond breaking energy has to be put in so the value is positive. For simple diatomic molecules where there are only two atoms the values can be known precisely. However or bonds such as the C–H bond there are many compounds containing a C–H bond and the value can di er slightly depending upon the surrounding atoms. The term **average bond enthalpy** is used. This is defined as the energy needed to break one mole o a bond in a gaseous molecule averaged over similar compounds.

I the bond enthalpy values are known or all the bonds in the reactants and products then the overall enthalpy change can be calculated.

Some bond enthalpies and average bond enthalpies All values in kJ mol⁻¹ H-H436 C = C614 C C 839 O=O 498 N N 945 C-C346 C-H 414 O-H463 N-H391 N-N158 Note that the O=O bond is stronger in O_2 , than the O–O bond in ozone, $O_{3'}$ (362 kJ mol⁻¹). This is

important as the ozone layer protects the Earth rom damaging ultraviolet radiation by absorbing both high and low energy uv light to break these bonds (see *Importance of the ozone layer* on page 33).



LIMITATIONS OF USING BOND ENTHALPIES

 $H = -(2830 - 2706) = -124 \text{ kJ mol}^{-1}$

Bond enthalpies can only be used on their own i all the reactants and products are in the gaseous state. I water were a liquid product in the above example then even more heat would be evolved since the enthalpy change o vaporization o water would also be needed to be included in the calculation.

In the above calculations some average bond enthalpies have been used. These have been obtained by considering a number o similar compounds containing the bond in question. In practice the energy o a particular bond will vary in di erent compounds. For this reason H values obtained rom using bond enthalpies will not necessarily be very accurate. Both these points are illustrated by the determination o the enthalpy change o combustion o methane by using bond enthalpies.

The equation or the reaction using bond enthalpies is:

$$H \xrightarrow{H} (g) + 20 = 0(g) \longrightarrow 0 = 0 = 0(g) + 20(g)$$

$$H \xrightarrow{H} (g) + 20 = 0(g) \xrightarrow{H} (g) \xrightarrow{$$

H reaction = (H products) – (H reactants)

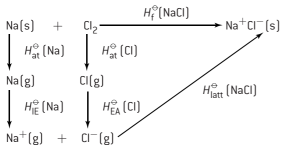
En	ergy taken in / kJ mol ⁻¹	Energy given out / kJ mol ⁻¹
4	C-H = 4 $414 = +1656$	2 $C=O=2$ $(-804) = -1608$
2	O=O=2 498 = + 996	4 $O-H = 4$ $(-463) = -1852$
	Total = + 2652	Total = -3408

The calculated enthalpy change or the reaction using bond enthalpies is there ore equal to -756 kJ mol^{-1} . However this is considerably di erent to the value o -891 kJ mol^{-1} given in Section 13 o the IB data booklet. It is to be expected that there will be a di erence as the definition o enthalpy o combustion is that the reactants and products should be in their normal states under standard conditions so we need to consider the extra 2 44 kJ mol^{-1} o energy given out when the two moles o gaseous water product turn to liquid water. This will now bring the enthalpy o combustion value to -844 kJ mol^{-1} . This is closer to -891 kJ mol^{-1} but there is still a di erence o more than 5%. This error is due to the act that the bond enthalpy or C=O in carbon dioxide is considerably di erent to the average C=O bond enthalpy.

🕕 Energy cyc es

BORN HABER CYCLES

Born Haber cycles are simply energy cycles or the ormation o ionic compounds. The enthalpy change o ormation o sodium chloride can be considered to occur through a number o separate steps.

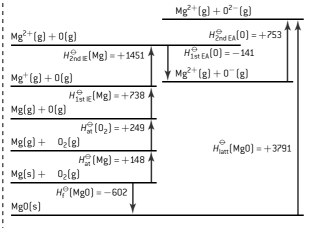


Using Hess law:

 $H^{\Theta}(\text{NaCl}) = H^{\Theta}_{at}(\text{Na}) + H^{\Theta}_{\text{IE}}(\text{Na}) + H^{\Theta}_{at}(\text{Cl}) + H^{\Theta}_{at}(\text{Cl})$ + $H^{\Theta}_{\text{EA}}(\text{Cl}) + H^{\Theta}_{\text{latt}}(\text{NaCl})$ Substituting the relevant values: $H^{\Theta}(\text{NaCl}) = +108 + 496 + 121 - 349 - 790$ = -414 kJ mol^{-1}

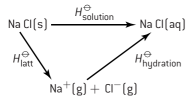
Note: it is the large lattice enthalpy that mainly compensates or the endothermic processes and leads to the enthalpy o ormation o ionic compounds having a negative value.

Sometimes Born Haber cycles are written as energy level diagrams with the arrows or endothermic processes in the opposite direction to the arrows or exothermic processes.



SOLUBILITY OF SALTS

It takes considerable energy to melt sodium chloride (melting point 801 C) due to the strong electrostatic attractions in its lattice and yet the lattice can easily be broken down by dissolving salt in water at room temperature. An energy cycle can be drawn to explain why.



The overall step is known as the enthalpy change o solution the enthalpy change when 1 mole o an ionic substance dissolves in water to give a solution o infinite dilution. This can be considered to proceed in two stages. The first involves the lattice enthalpy to break the lattice into gaseous ions, which will be highly endothermic and the second to hydrate the gaseous ions into

ENTHALPY OF ATOMIZATION

The standard enthalpy o atomization is the standard enthalpy change when one mole o gaseous atoms is ormed rom the element in its standard state under standard conditions. For diatomic molecules this is equal to hal the bond dissociation enthalpy.

 $\operatorname{Cl}_{_2}(g)$ $\operatorname{Cl}(g)$ $H_{_{\mathrm{at}}}^{\ominus} = +121 \text{ kJ mol}^{-1}$

LATTICE ENTHALPY

The lattice enthalpy relates either to the endothermic process o turning a crystalline solid into its gaseous ions or to the exothermic process o turning gaseous ions into a crystalline solid.

$$MX(s) \qquad M^{\scriptscriptstyle +}(g) + X^{\scriptscriptstyle -}(g)$$

The sign o the lattice enthalpy indicates whether the lattice is being ormed (-) or broken (+).

The size o the lattice enthalpy depends both on the size o the ions and on the charge carried by the ions.

The smaller the ion and the greater the charge, the higher the lattice enthalpy.

cation	n size in	creasing	anion s	size incr	easing
LiCl	NaCl	KCl	NaCl	NaBr	NaI
Lattice enthalpy / kJ mol ⁻¹					
864	790	720	790	754	705
		>			
charg	e on cat	ion increasing	charge	on anic	on increasing
MaCl	MgCl.		MgCl ₂	MgO	
maCI	0 2		0 2		
	0 2	alpy / kJ mol ⁻¹	0 2	0	

aqueous ions. This second step is known as the **hydration energy** and can be defined as the enthalpy change when 1 mole o gaseous ions dissolves in su ficient water to give an infinitely dilute solution. It is a highly exothermic process. Generally the smaller and more highly charged the ion the greater the hydration energy. In the case o sodium chloride the value or the sum o the hydration energies o the Na⁺ and Cl⁻ ions is very similar to the lattice enthalpy o NaCl and the small di erence o about 7 kJ mol⁻¹ can be made up by taking some heat rom the water so it dissolves with a slight lowering o temperature.

$$H_{\text{solution}}^{\ominus} = H_{\text{latt}}^{\ominus} + H_{\text{hydration}}^{\ominus} = +790 + (-783) = +7 \text{ kJ mol}^{-1}$$

Whether or not other salts are soluble in water depends upon the relative size o the lattice enthalpy compared with the hydration energy. The highly exothermic nature o hydration energy can explain why it is inadvisable to add water to sul ur trioxide. As sul ur trioxide is not ionic there is no strong lattice enthalpy to overcome and the hydration energy released as heat is so strong that the resulting sul uric acid can boil.

🕕 Entro y n s ont neity

ENTROPY

Entropy (*S*) re ers to the distribution o available energy among the particles in a system. The more ways the energy can be distributed the higher the entropy. This is sometimes equated to a measure o the disorder o a system. In nature, systems naturally tend towards an increase in entropy. An increase in entropy (disorder) can result rom:

mixing di erent types o particles, e.g. the dissolving o sugar in water

a change in state where the distance between the particles increases, e.g. liquid water steam

the increased movement o particles, e.g. heating a liquid or gas

increasing the number o particles, e.g. $2H_2O_2(l) = 2H_2O(l) + O_2(g)$.

The greatest increase in disorder is usually ound where the number o particles in the gaseous state increases.

The change in the disorder o a system is known as the entropy change, *S*. The more disordered the system becomes the more positive the value o *S* becomes. Systems which become more ordered will have negative *S* values.

SPONTANEITY

A reaction is said to be spontaneous i it causes a system to move rom a less stable to a more stable state. This will depend both upon the enthalpy change and the entropy change. These two actors can be combined and expressed as the Gibbs energy change G, o ten known as the ree energy change.

The standard ree energy change G^{\ominus} is defined as:

$$G^{\ominus} = H^{\ominus} - T S^{\ominus}$$

where all the values are measured under standard conditions. For a reaction to be spontaneous it must be able to do work, that is G° must have a negative value.

Note: the act that a reaction is spontaneous does not necessarily mean that it will proceed without any input o energy. For example, the combustion o coal is a spontaneous reaction and yet coal is stable in air. It will only burn on its own accord a ter it has received some initial energy so that some o the molecules have the necessary activation energy or the reaction to occur.

ABSOLUTE ENTROPY VALUES

The standard entropy o a substance is the entropy change per mole that results rom heating the substance rom 0 K to the standard temperature o 298 K. Unlike enthalpy, absolute values o entropy can be measured. The standard entropy change or a reaction can then be determined by calculating the di erence between the entropy o the products and the reactants.

 $S^{\ominus} = S^{\ominus}$ (products) $-S^{\ominus}$ (reactants)

e.g. or the ormation o ammonia

 $3H_{2}(g) + N_{2}(g) = 2NH_{3}(g)$

the standard entropies o $\,$ hydrogen, nitrogen and ammonia are respectively 131, 192 and 192 J $K^{_{-1}}$ mol $^{_{-1}}$.

There ore per mole o reaction

 $S^{\ominus} = 2$ 192 - [(3 131) + 192] = -201 J K^{-1} mol^{-1}

(or per mole o ammonia $S^{\Theta} = \frac{-201}{2} = -101 \text{ J } \text{K}^{-1} \text{ mol}^{-1}$)

FREE ENERGY, G^{\ominus} , and the position of Equilibrium

As a reaction proceeds, the composition o the reactants and products is continually changing and the ree energy will also be changing. The position o equilibrium corresponds to a maximum value o entropy and a minimum in the value o the Gibbs ree energy change. The reaction will not proceed any urther at this point, i.e. the rate o the orward reaction will equal the rate o the reverse reaction. The equilibrium composition o an equilibrium mixture thus depends upon the value o G^{\ominus} . From this it can also be deduced that the equilibrium constant or the reaction, $K_{e^{t}}$ will also depend upon the value o G^{\ominus} (see page 56).

💵 S ont neity o 🛛 re ction

POSSIBLE COMBINATIONS FOR FREE ENERGY CHANGES

Some reactions will always be spontaneous. I H^{\ominus} is negative or zero and S^{\ominus} is positive then G^{\ominus} must always have a negative value. Conversely i H^{\ominus} is positive or zero and S^{\ominus} is negative then G^{\ominus} must always be positive and the reaction will never be spontaneous.

For some reactions whether or not they will be spontaneous depends upon the temperature. I H^{\ominus} is positive or zero and S^{\ominus} is positive, then G^{\ominus} will only become negative at high temperatures when the value o T S^{\ominus} exceeds the value o H^{\ominus} .

Туре	H^{\ominus}	S^{\ominus}	$T S^{\ominus}$	$H^{\ominus} - T S^{\ominus}$	G^{\ominus}
1	0	+	+	(0) - (+)	_
2	0	-	_	(0) - (-)	+
3	-	+	+	(-) - (+)	-
4	+	_	_	(+) - (-)	+
5	+	+	+	(+) - (+)	- or +
6	_	_	_	(-) - (-)	+ or –

- **Type 1.** Mixing two gases. G^{\ominus} is negative so gases will mix o their own accord. Gases do not unmix o their own accord (Type 2) as G^{\ominus} is positive.
- **Type 3.** $(NH_4)_2 Cr_2 O_7(s) = N_2(g) + Cr_2 O_3(s) + 4H_2 O(g)$ The decomposition o ammonium dichromate is spontaneous at all temperatures.
- **Type 4.** $N_2(g) + 2H_2(g) = N_2H_4(g)$ The ormation o hydrazine rom its elements will never be spontaneous.

Type 5. $CaCO_3(s) = CaO(s) + CO_2(g)$ The decomposition o calcium carbonate is only spontaneous at high temperatures.

Type 6. $C_2H_4(g) + H_2(g) = C_2H_6(g)$ Above a certain temperature this reaction will cease to be spontaneous.

MULTIPLE CHOICE QUESTIONS ENERGETICS / THERMOCHEMISTRY

1. Which statement about the reaction below is correct?

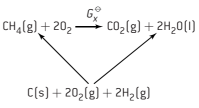
 $2NO(g) + O_{2}(g)$ $2NO_{2}(g)$ $H^{\ominus} = -114 \text{ kJ}$

- A. 114 kJ o energy are absorbed or every mole o NO reacted.
- B. 114 kJ o energy are released or every mole o NO reacted.
- C. 57 kJ o energy are absorbed or every mole o NO reacted.
- D. 57 kJ o energy are released or every mole o NO reacted.
- 2. When an aqueous solution o sul uric acid is added to an aqueous solution o potassium hydroxide the temperature increases. Which describes the reaction taking place?

	Туре	Sign of H^{\ominus}
Α.	Exothermic	+
В.	Exothermic	_
С.	Endothermic	—
D.	Endothermic	+

DETERMINING THE VALUE OF

The precise value o G^{\ominus} or a reaction can be determined rom G^{\ominus} values using an energy cycle, e.g. to find the standard ree energy o combustion o methane given the standard ree energies o ormation o methane, carbon dioxide, water, and oxygen.



By Hess law

$$\begin{array}{l} G_{x}^{\ominus} = [G_{x}^{\ominus}(\mathrm{CO}_{2}) + 2 \quad G^{\ominus}(\mathrm{H}_{2}\mathrm{O})] - [G^{\ominus}(\mathrm{CH}_{4}) \\ + 2 \quad G^{\ominus}(\mathrm{O}_{2})] \end{array}$$

Substituting the actual values

$$G_x^{\ominus} = [-394 + 2 \quad (-237)] - [-50 + 2 \quad 0]$$

= -818 kJ mol⁻¹

 G^{\ominus} values can also be calculated rom using the equation $G^{\ominus} = H^{\ominus} - T S^{\ominus}$. For example, in Type 5 in the adjacent list the values or H^{\ominus} and S^{\ominus} or the thermal decomposition o calcium carbonate are +178 kJ mol⁻¹ and +165.3 J K⁻¹ mol⁻¹ respectively. Note that the units o S^{\ominus} are di erent to those o H^{\ominus} .

At 25 C (298 K) the value or $G^{\ominus} = 178 - 298 \frac{165.3}{1000}$ = +129 kJ mol⁻¹

which means that the reaction is not spontaneous.

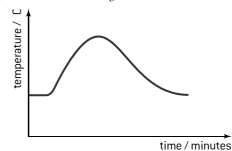
The reaction will become spontaneous when $T S^{\ominus} > H^{\ominus}$.

$$T \quad S^{\Theta} = H^{\Theta} \text{ when } T = \frac{H^{\Theta}}{S^{\Theta}} = \frac{178}{165.3/1000} = 1077 \text{K} (804 \text{ C})$$

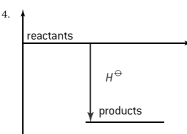
There ore above 804 C the reaction will be spontaneous.

Note: this calculation assumes that the entropy value is independent o temperature, which is not strictly true.

3. A student measured the temperature o a reaction mixture over time using a temperature probe. By considering the graph, which o the ollowing deductions can be made?



- I. The reaction is exothermic.
- II. The products are more stable than the reactants.
- III. The reactant bonds are stronger than the product bonds.
- A. I and II only C. II and III only
- B. I and III only
- D. I, II and III



What can be deduced about the relative stability of the reactants and products and the sign o H° , rom the enthalpy level diagram above?

Relative stability	Sign of	\pmb{H}^{\ominus}
A. Products more stable	_	
B. Products more stable	+	
C. Reactants more stable	_	
D. Reactants more stable	+	
Consider the ollowing reactions.		
$CH_4(g) + O_2(g) $ HCHO(l) + $H_2O(l)$		H^{\ominus}
$HCHO(l) + O_2(g) HCOOH(l)$		H^{\ominus}
$2HCOOH(l) + O_2(g) (COOH)_2(s) -$	$+ H_{2}O(l)$	H^{\ominus}
What is the enthalpy change o the reaction	n below?	
$2CH_4(g) + 3 O_2(g) (COOH)_2(s) + 3$	H ₂ O(l)	

x = x

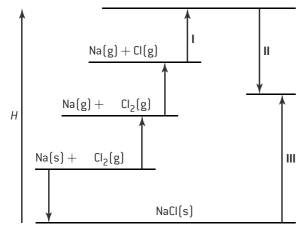
= y

= z

$$A. x + y + z$$
 $C. 2x + 2y + z$
 $B. 2x + y + z$
 $D. 2x + 2y + 2z$

5.

- 9. Which is a correct definition o lattice enthalpy?
 - A. It is the enthalpy change that occurs when an electron is removed rom 1 mol o gaseous atoms.
 - B. It is the enthalpy change that occurs when 1 mol o a compound is ormed rom its elements.
 - C. It is the enthalpy change that occurs when 1 mol o solid crystal changes into a liquid.
 - D. It is the enthalpy change that occurs when 1 mol o solid crystal is ormed rom its gaseous ions.
- 10. The diagram represents the Born Haber cycle or the lattice enthalpy o sodium chloride.



- 6. Which equation represents the H–F bond enthalpy?

7.
$$2 KHCO_{3}(s) \xrightarrow{H^{\ominus}} K_{2}CO_{3}(s) + CO_{2}(g) + H_{2}O(I)$$
$$+ 2HCI(aq) \xrightarrow{H_{1}^{\ominus}} H_{2}^{\ominus}$$
$$H_{2}^{\ominus} 2KCI(aq) + 2CO_{2}(g) + 2H_{2}O(I)$$

This cycle may be used to determine H^{\ominus} or the decomposition o potassium hydrogenearbonate. Which expression can be used to calculate H^{\ominus} ?

A. $H^{\ominus} = H_1^{\ominus} + H_2^{\ominus}$ C. $H^{\ominus} = \frac{1}{2} H_1^{\ominus} - H_2^{\ominus}$ B. $H^{\ominus} = H_1^{\ominus} - H_2^{\ominus}$ D. $H^{\ominus} = H_2^{\ominus} - H_1^{\ominus}$ 8. Which process is endothermic? A. HNO₃(aq) + NaOH(aq) NaNO₃(aq) + H₂O(l) B. Cl(g) + e⁻ Cl⁻(g) C. H₂O(l) H₂O(g) D. CH OW(h + 2O(h)) = 2CO(h) + 2H O(h)

D. $C_{_2}H_{_5}OH(l) + 3O_{_2}(g) = 2CO_{_2}(g) + 3H_{_2}O(l)$

What is the name o	the enthalpy chang	es I, II and III?
--------------------	--------------------	-------------------

	I	II	III
A.	ionization energy	electron a finity	lattice enthalpy
	o Na	o Cl	o NaCl
В.	lattice enthalpy	ionization energy	electron a finity
	o NaCl	o Na	o Cl
C.	electron a finity	ionization energy	lattice enthalpy
	o Cl	o Na	o NaCl
D.	01	lattice enthalpy o NaCl	electron a finity o Cl

- 11. Which reaction has the largest increase in entropy?
 - A. $H_2(g) + Cl_2(g) = 2HCl(g)$
 - B. $Al(OH)_{3}(s) + NaOH(aq)$ $Al(OH)_{4}^{-}(aq) + Na^{+}(aq)$
 - C. $Na_2CO_3(s) + 2HCl(aq) = 2NaCl(aq) + CO_2(g) + H_2O(l)$
 - D. $BaCl_2(aq) + Na_2SO_4(aq) = BaSO_4(s) + 2NaCl(aq)$
- 12. Which statements about entropy or the ollowing reaction at 298 K are correct?
 - $2NO(g) + O_2(g) = 2NO_2(g)$

I.
$$S^{\ominus}(O_2) = 0$$

II. $S^{\ominus} = 2S^{\ominus}(NO_2) - 2S^{\ominus}(NO) - S^{\ominus}(O_2)$

III. $S^{\ominus} < 0$

- A. I and II only C. II and III only
- B. I and III only D. I, II and III

SHORT ANSWER QUESTIONS ENERGETICS / THERMOCHEMISTRY

- In an experiment to measure the enthalpy change o combustion o ethanol, a student heated a copper calorimeter containing 100 cm³ o water with a spirit lamp and collected the ollowing data.
 - Initial temperature o water: 20.0 C
 - Final temperature o water: 55.0 C

Mass o ethanol burned: 1.78 g

- Density o water: 1.00 g cm⁻³
- a) Use the data to calculate the heat evolved (in kJ) when the ethanol was combusted.
- b) Calculate the enthalpy change o combustion per mole o ethanol.
- c) Suggest two reasons why the result is not the same as the value in the IB data booklet. [2]
- Ethanol is used as a component in uel or some vehicles. One uel mixture contains 10% by mass o ethanol in unleaded petrol (gasoline). This mixture is o ten re erred to as Gasohol E10.
 - a) Assume that the other 90% by mass o Gasohol E10 is octane. 1.00 kg o this uel mixture was burned.

 $CH_{2}CH_{2}OH(l) + 3O_{2}(g) = 2CO_{2}(g) + 3H_{2}O(l)$

$$H^{\ominus} = -1367 \text{ kJ mol}^{-1}$$

 $C_{_8}H_{_{18}}(l) + 12 O_{_2}(g) 8CO_{_2}(g) + 9H_{_2}O(l)$

 $H^{\ominus} = -5470 \text{ kJ mol}^{-1}$

[3]

- (i) Calculate the mass, in g, o ethanol and octane in 1.00 kg o the uel mixture. [1]
- (ii) Calculate the amount, in mol, o ethanol and octane in 1.00 kg o the uel mixture. [1]
- (iii) Calculate the total amount o energy, in kJ, released when 1.00 kg o the uel mixture is completely burned.
- b) I the uel blend was vaporized be ore combustion, predict whether the amount o energy released would be greater, less or the same. Explain your answer. [2]
- The data shown are rom an experiment to measure the enthalpy change or the reaction o aqueous copper(II) sul ate, CuSO₄ (aq) and zinc, Zn (s).

 $Cu^{_{2+}}(aq) + Zn (s) - Cu (s) + Zn^{_{2+}}(aq)$

50.0 cm³ o 1.00 mol dm⁻³ copper(II) sul ate solution was placed in a polystyrene cup and zinc powder was added a ter

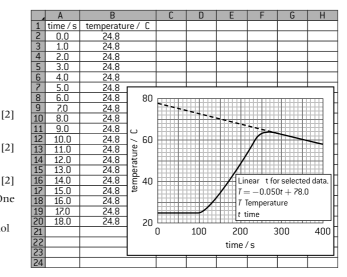
H

4. a) The standard enthalpy change o three combustion reactions is given below in kJ. $2C_{2}H_{4}(g) + 7O_{2}(g) = 4CO_{2}(g) + 6H_{2}O(l)$ $H^{\ominus} = -3120$ $2H_2(g) + O_2(g) = 2H_2O(l)$ $H^{\ominus} = -572$ $C_{2}H_{4}(g) + 3O_{2}(g) = 2CO_{2}(g) + 2H_{2}O(l)$ $H^{\ominus} = -1411$ Based on the above in ormation, calculate the standard change in enthalpy, H^{\ominus} , or the ollowing reaction. $C_{2}H_{2}(g) = C_{2}H_{4}(g) + H_{2}(g)$ [4] b) Predict, stating a reason, whether the sign o S^{\ominus} or the above reaction would be positive or negative. [2] c) Discuss why the above reaction is non-spontaneous

at low temperature but becomes spontaneous at

high temperatures.

100 seconds. The temperature time data was taken rom a data-logging so tware program. The table shows the initial 19 readings.



A straight line has been drawn through some o the data points. The equation or this line is given by the data-logging so tware as T = -0.050t + 78.0

The heat produced by the reaction can be calculated rom the temperature change, T, using the expression Heat change = Volume o $CuSO_4(aq)$ Specific heat capacity o H_2O T

- a) Describe two assumptions made in using this expression to calculate heat changes. [2]
- b) (i) Use the data presented by the data-logging so tware to deduce the temperature change, *T*, which would have occurred i the reaction had taken place instantaneously with no heat loss. [2]
 - (ii) State the assumption made in part b) (i). [1]
 - (iii) Calculate the heat, in kJ, produced during the reaction using the expression given in part a).
- c) The colour o the solution changed rom blue to colourless. Deduce the amount, in moles, o zinc which reacted in the polystyrene cup. [1]
- d) Calculate the enthalpy change, in kJ mol⁻¹, or this reaction.
- d) Using bond enthalpy values, calculate H^{\ominus} or the ollowing reaction.

$$_{2}H_{6}(g) = C_{2}H_{4}(g) + H_{2}(g)$$
 [3]

e) Suggest with a reason, why the values obtained in parts a) and d) are di erent. [1]

5. Synthesis gas is produced by the ollowing reaction.

 $CH_4(g) + H_2O(g) = 3H_2(g) + CO(g) = H^{\ominus} = +210 \text{ kJ}$

For this reaction $S^{\ominus} = +216 \text{ J K}^{-1}$.

С

- a) Explain why this reaction is not spontaneous at 298 K. [2]
- b) Calculate the temperature at which this reaction becomes spontaneous. [2]

[2]

6 CHEMICAL KINETICS

Rates reactin and cllisin the r

RATE OF REACTION

Chemical kinetics is the study o the actors a ecting the rate o a chemical reaction. The rate o a chemical reaction can be defined either as the increase in the concentration o one o the products per unit time or as the decrease in the concentration o one o the reactants per unit time. It is measured in mol dm 3 s 1 .

The change in concentration can be measured by using any property that di ers between the reactants and the products. Common methods include mass or volume changes when a gas is evolved, absorption using a spectrometer when there is a colour change, pH changes when there is a change in acidity, and electrical conductivity when there is a change in the ionic concentrations. Data loggers could be used or all these methods. A graph o concentration against time is then usually plotted. The rate at any stated point in time is then the gradient o the graph at that time. Rates o reaction usually decrease with time as the reactants are used up.

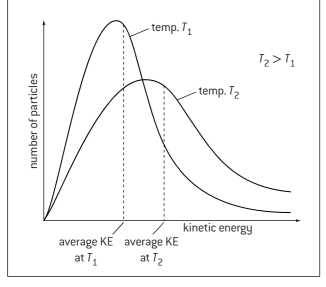
The reaction o hydrochloric acid with calcium carbonate can be used to illustrate three typical curves that could be obtained depending on whether the concentration o reactant, the volume o the product or the loss in mass due to the carbon dioxide escaping is ollowed.

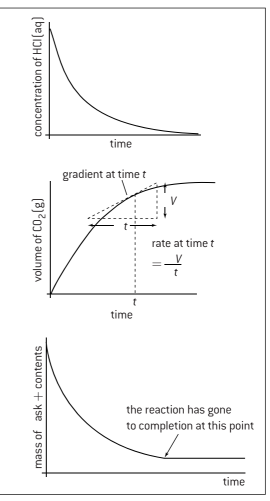
 $CaCO_{3}(s) + 2HCl(aq) = CaCl_{2}(aq) + H_{2}O(l) + CO_{2}(g)$

MAXWELL BOLTZMANN DISTRIBUTION

The moving particles in a gas or liquid do not all travel with the same velocity. Some are moving very ast and others much slower. The aster they move the more kinetic energy they possess. The distribution o kinetic energies is shown by a Maxwell Boltzmann curve.

As the temperature increases, the area under the curve does not change as the total number o particles remains constant. More particles have a very high velocity resulting in an increase in the average kinetic energy, which leads to a broadening o the curve.





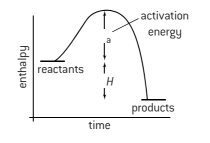
COLLISION THEORY

For a reaction between two particles to occur three conditions must be met.

The particles must collide.

They must collide with the appropriate geometry or orientation so that the reactive parts o the particles come into contact with each other.

They must collide with su ficient energy to bring about the reaction.

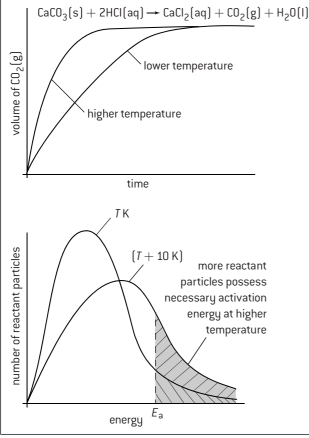


This minimum amount o energy required is known as the **activation energy**. Any actor that either increases the requency o the collisions or increases the energy with which they collide will make the reaction go aster.

act rs affecting the rate

TEMPERATURE

As the temperature increases, the particles will move faster so there will be more collisions per second. However, the main reason why an increase in temperature increases the rate is that more of the colliding particles will possess the necessary activation energy resulting in more successful collisions. As a rough rule of thumb an increase of 10 C doubles the rate of a chemical reaction.



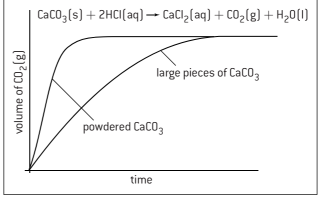
CATALYST

Catalysts increase the rate of a chemical reaction without themselves being chemically changed at the end of the reaction. They work essentially by bringing the reactive parts of the reactant particles into close contact with each other. This provides an alternative pathway for the reaction with a

reacti n

SUR ACE AREA

In a solid substance only the particles on the surface can come into contact with a surrounding reactant. If the solid is in powdered form then the surface area increases dramatically and the rate increases correspondingly.



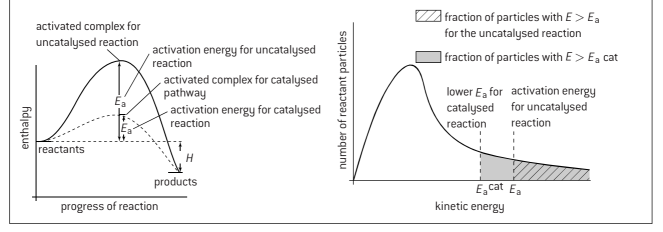
CONCENTRATION

The more concentrated the reactants the more collisions there will be per second per unit volume. As the reactants get used up their concentration decreases. This explains why the rate of most reactions gets slower as the reaction proceeds. (Some exothermic reactions do initially speed up if the heat that is given out more than compensates for the decrease in concentration.)

CaCO₃(s) + 2HCI(aq)
$$\rightarrow$$
 CaCI₂(aq) + CO₂(g) + H₂O(I)
1 mol dm ³ HCI
2 mol dm ³ HCI
time

Note: this graph assumes that calcium carbonate is the limiting reagent or that equal amounts (mol) of acid have been added.

lower activation energy. More of the reactants will possess this lower activation energy, so the rate increases.





RATE EXPRESSIONS

The rate o reaction between two reactants, A and B, can be ollowed experimentally. The rate will be ound to be proportional to the concentration o A raised to some power and also to the concentration o B raised to a power. I square brackets are used to denote concentration this can be written as rate $[A]^x$ and rate $[B]^y$. They can be combined to give the rate expression:

rate = $k[A]^x[B]^y$

where *k* is the constant o proportionality and is known as the **rate constant**.

x is known as the **order of the reaction** with respect to A.

y is known as the order o the reaction with respect to B.

The overall order o the reaction = x + y.

Note: the order o the reaction and the rate expression can only be determined experimentally. They cannot be deduced rom the balanced equation or the reaction.

UNITS OF RATE CONSTANT

The units o the rate constant depend on the overall order o the reaction.

First order: rate = k[A]

$$k = \frac{\text{rate}}{[A]} = \frac{\text{mol } dm^{-3} s^{-1}}{\text{mol } dm^{-3}} = s^{-1}$$

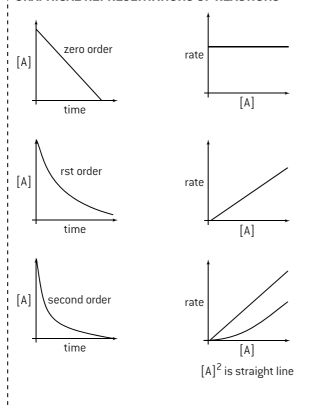
Second order: rate = $k[A]^2$ or k = [A][B]

$$k = \frac{\text{rate}}{[A]^2} = \frac{\text{mol } dm^{-3} s^{-1}}{(\text{mol } dm^{-3})^2} = dm^3 \text{ mol}^{-1} s^{-1}$$

Third order: rate = $k[A]^2[B]$ or rate = $k[A][B]^2$

$$k = \frac{\text{rate}}{[A]^2[B]} = \frac{\text{mol } dm^{-3} s^{-1}}{(\text{mol } dm^{-3})^3} = dm^6 \text{ mol}^{-2} s^{-1}$$

GRAPHICAL REPRESENTATIONS OF REACTIONS



DERIVING A RATE EXPRESSION BY INSPECTION OF DATA

Experimental data obtained rom the reaction between hydrogen and nitrogen monoxide at 1073 K:

 $2H_{_2}(g) + 2NO(g) = 2H_{_2}O(g) + N_{_2}(g)$

periment	Initial		Ini	Initial		Initial rate	
	concentration		n concer	concentration		of formation	
	of $H_2(g)$ /		of N	of NO(g) /		of $N_2(g)$ /	
	mol	dm-3	mol	dm-3	mol d	$m^{-3} s^{-1}$	
1	1	10 ³	6	10 3	3	10 ³	
2	2	10 3	6	10 3	6	10 3	
3	6	10 3	1	10 3	0.5	10 3	
4	6	10 3	2	10 3	2.0	10 3	

From experiments 1 and 2 doubling $[H_2]$ doubles the rate so rate $[H_2]$.

From experiments 3 and 4 doubling [NO] quadruples the rate so rate [NO]².

Rate expression given by rate = $k[H_2][NO]^2$.

The rate is first order with respect to hydrogen, second order with respect to nitrogen monoxide, and third order overall. The value o k can be ound by substituting the values rom any one o the our experiments:

$$k = \frac{\text{rate}}{[\text{H}][\text{NO}]^2} = 8.33 \quad 10^4 \text{ dm}^6 \text{ mol}^{-2}$$

Ex

For a first-order reaction the rate o change o concentration o A is equal to k[A]. This can be expressed as $\frac{d [A]}{dt} = k$ [A].

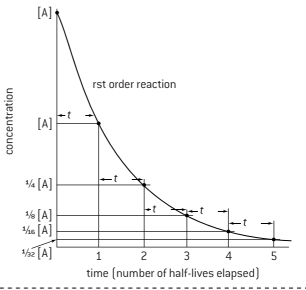
I this expression is integrated then $kt = \ln [A]_o \ln [A]$ where $[A]_o$ is the initial concentration and [A] is the concentration at time *t*. This expression is known as the integrated orm o the rate equation.

The hal -li e is defined as the time taken or the concentration o a reactant to all to hal o its initial value.

At $t [A] = [A]_{o}$ the integrated rate expression then becomes

 $kt = \ln [A]_{o} \ln [A]_{o} = \ln 2$ since $\ln 2 = 0.693$ this simplifies to $t = \frac{0.693}{k}$

From this expression it can be seen that the hal -li e o a firstorder reaction is independent o the original concentration o A, i.e. first-order reactions have a constant hal -li e.





Reaction mechanisms and activation energy

REACTION MECHANISMS

Many reactions do not go in one step. This is particularly true when there are more than two reactant molecules as the chances o a success ul collision between three or more particles is extremely small. When there is more than one step then each step will proceed at its own rate. No matter how ast the other steps are the overall rate o the reaction will depend only upon the rate o the slowest step. This slowest step is known as the rate-determining step.

e.g. consider the reaction between A and B to orm $A_{,B}$: $2A + B = A_{,B}$. A possible mechanism might be:

Step 1.
$$A + A$$
Step 2. $A + B$ $A = A$ $A = A$ Step 2. $A = A + B$ A_2B

However ast A A reacts with B the rate o production o A, B will only depend on how ast A A is ormed.

When the separate steps in a chemical reaction are analysed there are essentially only two types o processes. Either a single species can break down into two or more products by what is known as a unimolecular process, or two species can collide and interact by a **bimolecular process**.

In a bimolecular process the species collide with the necessary activation energy to give initially an activated complex. An activated complex is not a chemical substance which can be isolated, but consists o an association o the reacting particles in which bonds are in the process o being broken and ormed. An activated complex either breaks down to orm the products or reverts back to the original reactants.

The number o species taking part in any specified step in the reaction is known as the **molecularity**. In most cases the molecularity re ers to the slowest step, that is the rate-determining step.

In the reaction on the previous page, between nitrogen monoxide and hydrogen, the stoichiometry o the reaction involves two molecules o hydrogen and two molecules o nitrogen monoxide. Any proposed mechanism must be consistent with the rate expression. For third-order reactions, such as this, the rate-determining step will never be the first step. The proposed mechanism is:

ast NO(g) + NO(g)NO(g)slow $N_{2}O_{2}(g) + H_{2}(g)$ $N_{2}O(g) + H_{2}O(g)$ rate-determining step $N_2(g) + H_2O(g)$ $N_2O(g) + H_2(g)$ $2NO(g) + 2H_2(g)$ $N_{2}(g) + 2H_{2}O(g)$

Overall

I the first step was the slowest step the the rate expression would be rate = $k[NO]^2$ and the rate would be zero order with respect to hydrogen. The rate or the second step depends on [H,] and [N,O,]. However, the concentration o N,O, depends on the first step. So the rate expression or the second step becomes rate = $k[H_1][NO]^2$, which is consistent with the experimentally determined rate expression. The molecularity o the reaction is two, as two reacting species are involved in the rate-determining step.

ARRHENIUS EQUATION

The rate constant or a reaction is only constant i the temperature remains constant. As the temperature increases the reactants possess more energy and the rate constant increases. The relationship between rate constant and absolute temperature is given by the Arrhenius equation:

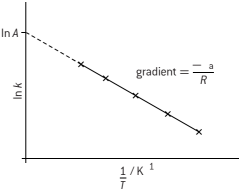
 $k = A e^{(-E_a/RT)}$

where E_1 is the activation energy and R is the gas constant. A is known as the requency actor (or pre-exponential actor) and is indicative o the requency o collisions with the correct orientation or the reaction to occur. This equation is o ten expressed in its logarithmic orm:

$$\ln k = \frac{-E_a}{RT} + \ln A$$

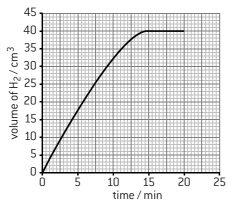
The equation can be used to determine both the requency actor and

the activation energy or the reaction. This can be done either by substitution using simultaneous equations or by plotting $\ln k$ against $\frac{1}{T}$ to give a straight line graph. The gradient o the graph will be equal to $\frac{1}{R}$ rom which the activation energy can be calculated. Extrapolating the graph back to the ln k axis will give an intercept with a value equal to ln A.

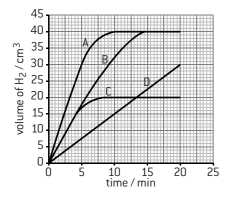


MULTIPLE CHOICE QUESTIONS CHEMICAL KINETICS

 A piece of zinc was added to aqueous nitric acid and the volume of hydrogen gas produced was measured every minute. The results are plotted on the graph below.



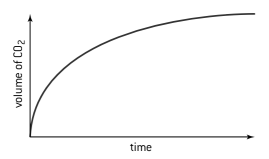
Which graph would you expect if the same mass of powdered zinc was added to nitric acid with the same concentration?



2. Which changes increase the rate of the reaction below?

 $C_4H_{10}(g) + Cl_2(g) = C_4H_9Cl(l) + HCl(g)$

- I. Increase of pressure
- II. Increase of temperature
- III. Removal of HCl(g)
- A. I and II only
- B. I and III only
- C. II and III only
- D. I, II and III
- 3. The reaction between excess calcium carbonate and hydrochloric acid can be followed by measuring the volume of carbon dioxide produced with time. The results of one such reaction are shown below. How does the rate of this reaction change with time and what is the main reason for this change?



- A. The rate increases with time because the calcium carbonate particles get smaller.
- B. The rate increases with time because the acid becomes more dilute.

- C. The rate decreases with time because the calcium carbonate particles get smaller.
- D. The rate decreases with time because the acid becomes more dilute.
- 4. Hydrochloric acid is reacted with large pieces of calcium carbonate, the reaction is then repeated using calcium carbonate powder. How does this change affect the activation enery and the collision frequency?

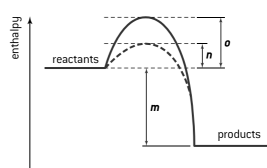
	Activation energy	Collision frequency		
A.	increases	increases		
В.	stays constant	increases		
С.	increases	stays constant		
D.	stays constant	stays constant		

5. Which statement is true about using sulfuric acid as a catalyst in the following reaction?

$$CH_3 CO CH_3(aq) + I_2(aq) CH_3 CO CH_2 I(aq)$$

+ HI (aq)

- I. The catalyst increases the rate of reaction.
- II. The catalyst lowers the activation energy for the reaction.
- III. The catalyst has been consumed at the end of the chemical reaction.
- A. I and II only
- B. I and III only
- C. II and III only
- D. I, II and III
- 6. Which are appropriate units for the rate of a reaction?
 - A. mol dm⁻³ s
 - B. mol dm⁻³ s⁻¹
 - C. mol dm⁻³
 - D. s
- The following enthalpy level diagram shows the effect of the addition of a catalyst to a chemical reaction. What do *m*, *n* and *o* represent?



progress of reaction

	т	п	0
A.	Н	E_{a} (without a catalyst)	E_{a} (with a catalyst)
В.	E _a (with a catalyst)	Н	E_{a} (without a catalyst)
C.	E _a (with a catalyst)	E_{a} (without a catalyst)	Н
D.	Н	E_{a} (with a catalyst)	E_{a} (without a catalyst)

MULTIPLE CHOICE QUESTIONS CHEMICAL KINETICS

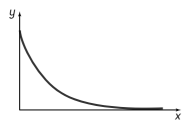
- 8. Consider the reaction between magnesium and hydrochloric acid. Which actors will a ect the reaction rate?
 - I. The collision requency o the reactant particles
 - II. The number or reactant particles with $E = E_a$
 - III. The number o reactant particles that collide with the appropriate geometry
 - A. I and II only
 - B. I and III only
 - C. II and III only
 - D. I, II and III
- 9. What is the best definition o rate of reaction?
 - A. The time it takes to use up all the reactants
 - B. The rate at which all the reactants are used up
 - C. The increase in concentration o a product per unit time
 - D. The time it takes or one o the reactants to be used up



11. The decomposition o N_2O_5 occurs according to the ollowing equation.

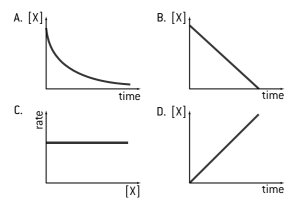
 $2N_2O_5(g) = 4NO_2(g) + O_2(g)$

The reaction is first order with respect to N_2O_3 . What combination o variables could the axes represent on the graph below?



	x-axis	y-axis
A.	time	[N ₂ O ₅]
В.	[N ₂ O ₅]	time
C.	[N ₂ O ₅]	rate o reaction
D.	rate o reaction	[N ₂ O ₅]

12. Which graph represents a reaction that is second order with respect to X or the reaction X products?



10. At 25 C, 200 cm³ o 1.0 mol dm⁻³ nitric acid is added to
5.0 g o magnesium powder. I the experiment is repeated using the same mass o magnesium powder, which conditions will result in the same initial reaction rate?

	Volume of HNO ₃ / cm ³	Concentration of HNO ₃ / mol dm ⁻³	Temperature / C	
A.	200	2.0	25	
В.	200	1.0	50	
C.	100	2.0	25	
D.	100	1.0	25	

13. Consider the reaction:

 $2NO(g) + Br_2(g) = 2NOBr(g)$

One suggested mechanism is:

 $NO(g) + Br_2(g) \Longrightarrow NOBr_2(g)$ fast

 $NOBr_2(g) + NO(g) \longrightarrow 2NOBr(g)$ slow

Which statements are correct?

- I. $NOBr_2(g)$ is an intermediate.
- II. The second step is the rate-determining step.
- III. rate = $k [NO]^2 [Br_2]$
- A. I and II only
- B. I and III only
- C. II and III only
- D. I, II and III
- 14. Consider the ollowing reaction.

2P+Q R+S

- This reaction occurs according to the ollowing mechanism.
- P + Q = X slow
- P + X = R + S fast
- What is the rate expression?
- A. rate = k[P]
- B. rate = k[P][X]
- C. rate = k[P] [Q]
- D. rate = $k[P]^2 [Q]$

SHORT ANSWER QUESTIONS CHEMICAL KINETICS

l.	a) (i)	Draw a graph that shows the distribution o
		molecular energies in a sample o a gas at two
		di erent temperatures, T_1 and T_2 such that
		T_2 is greater than T_1 .

(ii) Define the term activation energy.

[1]

[2]

- (iii) State and explain the e ect o a catalyst on the rate o an endothermic reaction.
- Magnesium is added to a solution o hydrochloric b) (i) acid. Sketch a graph o acid concentration on the y-axis against time on the x-axis to illustrate the progress o the reaction. [1]
 - (ii) Describe how the slope o the line changes with time.
 - (iii) Use the collision theory to state and explain the e ect o decreasing concentration on the rate o the reaction.
- 2. Hydrogen peroxide, H₂O₂ (aq), releases oxygen gas,
 - O_{2} (g), as it decomposes according to the equation:
 - $2H_{2}O_{2}(aq) = 2H_{2}O(l) + O_{2}(g)$

50.0 cm3 o hydrogen peroxide solution was placed in a boiling tube, and a drop o liquid detergent was added to create a layer o bubbles on the top o the hydrogen peroxide solution as oxygen gas was released. The tube was placed in a water bath at 75 C and the height o the bubble layer was



3. Hydrogen and nitrogen(II) oxide react together exothermically as ollows.

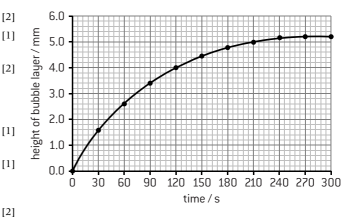
> $2H_2(g) + 2NO(g)$ $2H_{2}O(g) + N_{2}(g)$

The rate o this reaction was investigated in a series o experiments carried out at the same temperature, the results o which are shown below.

Experiment	Initial [H ₂ (g)] / mol dm ⁻³	Initial [NO(g)] / mol dm ⁻³	Initial rate of reaction / mol dm ⁻³ s ⁻¹	
1	2.0 10-3	4.0 10-3	4.0 10-3	
2	4.0 10 ⁻³	4.0 10-3	8.0 10-3	
3	6.0 10 ⁻³	4.0 10-3		
4	2.0 10-3	2.0 10-3	1.0 10-3	
5	2.0 10-3	1.0 10-3		

- a) Explain how the results rom Experiments 1 and 2 can be used to deduce that the order o reaction [1] with respect to hydrogen is 1.
- b) Deduce the order o reaction with respect to nitrogen(II) oxide, giving a reason or your answer. [2]
- c) Use your answers rom parts a) and b) to deduce the rate expression or the reaction. [1] d) Calculate the rate o reaction or each o
- Experiments 3 and 5. e) Use the results rom Experiment 1 to determine
- the value o, and the units or, the rate constant, k, or the reaction.

measured every thirty seconds. A graph was plotted o the height o the bubble layer against time.



- a) Explain why the curve reaches a maximum. [1]
- b) Use the graph to calculate the rate o decomposition o hydrogen peroxide at 120 s. [3]
- c) The experiment was repeated using solid manganese(IV) oxide, MnO, (s), as a catalyst.
 - (i) Draw a curve on the graph above to show how the height o the bubble layer changes with time when manganese(IV) oxide is present. [1]
 - (ii) Explain the e ect o the catalyst on the rate o decomposition o hydrogen peroxide. [2]
-) Suggest a mechanism or the reaction that is consistent with the rate expression. [2]
- g) The reaction is aster in the presence o a heterogeneous catalyst. Explain the meaning o the term heterogeneous as applied to a catalyst. Draw a labelled enthalpy level diagram that shows the e ect o the catalyst. [3]
- 4. a) The reaction between nitrogen(II) oxide and chlorine was studied at 263 K.

 $2NO(g) + Cl_{2}(g)$ 2NOCl(g)

It was ound that the orward reaction is first order with respect to Cl₂ and second order with respect to NO. The reverse reaction is second order with respect to NOCl.

(i) State the ra reaction.	(i) State the rate expression or the orward reaction.				
(ii) Predict the e ect on the rate o the orward reaction and on the rate constant i the concentration o NO is halved.					
Consider the olle	owing reaction	1.			
$NO_2(g) + C$	O(g) NO(g	$) + CO_{2}(g)$			
Possible reaction	mechanisms a	are:			
Above 775 K:	$NO_2 + CO$	$NO + CO_2$	slow		
Below 775 K:	2NO ₂ NO	+ NO ₃	slow		
	$NO_3 + CO$	$NO_2 + CO_2$	fast		
Based on the mechanisms, deduce the rate expressions					
above and below 775 K.					
State two situations when the rate o a chemical					

reaction is equal to the rate constant. [2]

b)

[2]

[2]

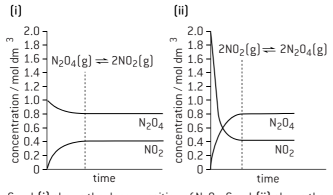
7 EQUILIBRIUM

The equilibrium law

DYNAMIC EQUILIBRIUM

A + B $\overbrace{reverse reaction}^{\text{orward reaction}} C + D$

Most chemical reactions do not go to completion. Once some products are ormed the reverse reaction can take place to re orm the reactants. In a closed system the concentrations o all the reactants and products will eventually become constant. Such a system is said to be in a state o **dynamic equilibrium**. The orward and reverse reactions continue to occur, but at equilibrium the rate o the orward reaction is equal to the rate o the reverse reaction.



Graph(i) shows the decomposition of N_2O_4 . Graph(ii) shows the reverse reaction starting with NO_2 . Once equilibrium is reached (shown by the dotted line), the composition of the mixture remains constant and is independent of the starting materials.

REACTION QUOTIENT AND EQUILIBRIUM CONSTANT

Consider the ollowing general reversible reaction in which w moles o A react with x moles o B to produce y moles o C and z moles o D.

 $wA + xB \quad yC + zD$

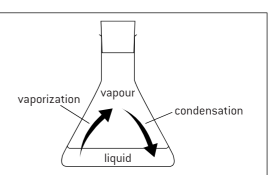
At any particular point in time the concentrations o A, B, C and D can be written as [A], [B], [C] and [D] respectively. The reaction quotient, Q, is defined as being

$$Q = \frac{[C]^{y} \quad [D]^{z}}{[A]^{w} \quad [B]^{y}}$$

As the reaction proceeds, the reaction quotient will change until the point o equilibrium is reached. At that point the concentrations o A, B, C and D remain constant and the reaction quotient is known as the equilibrium constant, *K*.

The equilibrium law states that or this reaction at a particular temperature

$$K_{c} = \frac{[C]_{eqm}^{y}}{[A]_{eqm}^{w}} [B]_{eqm}^{z}$$



Dynamic equilibrium also occurs when physical changes take place. In a closed ask, containing some water, equilibrium will be reached between the liquid water and the water vapour. The aster moving molecules in the liquid will escape rom the sur ace to become vapour and the slower moving molecules in the vapour will condense back into liquid. Equilibrium will be established when the rate o vaporization equals the rate o condensation.

 $\mathrm{H_{2}O}(l) \qquad \mathrm{H_{2}O}(g)$

CLOSED SYSTEM

A closed system is one in which neither matter nor energy can be lost or gained rom the system, that is, the macroscopic properties remain constant. I the system is open some o the products rom the reaction could escape and equilibrium would never be reached.

Examples

1

Formation o sul ur trioxide in the Contact process

$$2SO_{2}(g) + O_{2}(g) - 2SO_{3}(g)$$
$$\zeta_{c} = \frac{[SO_{3}]^{2}_{cqm}}{[SO_{2}]^{2}_{cam} - [O_{2}]_{cam}}$$

Formation o an ester rom ethanol and ethanoic acid $C_{1}H_{0}OH(l) + CH_{1}COOH(l) = CH_{1}COOC_{1}H_{0}(l) + H_{1}O(l)$

$$K_{c} = \frac{[CH_{3}COOC_{2}H_{5}]_{eqm}}{[C_{2}H_{5}OH]_{eqm}} \frac{[H_{2}O]_{eqm}}{[CH_{3}COOH]_{eqm}}$$

In both o these examples all the reactants and products are in the same phase. In the first example they are all in the gaseous phase and in the second example they are all in the liquid phase. Such reactions are known as *homogeneous reactions*. Another example o a homogeneous system would be where all the reactants and products are in the aqueous phase.

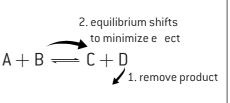
MAGNITUDE OF THE EQUILIBRIUM CONSTANT

Since the equilibrium expression has the concentration o products on the top and the concentration o reactants on the bottom it ollows that the magnitude o the equilibrium constant is related to the position o equilibrium. When the reaction goes nearly to completion K_c 1. I the reaction hardly proceeds then K_c 1. I the value or K_c lies between about 10^{-2} and 10^2 then both reactants and products will be present in the system in noticeable amounts. The value or K_c in the esterification reaction above is 4 at 100 °C. From this it can be in erred that the concentration o the products present in the equilibrium mixture is roughly twice that o the reactants.

Le Chatelier s principle and actors affecting the position o equilibrium

LE CHATELIER S PRINCIPLE

Provided the temperature remains constant the value for K_c must remain constant. If the concentration of the reactants is increased, or one of the products is removed from the equilibrium mixture then more of the reactants must react in order to keep K_c constant, i.e. the position of equilibrium will shift to the right (towards more products). This is the explanation for Le Chatelier s principle, which states that if a system at equilibrium is subjected to a small change the equilibrium tends to shift so as to minimize the effect of the change.



FACTORS AFFECTING THE POSITION OF EQUILIBRIUM

CHANGE IN CONCENTRATION

 $\label{eq:c2} C_2H_5OH(l)+CH_3COOH(l) \qquad CH_3COOC_2H_5(l)+H_2O(l)$

If more ethanoic acid is added the concentration of ethanoic acid increases so that at the point of addition:

 $K_{c} = \frac{[ester]}{[acid]} [water]$

To restore the system so that the equilibrium law is obeyed the equilibrium will move to the right, so that the concentration of ester and water increases and the concentration of the acid and alcohol decreases.

CHANGE IN PRESSURE

If there is an overall volume change in a gaseous reaction then increasing the pressure will move the equilibrium towards the side with less volume. This shift reduces the total number of molecules in the equilibrium system and so tends to minimize the pressure.

$2NO_2(g)$	$N_{2}O_{4}(g)$
brown	colourless
(2 vols)	(1 vol)

If the pressure is increased the mixture will initially go darker as the concentration of NO_2 increases then become lighter as the position of equilibrium is re-established with a greater proportion of N_2O_4 .

CHANGE IN TEMPERATURE

colourless

brown

In exothermic reactions heat is also a product. Taking the heat away will move the equilibrium towards the right, so more products are formed. The forward reaction in exothermic reactions is therefore increased by lowering the temperature

 $2NO_{2}(g) = N_{2}O_{4}(g) \qquad \qquad H^{\Theta} = -24 \text{ kJ mol}^{-1}$

 $H = -24 \text{ kJ mol}^{-1}$

Lowering the temperature will cause the mixture to become lighter as the equilibrium shifts to the right.

For an endothermic reaction the opposite will be true.

Unlike changing the concentration or pressure, a change in temperature will also change the value of K_c . For an exothermic reaction the concentration of the products in the equilibrium mixture decreases as the temperature increases, so the value of K_c will decrease. The opposite will be true for endothermic reactions.

e.g.
$$H_2(g) + CO_2(g)$$
 $H_2O(g) + CO(g)$ $H^{\Theta} = +41 \text{ kJ mol}^{-1}$

 T / K K_c

 298
 1.00
 10⁵

 500
 7.76
 10³

 700
 1.23
 10¹

 900
 6.01
 10¹

ADDING A CATALYST

A catalyst will increase the rate at which equilibrium is reached, as it will speed up both the forward and reverse reactions equally, but it will have no effect on the position of equilibrium and hence on the value of K_c .

MANIPULATING EQUILIBRIUM CONSTANTS

When a reaction is reversed the equilibrium constant for the reverse reaction will be the reciprocal of the equilibrium constant for the forward reaction, K_c . For example, for the reverse reaction of the Haber process, $2NH_3(g) = N_2(g) + 3H_2(g)$

$$K_{c} = \frac{[N_{2}] [H_{2}]^{3}}{[NH_{3}]^{2}} = \frac{1}{K_{c}}$$

If there are multiple steps in a reaction, each with its own equilibrium constant, then the equilibrium constants are multiplied to give the overall value of K_{c} . For example,

For the step $A + B = C$	$K_{c}^{1} = \frac{[C]}{[A] [B]}$
For the step $C + D = X + Y$	$K_{c}^{2} = \frac{[X] [Y]}{[C] [D]}$
For the overall reaction $A + B + D = X + Y$	$K_{c} = \frac{[X] [Y]}{[A] [B] [D]} = K_{c}^{1} K_{c}^{2}$

🖽 Equilibrium calculations

EQUILIBRIUM CALCULATIONS

The equilibrium law can be used either to find the value or the equilibrium constant, or to find the value o an unknown equilibrium concentration.

a) 23.0 g (0.50 mol) o ethanol was reacted with 60.0 g (1.0 mol) o ethanoic acid and the reaction allowed to reach equilibrium at 373 K. 37.0 g (0.42 mol) o ethyl ethanoate was ound to be present in the equilibrium mixture. Calculate K_{1} to the nearest integer at 373K.

	$C_{_2}H_{_5}OH(l)$	+	$CH_{3}COOH(l)$	$\mathrm{CH}_{_{3}}\mathrm{COOC}_{_{2}}\mathrm{H}_{_{5}}(\mathrm{l})$	+	$\mathrm{H_{2}O}(l)$
Initial amount / mol	0.50		1.00			
Equilibrium amount / mol	(0.50 0.42)		(1.00 0.42)	0.42		0.42
¹ Equilibrium concentration / mol dm ⁻³	(0.50 0.42)		(1.00 0.42)	0.42		0.42
(where V = total volume)	V		V	V		V

 $\frac{[\text{ester}] \quad [\text{water}]}{[\text{orid}]} = \frac{(0.421 \text{ v})}{(0.08/V)}$ (0.42/V) (0.42/V) = 4 (to the nearest integer) K =(0.58/V)

b) What mass o ester will be ormed at equilibrium i 2.0 moles o ethanoic acid and 1.0 moles o ethanol are reacted under the same conditions?

Let *x* moles o ester be ormed and let the total volume be $V dm^3$.

$$K_{c} = 4 = \frac{[\text{ester}] \quad [\text{water}]}{[\text{alcohol}] \quad [\text{acid}]} = \frac{x^{2}/V^{2}}{(1.0 \quad x)/V \quad (2.0 \quad x)/V} = \frac{x^{2}}{(x^{2} \quad 3x + 2)}$$

$$3x^{2} \quad 12x + 8 = 0$$

solve by substituting into the quadratic expression
$$x = \frac{-b \quad b^{2} \quad 4ac}{2a} \qquad x = \frac{12 \quad 144 \quad 96}{6}$$

solve by substituting into the quadratic expression

x = 0.845 or 3.15 (it can not be 3.15 as only 1.0 mol o ethanol was taken)

Mass o ester = 0.845 88.08 = 74.4 g (Note: IB Diploma Programme chemistry does not examine the use o the quadratic expression.)

c) 1.60 mol o hydrogen and 1.00 mol o iodine are allowed to reach equilibrium at a temperature o 704 K in a 4.00 dm³ ask, the amount o hydrogen iodide ormed in the equilibrium mixture is 1.80 mol. Determine the value o the equilibrium constant at this temperature.

			$H_{2}(g)$	$I_2(g)$	2HI(g)
Initi	al amount / mol		1.60	1.00	0
Equ	ilibrium amount / m	ol	0.70	0.10	1.80
Equ	ilibrium concentratio	on / mol dm ⁻³	0.175	0.025	0.450
**	$[HI(g)]^2$	0.450^{2}			

$$K_{\rm c} = \frac{[{\rm HI}({\rm g})]^2}{[{\rm H}_2({\rm g})] [{\rm I}_2({\rm g})]} = \frac{0.450^2}{0.175 \ 0.025} = 46.3 \text{ at } 704 \text{ K}$$

RELATIONSHIP BETWEEN FREE ENERGY CHANGE AND THE EQUILIBRIUM CONSTANT

The position o equilibrium corresponds to a maximum value o entropy and a minimum in the value o the Gibbs ree energy change. This means that the equilibrium constant, K_c and the Gibbs ree energy change, G^{\ominus} can both be used to measure the position o equilibrium in a reaction. They are related by the equation

 $G = -RT \ln K$

This can be illustrated by the dissociation o water according to the equation

H₂O(l) H⁺(aq) + OH⁻(aq) $H^{\Theta} = +55.8 \text{ kJ mol}^{-1}$

The relevant entropy values are:

	$H_{2}O(l)$	$H^+(aq)$	OH- (aq)
S^{\ominus} / J K ⁻¹ mol ⁻¹	+70.0	0	10.9

The change in entropy, $S^{\Theta} = (S^{\Theta} products) (S^{\Theta} reactants) = (-10.9) - (+70.0) = -80.8 \text{ J K}^{-1} \text{ mol}^{-1}$

 $G^{\ominus} = H^{\ominus}$ T $S^{\ominus} = 55.8$ 1000 - (298 -80.8) = +79900 J mol⁻¹

Using the expression
$$G = -RT \ln K$$

 $\ln K = \frac{1}{8.314} = -32.2$

 $K = e^{-32.2} = 1.00$ 10⁻¹⁴ at 298 K

This is the value or the equilibrium constant o water at 298 K, known as the ionic product constant or water (K), given in Section 2 o the IB data booklet. _____

MULTIPLE CHOICE QUESTIONS EQUILIBRIUM

- 1. Which statement is true about a chemical reaction at equilibrium?
 - A. The reaction has completely stopped.
 - B. The concentrations o the products are equal to the concentrations o the reactants.
 - C. The rate o the orward reaction is equal to the rate o the reverse reaction.
 - D. The concentrations o the products and reactants are constantly changing.
- 2. What is the equilibrium constant expression, *K_e*, or the ollowing reaction?

 $2NOBr(g) = 2NO(g) + Br_{2}(g)$ A. $K_{c} = \frac{[NO][Br_{2}]}{[NOBr]}$ B. $K_{c} = \frac{[NO]^{2}[Br_{2}]}{[NOBr]^{2}}$ C. $K_{c} = \frac{2[NO] + [Br_{2}]}{[2NOBr]}$ D. $K_{c} = \frac{[NOBr]^{2}}{[NO]^{2}[Br_{2}]}$

- 3. The ollowing are K_c values or a reaction, with the same starting conditions, carried out at di erent temperatures. Which equilibrium mixture has the highest concentration o products?
 - A. 1 10⁻²
 - B. 1
 - C. 1 10¹
 - D. 1 10²
- 4. What e ect will an increase in temperature have on the K_c value and the position o equilibrium in the ollowing reaction?

 $N_2(g) + 3H_2(g) = 2NH_3(g) = H = -92 kJ$

	K _c	Equilibrium position
A.	increases	shi ts to the right
B.	decreases	shi ts to the le t
C.	increases	shi ts to the le t
D.	decreases	shi ts to the right

14		
	-	
N		
	_	_

 0.50 mol o I₂(g) and 0.50 mol o Br₂(g) are placed in a closed flask. The ollowing equilibrium is established.

 $I_2(g) + Br_2(g) = 2IBr(g)$ The equilibrium mixture contains 0.80 mol o IBr(g). What is

the value o K_c ?

 A. 0.64
 C. 2.6

 B. 1.3
 D. 64

10. A 2.0 dm³ reaction vessel initially contains 4.0 mol o P and 5.0 mol o Q. At equilibrium 3 mol o R is present. What is the value o K_c or the ollowing reaction?

$$P(g) + Q(g)$$
 $R(g) + S(g)$
A. $\frac{2}{9}$ C. 4.5

B.
$$\frac{9}{20}$$
 D. 9

5. Consider the ollowing equilibrium reaction.

$$2SO_{3}(g) + O_{3}(g) = 2SO_{3}(g) \qquad H^{\Theta} = -197 \, k_{s}^{2}$$

Which change in conditions will increase the amount o SO, present when equilibrium is re-established?

- A. Decreasing the concentration o SO_2
- B. Increasing the volume
- C. Decreasing the temperature
- D. Adding a catalyst
- 6. The Haber process uses an iron catalyst to convert hydrogen gas, $H_2(g)$, and nitrogen gas, $N_2(g)$, to ammonia gas, $NH_3(g)$.

 $3H_{_2}(g) + N_{_2}(g) - 2NH_{_3}(g)$

- Which statements are correct or this equilibrium system?
 - I. The iron catalyst increases rates o the orward and reverse reactions equally.
 - II. The iron catalyst does not a ect the value o the equilibrium constant, K_c .
 - III. The iron catalyst increases the yield or ammonia gas, $\mathrm{NH}_{\sc c}(g).$
- A. I and II only
- B. I and III only
- C. II and III only
- D. I, II and III
- The ormation o nitric acid, HNO₃(aq), rom nitrogen dioxide, NO₂(g), is exothermic and is a reversible reaction.

 $4NO_2(g) + O_2(g) + 2H_2O(l) - 4HNO_3(aq)$

What is the e ect o a catalyst on this reaction?

- A. It increases the yield o nitric acid.
- B. It increases the rate o the orward reaction only.
- C. It increases the equilibrium constant.
- D. It has no e ect on the equilibrium position.
- 8. The value o K_c or the reaction $H_2(g) + Br_2(g) = 2HBr(g)$ is 4.0 10^{-2} . What is the value o the equilibrium constant or the reaction $2HBr(g) = H_2(g) + Br_2(g)$ at the same temperature?

A. 4.0	10 ⁻²	C. 25
B. 2.0	10 ⁻¹	D. 400

11. At 35 C $K_c = 1.6$ 10⁻⁵ mol dm⁻³ or the reaction 2NOCl(g) 2NO(g) + Cl₂(g)

Which relationship must be correct at equilibrium?

- A. [NO] = [NOCl] C. $[NOCl] < [Cl_2]$
- B. $2[NO] = [Cl_2]$ D. [NO] < [NOC]
- 12. Free energy change and the equilibrium constant are related by the equation $G = -RT \ln K_c$. Which combination is most likely or a reaction to go to completion at all temperatures?

	H^{\ominus}	S^{\ominus}	$K_{_{\rm c}}$
А		+	> 1
В	+	+	> 1
С	_	—	< 1
D		_	> 1

SHORT ANSWER QUESTIONS EQUILIBRIUM

1. Ethanol is manu actured by the hydration o ethene according to the equation below.

 $C_{2}H_{4}(g) + H_{2}O(g) = C_{2}H_{5}OH(g)$

a) State the expression or the equilibrium constant, *K_c*, or this reaction.

[1]

[1]

[3]

[4]

[1]

[2]

[1]

- b) Under certain conditions, the value o K_c or this reaction is 3.7 10^{-3} . When the temperature is increased the value is 4.9 10^{-4} .
 - (i) State what can be deduced about the position

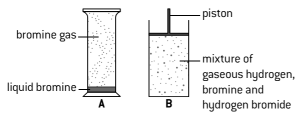
 o equilibrium at the higher temperature rom
 these values o K_c.
 - (ii)State what can be deduced about the sign*O H* or the reaction, explaining your choice.
- c) The process used to manu acture ethanol is carried out at high pressure. State and explain two advantages o using high pressure.
- 2. Ammonia is produced by the Haber process according to the ollowing reaction

 $N_2(g) + 3H_2(g) = 2NH_3(g)$ H is negative

- a) State the equilibrium expression or the above reaction. [1]
- b) Predict, giving a reason, the e ect on the positiono equilibrium when the pressure in the reactionvessel is increased. [2]
- c) State and explain the e ect on the value o K_c when the temperature is increased. [2]
- d) Explain why a catalyst has no e ect on the positiono equilibrium. [1]

H

Consider the two equilibrium systems involving bromine gas illustrated below.



- a) Formulate equations to represent the equilibria in A and B with Br,(g) on the le t-hand side in both equilibria. [2]
- b) (i) Describe what you would observe i a small amount o liquid bromine is introduced into A. [1]
 - (ii)Predict what happens to the position o equilibrium i a small amount o hydrogen is introduced into B.
- (iii)State and explain the e ect o increasing the pressure in **B** on the position o equilibrium.
- c) (i) Deduce the equilibrium constant expression,*K*, or the equilibrium in **B**. [1]
 - (ii) State the e ect o increasing $[H_2]$ in **B** on the value o K_2 .
- 6. Ammonia production is important in industry.
 - $N_2(g) + 3H_2(g) = 2NH_3(g)$ H = -92 kJ
 - a) The standard entropy values, S, at 298 K or $N_2(g)$, $H_2(g)$ and $NH_3(g)$ are 193, 131 and 192 J K⁻¹ mol⁻¹

3. Consider the ollowing equilibrium: $H^{\oplus} = -909 \,\mathrm{kJ}$ $4NH_{2}(g) + 5O_{2}(g)$ $4NO(g) + 6H_0O(g)$ a) Deduce the equilibrium constant expression, K_{c} , or the reaction. [1] b) Predict the direction in which the equilibrium will shi t when the ollowing changes occur. [4] (i) The volume increases. (ii) The temperature decreases. (iii)H₂O(g) is removed rom the system. (iv)A catalyst is added to the reaction mixture. c) Define the term *activation energy*. [1] 4. An example o a homogeneous reversible reaction is the reaction between hydrogen and iodine. $H_{2}(g) + I_{2}(g) = 2HI(g)$ a) Outline the characteristics o a homogeneous chemical system that is in a state o equilibrium. [2] b) Formulate the expression or the equilibrium constant, K. [1] c) Predict what would happen to the position o equilibrium and the value o K_c i the pressure is increased rom 1 atm to 2 atm. [2] d) The value o K_c at 500 K is 160 and the value o K_c at 700 K is 54. Deduce what this in ormation tells us about the enthalpy change o the orward reaction. [1] (e) Deduce the value o the equilibrium constant, *K* at 500 K or the reaction below: [1] $2HI(g) = H_{2}(g) + I_{2}(g)$

respectively. Calculate S^{\ominus} or the reaction as shown by the equation above. [2] b) Determine G^{\ominus} or the reaction at 298 K. [2]

- c) Describe and explain the e ect o increasing temperature on the spontaneity o the reaction. [2]
- d) Determine the value of the equilibrium constant at 298 K by using the value o G^{\ominus} that you obtained in b). [3]
- e) 0.20 mol o N(g) and 0.20 mol o H₂(g) were allowed to reach equilibrium in a 1 dm³ closed container at a temperature T₂ which is di erent to 298 K. At equilibrium the concentration o NH₃(g) was ound to be 0.060 mol dm⁻³. Determine the value o K_c at temperature T₃.
-) Comment on the two di erent values or *K_c* that you have obtained. [2]
- g) Describe how increasing the pressure a ects the yieldo ammonia. [2]
- h) In practice, typical conditions used in the Haber process are a temperature o 500 C and a pressure o 200 atmospheres. Suggest why these conditions are used rather than those that give the highest yield.
 [2]
- i) Iron is used as a catalyst in this manu acturing process. A catalyst has no e ect on the value o K_c or on the position o equilibrium. Suggest why a catalyst is used in this process. [1]

B ACIDS AND BASES

Theories and properties of acids and bases

THE IONIC THEORY

An acid was originally distinguished by its sour taste. Later it was said to be the oxide o a non-metal combined with water although hydrochloric acid does not fit into this definition. The ionic theory which is still commonly used today states that an acid is a substance which produces hydrogen ions, $H^+(aq)$, in aqueous solution, e.g.

HCl(aq) $H^+(aq) + Cl^-(aq)$

In aqueous solution hydrogen ions are hydrated to orm hydroxonium ions, $H_3O^+(aq)$. In the International Baccalaureate it is correct to write either $H^+(aq)$ or $H_3O^+(aq)$ to represent the hydrogen ions in an aqueous solution. Strictly speaking an acid gives a hydrogen ion concentration in aqueous solution greater than 1.0 10^{-7} mol dm⁻³. A base is a substance that can neutralize an acid. An alkali is a base that is soluble in water.

BRØNSTED LOWRY ACIDS AND BASES

A Br nsted Lowry acid is a substance that can *donate* a proton. A Br nsted Lowry base is a substance that can *accept* a proton. Consider the reaction between hydrogen chloride gas and water.

HCl(g)	$^+$	$H_{2}O(l)$	$H_{_3}O^+(aq)$	$^+$	Cl-(aq)
acid		base	acid		base

Under this definition both HCl and H_3O^+ are acids as both can donate a proton. Similarly both H_2O and Cl^- are bases as both can accept a proton. Cl^- is said to be the **conjugate base** o HCl and H_2O is the conjugate base o H_3O^+ . The conjugate base o an acid is the species remaining a ter the acid has lost a proton. Every base also has a conjugate acid, which is the species ormed a ter the base has accepted a proton. In the reaction with hydrogen chloride water is behaving as a base. Water can also behave as an acid.

$NH_{3}(g)$	$^+$	$H_{2}O(l)$	$\mathrm{NH}_{4}^{+}(\mathrm{aq})$	+	OH ⁻ (aq)
base		acid	acid		base

Substances such as water, which can act both as an acid and as a base, are described as amphiprotic.

Many acids, particularly organic acids, contain one or more non-acidic hydrogen atoms. The location o the acidic hydrogen atom(s) should be clearly identified. For example, ethanoic acid should be written as CH_3COOH rather than $C_2H_4O_2$ so that the conjugate base can be identified as the carboxylate anion CH_3COO^- rather than just $C_2H_3O_2^-$.

TYPICAL PROPERTIES OF ACIDS AND BASES

The typical reactions o acids are:

1. With indicators.

Acid base indicators can be used to determine whether or not a solution is acidic. Common indicators include:

Indicator	Colour in acidic solution	Colour in alkaline solution
litmus	red	blue
phenolphthalein	colourless	pink
methyl orange	red	yellow

2. Neutralization reactions with bases.

a) With hydroxides to orm a salt and water,

e.g. $CH_3COOH(aq) + NaOH(aq) = NaCH_3COO(aq) + H_2O(l)$

b) With metal oxides to orm a salt and water,

- e.g. $H_2SO_4(aq) + CuO(s) = CuSO_4(aq) + H_2O(l)$
- c) With ammonia to orm a salt.

e.g. $HCl(aq) + NH_{3}(aq) = NH_{4}Cl(aq)$

3. With reactive metals (those above copper in the activity series) to orm a salt and hydrogen, e.g.

 $2HCl(aq) + Mg(s) MgCl_2(aq) + H_2(g)$

4. With carbonates (soluble or insoluble) to orm a salt, carbon dioxide and water, e.g.

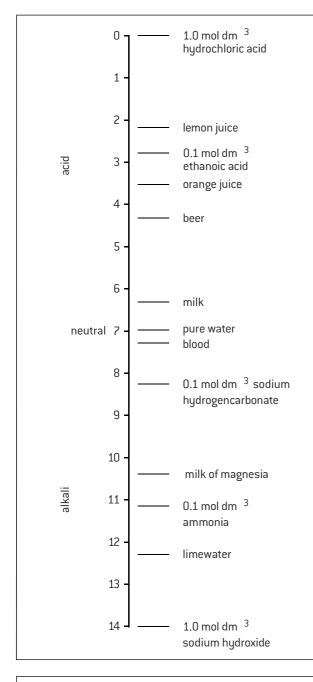
 $2\mathrm{HNO}_{_{3}}(\mathrm{aq}) + \mathrm{Na}_{_{2}}\mathrm{CO}_{_{3}}(\mathrm{aq}) \qquad 2\mathrm{NaNO}_{_{3}}(\mathrm{aq}) + \mathrm{CO}_{_{2}}(\mathrm{g}) + \mathrm{H}_{_{2}}\mathrm{O}(\mathrm{l})$

 $2HCl(aq) + CaCO_{3}(s)$ $CaCl_{2}(aq) + CO_{2}(g) + H_{2}O(l)$

5. With hydrogencarbonates to orm a salt, carbon dioxide and water, e.g.

 $HCl(aq) + NaHCO_3(aq) = NaCl(aq) + CO_2(g) + H_2O(aq)$

The pH scale



DETERMINATION OF pH

The pH o a solution can be determined by using a pH meter or by using universal indicator, which contains a mixture o indicators that give a range o colours at di erent pH values.

рН	•	[H ⁺]/ ol dm ⁻³	-)H ⁻]/ l dm ⁻³	Description	Colour of universal indicator
0	1		1	10^{-14}	very acidic	red
4	1	10^{-4}	1	10^{-10}	acidic	orange
7	1	10^{-7}	1	10^{-7}	neutral	green
10	1	10^{-10}	1	10^{-4}	basic	blue
14	1	10^{-14}	1		very basic	purple

THE pH SCALE

Pure water is very slightly dissociated:

 $H_2O(l) \longrightarrow H^+(aq) + OH^-(aq)$

At 25 C the equilibrium constant or this reaction is $1 - 10^{-14}$.

i.e. $K_w = [H^+(aq)][OH^-(aq)] = 1 \quad 10^{-14}.$

The concentration o the hydrogen ions (which is the same as the concentration o the hydroxide ions) equals $1 \quad 10^{-7}$ mol dm⁻³.

pH (which stands or **p**ower o **H**ydrogen) is defined as being equal to minus the logarithm to the base ten o the hydrogen ion concentration.

i.e. $pH = -log_{10}[H^+(aq)]$

In practice this means that it is equal to the power o ten o the hydrogen ion concentration with the sign reversed. The pH o pure water is thus 7.

Pure water is neutral, so the pH o any neutral solution is 7. I the solution is acidic the hydrogen ion concentration will be greater than 10^{-7} mol dm⁻³ and the pH will decrease. Similarly alkaline solutions will have a pH greater than 7.

The pH scale runs rom 0 to 14. Because it depends on the power o ten a change in one unit in the pH corresponds to a ten old change in the hydrogen ion concentration. A 0.1 mol dm^{-3} solution o a strong monoprotic acid will have a pH o 1, a 0.001 mol dm^{-3} solution o the same acid will have a pH o 3.

1	-4	I	1	1	1	I	I	1	l	5
Log ₁₀	scale 0.0001			ces be 0.1		n pow 10	ers o			ual 100 000
	10 ⁻⁴			10-1	10º I	10 ¹	10²	10 ³	104	10 ⁵

STRONG, CONCENTRATED AND CORROSIVE

In English the words strong and concentrated are o ten used interchangeably. In chemistry they have very precise meanings:

strong: completely dissociated into ions

concentrated: a high number o moles o solute per litre (dm³) o solution

corrosive: chemically reactive.

Similarly weak and dilute also have very di erent chemical meanings:

weak: only slightly dissociated into ions

dilute: a low number o moles o solute per litre o solution.

Strong and weak acids and bases and simple pH calculations

STRONG AND WEAK ACIDS AND BASES

A strong acid is completely dissociated (ionized) into its ions in aqueous solution. Similarly a strong base is completely dissociated into its ions in aqueous solution. Examples o strong acids and bases include:

Strong acids hydrochloric acid, HCl nitric acid, HNO₃ sul uric acid, H₂SO₄

Strong bases sodium hydroxide, NaOH

potassium hydroxide, KOH barium hydroxide, Ba(OH)₂

Note: because one mole o HCl produces one mole o hydrogen ions it is known as a **monoprotic** acid. Sul uric acid is known as a **diprotic** acid as one mole o sul uric acid produces two moles o hydrogen ions.

Weak acids and bases are only slightly dissociated (ionized) into their ions in aqueous solution.

Weak acids ethanoic acid, CH₃COOH carbonic acid (CO₂ in water), H₂CO₃ **Weak bases** ammonia, NH₃ aminoethane, C₂H₅NH₂

The di erence can be seen in their reactions with water:

 $\begin{array}{ll} \mbox{Strong acid: } HCl(g) + H_2O(l) & H_3O^+(aq) + Cl^-(aq) \\ \mbox{ reaction goes to completion} \end{array}$

Weak acid:

 $CH_{3}COOH(aq) + H_{2}O(l) \rightleftharpoons CH_{3}COO^{-}(aq) + H_{3}O^{+}(aq)$ equilibrium lies on the le t

i.e. a solution o hydrochloric acid consists only o hydrogen ions and chloride ions in water, whereas a solution o ethanoic acid contains mainly undissociated ethanoic acid with only very ew hydrogen and ethanoate ions.

Strong base: KOH(s) $\xrightarrow{H_2O(l)} K^+(aq) + OH^-(aq)$ Weak base: NH₃(g) + H₂O(l) $\xrightarrow{}$ NH₄⁺(aq) + OH⁻(aq) equilibrium lies on the le t

EXPERIMENTS TO DISTINGUISH BETWEEN STRONG AND WEAK ACIDS AND BASES

1. pH measurement

Because a strong acid produces a higher concentration o hydrogen ions in solution than a weak acid, with the same concentration, the pH o a strong acid will be lower than a weak acid. Similarly a strong base will have a higher pH in solution than a weak base, with the same concentration. The most accurate way to determine the pH o a solution is to use a pH meter.

$0.10 \text{ mol } dm^{-3} \text{ HCl}(aq) \qquad \text{pH}$
--

 $0.10 \text{ mol } dm^{-3} \text{ CH}_{3} \text{COOH}$ pH = 2.9

- Conductivity measurement Strong acids and strong bases in solution will give much higher readings on a conductivity meter than equimolar (equal concentration) solutions o weak acids or bases, because they contain more ions in solution.
- 3. Concentration measurement As the concentration o hydrogen ions is much greater, the rate o reaction o strong acids with metals, metal oxides, metal hydroxides, metal hydrogen carbonates and metal carbonates is greater than that o weak acids with the same concentration.

STRONG ACID AND BASE pH CALCULATIONS

For pure water the pH must be 7 at 25 C as the concentration o $H^+(aq)$ is equal to the concentration o $OH^-(aq)$ and $[H^+(aq)] = [OH^-(aq)] = 1 = 10^{-14}$. Strong acids are completely dissociated so, or example, the hydrogen ion concentration o 0.100 mol dm⁻³ hydrochloric acid, HCl(aq) will be 0.100 mol dm⁻³ as each mole o acid produces one mole o hydrogen ions when

it dissociates (ionizes). The pH o 0.100 mol dm⁻³ HCl(aq) will there ore be equal to $-\log_{10} (0.100) = 1$.

I the acid is diluted ten times the new hydrogen ion concentration will be 0.0100 mol dm⁻³ and the pH = $-\log_{10} (0.0100)$ or $-\log_{10} (1.00 \quad 10^{-2}) = 2$.

Sul uric acid is assumed or simplicity to be a strong diprotic acid. The hydrogen ion concentration o 0.0100 mol dm⁻³ H₂SO₄(aq) will there ore be 2 0.0100 or 2.00 10^{-2} mol dm⁻³ and the pH will equal $-\log_{10} (2.00 \ 10^{-2}) = 1.7$.

Note that pH is a measure o concentration so 10.0 cm³ o 0.100 mol dm⁻³ HCl(aq) will have the same pH as 100 cm³ o 0.100 mol dm⁻³ HCl(aq). Also note that $[H^+(aq)] = 10^{-pH}$ so i the pH o an acid is 3 then $[H^+(aq)] = 10^{-3} = 1.0$ 10^{-3} mol dm⁻³.

To calculate the pH o a strong base work out the hydroxide concentration first then calculate the hydrogen ion concentration rom the expression $[H^+(aq)] = [OH^-(aq)] = 1 = 10^{-14}$.

Example 1: To calculate the pH o 0.100 mol dm⁻³ NaOH(aq). $[OH^{-}(aq)] = 0.100 mol dm^{-3}$ Example 2: To calculate the pH o 0.100 mol dm⁻³ Ba(OH)₂(aq). $[OH^{-}(aq)] = 0.100 mol dm^{-3}$ $[OH^{-}(aq)] = 0.200 mol dm^{-3}$ $[H^{+}(aq)] = \frac{1}{0.100} = 1$ $10^{-13} mol dm^{-3}$ $pH = -log_{10} (1.00 - 10^{-13}) = 13.$ $PH = -log_{10} (5.00 - 10^{-14}) = 13.3$

Acid deposition

OXIDES OF SULFUR SO

Sul ur dioxide occurs naturally rom volcanoes. It is produced industrially rom the combustion o sul ur-containing ossil uels and the smelting o sulfide ores.

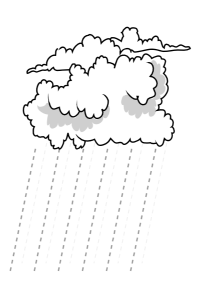
$$\begin{split} S(s) + O_{_2}(g) & SO_{_2}(g) \\ \label{eq:solution} \mbox{In the presence o sunlight sul ur} \\ \mbox{dioxide is oxidized to sul ur trioxide}. \end{split}$$

 $SO_{2}(g) + \frac{1}{2}O_{2}(g) = SO_{3}(g)$

The oxides can react with water in the air to orm sul urous acid and sul uric acid:

 $\begin{array}{lll} SO_2(g) + H_2O(l) & H_2SO_3(aq)\\ and \\ SO_3(g) + H_3O(l) & H_3SO_4(aq) \end{array}$

ACID DEPOSITION



OXIDES OF NITROGEN NO_v

Nitrogen oxides occur naturally rom electrical storms and bacterial action. Nitrogen monoxide is produced in the internal combustion engine and in jet engines. $N_{2}(g) + O_{2}(g) = 2NO(g)$ Oxidation to nitrogen dioxide occurs in the air. $2NO(g) + O_{2}(g) = 2NO_{2}(g)$ The nitrogen dioxide then reacts with water to orm nitric acid and nitrous acid: $2NO_3(g) + H_2O(l) = HNO_3(aq) +$ HNO (aq) or is oxidized directly to nitric acid by oxygen in the presence o water: $4NO_{2}(g) + O_{2}(g) + 2H_{2}O(l)$ 4HNO₁(aq)

Pure rainwater is naturally acidic with a pH o 5.65 due to the presence o dissolved carbon dioxide. Carbon dioxide itsel is *not* responsible or acid rain since acid rain is defined as rain with a pH less than 5.6. It is the oxides o sul ur and nitrogen present in the atmosphere which are responsible or **acid deposition** the process by which acidic particles, gases and precipitation leave the atmosphere. Wet deposition, due to the acidic oxides dissolving and reacting with water in the air, is known as acid rain and includes og, snow and dew as well as rain. Dry deposition includes acidic gases and particles.

VEGETATION

Increased acidity in the soil leaches important nutrients, such as Ca^{2+} , Mg^{2+} and K^+ . Reduction in Mg²⁺ can cause reduction in chlorophyll and consequently lowers the ability o plants to photosynthesize. Many trees have been seriously a ected by acid rain. Symptoms include stunted growth, thinning o tree tops, and yellowing and loss o leaves. The main cause is the aluminium leached rom rocks into the soil water. The Al³⁺ ion damages the roots and prevents the tree rom taking up enough water and nutrients to survive.

LAKES AND RIVERS Increased levels o

aluminium ions in water can kill fish. Aquatic li e is also highly sensitive to pH. Below pH 6 the number o sensitive fish, such as salmon and minnow, decline as do insect larvae and algae. Snails cannot survive a pH less than 5.2 and below pH 5.0 many microscopic animal species disappear. Below pH 4.0 lakes are e ectively dead. The nitrates present in acid rain can also lead to eutrophication.

BUILDINGS

Stone, such as marble, that contains calcium carbonate is eroded by acid rain. With sul uric acid the calcium carbonate reacts to orm calcium sul ate, which can be washed away by rainwater thus exposing more stone to corrosion. Salts can also orm within the stone that can cause the stone to crack and disintegrate.

$$\begin{split} & \mathsf{CaCO}_{_3}(s) + \mathrm{H_2SO}_{_4}(aq) \\ & \mathsf{CaSO}_{_4}(aq) + \mathrm{CO}_{_2}(g) + \mathrm{H_2O}(l) \end{split}$$

HUMAN HEALTH

The acids ormed when NO₂ and SO₂ dissolve in water irritate the mucous membranes and increase the risk o respiratory illnesses, such as asthma, bronchitis and emphysema. In acidic water there is more probability o poisonous ions, such as Cu²⁺ and Pb²⁺, leaching rom pipes and high levels o aluminium in water may be linked to Alzheimer s disease.

METHODS TO LOWER OR COUNTERACT THE EFFECTS OF ACID RAIN

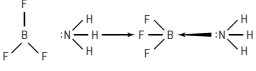
- 1. Lower the amounts o NO_x and SO_x ormed, e.g. by improved engine design, the use o catalytic converters, and removing sul ur be ore, during, and a ter combustion o sul ur-containing uels.
- 2. Switch to alternative methods o energy (e.g. wind and solar power) and reduce the amount o uel burned, e.g. by reducing private transport and increasing public transport and designing more e ficient power stations.
- 3. Liming o lakes adding calcium oxide or calcium hydroxide (lime) neutralizes the acidity, increases the amount o calcium ions and precipitates aluminium rom solution. This has been shown to be e ective in many, but not all, lakes where it has been tried.



LEWIS ACIDS AND BASES

Br nsted Lowry bases must contain a non-bonding pair o electrons to accept the proton. The Lewis definition takes this urther and describes bases as substances which can *donate* a pair o electrons, and acids as substances which can *accept* a pair o electrons. In the process a coordinate (both electrons provided by one species) covalent bond is ormed between the base and the acid.

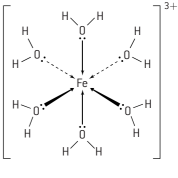
The Lewis theory is all-embracing, so the term Lewis acid is usually reserved or substances which are not also Br nsted Lowry acids. Many Lewis acids do not even contain hydrogen.



acid bas

BF₃ is a good Lewis acid as there are only six electrons around the central boron atom which leaves room or two more. Other common Lewis acids are aluminium chloride, AlCl₃, and also transition metal ions in aqueous solution which can accept a pair o electrons rom each o six surrounding water molecules, e.g. $[Fe(H_2O)_6]^{3+}$.

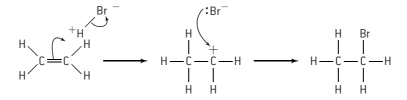
Note that hydrated transition metal ions, such as $[Fe(H_2O)_6]^{3+}$ are acidic in solution as the +3 charge is spread over a very small ion which gives the ion a high charge density. The non-bonded pair o electrons on one o the water molecules surrounding the ion will be strongly attracted to the ion and the water molecule will lose a hydrogen ion in the process. This process can continue until iron(III) hydroxide is ormed. The equilibrium can be urther moved to the right by adding hydroxide ions, OH⁻(aq) or back to the let by adding hydrogen ions, H⁺(aq) which exemplifies the ion's amphoteric nature.



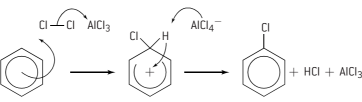
$$[Fe(H_{2}0)_{6}]^{3+} \xrightarrow{-H^{+}}_{H^{+}} [Fe(H_{2}0)_{5}0H]^{2+} \xrightarrow{-H^{+}}_{H^{+}} [Fe(H_{2}0)_{4}(0H)_{2}]^{+} \xrightarrow{-H^{+}}_{H^{+}} Fe(H_{2}0)_{3}(0H)_{3}$$
$$0H^{-} \iint_{H^{+}} H^{+} [Fe(H_{2}0)_{2}(0H)_{4}]^{-}$$

The Lewis acid and base concept is also used in organic chemistry particularly to identi y reacting species and in the use o curly arrows to explain the movement o pairs o electrons in organic reaction mechanisms.

For example the addition o hydrogen bromide to an alkene proceeds by an electrophilic addition mechanism (see page 90). The ⁺ hydrogen atom o the hydrogen bromide molecule acts as the electrophile and accepts a pair o electrons rom the double bond o the alkene. Hence, the electrophile is acting as a Lewis acid and the alkene is acting as a Lewis base. Since a curly arrow shows the movement o a pair o electrons the arrow always originates rom the Lewis base and the head o the curly arrow always points towards the Lewis acid. A second Lewis acid base reaction occurs when the bromide ion (acting as the Lewis base) donates a pair o electrons to the positive carbon atom in the carbocation intermediate to orm the brominated addition product.



Another good example o Lewis acids in organic chemistry is the unction o halogen carriers as catalysts in the electrophilic reactions o benzene (see page 88). Benzene is electron-rich due to its delocalized bond and can react with chlorine to orm chlorobenzene. A halogen carrier such as aluminium chloride is added to provide a positive chloride ion which acts as the electrophile. So in this reaction aluminium chloride (Lewis acid) and chlorine (Lewis base) undergo a Lewis acid base reaction and then a second Lewis acid base reaction occurs between the positive chloride ion (Lewis acid) and the benzene molecule (Lewis base). A third Lewis acid base reaction occurs when the hydrogen atom (Lewis acid) is removed rom the intermediate to orm hydrogen chloride and regenerate the aluminium chloride catalyst.



🕕 Calculations involving pH, pOH and pK

THE IONIC PRODUCT OF WATER

Pure water is very slightly ionized:

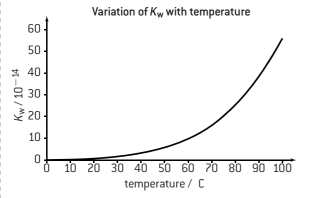
 $\mathrm{H}_{2}\mathrm{O}(\mathrm{l}) \rightleftharpoons \mathrm{H}^{+}(\mathrm{aq}) + \mathrm{OH}^{-}(\mathrm{aq}) \qquad H^{\ominus} = +57.3 \mathrm{ kJ mol}^{-1}$

 $K_{c} = \frac{[\mathrm{H}^{+}(\mathrm{aq})] [\mathrm{OH}^{-}(\mathrm{aq})]}{[\mathrm{H}_{2}\mathrm{O}(\mathrm{l})]}$

Since the equilibrium lies far to the left the concentration of water can be regarded as constant so

 $K_{\rm w} = [{\rm H^+}({\rm aq})]$ [OH⁻(aq)] = 1.00 10⁻¹⁴ at 298 K, where $K_{\rm w}$ is known as the ionic product of water.

The dissociation of water into its ions is an endothermic process, so the value of K_w will increase as the temperature is increased.



For pure water $[H^+(aq)] = [OH^-(aq)]$
$= 1.00 10^{-7} \text{ mol } dm^{-3} \text{ at } 298 \text{ K}$
From the graph the value for $K_{\rm w} = 1.00 10^{-13}$ at 334 K (61 C)
At this temperature $[H^+(aq)] = 1.00 10^{-13}$ = 3.16 10^{-7} mol dm ⁻³

pH, pOH AND pK __ FOR STRONG ACIDS AND BASES

As stated earlier in this chapter the pH of a solution depends only upon the hydrogen ion concentration and is independent of the volume of the solution.

_ _ _ _ _ _ _

 $pH = -log_{10} [H^+(aq)]$

For strong monoprotic acids the hydrogen ion concentration will be equal to the concentration of the acid and will be twice the value of the acid concentration for strong diprotic acids.

The use of the logarithmic scale can be extended to other values, e.g. pOH and pK_{w} .

 $pOH = -log_{10} [OH^-(aq)] and pK_w = -log_{10} K_w$

 $K_{\rm w} = [\mathrm{H^{\scriptscriptstyle +}}(\mathrm{aq})] \quad [\mathrm{OH^{\scriptscriptstyle -}}(\mathrm{aq})]$

If logarithms to the base ten are taken then

 $\log_{10} K_w = \log_{10} [H^+(aq)] + \log_{10} [OH^-(aq)]$ which can also be written as

 $-\log_{10}K_{w} = -\log_{10} [H^{+}(aq)] - \log_{10} [OH^{-}(aq)]$

This leads to the useful expression

$$pK_w = pH + pOH$$

At 25 C $K_{w} = 10^{-14}$ and pH + pOH = 14

This expression gives another way of calculating the pH of a strong base since the pOH can be determined directly from the hydroxide ion concentration then subtracted from 14. For example, to determine the pH of $4.00 - 10^{-3}$ mol dm⁻³ Ba(OH),

 $[OH^{-}(aq)] = 2$ 4.00 $10^{-3} = 8.00$ 10^{-3} mol dm⁻³

 $pOH = -log_{10} 8.00 \quad 10^{-3} = 2.10$

pH = 14 - 2.10 = 11.9

🕕 Calculations with weak acids and bases

WEAK ACIDS

The dissociation of a weak acid HA in water can be written:

 $HA(aq) \rightarrow H^+(aq) + A^-(aq)$

The equilibrium expression for this reaction is:

 $K_{a} = \frac{[\mathrm{H}^{+}] \quad [\mathrm{A}^{-}]}{[\mathrm{HA}]}$ where K_{a} is known as the acid dissociation constant

For example, to calculate the pH of 0.10 mol dm⁻³ CH₃COOH given that $K_a = 1.8 \quad 10^{-5}$ mol dm⁻³ at 298 K:

 $CH_{3}COOH(aq) \iff CH_{3}COO^{-}(aq) + H^{+}(aq)$ Initial concentration / mol dm⁻³ 0.10 Equilibrium concentration / mol dm⁻³ (0.10 - x) x x K = $\frac{[CH_{3}COO^{-}] [H^{+}]}{x^{2}} = \frac{x^{2}}{x^{2}}$

$$\begin{aligned} & \text{[CH}_{3}\text{COOH]} & (0.10 - x) \\ &= 1.8 \quad 10^{-5} \text{ mol dm}^{-3} \\ & x^{2} + (1.8 \quad 10^{-5}x) - 1.8 \quad 10^{-6} = 0 \\ & \text{by solving the quadratic equation} \end{aligned}$$

x = 1.33 10⁻³ mol dm⁻³

 $pH = -\log_{10} 1.33 \quad 10^{-3} = 2.88$

If the acids are quite weak the equilibrium concentration of the acid can be assumed to be the same as its initial concentration. Provided the assumption is stated it is usual to simplify the expression in calculations to avoid a quadratic equation. In the above example:

$$K_{a} = \frac{[CH_{3}COO^{-}] [H^{+}]}{[CH_{3}COOH]} \frac{[H^{+}]^{2}}{0.10}$$
$$= 1.8 \quad 10^{-5} \text{ mol } dm^{-3}$$
$$[H^{+}] = 1.8 \quad 10^{-6} = 1.34 \quad 10^{-3} \text{ mol } dm^{-3}$$

pH = 2.87

Examples of other weak acid calculations

 The pH of a 0.020 mol dm⁻³ solution of a weak acid is 3.9. Find the K₃ of the acid.

$$K_{a} = \frac{[\mathrm{H}^{+}]^{2}}{(0.020 - [\mathrm{H}^{+}])} \frac{10^{-3.9}}{0.020}$$

 $= 7.92 \quad 10^{-7} \text{ mol } dm^{-3}$

2. An acid whose K_a is 4.1 10^{-6} mol dm⁻³ has a pH of 4.5. Find the concentration of the acid.

$$[\text{HA}] = \frac{[\text{H}^+]^2}{K_{\text{a}}} = \frac{10^{-4.5}}{4.1} \frac{10^{-4.5}}{10^{-6}}$$

$$= 2.44 \quad 10^{-4} \text{ mol } dm^{-3}$$

Note that the weaker the weak acid the **smaller** the value of K_a and the **larger** the value of pK_a . Thus ethanoic acid $(pK_a = 4.76)$ is a weaker acid than methanoic acid $(pK_a = 3.75)$.

WEAK BASES

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The reaction of a weak base can be written:
```

 $B(aq) + H_2O(l) \implies BH^+(aq) + OH^-(aq)$

Since the concentration of water is constant:

$$K_{\rm b} = \frac{[{\rm B}{\rm H}^+] [{\rm O}{\rm H}^-]}{[{\rm B}]}$$

where K_{b} is the base dissociation constant If one considers the reverse reaction of BH⁺ acting as an acid to give B and H⁺ then:

 $K_{a} = \frac{[B] [H^{+}]}{[BH^{+}]}$

then

$$\begin{split} K_{\mathrm{a}} \quad K_{\mathrm{b}} &= \frac{[\mathrm{B}] \quad [\mathrm{H}^+]}{[\mathrm{B}\mathrm{H}^+]} \quad \frac{[\mathrm{B}\mathrm{H}^+] \quad [\mathrm{O}\mathrm{H}^-]}{[\mathrm{B}]} \\ &= [\mathrm{H}^+] \quad [\mathrm{O}\mathrm{H}^-] = K_{\mathrm{w}} \end{split}$$

since $pK_a = -\log_{10} K_a$; $pK_b = -\log_{10} K_b$ and $pK_w = -\log_{10} K_w$ = 14 this can also be expressed as:

$$pK_{a} + pK_{b} = 14$$

Examples of calculations

1. The $K_{\rm b}$ value for ammonia is 1.8 10^{-5} mol dm⁻³. Find the pH of a 1.00 10^{-2} mol dm⁻³ solution.

Since $[NH_4^+] = [OH^-]$ then

$$K_{\rm b} = \frac{[\rm OH^{-}]^2}{[\rm NH_3]} - \frac{[\rm OH^{-}]^2}{1.00 \ 10^{-2}} = 1.8 \ 10^{-2}$$

 $[OH^{-}] = 1.8 \quad 10^{-7} = 4.24 \quad 10^{-4} \text{ mol } dm^{-3}$

$$pOH = -log_{10} 4.24$$
 $10^{-4} = 3.37$
 $pH = 14 - 3.37 = 10.6$

2. The pH of a 3.00 10^{-2} mol dm⁻³ solution of weak base is 10.0. Calculate the p $K_{\rm b}$ value of the base.

$$pH = 10.0 \text{ so } pOH = 4.0$$

$$K_{\rm b} = \frac{10^{-4}}{3.00} \frac{10^{-4}}{10^{-2}} = 3.33 \quad 10^{-7} \text{ mol } \mathrm{dm}^{-3}$$

 $pK_{b} = 6.48$

 The value for the pK_a of methylamine (aminomethane) is 10.66. Calculate the concentration of an aqueous solution of methylamine with a pH of 10.8.

$$pK_{b} = 14 - 10.66 = 3.34; pOH = 14 - 10.8 = 3.2$$

[OH⁻¹² 10^{-3.2} 10^{-3.2}

$$[CH_{3}NH_{2}] = \frac{[011]_{1}}{K_{b}} = \frac{10^{-1}10^{-1}}{10^{-3.34}}$$
$$= 8.71 \quad 10^{-4} \text{ mol } dm^{-3}$$

Note that the weaker the weak base the **smaller** the value of $K_{\rm b}$ and the **larger** the value of $\mathbf{p}K_{\rm b}$. Thus ammonia $(\mathbf{p}K_{\rm b} = 4.75)$ is a weaker base than methylamine $(\mathbf{p}K_{\rm b} = 3.34)$.

💷 Salt hydrolysis and buffer solutions

SALT HYDROLYSIS

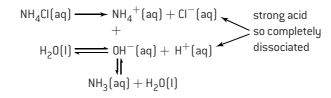
Sodium chloride is neutral in aqueous solution. It is the salt o a strong acid and a strong base so its ions remain completely dissociated in solution. Salts made rom a weak acid and a strong base, such as sodium ethanoate, are alkaline in solution. This is because the ethanoate ions will combine with hydrogen ions rom water to orm mainly undissociated ethanoic acid, leaving excess hydroxide ions in solution.

$$NaCH_{3}COD(aq) \longrightarrow Na^{+}(aq) + CH_{3}COD^{-}(aq)$$

$$+ H_{2}O(I) \longrightarrow OH^{-}(aq) + H^{+}(aq)$$

strong base so completely dissociated CH₃COOH(aq)

Similarly salts derived rom a strong acid and a weak base will be acidic in solution.



BUFFER SOLUTIONS

A bu er solution resists changes in pH when small amounts o acid or alkali are added to it.

An acidic bu er solution can be made by mixing a weak acid together with the salt o that acid and a strong base. An example is a solution o ethanoic acid and sodium ethanoate. The weak acid is only slightly dissociated in solution, but the salt is ully dissociated into its ions, so the concentration o ethanoate ions is high.

 $NaCH_3COO(aq) = Na^+(aq) + CH_3COO^-(aq)$

 $CH_2COOH(aq) \rightleftharpoons CH_2COO^-(aq) + H^+(aq)$

I an acid is added the extra H⁺ ions coming rom the acid are removed as they combine with ethanoate ions to orm undissociated ethanoic acid, so the concentration o H⁺ ions remains unaltered.

 $CH_{2}COO^{-}(aq) + H^{+}(aq) \Longrightarrow CH_{2}COOH(aq)$

I an alkali is added the hydroxide ions rom the alkali are removed by their reaction with the undissociated acid to orm water, so again the H⁺ ion concentration stays constant.

 $CH_3COOH(aq) + OH^-(aq) = CH_3COO^-(aq) + H_2O(l)$

In practice acidic bu ers are o ten made by taking a solution o a strong base and adding excess weak acid to it, so that the solution contains the salt and the unreacted weak acid.

 $NaOH(aq) + CH_2COOH(aq)$ $NaCH_{2}COO(aq) + H_{2}O(l) + CH_{2}COOH(aq)$ limiting reagent salt

excess weak acid

bu er solution

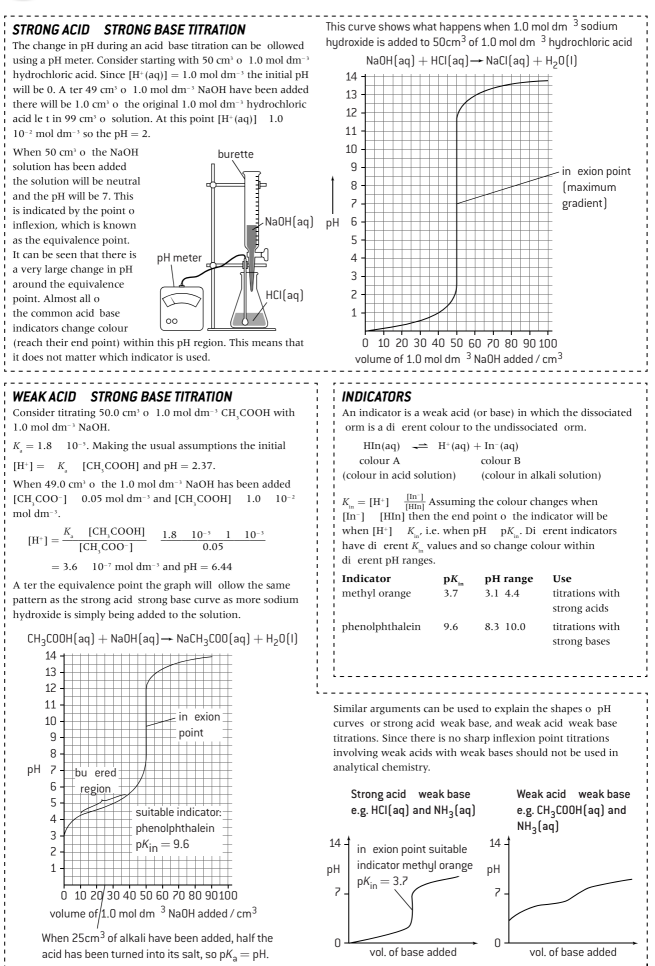
An alkali bu er with a fixed pH greater than 7 can be made rom a weak base together with the salt o that base with a strong acid. An example is ammonia with ammonium chloride.

 $NH_4Cl(aq) = NH_4^+(aq) + Cl^-(aq)$

 $NH_{a}(aq) + H_{a}O(l) \implies NH_{a}^{+}(aq) + OH^{-}(aq)$

I H⁺ ions are added they will combine with OH⁻ ions to orm water and more o the ammonia will dissociate to replace them. I more OH⁻ ions are added they will combine with ammonium ions to orm undissociated ammonia. In both cases the hydroxide ion concentration and the hydrogen ion concentration remain constant.

🕕 Titration curves and indicators



MULTIPLE CHOICE QUESTIONS ACIDS AND BASES

- Which statement about hydrochloric acid is alse?
 A. It can react with copper to give hydrogen
 - B. It can react with sodium carbonate to give carbon dioxide
 - C. It can react with ammonia to give ammonium chloride
 - D. It can react with copper oxide to give water
- 2. 1.00 cm³ o a solution has a pH o 3. 100 cm³ o the same solution will have pH o :
 - A. 1 C. 5
 - B. 3 D. Impossible to calculate rom the data given.
- 3. Which statement(s) is/are true about separate solutions o a strong acid and a weak acid both with the same concentration?
 - I. They both have the same pH.
 - II. They both have the same electrical conductivity.
 - A. I and II C. II only
 - B. I only D. Neither I nor II
- Identi y the correct statement about 25 cm³ o a solution o 0.1 mol dm⁻³ ethanoic acid CH₃COOH.
 - A. It will contain more hydrogen ions than 25 cm³ o
 0.1 mol dm⁻³ hydrochloric acid.
 - B. It will have a pH greater than 7.
 - C. It will react exactly with 25 cm 3 o $\,$ 0.1 mol dm $^{-3}$ sodium hydroxide.
 - D. It is completely dissociated into ethanoate and hydrogen ions in solution.
- 5. What is the pH o 1.0 10⁻⁴ mol dm⁻³ sul uric acid, H₂SO₄(aq)?
 A. -4
 C. 4

B. between 3 and 4D. between 4 and 5

- HL
- 11. During the titration o a known volume o a strong acid with a strong base:
 - A. there is a steady increase in pH
 - B. there is a sharp increase in pH around the end point
 - C. there is a steady decrease in pH
 - D. there is a sharp decrease in pH around the end point.
- 12. Three acids, HA, HB, and HC have the ollowing K_a values K_a (HA) = 1 10^{-5} K_a (HB) = 2 10^{-5}
 - $K_{a}(HC) = 1 10^{-6}$

What is the correct order o increasing acid strength (weakest first)?

А. НА, НВ, НС	С.	НС, НА, НВ
B. HC, HB, HA	D.	НВ, НА, НС

- D. IIC, IID, IIA D. IID, IIA
- 13. Which o the ollowing reagents could not be added together to make a bu er solution?
 - A. NaOH(aq) and CH₃COOH(aq)
 - B. NaCH₃COO(aq) and CH₃COOH(aq)
 - C. NaOH(aq) and NaCH₃COO(aq)
 - D. $NH_4Cl(aq)$ and $NH_3(aq)$

- NH₃(aq), HCl(aq), NaOH(aq), CH₃COOH(aq) When 1.0 mol dm⁻³ solutions o the substances above are arranged in order o **decreasing** pH the order is:
 - A. NaOH(aq), $NH_{3}(aq)$, $CH_{3}COOH(aq)$, HCl(aq)
 - B. $NH_3(aq)$, NaOH(aq), HCl(aq), $CH_3COOH(aq)$
 - C. $CH_{3}COOH(aq)$, HCl(aq), NaOH(aq), $NH_{3}(aq)$
 - D. HCl(aq), CH₃COOH(aq), NH₃(aq), NaOH(aq)
- 7. A solution with a pH o 8.5 would be described as:
 - A. very basic C. slightly acidic
 - B. slightly basic D. very acidic
- 8. Which statement is true about two solutions, one with a pH o 3 and the other with a pH o 6?
 - A. The solution with a pH o 3 is twice as acidic as the solution with a pH o 6
 - B. The solution with a pH o 6 is twice as acidic as the solution with a pH o 3
 - C. The hydrogen ion concentration in the solution with a pH o 6 is one thousand times greater than that in the solution with a pH o 3
 - D. The hydrogen ion concentration in the solution with a pH o 3 is one thousand times greater than that in the solution with a pH o 6
- Which o the ollowing is not a conjugate acid base pair?
 A. HNO₃/NO₃⁻
 - B. H,SO,/HSO
 - C. NH₃/NH₂⁻
 - D. H, O⁺/OH
- 10. Which gas cannot lead to acid deposition?
 - A. CO₂ C. NO B. SO, D. NO,
- 14. When 1.0 cm³ o a weak acid solution is added to 100 cm³ o a bu er solution:
 - A. the volume o the resulting mixture will be 100 cm³
 - B. there will be almost no change in the pH o the solution
 - C. the pH o the solution will increase noticeably
 - D. the pH o the solution will decrease noticeably.
- 15. Which species cannot act as a Lewis acid?

A. NH ₃	C. Fe ²⁺
B. BF ₃	D. AlCl ₃

- 16. Which salt does not orm an acidic solution in water?
 - A. MgCl₂ C. FeCl₃
 - B. Na₂CO₃ D. NH₄NO₃
- 17. An indicator changes colour in the pH range 8.3 10.0. This indicator should be used when titrating a known volume o :
 - A. a strong acid with a weak base
 - B. a weak acid with a weak base
 - C. a weak base with a strong acid
 - D. a weak acid with a strong base.

SF

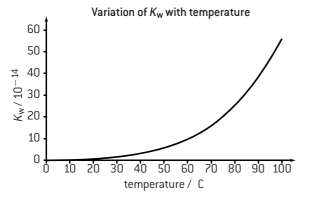
HOR	T ANSWER QUESTIONS ACIDS AND BASES	
a)	(i) De ne a Br nsted Lowry base.	[1]
	(ii) Deduce the two acids and their conjugate bases in the ollowing reaction:	
	$H_2O(l) + HCl(aq) \implies H_3O^+(aq) + Cl^-(aq)$	[2]
b)	Ethanoic acid, CH ₃ COOH, is a weak acid.	
	(i) Explain the di erence between a strong acid and a weak acid.	[2]
	(ii) State the equation or the reaction o ethanoic acid with aqueous ammonia.	[1]
	(iii) Compare and contrast the reactions o $$ 1.00 mol dm^- 3 hydrochloric acid and 1.00 mol dm^- 3 ethanoic acid	
	with excess magnesium metal.	[4]
a)	10 cm ³ o 5.00 10^{-3} mol dm ⁻³ sul uric acid, H ₂ SO ₄ (ac is added to an empty volumetric flask.	l) <i>,</i>
	(i) Calculate the pH o the sul uric acid solution (assume it is a completely strong diprotic acid).	[2]
	(ii) Determine the pH o the diluted solution i the total volume is made up to 100 cm³ with	
	distilled water.	[2]
b)	State the equation or the reaction o sul uric acid with sodium hydroxide solution.	[1]
c)	The diluted solution in (a)(ii) is used to titrate 25.0 cm^3	

- o 1.00 10⁻⁴ mol dm⁻³ sodium hydroxide solution.
- (i) Describe what will be observed when the end point is reached i phenolphthalein is used as the indicator or this titration. [2]

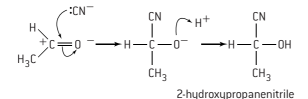
1.

2.

5. The graph below shows how the ionic product o water, K_{w} varies with temperature.



- a) State the equation or the dissociation o water and deduce rom the graph whether the reaction is exothermic or endothermic. [2]
- b) Determine the hydrogen ion concentration and the hydroxide ion concentration in pure water at 90 C and hence deduce the pH o pure water at this temperature. [3]
- 6. a) Explain why a nucleophile can also be described as a Lewis base. [1]
 - b) The diagram below shows the mechanism or the nucleophilic addition reaction between cyanide ions and ethanal to orm 2-hydroxypropanenitrile.



]		(ii) Determine the volume o acid needed to reach the equivalence point o this titration.	[2]
]	3.	When hydrochloric acid is added to a solution o sodium hydrogencarbonate, NaHCO ₃ (aq) carbon dioxide is evolved	l.
		a) State the equation or this reaction.	[2]
1		b) The hydrogencarbonate ion can act either as an acid or a base according to Br nsted Lowry theory.	
1		(i) Deduce the ormula o the conjugate base i it is behaving as an acid.	[1]
		(ii) Deduce the ormula o the acid i it is behaving as a conjugate base.	[1]
]		 c) State the equations or the reaction o hydrochloric acid with (i) copper(II) oxide CuO and (ii) sodium carbonate, Na₂CO₃. 	[2]
1	4.	a) Explain why rain water with a pH o 6 is not classi ed as acid rain even though its pH is less than 7?	[2]
		b) State the equation or the ormation o nitrogen(II) oxide, NO(g), in an internal combustion engine and	
]		describe with equations how it is converted into nitric acid in the atmosphere.	[4]
]		c) Explain why marble statues become corroded by acid rain.	[2]
		 d) Outline why adding calcium hydroxide (lime) to lakes can reduce the e ects o acid deposition. 	[2]

- (i) Identi y the nucleophile in this reaction. [1]
- (ii) Explain the mechanism in terms o Lewis acid base theory. [3]

[3]

- c) Explain why the reaction between a transition metal ion and six monodentate ligands to orm an octahedral complex ion is an example o a Lewis acid base reaction.
- 7. a) (i) State the equation or the reaction between propanoic acid, C,H,COOH(aq) and water and deduce the equilibrium expression. [2]
 - (ii) Calculate the pH o a $2.00 mtext{ } 10^{-3} ext{ mol dm}^{-3}$ solution o propanoic acid ($pK_a = 4.87$). [3]
 - (iii) State any assumptions you have made in arriving at your answer to (a)(ii). [2]
 - b) 25.0 cm³ o 2.00 10⁻³ mol dm⁻³ sodium hydroxide solution, NaOH(aq) was added to 50 cm3 o 2.00 10⁻³ mol dm⁻³ propanoic acid.
 - (i) Identi y all the chemical species present in the resulting solution. [2]
 - (ii) Explain how the resulting solution can unction as a bu er solution is a small amount o alkali is added. [2]
- 8. A particular indicator is a weak acid and can be represented as HIn. The K_1 or HIn is 3 10^{-10} . HIn(aq) is colourless and In⁻(aq) is pink in aqueous solution.
 - a) Identi y the colour this indicator will show in strongly alkaline solution. [1]
 - b) Explain whether or not this indicator would be suitable to use when titrating hydrochloric acid with ammonia solution. [2]
 - Explain why no acid base indicator is suitable to use C) when titrating ethanoic acid with ammonia solution. [1]

9 REDOX PROCESSES R d [1]

DEFINITIONS OF OXIDATION AND REDUCTION

Oxidation used to be narrowly defined as the addition o oxygen to a substance. For example, when magnesium is burned in air the magnesium is oxidized to magnesium oxide.

 $2Mg(s) + O_2(g) = 2MgO(s)$

The electronic configuration o magnesium is $[Ne]3s^2$. During the oxidation process it loses two electrons to orm the Mg^{2+} ion with the electronic configuration o [Ne]. **Oxidation** is now defined as the *loss of one or more electrons from a substance*. This is a much broader definition, as it does not necessarily involve oxygen. Bromide ions, or example, are oxidized by chlorine to orm bromine.

 $2Br^{-}(aq) + Cl_2(aq) = Br_2(aq) + 2Cl^{-}(aq)$

I a substance loses electrons then something else must be gaining electrons. *The gain of one or more electrons* is called **reduction**. In the first example oxygen is reduced as it is gaining two electrons rom magnesium to orm the oxide ion O^{2-} . Similarly, in the second example chlorine is reduced as each chlorine atom gains one electron rom a bromide ion to orm a chloride ion.

Since the processes involve the trans er o electrons oxidation and reduction must occur simultaneously. Such reactions are known as **redox reactions**. In order to distinguish between the two processes hal -equations are o ten used:

$2Mg(s) = 2Mg^{2+}(s) + 4e^{-}$	OXIDATION	$2Br^{-}(aq)$ $Br_{2}(aq) + 2e^{-}$
$O_{_2}(g) + 4e^- 2O^{_2-}(s)$	REDUCTION	$\operatorname{Cl}_2(\operatorname{aq}) + 2e^- 2\operatorname{Cl}^-(\operatorname{aq})$
$2Mg(s) + O_2(g) = 2MgO(s)$	OVERALL REDOX EQUATION	$2Br^{-}(aq) + Cl_2(aq) = Br_2(aq) + 2Cl^{-}(aq)$

Understanding that magnesium must lose electrons and oxygen must gain electrons when magnesium oxide MgO is ormed rom its elements is a good way to remember the definitions o oxidation and reduction. Some students pre er to use the mnemonic OILRIG: **O**xidation **I**s the **L**oss o electrons, **R**eduction **I**s the **G**ain o electrons.

RULES FOR DETERMINING OXIDATION STATES

It is not always easy to see how electrons have been trans erred in redox processes. Oxidation states can be a use ul tool to identi y which species have been oxidized and which reduced. Oxidation states are assigned according to a set o rules:

1. In an ionic compound between two elements the oxidation state o each element is equal to the charge carried by the ion, e.g.

 $\label{eq:constraint} \begin{array}{ll} Na^{_{+}}Cl^{_{-}} & Ca^{2+}Cl^{_{-2}} \\ (Na=+1;\,Cl=-1) & (Ca=+2;\,Cl=-1) \end{array}$

2. For covalent compounds assume the compound is ionic with the more electronegative element orming the negative ion, e.g.

 $\begin{array}{c} \text{CCl}_{_4} & \text{NH}_{_3} \\ \text{(C}=+4; \text{Cl}=-1) & \text{(N}=-3; \text{H}=+1) \end{array}$

3. The algebraic sum o all the oxidation states in a compound = zero, e.g.

$$CCl_{4} [(+4) + 4 \quad (-1) = 0];$$

H₂SO₄ [2 (+1) + (+6) + 4 (-2) = 0]

4. The algebraic sum o all the oxidation states in an ion = the charge on the ion, e.g.

$$\begin{split} & \text{SO}_4^{\ 2^-} \ [(+6) + 4 \quad (-2) = -2]; \\ & \text{MnO}_4^{\ -} \ [(+7) + 4 \quad (-2) = -1]; \\ & \text{NH}_4^{\ +} \ [(-3) + 4 \quad (+1) = +1] \end{split}$$

- Elements not combined with other elements have an oxidation state o zero, e.g. O₂; P₄; S₈.
- Oxygen when combined always has an oxidation state o −2 except in peroxides (e.g. H₂O₂) when it is −1.
- 7. Hydrogen when combined always has an oxidation state o +1 except in certain metal hydrides (e.g. NaH) when it is -1.

Many elements can show di erent oxidation states in di erent compounds, e.g. nitrogen in:

NH ₃	$N_{_2}H_{_4}$	N_2	$N_{2}O$	NO	NO_{2}	NO_{3}^{-}
(-3)	(-2)	(0)	(+1)	(+2)	(+4)	(+5)

When elements show more than one oxidation state the oxidation number is represented by using Roman numerals when naming the compound,

e.g. FeCl₂ iron(II) chloride; FeCl₃ iron(III) chloride K₂Cr₂O₇ potassium dichromate(VI); KMnO₄ potassium manganate(VII) Cu₃O copper(I) oxide; CuO copper(II) oxide.

OXIDIZING AND REDUCING AGENTS

A substance that readily oxidizes other substances is known as an oxidizing agent. Oxidizing agents are thus substances that readily accept electrons. Usually they contain elements that are in their highest oxidation state, e.g. O_2 , Cl_2 , F_2 , SO_3 (SO_4^{2-} in solution), MnO_4^{-} , and $Cr_2O_7^{2-}$. Reducing agents readily donate electrons and include H₂, Na, C, CO, and SO₂ (SO₂²⁻ in solution), $Cr_{2}O_{2}^{2}(aq)$ + $3SO_{3}^{2-}(aq) + 8H^{+}(aq) = 2Cr^{3+}(aq) + 3SO_{4}^{2-}(aq) + 4H_{2}O(l)$ e.g. (+4)(+6)(+6)(+3) (orange) (green) (oxidizing agent) (reducing agent)

Rd

[2]

 $Mg^{2+}(aq) + 2e^{-}$ e.g. Mg(s) (0)(+2)When an element is reduced its oxidation state decreases, e.g. $SO_4^{2-}(aq) + 2H^+(aq) + 2e^ SO_{2^{2}}(aq) + H_{2}O(l)$ (+6)(+4)The change in the oxidation state will be equal to the number o electrons involved in the hal -equation. Using oxidation states makes it easy to identi y whether or not a reaction is a redox reaction. Redox reactions (change in oxidation states) $CuO(s) + H_2(g) = Cu(s) + H_2O(l)$

OXIDATION AND REDUCTION IN TERMS OF OXIDATION STATES

When an element is oxidized its oxidation state *increases*.

(+2)(0) (0) (+1) $5Fe^{2+}(aq) + MnO_4^{-}(aq) + 8H^{+}(aq) = 5Fe^{3+}(aq) + Mn^{2+}(aq) + 4H_2O(l)$ (+2)(+7)(+3) (+2)

Not redox reactions (no change in oxidation states)

precipitation $Ag^+(aq) + Cl^-(aq) = AgCl(s)$ (+1)(-1)(+1) (-1)neutralization HCl(aq) + NaOH(aq) $NaCl(aq) + H_2O(l)$ (+1)(-1)(+1)(-2)(+1)(+1)(-1)(+1)(-2)

Note: reactions where an element is uncombined on one side o the equation and combined on the other side *must* be redox reactions since there must be a change in oxidation state,

e.g. $Mg(s) + 2HCl(aq) = MgCl_2(aq) + H_2(g)$

BALANCING REDOX EQUATIONS

To obtain the overall redox equation the number o electrons in the oxidation hal -equation must balance the number o electrons in the reduction hal -equation. For many redox equations this is straight orward. For example, the oxidation o magnesium by silver(I) ions. Write the two hal -equations, then double the silver hal -equation so that both hal -equations involve two electrons and then simply add them together:

 $2Ag^{+}(aq) + 2e^{-}$ 2Ag(s)

 $Mg(s) = Mg^{2+}(aq) + 2e^{-2}$

Overall $2Ag^{+}(aq) + Mg(s) = 2Ag(s) + Mg^{2+}(aq)$

It is less straight orward when there is a change in the number o oxygen atoms in a compound or ion. The rule is that if oxygen atoms need to be accounted for then water is used and if hydrogen atoms need to be accounted for then hydrogen ions, $H^+(aq)$, are used. Consider the oxidation o sulfite ions, $SO_3^{2-}(aq)$ to sul ate ions, $SO_4^{2-}(aq)$ by purple permanganate(VII) ions, $MnO_a^{-}(aq)$, which are reduced to very pale pink (virtually colourless) $Mn^{2+}(aq)$ ions in the process.

 $SO_{2^{-}}(aq)$ $SO_4^{2-}(aq)$. Add the extra O atom by adding water and ensure the charges on both sides are equal so the balanced hal equation becomes: $SO_{3^{2-}}(aq) + H_{2}O(l) = SO_{4^{-2-}}(aq) + 2H^{+}(aq) + 2e^{-2}$

 $MnO_{-}(aq) = Mn^{2+}(aq)$. Add 8 H⁺ ions to remove the oxygen atoms as water and ensure the charges on both sides are equal so the balanced hal -equation becomes: $MnO_{-}(aq) + 8H^{+}(aq) + 5e^{-} Mn^{2+}(aq) + 4H_{2}O(l)$.

Note that the number o electrons can also be obtained rom the change in oxidation states. S goes rom +4 to +6 so two electrons are involved and Mn goes rom +7 to +2 so five electrons are involved.

To ensure both hal -equations have the same number o electrons multiply the sulfite equation by 5 and the permanganate(VII) equation by 2 then obtain the overall redox equation by adding them together to cancel out the ten electrons and simpli y the water and hydrogen ions.

 $2MnO_4^{-}(aq) + 16H^+(aq) + 10e^- + 5SO_3^{-2-}(aq) + 5H_3O(l) \\ 2Mn^{2+}(aq) + 8H_3O(l) + 5SO_4^{-2-}(aq) + 10H^+(aq) + 10e^- + 5SO_3^{-2-}(aq) + 5H_3O(l) \\ - 2MnO_4^{-}(aq) + 8H_3O(l) + 5SO_4^{-2-}(aq) + 10H^+(aq) + 10H^+(ad) + 10H^+(ad) + 10H^+(ad) + 1$ Which simplifies to:

 $2MnO_{4^{-}}(aq) + 6H^{+}(aq) + 5SO_{3^{-}}(aq) = 2Mn^{2+}(aq) + 3H_{2}O(l) + 5SO_{4^{-}}(aq)$ This means that acid (H⁺ ions) needs to be present or the reaction to take place.

A vy

ACTIVITY SERIES

Lithium, sodium and potassium all react with cold water to give similar products but the reactivity increases down the group.

h d

 $2M(s) + 2H_2O(l)$ $2M^+(aq) + 2OH^-(aq) + H_2(g)$ (M = Li, Na or K)

Slightly less reactive metals react with steam and will give hydrogen with dilute acids, e.g.

dW k

 $Mg(s) + 2H_2O(g) = Mg(OH)_2 + H_2(g)$

Mg(s) + 2HCl(aq) $Mg^{2+}(aq) + 2Cl^{-}(aq) + H_{2}(g)$

In all o these reactions the metal is losing electrons that is, it is being oxidized and in the process it is acting as a reducing agent. An activity series o reducing agents can be deduced by considering the reactivity o metals with water and acids, and the reactions o metals with the ions o other metals.

The series used by the IB can be ound in Section 25 o the IB Chemistry data booklet. A simplified series ollows the order:

 $\mathrm{Li} > \mathrm{K} > \mathrm{Ca} > \mathrm{Na} > \mathrm{Mg} > \mathrm{Al} > \mathrm{Zn} > \mathrm{Fe} > \mathrm{Sn} > \mathrm{Pb} > \mathrm{H} > \mathrm{Cu} > \mathrm{Ag} > \mathrm{Au}$

Generally the more readily the metal ion loses its outer electron(s) the more reactive it is although note that lithium is actually higher than potassium in the activity series as the redox reactions also involve the hydration o the ions ormed. Metals higher in the series can displace metal ions lower in the series rom solution, e.g. zinc can react with copper ions to orm zinc ions and precipitate copper metal.

 $Zn(s) + Cu^{\scriptscriptstyle 2+}(aq) \qquad Zn^{\scriptscriptstyle 2+}(aq) + Cu(s)$

 $Zn(s) = Zn^{2+}(aq) + 2e^{-}$ Zn loses electrons in pre erence to Cu

 $Cu^{2+}(aq) + 2e^{-}$ Cu(s) Cu^{2+} gains electrons in pre erence to Zn^{2+} This also explains why only metals above hydrogen can react with acids (displace hydrogen ions) to produce hydrogen gas, e.g.

 $Zn(s) + 2H^{+}(aq) = Zn^{2+}(aq) + H_{2}(g)$

WINKLER METHOD

One application o a redox process is the Winkler method which is used to measure Biological Oxygen Demand (BOD). BOD is a measure o the dissolved oxygen (in ppm) required to decompose the organic matter in water biologically over a set time period, which is usually five days. Polluted water with a high BOD without the means o replenishing oxygen will not sustain aquatic li e.

The sample o the water is saturated with oxygen so the initial concentration o dissolved oxygen is known. A measured volume o the sample is then incubated at a fixed temperature or five days while microorganisms in the water oxidize the organic material. An excess o a manganese(II) salt is then added to the sample.

Under alkaline conditions manganese(II) ions are oxidized to manganese(IV) oxide by the remaining oxygen.

 $2Mn^{2+}(aq) + 4OH^{-}(aq) + O_{2}(aq) = 2MnO_{2}(s) + 2H_{2}O(l)$ Potassium iodide is then added which is oxidized by the manganese(IV) oxide in acidic solution to orm iodine.

 $MnO_{2}(s) + 2I^{-}(aq) + 4H^{+}(aq) \qquad Mn^{2+}(aq) + I_{2}(aq) + 2H_{2}O(l)$

The iodine released is then titrated with standard sodium thiosul ate solution.

 $I_2(aq) + 2S_2O_3^{2-}(aq) = S_4O_6^{2-}(aq) + 2I^{-}(aq)$

By knowing the amount (in mol) o iodine produced the amount o oxygen present in the sample o water can be calculated and hence its concentration.

Wkd xp

100 cm³ o a sample o water was treated with an excess o alkaline manganese(II) sulphate, $MnSO_4(aq)$. A ter all the dissolved oxygen had reacted the solution was then acidified

The series can be extended or oxidizing agents. The most reactive oxidizing agent will be the species that gains electrons the most readily. For example, in group 17

 $I^{-}(aq) = e^{-} + I_{2}(aq)$ $Br^{-}(aq) = e^{-} + Br_{2}(aq)$ $Cl^{-}(aq) = e^{-} + Cl_{2}(aq)$ $F^{-}(aq) = e^{-} + F_{2}(aq)$ VOxidizing agents lower in the series gain electrons rom species higher in the series, e.g. $Cl_{2}(aq) + 2Br^{-}(aq) = 2Cl^{-}(aq) + Br_{2}(aq)$

and excess potassium iodide. KI(aq) added. The iodine released was titrated with sodium thiosul ate solution, Na₂S₂O₃(aq), using starch as an indicator. It was ound that 6.00 cm³ o $1.00 - 10^{-2}$ mol dm⁻³ Na₂S₂O₃(aq) was required to react with all the iodine. Determine the concentration o dissolved oxygen in parts per million (ppm).

Step 1. Calculate the amount o sodium thiosul ate used in the titration.

Amount o $S_2O_3^{2-}(aq) = \frac{6.00}{1000}$ 1.00 $10^{-2} = 6.00$ 10^{-5} mol **Step 2.** Calculate the amount o iodine that reacted with the sodium thiosul ate.

$$\begin{split} &I_{_2}(aq) + 2S_{_2}O_{_3}{}^{_2-}(aq) \qquad S_{_4}O_{_6}{}^{_2-}(aq) + 2I^{_-}(aq) \\ &\text{Amount o } I_{_3}(aq) = \qquad 6.00 \quad 10^{-5} = 3 \quad 10^{-5} \text{ mol} \end{split}$$

Step 3. Calculate the amount o MnO_2 produced by the oxidation o the Mn^{2+} ions.

$$\begin{split} MnO_{_2}(s) &+ 2I^{_-}(aq) + 4H^{_+}(aq) \qquad Mn^{_{2+}} + I_{_2}(aq) + 2H_{_2}O(l) \\ Amount \ o \ MnO_{_2} &= 3 \qquad 10^{_{-5}} \ mol \end{split}$$

Step 4. Calculate the mass o oxygen dissolved in the 100 cm³ sample o water.

$$\begin{split} & 2Mn^{2_+}(aq) + 4OH^-(aq) + O_2(aq) \qquad 2MnO_2(s) + 2H_2O(l) \\ & Amount \ o \ O_2 = 3 \ 10^{-5} = 1.5 \ 10^{-5} \ mol \\ & Mass \ o \ O_2 = 32.00 \ 1.5 \ 10^{-5} = 4.8 \ 10^{-4} \ g \end{split}$$

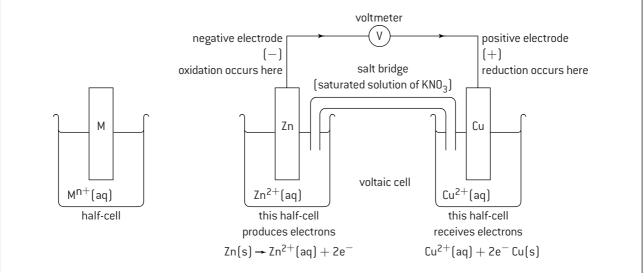
Step 5. Calculate the concentration o dissolved oxygen in parts per million (ppm). (This is the same as the mass in mg dissolved in 1 dm³ o water.)

 100 cm^3 contains 4.8 $10^{-4} \text{ g o } O_2$ so concentration = 4.8 $10^{-3} \text{ g dm}^{-3} = 4.8 \text{ ppm}.$

SIMPLE VOLTAIC CELLS

A hal -cell is simply a metal in contact with an aqueous solution o its own ions. A voltaic cell consists o two di erent hal -cells, connected together to enable the electrons trans erred during the redox reaction to produce energy in the orm o electricity. The cells are connected by an external wire and by a salt bridge, which allows the ree movement o ions.

A good example o a voltaic cell is a zinc hal -cell connected to a copper hal -cell. Because zinc is higher in the activity series the electrons will flow rom the zinc hal -cell towards the copper hal -cell. To complete the circuit and to keep the hal -cells electrically neutral, ions will flow through the salt bridge. The voltage produced by a voltaic cell depends on the relative di erence between the two metals in the activity series. Thus the voltage rom a $Mg(s)/Mg^{2+}(aq)$ hal -cell connected to a $Cu(s)/Cu^{2+}(aq)$ hal -cell will be greater than that obtained rom a $Zn(s)/Zn^{2+}(aq)$ hal -cell connected to a $Fe(s)/Fe^{2+}(aq)$ hal -cell.



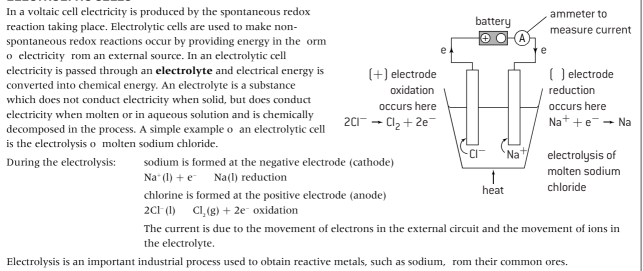
CONVENTION FOR WRITING CELLS

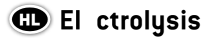
By convention in a cell diagram the hal -cell undergoing oxidation is placed on the let o the diagram and the hal -cell undergoing reduction on the right o the diagram. The two aqueous solutions are then placed either side o the salt bridge e.g.

 $Zn(s)/Zn^{2+}(aq) = Cu^{2+}(aq)/Cu(s)$

The words cathode and anode can also be used to describe the electrodes. The anode is where oxidation occurs so or a voltaic cell it is the negative electrode but or an electrolytic cell (see below), where the electricity is provided rom an external source, the anode is the positive electrode, which can cause some con usion.

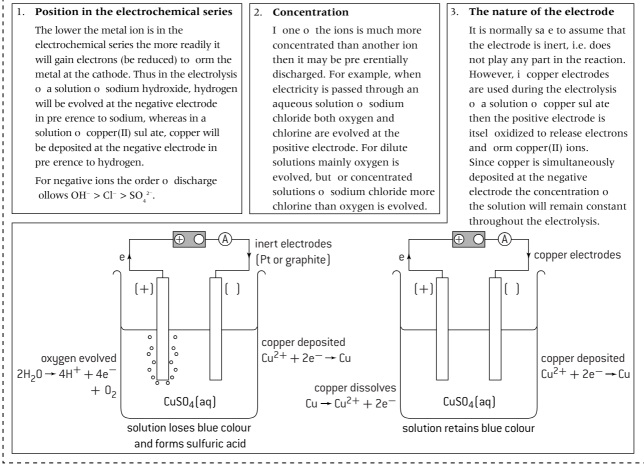
ELECTROLYTIC CELLS





FACTORS AFFECTING THE DISCHARGE OF IONS DURING ELECTROLYSIS

During the electrolysis o molten salts there are only usually two ions present, so the cation will be discharged at the negative electrode (cathode) and the anion at the positive electrode (anode). However, or aqueous electrolytes there will also be hydrogen ions and hydroxide ions rom the water present. There are three main actors that influence which ions will be discharged at their respective electrodes.



FACTORS AFFECTING THE QUANTITY OF PRODUCTS DISCHARGED DURING ELECTROLYSIS

The amount o substance deposited will depend on:

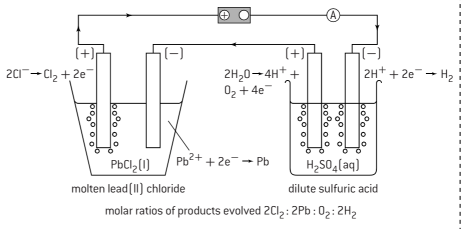
1. The number o electrons flowing through the system, i.e. the amount o charge passed. This in turn depends on the current and the time or which it flows. I the current is doubled then twice as many electrons pass through the system and twice as much product will be ormed. Similarly i the time is doubled twice as many electrons will pass through the system and twice as much product will be ormed.

charge = current time (1 coulomb = 1 ampere 1 second)

 The charge on the ion. To orm one mole o sodium in the electrolysis o molten sodium chloride requires one mole o electrons to flow through the cell. However, the ormation o one mole o lead during the electrolysis o molten lead(II) bromide requires two moles o electrons.

- $Na^+(l) + e^- Na(l)$
- $Pb^{2+}(l) + 2e^{-}$ Pb(l)

I cells are connected in series then the same amount o electricity will pass through both cells and the relative amounts o products obtained can be determined.

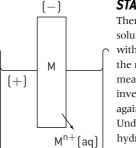


🚥 El ctroplating and standard 🛛 ctrod pot ntials

ELECTROPLATING

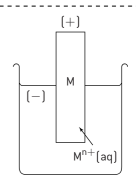
Electrolysis can also be used in industry to coat one metal with a thin layer o another metal. This process is known as **electroplating**.

For example, in copper plating the negative electrode (cathode) is made rom the metal to be copper plated. It is placed into a solution o copper(II) sul ate together with a positive electrode (anode) made rom a piece o copper. As electricity is passed through the solution the copper anode dissolves in the solution to orm $Cu^{2+}(aq)$ ions and the $Cu^{2+}(aq)$ ions in solution are deposited onto the cathode. By making the anode o impure copper and the cathode rom a small piece o pure copper this process can also be used to puri y impure copper. This is an important industrial process as one o the main uses o copper is or electrical wiring, where purity is important since impure copper has a much higher electrical resistance.



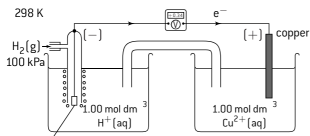
STANDARD ELECTRODE POTENTIALS

There are two opposing tendencies in a hal -cell. The metal may dissolve in the solution o its own ions to leave the metal with a negative potential compared with the solution, or the metal ions may deposit on the metal, which will give the metal a positive potential compared with the solution. It is impossible to measure this potential, as any attempt to do so inter eres with the system being investigated. However, the electrode potential o one hal -cell can be compared against another hal -cell. The hydrogen hal -cell is normally used as the standard. Under standard conditions o 100 kPa pressure, 298 K, and 1.0 mol dm⁻³ hydrogen ion concentration the standard electrode potential o the hydrogen electrode is assigned a value o zero volts.

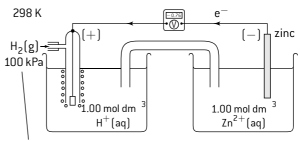


When the hal -cell contains a metal above hydrogen in the reactivity series

electrons flow rom the hal -cell to the hydrogen electrode, and the electrode potential is given a negative value. I the hal cell contains a metal below hydrogen in the reactivity series electrons flow rom the hydrogen electrode to the hal -cell, and the electrode potential has a positive value. The standard electrode potentials are arranged in increasing order to orm the electrochemical series.



The platinum electrode is coated with nely divided platinum, which serves as a catalyst for the electrode reaction.

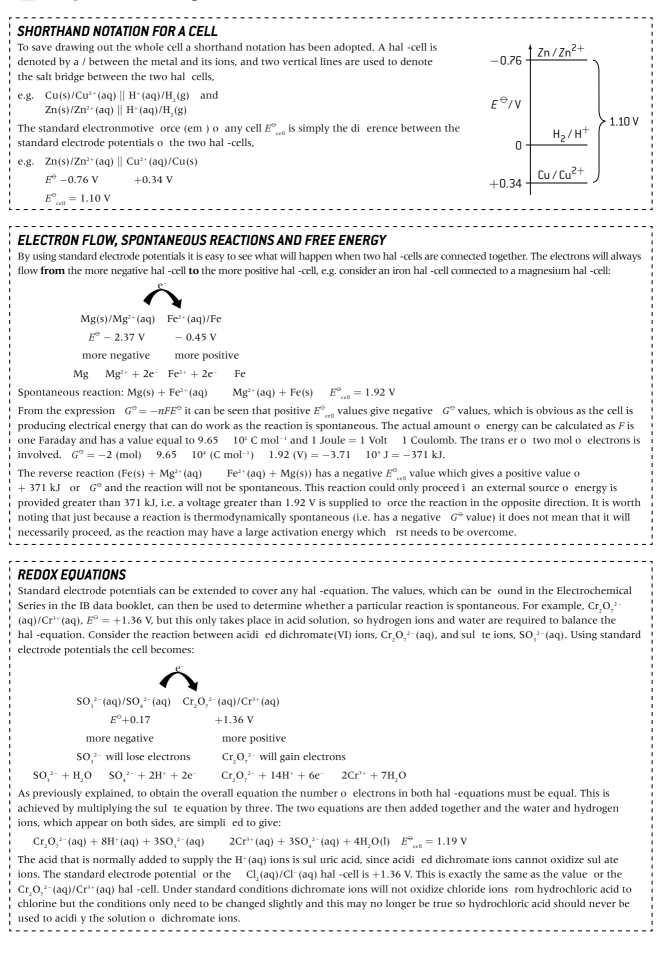


pressure just above 100 kPa so H_2 can escape from electrode

ELECTROCHEMICAL SERIES

ELECTROCHEMICAL SERIES					
(A more complete series can be	ound in the IB data booklet)				
Couple	E^{\ominus} / V				
$K(s)/K^+(aq)$	-2.93				
$Ca(s)/Ca^{2+}(aq)$	-2.87				
Na(s)/Na ⁺ (aq)	-2.71				
$Mg(s)/Mg^{2+}(aq)$	-2.37				
$Al(s)/Al^{3+}(aq)$	-1.66				
$Zn(s)/Zn^{2+}(aq)$	-0.76				
$Fe(s)/Fe^{2+}(aq)$	-0.45				
$H_{_2}(g)/H^+(aq)$	0.00				
$Cu(s)/Cu^{2+}(aq)$	+0.34				
$I^{-}(aq)/I_{2}(aq)$	+0.54				
$Ag(s)/Ag^{+}(aq)$	+0.80				
$Br^{-}(aq)/Br_{2}(aq)$	+1.09				
$Cl^{-}(aq)/Cl_{2}(aq)$	+1.36				
$F^{-}(aq)/F_{2}(aq)$	+2.87				

Spontaneity of electrochemical reaction



MULTIPLE CHOICE QUESTIONS REDOX PROCESSES

1. $5Fe^{2_{+}}(aq) + MnO_{4_{-}}(aq) + 8H^{+}(aq) = 5Fe^{3_{+}}(aq) + Mn^{2_{+}}(aq) + 4H_{2}O(l)$

In the equation above:

- **A.** $Fe^{2+}(aq)$ is the oxidizing agent
- **B.** H^+ (aq) ions are reduced
- **C.** $Fe^{2+}(aq)$ ions are oxidized
- **D.** $MnO_4^{-}(aq)$ is the reducing agent
- **2.** The oxidation states o nitrogen in NH₃, HNO₃ and NO₂ are, respectively
 - A. -3, -5, +4C. -3, +5, -4B. +3, +5, +4D. -3, +5, +4
- **3.** Which one o the ollowing reactions is **not** a redox reaction?
 - **A.** $Ag^+(aq) + Cl^-(aq) = AgCl(s)$
 - **B.** $2Na(s) + Cl_2(g) = 2NaCl(s)$
 - **C.** $Mg(s) + 2HCl(aq) MgCl_2(aq) + H_2(g)$
 - **D.** $Cu^{2+}(aq) + Zn(s) = Cu(s) + Zn^{2+}(aq)$
- 4. Which substance does not have the correct ormula?
 - **A.** iron(III) sul ate $Fe_2(SO_4)_3$
 - **B.** iron(II) oxide Fe₂O
 - **C.** copper(I) sul ate Cu_2SO_4
 - **D.** copper(II) nitrate $Cu(NO_3)_2$
- 5. For which conversion is an oxidizing agent required?
 - **A.** $2H^+(aq) = H_2(g)$ **C.** $SO_3(g) = SO_4^{-2-}(aq)$
 - **B.** $2Br^{-}(aq) = Br_{2}(aq)$ **D.** $MnO_{2}(s) = Mn^{2+}(aq)$
- **6.** Ethanol can be oxidized to ethanal by an acidic solution o dichromate(VI) ions.

 $\begin{array}{rl} C_{2}H_{5}OH(aq) + & H^{+}(aq) + & Cr_{2}O_{7}^{2-}(aq) \\ CH3CHO(aq) + & Cr^{3+}(aq) + & H_{2}O(l) \end{array}$

The sum o all the coe cients in the balanced equation is:

A. 24 **B.** 26 **C.** 28 **D.** 30

- When an Fe(s)/Fe²⁺(aq) hal -cell is connected to a Cu(s)/ Cu²⁺(aq) hal -cell by a salt bridge and a current allowed to flow between them
 - A. the electrons will flow rom the copper to the iron.
 - **B.** the salt bridge allows the flow o ions to complete the circuit.
 - **C.** the salt bridge allows the flow o electrons to complete the circuit.
 - **D.** the salt bridge can be made o copper or iron.
- **8.** During the electrolysis o molten sodium chloride using platinum electrodes
 - **A.** sodium is ormed at the negative electrode.
 - **B.** chlorine is ormed at the negative electrode.
 - **C.** sodium is ormed at the positive electrode.
 - **D.** oxygen is ormed at the positive electrode.
- 9. Which statement is true?
 - **A.** Lead chloride is ionic so solid lead chloride will conduct electricity.
 - **B.** When a molten ionic compound conducts electricity ree electrons pass through the liquid.
 - **C.** When liquid mercury conducts electricity mercury ions move towards the negative electrode.
 - **D.** During the electrolysis o a molten salt reduction will always occur at the negative electrode.
- **10.** The ollowing in ormation is given about reactions involving the metals X, Y and Z and solutions o their sul ates.
 - X(s) / YSO (aq) no reaction

$$Z(s) + YSO_4(aq) = Y(s) + ZSO_4(aq)$$

When the metals are listed in decreasing order o reactivity (most reactive rst), what is the correct order?

А.	Z > Y > X	C. $Y > X > Z$
B.	X > Y > Z	D. $Y > Z > X$

H

- **11.** Which are correct or a spontaneous reaction occurring in a voltaic cell?
 - I. E^{\ominus} or the cell has a positive value
 - II. G^{\ominus} or the reaction has a negative value
 - III. The reaction must occur under standard conditions

A. I and II only	C. II and III only
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- **B.** I and III only **D.** I, II and III
- 12. When the same quantity o electricity was passed through a dilute solution o sodium hydroxide and through a molten solution o lead bromide, 0.100 mol o lead was produced. What amount (in mol) o hydrogen gas was evolved?

А.	0.014	С.	0.100
B.	0.050	D.	0.200

Use the following information to answer questions 13 and 14.

$\operatorname{Sn}^{_{2+}}(\operatorname{aq}) + 2e^{}$	Sn(s)	$E^{\oplus} = -0.14 \mathrm{V}$
$\operatorname{Sn}^{\scriptscriptstyle 4+}(\operatorname{aq}) + 2e^-$	$\operatorname{Sn}^{\scriptscriptstyle 2+}(\operatorname{aq})$	$E^{\ominus} = +0.15 \mathrm{V}$
$\mathrm{Fe}^{_{2+}}(\mathrm{aq}) + 2\mathrm{e}^{_{-}}$	Fe(s)	$E^{\ominus} = -0.44 \mathrm{V}$
$Fe^{_{3+}}(aq) + e^{_{-}}$	$Fe^{2+}(aq)$	$E^{\odot} = +0.77 \mathrm{V}$

- 13. Under standard conditions which statement is correct?
 - **A.** $Sn^{2+}(aq)$ can reduce $Fe^{3+}(aq)$.
 - **B.** Fe(s) can oxidize $Sn^{2+}(aq)$.
 - **C.** Sn(s) can reduce Fe(s).
 - **D.** $Fe^{3+}(aq)$ can reduce $Sn^{4+}(aq)$.
- **14.** When a hal -cell o $Fe^{2+}(aq)/Fe^{3+}(aq)$ is connected by a salt bridge to a hal -cell o $Sn^{2+}(aq)/Sn^{4+}(aq)$ under standard conditions and a current allowed to flow in an external circuit the total em o the spontaneous reaction will be:

A. +0.92V **B.** -0.92V **C.** +0.62V **D.** -0.62V

SHORT ANSWER QUESTIONS REDOX PROCESSES

1. The data below is rom an experiment used to determine the percentage o iron present in a sample o iron ore. This sample was dissolved in acid and all o the iron was converted to Fe^{2+} . The resulting solution was titrated with a standard solution o potassium manganate(VII), KMnO₄. This procedure was carried out three times. In acidic solution, MnO_4^{-} reacts with Fe^{2+} ions to orm Mn^{2+} and Fe^{3+} and the end point is indicated by a slight pink colour.

Titre	1	2	3
Initial burette reading / cm ³	1.00	23.60	10.00
Final burette reading / cm ³	24.60	46.10	32.50

Mass of iron ore / g	3.682	10^{-1}
Concentration of KMnO ₄ solution / mol dm ⁻³	2.152	10^{-2}

a) Deduce the balanced redox equation or this reaction in **acidic** solution.

[2]

[1]

[2]

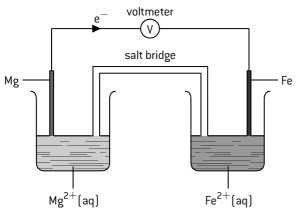
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- b) Identi y the reducing agent in the reaction.
- c) Calculate the amount, in moles, o MnO₄⁻ used in the titration.
- d) Calculate the amount, in moles, o Fe present in the $3.682 \quad 10^{-1}$ g sample o iron ore.
- e) Determine the percentage by mass o Fe present in the 3.682 10^{-1} g sample o iron ore.
- 2. Chemical energy can be converted to electrical energy in the voltaic cell below.



a) Explain how the diagram con rms that magnesium is above iron in the activity series.

Α

5. Consider the ollowing two hal -equations:

$$\begin{array}{ll} g^{+}(aq) + e^{-} & Ag(s) & E^{\ominus} = +0.80 \text{ V} \\ Cl_{2}(g) + e^{-} & Cl^{-}(aq) & E^{\ominus} = +1.36 \text{ V} \end{array}$$

- a) Deduce the overall equation or the spontaneous reaction that will occur when a silver hal -cell is connected to a chlorine hal -cell.
- b) Identi y the direction o electron flow in the external circuit when the cell is operating. [1]
- c) Determine the cell potential, E_{cell}^{\ominus} . [1]
- d) Calculate the ree energy produced by the cell when it is operating under standard conditions. [2]

- b) Identi y the positive electrode (anode) o the cell. [1]
- c) (i) State the hal -equation or the reaction occurring at the iron electrode. [1]
 - (ii) State the overall equation or the reaction when the cell is producing electricity. [2]
- d) Deduce whether the voltage produced by the cell would be greater or less i the iron hal -cell is replaced by a copper hal -cell. [2]
- 3. a) Molten sodium chloride can be electrolysed using graphite electrodes.
 - (i) Draw the essential components o this electrolytic cell and identi y the products that orm at each electrode.[2]
 - (ii) State the hal -equations or the oxidation and reduction processes and deduce the overall cell reaction, including state symbols.
 - b) Explain why solid sodium chloride does not conduct electricity. [1]

[2]

[2]

- c) Using another electrolysis reaction, aluminium can be extracted rom its ore, bauxite, which contains Al₂O₃. State **one** reason why aluminium is o ten used instead o iron in many engineering applications. [1]
- 4. Iodine reacts with thiosul ate ions to orm the tetrathionate ion according to the equation

 $I_2(aq) + 2S_2O_3^{2-}(aq) = S_4O_6^{2-}(aq) + 2I^-(aq)$

- a) Show that this reaction is a redox reaction.
- b) The thiosul ate ion has a structure similar to the sul ate ion, SO₄²⁻, except that one o the outer oxygen atoms has been replaced by a sul ur atom.



thiosulfate ion

- (i) Comment on the di erence in the oxidation state o sul ur in the thiosul ate ion compared with the sul ate ion. [2]
 (ii) Explore the concept o oxidation state using
- the tetrathionate ion, $S_4O_6^{2-}$, as an example. [3]
- c) Discuss whether the complete combustion o carbon in oxygen to orm carbon dioxide can be described as the oxidation o carbon according to all the di erent de nitions o oxidation. [4]
- a) Identi y the products that will be obtained at the positive (anode) and negative (cathode) electrodes when (i) a dilute solution and (ii) a concentrated solution o sodium chloride undergoes electrolysis. [3]
 - b) Explain why the electrolysis o a dilute solution o sul uric acid or a dilute solution o sodium hydroxide are both sometimes described as the electrolysis o water. [3]
 - c) Describe how you could use electrolysis to coat a spoon made o steel with a thin layer o silver. [2]

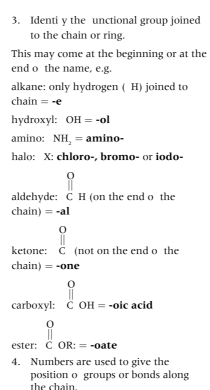
10 ORGANIC CHEMISTRY

Fundamentals of organic chemistry

NAMING ORGANIC COMPOUNDS Organic chemistry is concerned with the compounds o carbon. Since there are more compounds o carbon known than all the other elements put together, it is help ul to have a systematic way o naming them. 1. Identi y the longest carbon chain. 1 carbon = meth-2 carbons = eth-3 carbons = **prop-**4 carbons = **but-**5 carbons = **pent-** $6 \text{ carbons} = \mathbf{hex}$ -7 carbons = **hept-**8 carbons = **oct-**2. Identi y the type o bonding in the chain or ring All single bonds in the carbon chain = -an-One double bond in the carbon chain = -en-One triple bond in the carbon chain = -yn-

CLASS OF ORGANIC COMPOUND

Di erent compounds that all contain the same unctional group are divided into classes. Sometimes the name o the class is the same as the unctional group, but sometimes it is di erent. For example, the name o the class is the same or all esters that contain the ester functional group, (COOR), but it is di erent or all alcohols that contain the hydroxyl, (**OH**), **functional group**. The carbonyl unctional group is CO where the carbon atom is joined by a double bond to the oxygen atom. Several classes o compounds contain this unctional group as it depends upon what else is bonded to the carbon atom. For example, i H is bonded to the carbon atom then they are known as aldehydes (COH), whereas i an alkyl group is bonded to the carbon atom then they are known as ketones (COR). The COOH unctional group is known as the carboxyl group and the class o compounds containing this group is called carboxylic acids. Compounds containing an amino group are known as amines and compounds containing a halo group are known as halogenoalkanes (or more specifically chloroalkanes, bromoalkanes or iodoalkanes).



HOMOLOGOUS SERIES

The alkanes orm a series o compounds all with the general ormula $C_nH_{2n+2'}$ e.g. methane CH_4 ethane C_2H_6 propane C_3H_8 butane C_4H_{10} I one o the hydrogen atoms is

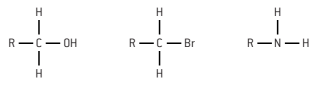
To ne of the hydrogen atoms is removed what is let is known as an alkyl radical R (e.g methyl CH_3 ; ethyl C_2H_5). When other atoms or groups are attached to an alkyl radical they can orm a di erent series o compounds. These atoms or groups attached are known as unctional groups and the series ormed are all homologous series.

Homologous series have the same general ormula with the neighbouring members o the series di ering by CH₂; or example the general ormula o alcohols is $C_n H_{2n+1}$ OH. The chemical properties o the individual members o a homologous series are similar and they show a gradual change in physical properties.

CLASSIFICATION OF ALCOHOLS, HALOGENOALKANES AND AMINES

Alcohols and halogenoalkanes may be classified according to how many R groups are bonded to the carbon atom containing the unctional group. Similar logic applies to amines but now it is the number o R groups attached to the nitrogen atom o the amino unctional group.

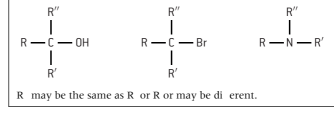
primary (one R group bonded to the C or N atom)



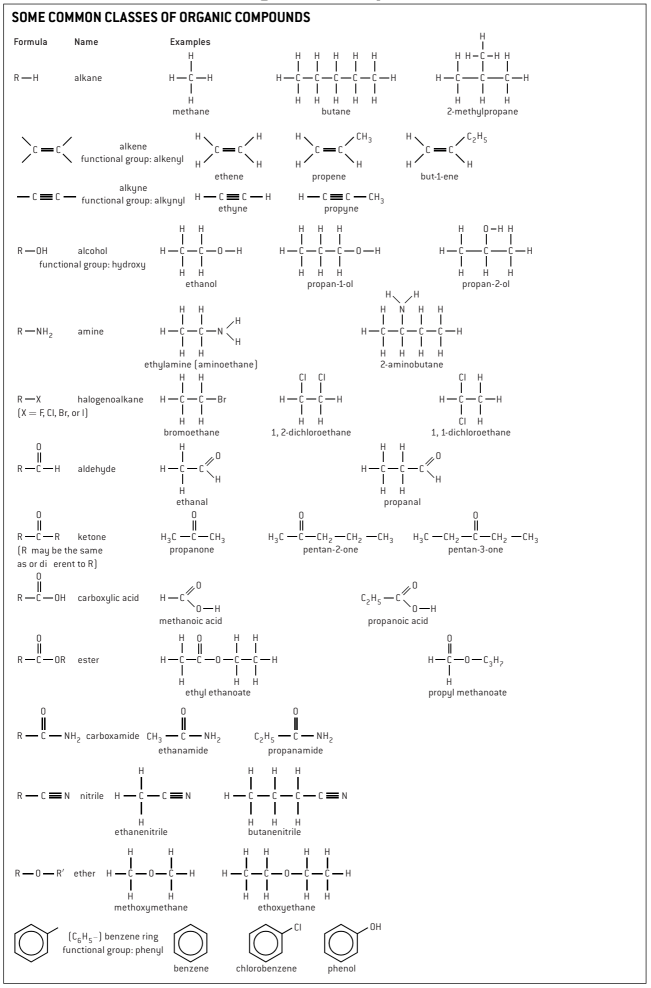
secondary (two R groups bonded to the C or N atom) R may be the same as R or di erent

$$\begin{array}{cccc} H & H & H \\ I & I \\ R - C - OH & R - C - Br & R - N - R' \\ I & I \\ R' & R' \end{array}$$

tertiary (three R groups bonded to the C or N atom)



Common classes of organic compounds

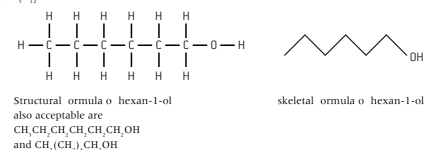


Structural formulas

STRUCTURAL FORMULAS

The di erence between the empirical, molecular and structural ormulas o a compound has already been covered in Topic 1 Stoichiometric relationships. Because the physical and chemical properties o a compound are determined by the unctional group and the arrangement o carbon and other atoms within the molecule, the structural ormulas or organic compounds are o ten used.

These may be shown in a variety o di erent ways but all ways should show unambiguously how the atoms are bonded together. When drawing ull structural ormulas with lines representing bonds and the symbol o the element representing atoms, all the hydrogen atoms must also be included in the diagram. This is because the skeletal ormula does not include any symbols or the elements and the end o a line represents a methyl group unless another atom is shown attached. Although you should understand skeletal ormulas they are not normally used or simple ormulas except where benzene is involved. Note that unless specifically asked or, Lewis structures, showing all the valence electrons, are not necessary. The bonding must be clearly indicated. Structures may be shown using lines as bonds or in their shortened orm, e.g. CH₂CH₂CH₂CH₂CH₂CH₂ or CH₂ (CH₂), CH₂ or pentane but the molecular ormula, C_sH₁₂ will not su fice.



STRUCTURAL FORMULA OF BENZENE

The simplest aromatic compound (arene) is benzene, C,H,. The Kekul structure o benzene (cyclohexa-1,3,5-triene) consists o three double bonds.

There is both physical and chemical evidence to support the act that benzene does not contain three separate double bonds but exists as a resonance hybrid structure with delocalized electrons. The two resonance hybrid orms are shown (right).

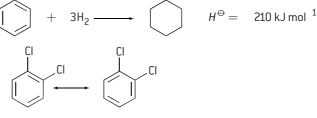
- 1. The C C bond lengths are all the same and have a value o 0.140 nm which lies between the values or C C (0.154 nm) and C=C (0.134 nm). $+ H_2 \xrightarrow{\text{Ni}}$
- 2. The enthalpy o hydrogenation o cyclohexene is -120 kJ mol⁻¹.

I benzene simply had the cyclohexa-1,3,5-triene structure with three double bonds the enthalpy change o hydrogenation o benzene would be expected to be equal to 3 times the enthalpy change o hydrogenation o cyclohexene, i.e. -360 kJ mol⁻¹.

However the experimentally determined value or benzene is 210 kJ mol⁻¹.

The di erence o 150 kJ mol⁻¹ is the extra energy associated with the delocalization.

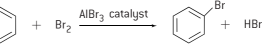
3. Only one isomer exists or 1,2- disubstituted benzene compounds. I there were simply alternate double bonds then two isomers would exist.



The two isomers of 1, 2- dichlorobenzene which would be expected if benzene has the cyclohexa-1, 3, 5-triene structure

4. I benzene had three normal double bonds it would be expected to readily undergo addition reactions. In act it will only undergo addition reactions with di ficulty and more

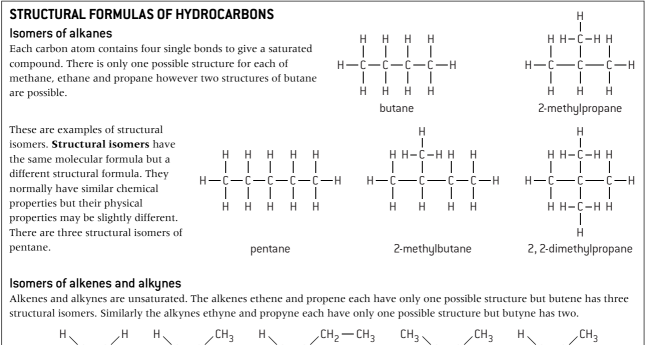
commonly undergoes substitution reactions. For example, with bromine it orms bromobenzene and hydrogen bromide rather than 1,2-dibromobenzene.

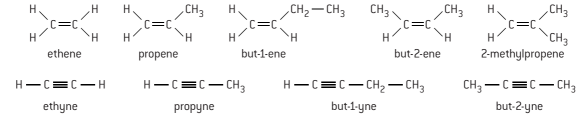


The actual bonding in benzene is best described by the delocalization o electrons. For this reason benzene is o ten represented by a hexagonal ring with a circle in the middle o it.

120 kJ mol ¹

Structural isomers

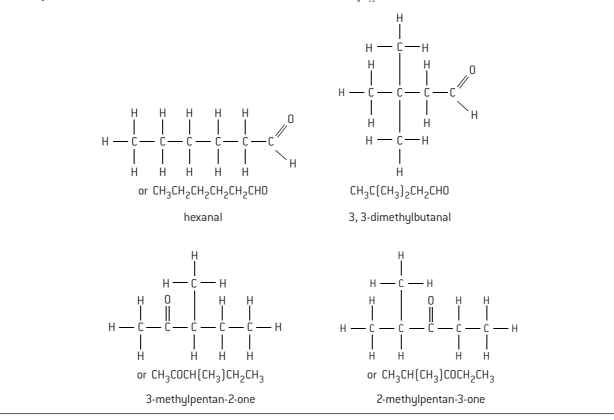




NAMING STRUCTURAL ISOMERS

The naming system explained on page 79 is known as the IUPAC (International Union of Pure and Applied Chemistry) system. The IUPAC names to distinguish between structural isomers of alkanes, alkenes, alkenes, alcohols, ethers, halogenoalkanes, aldehydes, ketones, esters and carboxylic acids, each containing up to six carbon atoms, are required.

For example, four different structural isomers with the molecular formula C₁H₁₀O are shown.

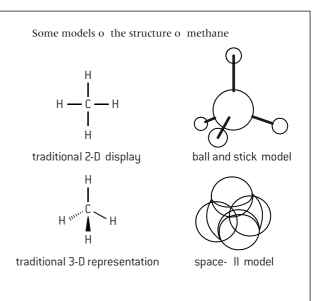


3-D models of structural formulas

3-D REPRESENTATIONS

To help distinguish between isomers and because the chemical and physical properties o molecules depend upon their shape and the types o bonds and unctional groups they contain it is help ul to be able to visualize the structures in three dimensions. Twodimensional display diagrams o alkanes, or examples, wrongly suggest that the H C H bond angles are 90 instead o 109.5 . The traditional way in which chemists have approached this is to use a ull wedge to show an atom coming out o the two dimensional page and a dotted line to show it going behind the page.

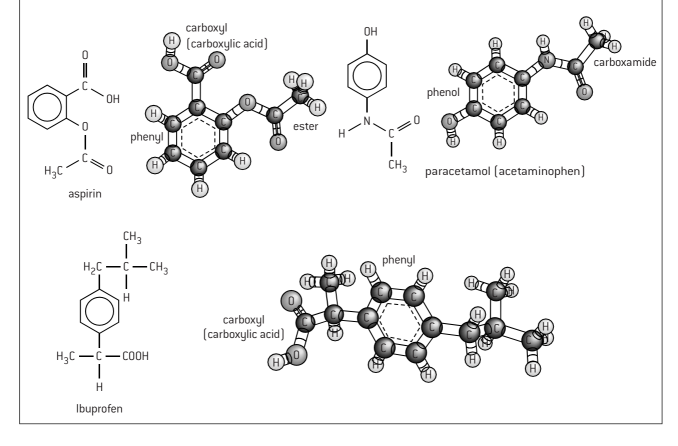
The very best way to see molecular structures in three dimensions is to use a molecular modelling kit and build the models or yoursel. This can be particularly help ul when distinguishing between isomers and understanding bond angles. Two common ways to represent molecules in 3-D are ball and stick models or space-fill models. You can also see virtual 3-D representations using apps on your laptop, smart phone or tablet, which are also able to show the molecules rotating. However it is easy to orget that these are still being displayed on a 2-D screen so are not truly three dimensional.



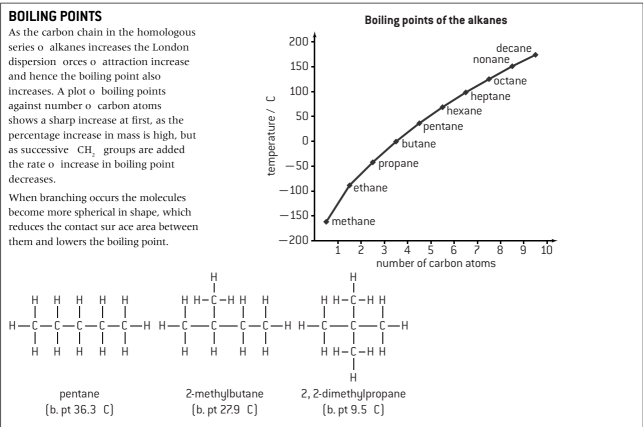
IDENTIFYING FUNCTIONAL GROUPS IN MOLECULES FROM VIRTUAL 3-D MODELS

As well as recognizing di erent unctional groups within more complex molecules rom their structural ormulas you should also be able to recognize them rom virtual 3-D representations.

The ollowing three molecules, aspirin, paracetamol (acetaminophen) and ibupro en, are all mild pain killers. As you can see they all contain a phenyl group, although this is a phenol in paracetamol as the phenyl group is directly bonded to an hydroxyl group. Aspirin and ibupro en contain a carboxyl group (carboxylic acid), aspirin also contains an ester group and paracetamol contains a carboxamide group. I you look at past papers or in the IB data booklet you can find other molecules such as morphine and heroin where you can do a similar exercise.



Properties o different homologous series



Other homologous series show similar trends but the actual temperatures at which the compounds boil will depend on the types o attractive orces between the molecules. The volatility o the compounds also ollows the same pattern. The lower members o the alkanes are all gases as the attractive orces are weak and the next ew members are volatile liquids. Methanol, the first member o the alcohols is a liquid at room temperature, due to the presence o hydrogen bonding. Methanol is classed as volatile as its boiling point is 64.5 C but when there are our or more carbon atoms in the chain the boiling points exceed 100 C and the higher alcohols have low volatility.

Compound	Formula	Mr	Class of compound	Strongest type of attraction	B. pt / C
butane	$C_{4}H_{10}$	58	alkane	London dispersion orces	-0.5
but-1-ene	C_4H_8	56	alkene	London dispersion orces	-6.5
but-1-yne	C_4H_6	54	alkyne	London dispersion orces	8.1
methyl methanoate	HCOOCH ₃	60	ester	dipole dipole	31.5
propanal	CH ₃ CH ₂ CHO	58	aldehyde	dipole dipole	48.8
propanone	CH ₃ COCH ₃	58	ketone	dipole dipole	56.2
aminopropane	$\mathrm{CH_{_3}CH_{_2}CH_{_2}NH_{_2}}$	59	amine	hydrogen bonding	48.6
propan-1-ol	$\rm CH_{_3}\rm CH_{_2}\rm CH_{_2}\rm OH$	60	alcohol	hydrogen bonding	97.2
ethanoic acid	CH ₃ COOH	60	carboxylic acid	hydrogen bonding	118

SOLUBILITY IN WATER

Whether or not an organic compound will be soluble in water depends on the polarity o the unctional group and on the chain length. The lower members o alcohols, amines, aldehydes, ketones and carboxylic acids are all water soluble. However, as the length o the non-polar hydrocarbon chain increases the solubility in water decreases. For example, ethanol and water mix in all proportions, but hexan-1-ol is only slightly soluble in water. Compounds with non-polar unctional groups, such as alkanes and alkenes, do not dissolve in water but are soluble in other non-polar solvents. Propan-1-ol is a good solvent because it contains both polar and non-polar groups and can to some extent dissolve both polar and non-polar substances.

Alkanes

LOW REACTIVITY OF ALKANES

Because o the relatively strong C C and C H bonds and because they have low polarity, alkanes tend to be quite unreactive. They only readily undergo combustion reactions with oxygen and substitution reactions with halogens in ultraviolet light.

COMBUSTION

Alkanes are hydrocarbons compounds that contain carbon and hydrogen only. All hydrocarbons burn in a plenti ul supply o oxygen to give carbon dioxide and water. The general equation or the combustion o any hydrocarbon is:

$$C_{x}H_{y} + (x + \frac{y}{4})O_{2} = xCO_{2} + \frac{y}{2}H_{2}O$$

Although the C C and C H bonds are strong the C=O and O H bonds in the products are even stronger so the reaction is very exothermic and much use is made o the alkanes as uels.

e.g. natural gas (methane)

 $CH_{_4}(g) + 2O_{_2}(g)$ $CO_{_2}(g) + 2H_{_2}O(l)$ $H^{\ominus} = -890.4 \text{ kJ mol}^{_{-1}}$

gasoline (petrol)

 $C_{_8}H_{_{18}}(l)\,+\,12\quad O_{_2}(g)\,-\,8CO_{_2}(g)\,+\,9H_{_2}O(l)$

 $\mathrm{H}^{\ominus} = -5512 \; kJ \; mol^{_{-1}}$

I there is an insu ficient supply o oxygen then incomplete combustion occurs and carbon monoxide and carbon are also produced as products.

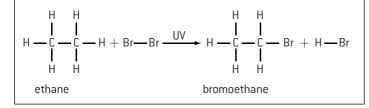
SUBSTITUTION REACTIONS

Alkanes can react with chlorine (or other halogens) in the presence o ultraviolet light to orm hydrogen chloride and a substituted alkane, e.g. methane can react with chlorine to orm chloromethane and ethane can react with bromine to orm bromoethane.

$$H \xrightarrow[H]{C} H + CI \xrightarrow{UV} H \xrightarrow{H} CI \xrightarrow{UV} H \xrightarrow{H} CI + H \xrightarrow{CI} CI$$

methane

chloromethane



MECHANISM OF CHLORINATION OF METHANE

The mechanism o an organic reaction describes the individual steps. When chemical bonds break they may break **heterolytically** or **homolytically**. In heterolytic fission both o the shared electrons go to one o the atoms resulting in a negative and a positive ion. In homolytic fission each o the two atoms orming the bond retains one o the shared electrons resulting in the ormation o two **free radicals**. The bond between two halogen atoms is weaker than the C H or C C bond in methane and can break homolytically in the presence o ultraviolet light.

 $Cl_2 = Cl + Cl$

This stage o the mechanism is called **initiation**.

Free radicals contain an unpaired electron and are highly reactive. When the chlorine ree radicals come into contact with a methane molecule they combine with a hydrogen atom to produce hydrogen chloride and a methyl radical.

 $H_{,C} H + Cl = H_{,C} + HCl$

Since a new radical is produced this stage o the mechanism is called **propagation**. The methyl ree radical is also extremely reactive and reacts with a chlorine molecule to orm the product and regenerate another chlorine radical. This is a urther propagation step and enables a chain reaction to occur as the process can repeat itsel .

$$CH_3 + Cl_2 = CH_3 Cl + Cl$$

In theory a single chlorine radical may cause up to 10 000 molecules o chloromethane to be ormed. **Termination** occurs when two radicals react together.

Cl + Cl = C	2l ₂	
$CH_{_3} + Cl$	CH ₃ Cl	termination
$CH_3 + CH_3$	C ₂ H ₆	

Further substitution can occur when chlorine radicals react with the substituted products. For example:

The substitution can continue even urther to produce trichloromethane and then tetrachloromethane.

The overall mechanism is called **free radical substitution**. [Note that in this mechanism hydrogen radicals H are not ormed.]

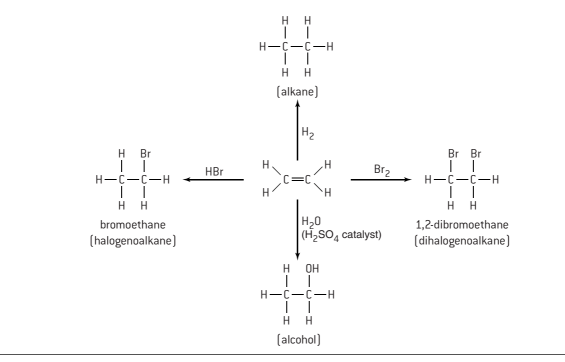
Alkenes

ADDITION REACTIONS

The bond enthalpy of the C=C double bond in alkenes has a value of 612 kJ mol⁻¹. This is less than twice the average value of 348kJ mol⁻¹ for the C C single bond and accounts for the relative reactivity of alkenes compared to alkanes. The most important reactions of alkenes are addition reactions. Reactive molecules are able to add across the double bond. The double bond is said to be **unsaturated** and the product, in which each carbon atom is bonded by four single bonds, is said to be **saturated**.



Addition reactions include the addition of hydrogen, bromine, hydrogen halides and water.



USES OF ADDITION REACTIONS

1. Bromination

Pure bromine is a red liquid but it has a distinctive yellow/orange colour in solution. When a solution of bromine is added to an alkene the product is colourless. This decolourization of bromine solution provides a useful test to indicate the presence of an alkene group.

2. Hydration

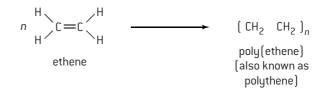
Ethene is an important product formed during the cracking of oil. Although ethanol can be made from the fermentation of starch and sugars, much industrial ethanol is formed from the addition of steam to ethene.

3. Hydrogenation

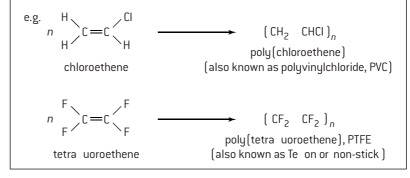
The addition of hydrogen to unsaturated vegetable oils is used industrially to make margarine. Hydrogenation reduces the number of double bonds in the polyunsaturated vegetable oils present in the margarine, which causes it to become a solid at room temperature.

ADDITION POLYMERIZATION

Under certain conditions ethene can also undergo addition reactions with itself to form a long chain polymer containing many thousands (typically 40 000 to 800 000) of carbon atoms.



These addition reactions can be extended to other substituted alkenes to give a wide variety of different addition polymers.



Alcohols

COMBUSTION

Ethanol is used both as a solvent and as a uel. It combusts completely in a plenti ul supply o oxygen to give carbon dioxide and water.

 $C_{2}H_{2}OH(l) + 3O_{2}(g) = 2CO_{2}(g) + 3H_{2}O(l) = H^{\ominus} = -1371 \text{ kJ mol}^{-1}$

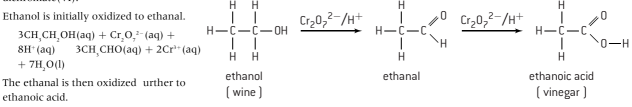
Ethanol is already partially oxidized so it releases less energy than burning an alkane o comparable mass. However, it can be obtained by the ermentation o biomass so in some countries it is mixed with petrol to produce gasohol which decreases the dependence on crude oil.

The general equation or an alcohol combusting completely in oxygen is:

 $C_n H_{(2n+1)} OH + (2n \ 1)O_2$ $nCO_2 + (n+1)H_2 O$

OXIDATION OF ETHANOL

Ethanol can be readily oxidized by warming with an acidi ed solution o potassium dichromate(VI). During the process the orange dichromate(VI) ion $\operatorname{Cr}_2 \operatorname{O}_7^{2-}$ is reduced rom an oxidation state o +6 to the green Cr^{3+} ion. Use is made o this in simple breathalyser tests, where a motorist who is suspected o having exceeded the alcohol limit blows into a bag containing crystals o potassium dichromate(VI).



 $3CH_{2}CHO(aq) + Cr_{2}O_{2}^{2-}(aq) + 8H^{+}(aq) = 3CH_{2}COOH(aq) + 2Cr^{3+}(aq) + 4H_{2}O(l)$

Unlike ethanol (b. pt 78.5 C) and ethanoic acid (b. pt 118 C) ethanal (b. pt 20.8 C) does not have hydrogen bonding between its molecules, and so has a lower boiling point. To stop the reaction at the aldehyde stage the ethanal can be distilled rom the reaction mixture as soon as it is ormed. I the complete oxidation to ethanoic acid is required, then the mixture can be heated under reflux so that none o the ethanal can escape.

OXIDATION OF ALCOHOLS

Ethanol is a primary alcohol, that is the carbon atom bonded to the OH group is bonded to two hydrogen atoms and one alkyl group. The oxidation reactions o alcohols can be used to distinguish between primary, secondary and tertiary alcohols.

All **primary alcohols** are oxidized by acidi ed potassium dichromate(VI), rst to aldehydes then to carboxylic acids.

$$\begin{array}{c} \stackrel{\stackrel{\scriptstyle }{\overset{\scriptstyle }}{\underset{\scriptstyle H}{\overset{\scriptstyle }}} \\ R - \overset{\scriptstyle C}{\underset{\scriptstyle H}{\overset{\scriptstyle }}} \\ OH \end{array} \xrightarrow{ Cr_2 O_7^{2-}/H^+ } R - C \overset{\scriptstyle \mathcal{C}}{\underset{\scriptstyle H}{\overset{\scriptstyle O}{\overset{\scriptstyle }}} \xrightarrow{ Cr_2 O_7^{2-}/H^+ } R - C \overset{\scriptstyle \mathcal{C}}{\underset{\scriptstyle OH}{\overset{\scriptstyle O}{\overset{\scriptstyle }}}$$

primary alcohol

Н

aldehyde

carboxylic acid

Secondary alcohols are oxidized to ketones, which cannot undergo urther oxidation.

$$\begin{array}{c} H & 0 \\ R - C - 0H & \frac{Cr_2 O_7^{2-} / H^+}{R} & \prod_{l=1}^{N} R - C - R \\ R & R \end{array}$$

secondary alcohol

ketone

Tertiary alcohols cannot be oxidized by acidi ed dichromate(VI) ions as they have no hydrogen atoms attached directly to the carbon atom containing the -OH group. It is not true to say that tertiary alcohols can never be oxidized, as they burn readily, but when this happens the carbon chain is destroyed.

tertiary alcohol

Substitution and condensation reactions

SUBSTITUTION REACTIONS OF HALOGENOALKANES

Because o the greater electronegativity o the halogen atom compared with the carbon atom halogenoalkanes have a polar bond. Reagents that have a non-bonding pair o electrons are attracted to the electron-de cient carbon atom in halogenoalkanes and a substitution reaction occurs. Such reagents are called nucleophiles.

$$Nu^{-}$$
: $C^{+}_{Br^{-}}$

| A double-headed curly arrow represents the movement o a pair o electrons. It shows where they come rom and where they move to.

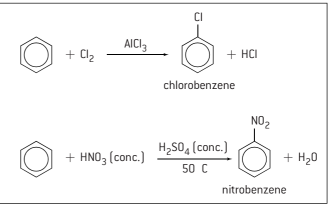
These reactions are use ul in organic synthesis as a wide variety o di erent compounds can be made by varying the nucleophile. For example, with warm dilute sodium hydroxide solution the product is an alcohol, with ammonia the product is an amine and with cyanide ions, CN, the product is a nitrile. This last reaction is particularly use ul as it provides a way to increase the number o atoms in the carbon chain.

Reaction o bromoethane with warm dilute sodium hydroxide solution:

 $C_{2}H_{5}Br(l) + NaOH(aq) = C_{2}H_{5}OH(aq) + NaBr(aq)$

SUBSTITUTION REACTIONS OF BENZENE

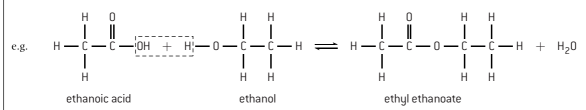
The extra stability provided by the delocalization o the electrons in the benzene ring means that benzene, unlike simple alkenes, does not readily undergo addition reactions. However benzene does undergo substitution reactions. Benzene has a high electron density so reacts with electrophiles. Electrophiles are electron-de cient species o ten ormed *in situ* that can accept electron pairs. Two examples o electrophilic substitution reactions o benzene are the reaction with chlorine in the presence o aluminium chloride (the electrophile is Cl⁺) to orm chlorobenzene and the reaction with nitric acid in the presence o sul uric acid (the electrophile is NO₂⁺) to orm nitrobenzene.



CONDENSATION REACTION BETWEEN AN ALCOHOL AND A CARBOXYLIC ACID

Alcohols can undergo a nucleophilic substitution reaction with carboxylic acids but this is more normally called esteri cation and is an example o a condensation reaction. A condensation reaction involves the reaction between two molecules to produce a larger molecule with the elimination o a small molecule such as water or hydrogen chloride.

Alcohols react with carboxylic acids in the presence o a small amount o concentrated sul uric acid, which acts as a catalyst, to orm an ester.



Most esters have a distinctive, pleasant ruity smell and are used both as natural and articial flavouring agents in ood. For example, ethyl methanoate HCOOCH₂CH₃ is added to chocolate to give it the characteristic flavour or rum tru fle. Esters are also used as solvents in per umes and as plasticizers (substances used to modi y the properties or polymers by making them more flexible.)

Many drugs contain one or more ester groups e.g. aspirin and heroin.

Fats and oils are natural triesters ormed rom the reaction between glycerol (propane-1,2-3-triol) and three atty acids. In a process known as transesteri cation they can react with alcohols in the presence o sodium hydroxide (which acts as a catalyst) to orm alkyl esters, which can be used as bio uel.

CH ₂ — OCOR ₁			СН ₂ — ОН	R ₁ — COOCH ₃
CH OCOR ₂ +	3 HOCH ₃ =	NaOH	и СН —— ОН +	R ₂ — COOCH ₃
CH ₂ — OCOR ₃			СН ₂ — ОН	R ₃ — COOCH ₃
triglyceride (fat or oil)	methanol		glycerol	methyl esters (biofuel)

🕕 Nucleophilic substitution

MECHANISM OF NUCLEOPHILIC SUBSTITUTION

Primary halogenoalkanes (one alkyl group attached to the carbon atom bonded to the halogen)

the carbon atom bonded to the halogen)

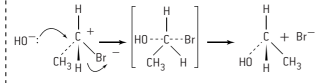
For example, the reaction between bromoethane and warm dilute sodium hydroxide solution.

 $C_2H_5Br + OH^ C_2H_5OH + Br^-$

The experimentally determined rate expression is:

rate = $k[C_{2}H_{5}Br][OH^{-}]$

The proposed mechanism involves the ormation o a transition state which involves both o the reactants.



Because the molecularity o this single-step mechanism is two it is known as an $S_{_N}^2$ mechanism (bimolecular nucleophilic substitution).

Note that the $S_{_N}^2$ mechanism is stereospecific with an inversion o configuration at the central carbon atom.

Tertiary halogenoalkanes (three alkyl groups attached to the carbon atom bonded to the halogen)

For example, the reaction between 2-bromo-2-methylpropane and warm dilute sodium hydroxide solution.

$$C(CH_3)_3 Br + OH^- \longrightarrow C(CH_3)_3 OH + Br^-$$

The experimentally determined rate expression or this

reaction is: rate = $k[C(CH_3)_3Br]$

A two-step mechanism is proposed that is consistent with this rate expression.

$$(CH_3)_3 Br \longrightarrow C(CH_3)_3^+ + Br$$

 $C(CH_3)_3^+ + OH^- \xrightarrow{fast} C(CH_3)_3 OH$

In this reaction it is the first step, the heterolytic fission o the C Br bond, that is the rate-determining step. The molecularity o this step is one and the mechanism is known as $S_{N}1$ (unimolecular nucleophilic substitution).

The mechanism or the hydrolysis o secondary halogenoalkanes (e.g 2-bromopropane $CH_3CHBrCH_3$) is more complicated as they can proceed by either $S_N 1$ or $S_N 2$ pathways or a combination o both.

CHOICE OF SOLVENT

Whether or not the reaction proceeds by an $S_N 1$ or $S_N 2$ mechanism also depends upon the solvent. Protic solvents which are polar, such as water or ethanol, avour the $S_N 1$ mechanism as they support the breakdown o halogenoalkanes into carbocations and halide ions whereas aprotic solvents which are less polar such as ethoxyethane, $C_2H_5OC_2O_5$, avour an $S_N 2$ mechanism involving a transition state.

FACTORS AFFECTING THE RATE OF NUCLEOPHILIC SUBSTITUTION

The nature of the nucleophile

The nature of the halogen

The e ectiveness o a nucleophile depends on its electron density. Anions tend to be more reactive than the corresponding neutral species. This explains why the hydroxide ion is a much better nucleophile than water. For both $S_{N}1$ and $S_{N}2$ reactions iodoalkanes react aster than bromoalkanes, which in turn react aster than chloroalkanes. This is due to the relative bond enthalpies as the C I bond is much weaker than the C Cl bond and there ore breaks more readily.

Bond enthalpy / kJ mol⁻¹

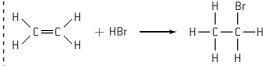
C I 228 C Br 285 C Cl 324

The nature of the halogenoalakane

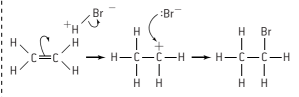
Tertiary halogenoalkanes react aster than secondary halogenoalkanes which in turn react aster than primary halogenoalkanes. The $S_n 1$ route which involves the ormation o an intermediate carbocation is aster than the $S_n 2$ route which involves a transition state with a relatively high activation energy.

💷 Electrophilic addition reactions (1)

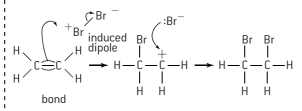
ELECTROPHILIC ADDITION TO SYMMETRIC ALKENES Ethene readily undergoes addition reactions. With hydrogen bromide it orms bromoethane.



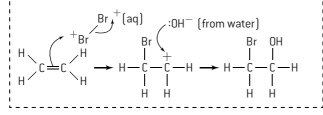
The reaction can occur in the dark which suggests that a ree radical mechanism is not involved. The double bond in the ethene molecule has a region o high electron density above and below the plane o the molecule. Hydrogen bromide is a polar molecule due to the greater electronegativity o bromine compared with hydrogen. The hydrogen atom (which contains a charge o +) rom the H Br is attracted to the double bond and the H Br bond breaks, orming a bromide ion. At the same time the hydrogen atom adds to one o the ethene carbon atoms leaving the other carbon atom with a positive charge. A carbon atom with a positive charge is known as a **carbocation**. The carbocation then combines with the bromide ion to orm bromoethane. Because the hydrogen bromide molecule is attracted to a region o electron density it is described as an **electrophile** and the mechanism is described as electrophilic addition.

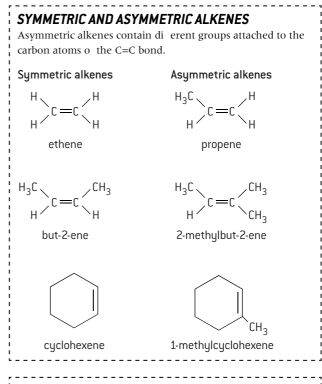


Electrophilic addition also takes place when bromine adds to ethene in a non-polar solvent to give 1,2-dibromoethane. Bromine itsel is non-polar but as it approaches the double bond o the ethene an induced dipole is ormed by the electron cloud.



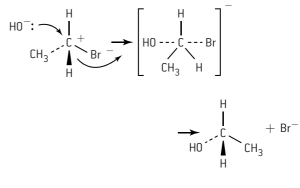
Evidence or this mechanism is that when bromine water is reacted with ethene the main product is 2-bromoethanol not 1,2-dibromoethane. This suggests that hydroxide ions rom the water add to the carbocation in pre erence to bromide ions.





CURLY ARROWS AND FISH HOOKS

Curly arrows are used to show the movement o a pair o electrons as or example in the electrophilic addition reaction o hydrogen bromide with ethene shown in the opposite box. The tail o the arrows shows the origin o the electron pair and the head where the electron pair ends up. In $S_{_N}2$ reactions such as the nucleophilic substitution reaction o bromoethane by hydroxide ions it is best to use three-dimensional representations to clari y the stereochemistry as the arrow head should go to the opposite side o the + carbon atom ormed due to the polar C Br bond (see page 89).



When single electrons are trans erred as in the ree radical substitution reactions o alkanes with halogens in ultraviolet light as described on page 85 then single-headed arrows known as fish hooks are used. Thus the initiation step which involves the homolytic fission o the chlorine to chlorine bond by ultraviolet light can be represented as:

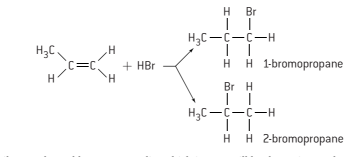




💵 Electrophilic addition reactions (2)

MARKOVNIKOV S RULE

When hydrogen halides add to asymmetric alkenes two products are possible depending upon which carbon atom the hydrogen atom bonds to. For example, the addition o hydrogen bromide to propene could produce 1-bromopropane or 2-bromopropane.

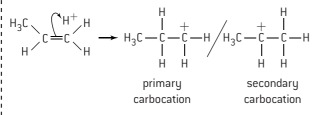


Markovnikov s rule enables you to predict which isomer will be the major product. It states that the hydrogen halide will add to the carbon atom that already contains the most hydrogen atoms bonded to it. Thus in the above example 2-bromopropane will be the major product.

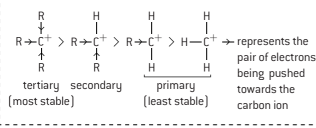
EXPLANATION OF MARKOVNIKOV S RULE

Markovnikov s rule enables the product to be predicted but it does not explain why. It can be explained by considering the nature o the possible intermediate carbocations ormed during the reaction.

When hydrogen ions react with propene two di erent carbocation intermediates can be ormed.

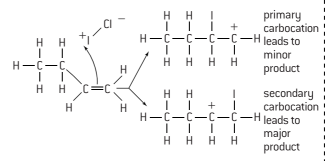


The first one has the general ormula RCH,⁺ and is known as a primary carbocation. The second one has two R groups attached to the positive carbon ion R₂CH⁺ and is known as a secondary carbocation. A tertiary carbocation has the general ormula R₂C⁺. The R-groups (alkyl groups) tend to push electrons towards the carbon atom they are attached to which tends to stabilize the positive charge on the carbocation. This is known as a **positive inductive effect**. This e ect will be greatest with tertiary carbocations and smallest with primary carbocations.



Thus in the above reaction the secondary carbocation will be pre erred as it is more stable than the primary carbocation. This secondary carbocation intermediate leads to the major product, 2-bromopropane.

Understanding this mechanism enables you to predict what will happen when an interhalogen adds to an asymmetric alkene even though no hydrogen atoms are involved. Consider the reaction o iodine chloride ICl with but-1-ene. Since iodine is less electronegative than chlorine the iodine atom will act as the electrophile and add first to the alkene.



The major product will thus be 2-chloro-1-iodobutane.

Electrophilic substitution reactions and reduction reactions

NITRATION OF BENZENE

Benzene reacts with a mixture o concentrated nitric acid and concentrated sul uric acid when warmed at 50 C to give nitrobenzene and water. Note that the temperature should not be raised above 50 C otherwise urther nitration to dinitrobenzene will occur.

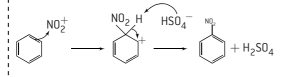
$$(\bigcirc + HNO_3 \text{ (conc.)} \xrightarrow{H_2SO_4 \text{ (conc.)}} (\bigcirc + H_2O_3 \text{ (conc.)} + H_2O_3 \text{ (conc.)}$$

The electrophile is the **nitryl cation** NO_2^+ (also called the nitronium ion). The concentrated sul uric acid acts as a catalyst. Its unction is to protonate the nitric acid which then loses water to orm the electrophile. In this reaction nitric acid is acting as a base in the presence o the more acidic sul uric acid.

The NO_2^+ is attracted to the delocalized bond and attaches to one o the carbon atoms. This requires considerable activation energy as the delocalized bond is partially broken. The positive charge is distributed over the remains o the bond in the intermediate. The intermediate then loses a proton and energy is evolved as the delocalized bond is re ormed. The proton can recombine with the hydrogensul ate ion to regenerate the catalyst.

 $(H^{1})^{+} (H^{2})^{+} (H^{2})^{+} (H^{2})^{-} (H^{$

Although it is more correct to draw the intermediate as a partially delocalized bond it can sometimes be convenient to show benzene as i it does contain alternate single and double carbon to carbon bonds. In this model the positive charge is located on a particular carbon atom.



REDUCTION REACTIONS

a) Reduction of carbonyl compounds

There are several reducing agents that can be used to reduce carbonyl compounds. Typical among these are lithium tetrahydridoaluminate (also known as lithium aluminium hydride), LiAlH, and sodium tetrahydridoborate (also known as sodium borohydride), NaBH₄. Both e ectively provide a source o H ions, which act as the reducing agent undergoing a nucleophilic addition reaction with the electronde cient carbon atom o the carbonyl group. Sodium tetrahydridoborate can be used in the presence o protic solvents such as water or ethanol but is ine ectual at reducing carboxylic acids. The stronger reducing agent lithium aluminium hydride must initially be used in aprotic solvents such as ether as it reacts with water, then the reaction is acidi ed to obtain the product. Another reducing agent that can be used is hydrogen itsel in the presence o a nickel, platinum or palladium catalyst.

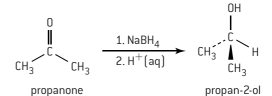
Aldehydes are reduced to primary alcohols.

e.g.

$$CH_{3}CH_{2}C \xrightarrow{0}_{H} \xrightarrow{1. NaBH_{4}} CH_{3}CH_{2}CH_{2}OH$$

propanal

Ketones are reduced to secondary alcohols. e.g.

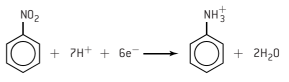


Carboxylic acids are reduced to primary alcohols according to the general equation:

b) Reduction of nitrobenzene

The reduction of nitrobenzene to phenylamine is usually carried out in two stages.

Stage 1. Nitrobenzene is refluxed with a mixture o tin and concentrated hydrochloric acid. The tin provides the electrons by acting as the reducing agent and the product is the phenylammonium ion.



Stage 2. The addition o sodium hydroxide solution releases the ree amine.





SYNTHESIS AND RETRO-SYNTHESIS

The raw materials or many organic compounds originate rom coal, crude oil or natural gas. The challenge or synthetic organic chemists is to devise reaction pathways to make new compounds rom simple starting materials using as ew steps as possible, each with the highest yield possible. The more steps there are in an organic synthesis, then the lower the nal yield is likely to be, because some material will be lost during each step. For example, but-2-ene can be obtained by cracking some o the higher boiling point ractions o crude oil. One way o obtaining butanone rom but-2-ene in a two-step process would be, rstly, to hydrate the but-2-ene by heating with steam in the presence o a sul uric acid catalyst to produce butan-2-ol. This secondary alcohol could then be oxidized with a warm acidi ed solution o potassium dichromate(VI).

This sub-topic requires organic synthesis using up to our di erent steps involving reactions that have already been covered. Rather than looking at the speci c compounds it is o ten help ul to look at the unctional groups involved and consider the reactions o these groups. For some syntheses it is help ul to work orward rom the reactants but it can be equally pro table to work backwards rom the product to nd the simplest and cheapest starting materials a process known as retro-synthesis. In the above example butanone is a ketone and ketones are ormed by oxidizing secondary alcohols. It can then be seen that alcohols are made by hydrating alkenes. The ollowing is a summary o the reactions covered in the Core and AHL organic chemistry.

Combustion o hydrocarbons and alcohols

Substitution o alkanes with halogens

Addition reactions o alkenes with H₂, H₂O, HX, X₂, interhalogens and polymerization

Substitution o halogenoalkanes with sodium hydroxide

Substitution o benzene to orm nitrobenzene

Reduction o nitrobenzene to orm phenylamine

Oxidation o alcohols and aldehydes

Reduction o aldehydes, ketones and carboxylic acids Esteri cation

When devising syntheses, the reagents and all necessary experimental conditions should be given. O ten more than one solution will be possible as the same product may be obtained by di erent routes. For example, bromoethane can be prepared rom ethene either by adding hydrogen bromide directly to ethene or by rst converting the ethene into ethane then reacting it with bromine in ultraviolet light.

WORKED EXAMPLES OF SYNTHESES

Starting with benzene, suggest a two-step synthesis o phenylamine.
 Working backwards, phenylamine can be obtained by reducing nitrobenzene, which can be obtained rom benzene by electrophilic substitution.

Step 1. React benzene with a mixture o concentrated nitric acid and concentrated sul uric acid at 50 C to orm nitrobenzene.

$$C_6H_6 + HNO_3(conc) \xrightarrow{H_2SO_4(conc)} C_6H_5NO_2 + H_2C$$

Step 2. Reflux nitrobenzene with tin and concentrated hydrochloric acid, then react with sodium hydroxide solution.

$$C_6H_5NO_2 \xrightarrow{\text{Sn/HCl, reflux}} C_6H_5NH_2$$

2. Starting with propane, suggest a our-step synthesis o propyl propanoate.

Propane reacts with halogens to orm halogenoalkanes, which can easily be converted into alcohols which are required to make an ester. A primary alcohol can also be oxidized to a carboxylic acid which will also be required to make the ester.

Step 1. React propane with bromine in ultraviolet light to give 1-bromopropane (other halogenated products will need to be discarded).

$$CH_3CH_2CH_3 + Br_2 \xrightarrow{UV} CH_3CH_2CH_2Br + HB$$

Step 2. React 1-bromopropane with warm dilute aqueous sodium hydroxide solution to give propan-1-ol.

CH,CH,CH,Br + NaOH <u>warm</u> → CH,CH,CH,OH + NaBr

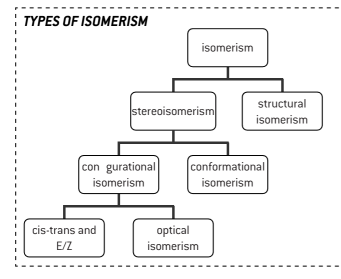
Step 3. Keep hal o the propan-1-ol and reflux the remainder with warm excess acidi ed potassium dichromate(VI) solution to convert it into propanoic acid.

 $CH_{2}CH_{2}CH_{2}OH + \frac{Cr_{2}O_{7}^{2}/H^{+}, reflux}{\longrightarrow} CH_{2}CH_{2}CH_{2}OH$

Step 4. Warm propan-1-ol and propanoic acid in the presence o a ew drops o concentrated sul uric acid as a catalyst to make the ester.

$$CH_{3}CH_{2}CH_{2}OH + CH_{3}CH_{2}COOH \xrightarrow{H^{+}} CH_{3}CH_{2}COOCH_{2}CH_{2}CH_{3} + H_{2}O$$

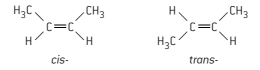
💷 Stereoisomerism (1)



Isomers are compounds that are composed o the same elements in the same proportions but di er in properties because o di erences in the arrangement o atoms. Structural isomers share the same molecular ormula but have di erent structural ormulas. That is, their atoms are bonded in di erent ways. Stereoisomers have the same structural ormula but di er in their spatial arrangement. Stereoisomers can be sub-divided into two classes con ormational isomers, which interconvert by rotation about a bond, and configurational isomers that interconvert only by breaking and re orming a bond. The IB syllabus on stereoisomerism is concerned with configurational isomers which can be urther sub-divided into cis trans and E/Zisomers (which together used to be known by the now obsolete term o geometrical isomers) and optical isomers.

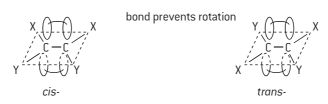
CIS TRANS ISOMERISM

Both cis trans and E/Z isomerism occur when rotation about a bond is restricted or prevented. The classic examples o cis trans isomerism occur with asymmetric non-cyclic alkenes o the type $R_1R_2C=CR_1R_2$. A cis-isomer is one in which the substituents are on the same side o the double bond. In a trans-isomer the substituents are on opposite sides o the double bond. For example, consider cis-but-2-ene and trans-but-2-ene.

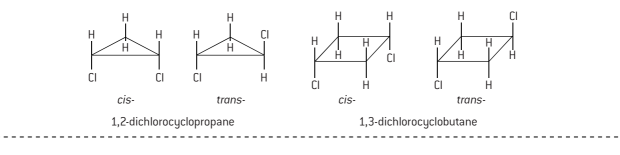


When there is a single bond between two carbon atoms ree rotation about the bond is possible. However, the double bond in an alkene is made up o a and a bond. The bond is ormed rom the combination o two p orbitals, one rom each o the carbon atoms. These two p orbitals must be in the same plane to combine. Rotating the bond would cause the bond to break so no rotation is possible.

Cis-trans isomerism will always occur in alkenes when the two groups X and Y attached to each o the two carbon atoms are di erent.



Cis trans isomerism can also occur in disubstituted cycloalkanes. The rotation is restricted because the C C single bond is part o a ring system. Examples include 1,2-dichlorocyclopropane and 1,3-dichlorocyclobutane.





E/Z ISOMERISM

Cis trans isomerism is a restricted form of E/Z isomerism as it only occurs when the two substituents R_1 and R_2 (or X and Y) occur on either side of the carbon to carbon double bond. E/Z isomerism covers every case where free rotation around a C=C double bond is not possible, i.e. for $R_1R_2C=CR_3R_4$ where $(R_1 \quad R_2, R_3 \quad R_4)$ and where neither R_1 nor R_2 need be different from R_3 or R_4 .

E/Z terminology is quite easy to apply and depends on what are known as the Cahn Ingold Prelog (CIP) rules for determining the priority of the atoms or groups attached to the two carbon atoms of the double bond. Each of these two carbon atoms on either side of the double bond is considered separately. In simple terms the higher the atomic number of the attached atoms to each carbon atom the higher the priority. Consider 1,2-dichloroethene. Chlorine has a higher atomic number than hydrogen so has a higher priority. If the two atoms/ groups lie on the same side the isomer is *Z* and if they lie on opposite sides they are *E*. In this simple case *Z* is the cisform and *E* is the trans- form.

However *Z* does not always equate to cis. If you consider 2-chlorobut-2-ene then the carbon atom of the methyl group has priority over the hydrogen atom on one side of the double bond but because chlorine has a higher atomic number than carbon the chlorine atom has priority over the methyl group on the other side of the double bond. Now the cis-isomer is the E isomer and the trans-isomer is the *Z* isomer.

E/Z and not cis trans must be used when the groups are all different. Consider 2-bromo-1-chloro-2-iodoethene. It is not obvious which would be the cis- and trans- forms. However using the Cahn Ingold Prelog rules the *E* and *Z* forms can easily be determined.

Z and *E* both come from German words. *Z* is from *zusammen* (together) and *E* is from *entgegen* (opposite). One easy way to remember the correct application is that *E* could stand for **e**nemy, and enemies are on opposite sides. Cl has a higher priority than H on the left-hand side and on the right-hand side.

Both highest priorities lie on the same side so *Z* isomer.



cis-1, 2-dichloroethene [*Z*]-1, 2-dichloroethene

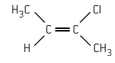
Highest priorities lie on opposite side so isomer.

)c=c

trans-1, 2-dichloroethene []-1, 2-dichloroethene

 CH_3 – has a higher priority than H on the left-hand side and Cl has a higher priority than CH_3 – on the right-hand side.

Both highest priorities lie on the same side so *Z* isomer.



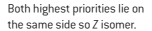
trans-2-chlorobut-2-ene (Z)-2-chlorobut-2-ene

Highest priorities lie on opposite side so isomer.

 $H_{3^{L}}$ C = C

cis-2-chlorobut-2-ene ()-2-chlorobut-2-ene

Cl has a higher priority than H on the left-hand side and I has a higher priority than Br on the right-hand side.



(Z)-2-bromo-1-chloro-2-iodoethene

Highest priorities lie on opposite side so isomer.

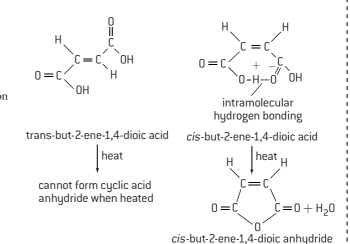


()-2-bromo-1-chloro-2-iodoethene

PHYSICAL AND CHEMICAL PROPERTIES OF E/Z AND CIS TRANS ISOMERS

The chemical properties of E/Z and cis trans isomers tend to be similar but their physical properties are different. For example, the boiling point of cis-1,2-dichloroethene is 60.3 C whereas trans-1,2-dichloroethene boils at the lower temperature of 47.5 C. Sometimes there can be a marked difference in both chemical and physical properties. This tends to occur when there is some sort of chemical interaction between the substituents. cis-but-2-ene-1,4-dioic acid melts with decomposition at 130 131 C. However, trans-but-2-ene,1,4-dioic acid does not melt until 286 C. In the cisisomer the two carboxylic acid groups are closer together so that intramolecular hydrogen bonding is possible between them. In the trans- isomer they are too far apart to attract each other so there are stronger intermolecular forces of attraction between different molecules, resulting in a higher melting point.

The cis- isomer reacts when heated to lose water and form a cyclic acid anhydride. The trans- isomer cannot undergo this reaction.

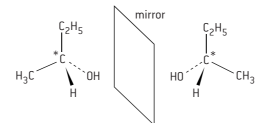




💵 Stereoisomerism (3)

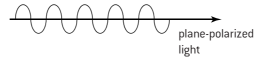
OPTICAL ISOMERISM

Optical isomerism is shown by all compounds that contain at least one asymmetric or chiral carbon atom within the molecule, that is, one that contains our di erent atoms or groups bonded to it, also known as a stereocentre. The two isomers are known as enantiomers and are mirror images o each other. Examples include butan-2-ol, CH₃CH(OH)C₃H₅, 2-hydroxypropanoic acid (lactic acid), CH₃CH(OH)COOH and all amino acids (except glycine, NH₂CH,COOH).



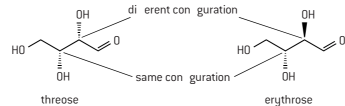
enantiomers of butan-2-ol (* asymmetric carbon/chiral carbon/stereocentre)

The two di erent isomers are optically active with plane-polarized light. Normal light consists o electromagnetic radiation which vibrates in all planes. When it is passed through a polarizing filter the waves only vibrate in one plane and the light is said to be plane-polarized.



The two enantiomers both rotate the plane o plane-polarized light. One o the enantiomers rotates it to the le t and the other rotates it by the same amount to the right. Apart rom their behaviour towards plane-polarized light enantiomers have identical physical properties. Their chemical properties are identical too except when they interact with other optically active substances. This is o ten the case in the body where the di erent enantiomers can have completely di erent physiological e ects. For example one o the enantiomers o the amino acid asparagine H,NCH(CH,CONH,)COOH tastes bitter whereas the other enantiomer tastes sweet.

I a molecule contains two or more stereocentres then several di erent stereoisomers are possible. They are known as enantiomers i they are mirror images and as diastereomers i they are not mirror images o each other. Diastereomerism occurs when two or more stereoisomers o a compound have di erent configurations at one or more (but not all) o the equivalent stereocentres. This is particularly important with many sugars and some amino acids. Diastereomers have di erent physical properties and di erent chemical reactivity. e.g.

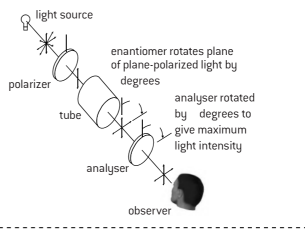


diastereomers of 2, 3, 4-trihydroxybutanal

POLARIMETRY

The optical activity o enantiomers can be detected and measured by an instrument called a polarimeter. It consists o a light source, two polarizing lenses, and between the lenses a tube to hold the sample o the enantiomer dissolved in a suitable solvent.

When light passes through the first polarizing lens (polarizer) it becomes plane-polarized. That is, it is vibrating in a single plane. With no sample present the observer will see the maximum intensity o light when the second polarizing lens (analyser) is in the same plane. Rotating the analyser by 90 will cut out all the light. When the sample is placed between the lenses the analyser must be rotated by degrees, either clockwise (dextrorotatory) or anticlockwise (laevorotatory) to give light o maximum intensity. The two enantiomers rotate the plane o plane-polarized light by the same amount but in opposite directions. I both enantiomers are present in equal amounts the two rotations cancel each other out and the mixture appears to be optically inactive. Such a mixture is known as a racemic mixture or racemate.



MULTIPLE CHOICE QUESTIONS ORGANIC CHEMISTRY

same homologous series?A. CH,COOHC. C,H,CHOA. CH,COOH and HCOOCH,B. CH,COOH and C,C,H,CHOD. HCOOCH,C. C,L4, and C,H,B. CH,OH and C,H,OHD. HCOOCH,B. CH,OH and C,H,OHD. C,H,CH and C,H,CLC. C,H,CHOD. C,H,CI and C,H,CIC. C,H,CHOD. HCOOCH,D. C,H,CI and C,H,CIA. CH,OOH and C,H,OHC. C, C,H,CHOJ. C, J, CI and C,H,CIB. 2C. ethanoi is partially oxidized by an acidified solution potassium dichromate(VI), the product that can be obtained by distillation as soon as it is ormet is:A. 1B. 2C. ethanoic acidC. 3D. 4T. Which ormula is that o a secondary halogenoalkane?A. 2. Mapping IUPAC rules what is the name oH3C-C-CH3? CH2 CH3C. C,CH3/2CH2CH3BrA. 2-methyl-2-ethylpropaneC. 2,2-dimethylbutane B. hexaneB. ethereD. (CH3/3CBrB. hexaneD. 2-methylpentaneC. nucleophilic substitutionA. It involves heterolytic fission and Cl ⁻ ions. B. It involves heterolytic fission and Cl radicals. C. It involves homolytic fission and Cl radicals.Y. Which alcohol cannot be oxidized by an acidified solution potassium dichromate(VI)?A. CH,COHC. (CH3/3CHCH2OH CH2CH2CH2OHA. (CH3/3CHCH2OH CH2CH2CH2OHC. C, CH3/3CHCH2OH	1.	Which o the ollowing two comp	ounds both belong to the	5.	Which compound is an ester?	
C. C_2H_4 and C_2H_6 C. C_2H_4 and C_2H_6 C. C_2H_4 and C_2H_5 CHB. CH_3OH and C_2H_5OH D. C_2H_5CI and $C_2H_4CI_5$ C. C_3H_{12} exist?C. Mow many di erent isomers o C_5H_{12} exist?A. 1 B. 2 C. 3 D. 4 C. ethanoic acidA. 1 B. 2 C. ethanoic acidC. 3 D. 4 C. ethanoic acidB. Applying IUPAC rules what is the name o $H_3C - C - CH_3$?C. $CH_5CH_2CH_2Br$ C. $(CH_3)_2CHCH_2Br$ B. Applying IUPAC rules what is the name o $H_3C - C - CH_3$?C. $CH_5CH_2CH_2CH_3Br$ C. $(CH_3)_5CHCH_2Br$ B. Applying IUPAC rules what is the name o $H_3C - C - CH_3$?C. $CH_5CH_2CH_2CH_3Br$ C. $(CH_3)_5CHCH_2Br$ B. Applying IUPAC rules what is the name o $H_3C - C - CH_3$?C. $CH_5CH_2CH_2CH_3$ D. $(CH_3)_5CHCH_2Br$ B. Applying IUPAC rules what is the name o $H_3C - C - CH_3$?C. $CH_5CH_2CH_2CH_3$ D. $(CH_3)_5CHCH_2Br$ B. Applying IUPAC rules what is the name o $H_3C - C - CH_3$?C. $CH_5CH_2CH_2CH_3$ D. $(CH_3)_5CHCH_2Br$ B. Applying IUPAC rules what is the name o $H_3C - C - CH_3$?C. $CH_5CH_2CH_2GH_3$ A. $CH_5CH_2CH_2CH_3$ A. 2-methyl-2-ethylpropaneC. $2,2$ -dimethylbutaneB. esterificationC. $C. (CH_3)_5CBr$ B. hexaneD. 2 -methylpentaneA. ree radical substitutionD. additionA. It involves heterolytic fission and Cl ⁻ ions.B. It involves heterolytic fission and Cl ⁻ ions.P. $CH CH CH CH CH_3CH_2CH_2CH_2CH_2CH_2CH_2CH_2CH_2CH_2CH_2$		same homologous series?			A. CH ₃ COOH	C. C ₂ H ₅ CHO
B. $CH_{3}OH$ and $C_{2}H_{3}OH$ Initiation $C_{2}H_{3}OH$ Initiation $C_{2}H_{3}OH$ D. $C_{2}H_{3}Cl$ and $C_{2}H_{4}Cl_{2}$ Initiation $C_{2}H_{3}OH$ Initiation $C_{2}H_{3}OH$ 2. How many dimension of $C_{3}H_{12}$ exist?A. 1B. 2C. 3D. 4C. ethanolic acidB. AD. 4C. ethanolic acidC. 3D. 4C. ethanolic acidC. 3D. 4C. ethanolic acidC. 3D. 4C. ethanolic acidA. Applying IUPAC rules what is the name o $H_{3}C - C - CH_{3}$? CH_{2} CH_{3} C. $C(H_{3})_{2}CHCH_{2}Br$ A. 2-methyl-2-ethylpropaneC. 2,2-dimethylbutaneB. hexaneD. 2-methylpentane4. Which statement is correct about the reaction between methane and chlorine?D. 2-methylpentane4. Which statement is correct about the reaction between methane and chlorine?P. etherolytic fission and Cl ⁻ ions.B. It involves heterolytic fission and Cl ions.P. Which alcohol cannot be oxidized by an acidified solution potassium dichromate(VI)?9. Which alcohol cannot be oxidized by an acidified solution potassium dichromate(VI)?9. CH		A. CH ₃ COOH and HCOOCH ₃			B. CH ₃ OC ₂ H ₅	D. HCOOCH ₃
b. $C_2H_3Cl and C_2H_4Cl_2$ How many di erent isomers o C_3H_{12} exist? A. 1 B. 2 C. 3 Applying IUPAC rules what is the name o $H_3C - C - C - CH_3$? A. 2-methyl-2-ethylpropane B. hexane M. 2-methyl-2-ethylpropane C. 2,2-dimethylbutane B. hexane M. 2 - methyl-2-ethylpropane C. 2,2-dimethylbutane B. hexane M. 2 - methylpentane C. 1 tinvolves heterolytic fission and Cl ⁻ ions. B. It involves heterolytic fission and Cl ⁻ ions. B. It involves heterolytic fission and Cl radicals. C. It involves homolytic fission and Cl ions. B. It involves heterolytic fission and Cl ions. B. It involves heterolytic fission and Cl ions. B. It involves heterolytic fission and Cl ions. B. CH		C. C_2H_4 and C_2H_6		6.	When ethanol is partially oxidized	by an acidified solution o
D: $C_2 \Pi_3$ of thic $C_2 \Pi_4$ or M_4 or M_2 A. ethanalC. ethanoic acid2. How many di erent isomers o $C_3 H_{12}$ exist?A. ethanalC. ethanoic acidA. 1B. 2B. etheneD. ethane-1,2-diolC. 3D. 4CH3FH33. Applying IUPAC rules what is the name o $H_3 C - C - C H_3$? $CH2C. (CH_3)_2 CHCH_2 BrA. 2-methyl-2-ethylpropaneC. 2,2-dimethylbutaneB. hexaneD. 2-methylpentane4. Which statement is correct about the reaction betweenmethane and chlorine?D. 2-methylpentane4. It involves heterolytic fission and Cl- ions.D. additionB. It involves heterolytic fission and Cl ions.P. Which alcohol cannot be oxidized by an acidified solutionpotassium dichromate(VI)?A. CH, CH, CH, CH, CH, OHD. CH, CHOHCH, CH$		B. $CH_{3}OH$ and $C_{2}H_{5}OH$				
2.How many uncertain isoners of $C_3 M_{12}$ exist?A. 1B. 2C. 3D. 4B. etheneD. ethane-1,2-diolC. 3D. 4J. 4 $H_3C - C - CH_3$?CH2LLCH2LLCH3CH2A. 2-methyl-2-ethylpropaneC. 2,2-dimethylbutaneB. hexaneD. 2-methylpentane4. Which statement is correct about the reaction betweenmethane and chlorine?A. It involves heterolytic fission and Cl ⁻ ions.B. It involves heterolytic fission and Cl radicals.C. It involves homolytic fission and Cl ions.		D. C_2H_5Cl and $C_2H_4Cl_2$			by distillation as soon as it is orme	d is:
A. 1D. 2C. 3D. 4S. Applying IUPAC rules what is the name o $H_3C - C - CH_3$? CH_2 CH_3 A. 2-methyl-2-ethylpropaneC. 2,2-dimethylbutane B. hexaneB. hexaneD. 2-methylpentane4. Which statement is correct about the reaction between methane and chlorine?A. It involves heterolytic fission and Cl ⁻ ions. B. It involves heterolytic fission and Cl ions.B. It involves heterolytic fission and Cl ions.C. It involves homolytic fission and Cl ions.	2.	How many di erent isomers o C_{5}	H ₁₂ exist?		A. ethanal	C. ethanoic acid
C. JC. GH_3 A. $CH_3CH_2CH_2CH_2Br$ C. $(CH_3)_2CHCH_2Br$ 3. Applying IUPAC rules what is the name o $H_3C - C - CH_3$? CH_2 CH_2 A. $CH_3CH_2CH_2CH_2Br$ C. $(CH_3)_2CHCH_2Br$ 3. Applying IUPAC rules what is the name o $H_3C - C - CH_3$? CH_2 CH_2 B. $CH_3CH_2CH_2CH_3$ D. $(CH_3)_3CBr$ 4. 2-methyl-2-ethylpropaneC. 2,2-dimethylbutane CH_3 A. ree radical substitutionB. esterification5. hexaneD. 2-methylpentaneC. nucleophilic substitutionD. addition6. It involves heterolytic fission and Cl radicals. C. It involves homolytic fission and Cl ions.9. Which alcohol cannot be oxidized by an acidified solution potassium dichromate(VI)?7. It involves homolytic fission and Cl ions.9. CH (CH CH CH OHC. $(CH_3)_2CHCH_2OH$ P. CH CHOCH CH		A. 1	B. 2		B. ethene	D. ethane-1,2-diol
 A. CH₃CH₂CH₂CH₂BF C. (CH₃)₂CHCH₂BF C. (CH₃)₂CHCH₂OH D. (CH₃)₂CHCH₂OH D. (CH₃)₂CHCH₂OH D. (CH₃)₂CHCH₂OH 		C. 3	D. 4	7.	Which ormula is that o a seconda	ry halogenoalkane?
 A. 2-methyl-2-ethylpropane A. 2-methyl-2-ethylpropane C. 2,2-dimethylbutane B. hexane D. 2-methylpentane Which statement is correct about the reaction between methane and chlorine? A. It involves heterolytic fission and Cl⁻ ions. B. It involves heterolytic fission and Cl radicals. C. It involves homolytic fission and Cl ions. D. CH CH					A. $CH_3CH_2CH_2CH_2Br$	C. (CH ₃) ₂ CHCH ₂ Br
I CH3an example o :A. 2-methyl-2-ethylpropaneC. 2,2-dimethylbutaneA. ree radical substitutionB. hexaneD. 2-methylpentaneB. esterification4. Which statement is correct about the reaction between methane and chlorine?C. nucleophilic substitutionA. It involves heterolytic fission and Cl ⁻ ions.9. Which alcohol cannot be oxidized by an acidified solution potassium dichromate(VI)?B. It involves homolytic fission and Cl radicals. C. It involves homolytic fission and Cl ions.9. CH (CH CH OHC. It involves homolytic fission and Cl ions.9. CH (CH CH OHC. It involves homolytic fission and Cl ions.9. CH (CH CH OHC. It involves homolytic fission and Cl ions.9. CH (CH CH OHC. It involves homolytic fission and Cl ions.9. CH (CH CH OHC. It involves homolytic fission and Cl ions.9. CH (CH CH OHC. It involves homolytic fission and Cl ions.9. CH (CH CH OHC. It involves homolytic fission and Cl ions.9. CH (CH CH OHC. It involves homolytic fission and Cl ions.9. CH (CH CH OHC. It involves homolytic fission and Cl ions.9. CH (CH CH OHC. It involves homolytic fission and Cl ions.9. CH (CH CH OHC. It involves homolytic fission and Cl ions.9. CH (CH CH OHC. It involves homolytic fission and Cl ions.9. CH (CH CH OHC. It involves homolytic fission and Cl ions.9. CH (CH CH OHC. It involves homolytic fission and Cl ions.9. CH (CH CH OHC. It involves homolytic fission and Cl ions.9. CH (CH CH CH OHC. It involves homolytic fission and	3.	Applying IUPAC rules what is the	name o H ₃ C—C—CH _{3?}		B. CH ₃ CHBrCH ₂ CH ₃	D. (CH ₃) ₃ CBr
LH3 A. 2-methyl-2-ethylpropane C. 2,2-dimethylbutane B. hexane D. 2-methylpentane B. esterification 4. Which statement is correct about the reaction between methane and chlorine? D. 2-methylpentane A. It involves heterolytic fission and Cl ⁻ ions. D. addition B. It involves heterolytic fission and Cl radicals. P. Which alcohol cannot be oxidized by an acidified solution potassium dichromate(VI)? A. (CH ₃) ₃ COH C. (CH ₃) ₂ CHCH ₂ OH B. CH CH CH CH CH OH D. CH CHOHCH CH			CH ₂	8.	The reaction between bromine and	l ethene in the dark is
 A. 2-methyl-2-ethylpropane B. hexane C. 2,2-dimethylbutane B. hexane D. 2-methylpentane C. 12-methylpentane Mhich statement is correct about the reaction between methane and chlorine? A. It involves heterolytic fission and Cl⁻ ions. B. It involves heterolytic fission and Cl radicals. C. It involves homolytic fission and Cl ions. A. ree radical substitution B. esterification C. nucleophilic substitution D. addition Which alcohol cannot be oxidized by an acidified solution potassium dichromate(VI)? A. (CH₃)₃COH C. (CH₃)₂CHCH₂OH B. CH CH CH CH CH OH C. HOHCH CH 			CH3		an example o :	
 B. hexane B. hexane C. nucleophilic substitution C. nucleophilic substitution D. addition D. addition C. nucleophilic substitution D. addition S. It involves heterolytic fission and Cl⁻ ions. E. It involves heterolytic fission and Cl radicals. C. It involves homolytic fission and Cl ions. 		A. 2-methyl-2-ethylpropane	-		A. ree radical substitution	
 4. Which statement is correct about the reaction between methane and chlorine? A. It involves heterolytic fission and Cl⁻ ions. B. It involves heterolytic fission and Cl radicals. C. It involves homolytic fission and Cl ions. C. It involves homolytic fission and Cl ions. D. addition 9. Which alcohol cannot be oxidized by an acidified solution potassium dichromate(VI)? A. (CH₃)₃COH C. (CH₃)₂CHCH₂OH B. CH CH CH CH CH OH C. HCHOHCH CH 		B. hexane	D. 2-methylpentane		B. esterification	
methane and chlorine?D. additionA. It involves heterolytic fission and Cl ⁻ ions.9. Which alcohol cannot be oxidized by an acidified solution potassium dichromate(VI)?B. It involves heterolytic fission and Cl radicals. C. It involves homolytic fission and Cl ions.9. Which alcohol cannot be oxidized by an acidified solution potassium dichromate(VI)?A. (CH ₃) ₃ COHC. (CH ₃) ₂ CHCH ₂ OH CH CH CH CH CH CH CH	4.	Which statement is correct about	the reaction between		C. nucleophilic substitution	
B. It involves heterolytic fission and Cl radicals. potassium dichromate(VI)? C. It involves homolytic fission and Cl ions. A. (CH ₃) ₃ COH C. (CH ₃) ₂ CHCH ₂ OH B. CH D. CH CHOHCH CH					D. addition	
C. It involves homolytic fission and Cl ions. A. (CH ₃) ₃ COH C. (CH ₃) ₂ CHCH ₂ OH C. (CH ₃) ₂ CHCH ₂ OH C. (CH ₃) ₂ CHCH ₂ OH		A. It involves heterolytic fission a	nd Cl⁻ ions.	9.	Which alcohol cannot be oxidized	by an acidified solution o
		B. It involves heterolytic fission a	nd Cl radicals.		potassium dichromate(VI)?	
D. It involves homolytic fission and Cl radicals. B. $CH_3CH_2CH_2CH_2OH$ D. $CH_3CHOHCH_2CH_3$		C. It involves homolytic fission an	ld Cl ions.		A. (CH ₃) ₃ COH	C. $(CH_3)_2CHCH_2OH$
		D. It involves homolytic fission an	d Cl radicals.		B. CH ₃ CH ₂ CH ₂ CH ₂ OH	D. CH ₃ CHOHCH ₂ CH ₃
	(

10. Which statement(s) is/are true about the reactions o halogenoalkanes with warm dilute sodium hydroxide solution?

I. CH₃I reacts aster than CH₃F

- II. (CH₃)₃CBr reacts aster than CH₃Br
- III. $(CH_3)_3$ CBr and $(CH_3)_3$ CCl both react by S_N^{1} mechanisms.
- A. I and II only C. II and III only
- B. I and III only D. I, II and III
- 11. The product rom the reaction o iodine monochloride, ICl, with pent-1-ene is:

A. CH ₃ CH ₂ CHICHClCH ₃	C. CH ₃ CH ₂ CH ₂ CHClCH ₂ I
B. CH ₃ CH ₂ CH ₂ CHICH ₂ Cl	D. CH ₃ CH ₂ CHClCHICH ₃

- 12. Ketones are reduced by sodium borohydride, NaBH₄, to give:
 - A. a primary alcohol C. a tertiary alcohol
 - B. a secondary alcohol D. a carboxylic acid
- 13. During the conversion o nitrobenzene to phenylamine (aniline) by tin and concentrated hydrochloric acid the organic intermediate is:
 - A. the phenylammonium ion C. phenol
 - B. dinitrobenzene D. chlorobenzene
- 14. Which can exist as enantiomers and diastereomers?

I. H₂N — CH — COOH H₃C — CH — OH

II. HO
$$\longrightarrow$$
 CH \longrightarrow CH \longrightarrow CH₂ \longrightarrow NH \longrightarrow CH₃
HO \longrightarrow OH

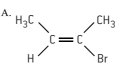
III. CH, CH(OH)CH, CH(OH)CHO

A. I and II only	C. II and III only
B. I and III only	D. I, II and III

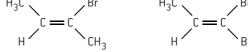
- 15. How many our-membered ring isomers are there o dichlorocyclobutane, C₄H₆Cl₂?
 - A. 3 C. 5 B. 4 D. 6

16. Which is an *E* isomer?

В







D

- 17. An organic compound is prepared rom ethanol by a our-step synthesis. Each step gives a yield o 70%. What will be the yield o the organic compound based on the initial amount o ethanol?
 - A. 24%C. 34%B. 28%D. 70%

SHORT ANSWER QUESTIONS ORGANIC CHEMISTRY

- 1. Alkenes are important starting materials or a variety o products.
 - a) State and explain the trend o the boiling points o the first five members o the alkene homologous series.
 - b) Describe two eatures o a homologous series. [2]

[3]

[5]

[1]

[1]

[1]

[1]

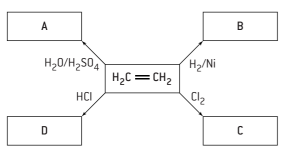
[1]

[4]

[6]

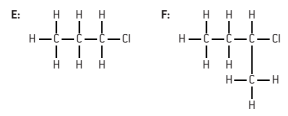
[8]

c) Below is a schematic diagram representing some reactions o ethene. The letters A D represent the organic compounds ormed rom the reactants and catalysts shown.



Deduce the structural ormulas o compounds **A**, **B**, **C** and **D** and state the IUPAC name o compound **C**.

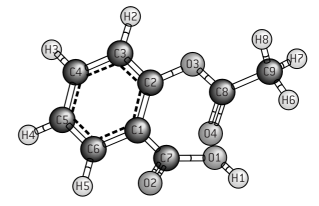
- d) Describe a chemical test that could be used to distinguish between pent-1-ene and pentane. [2]
- e) State and explain whether the ollowing molecules are primary, secondary or tertiary halogenoalkanes. [4]





- 3. Consider alkenes with the molecular ormula C₅H₉Br. Give structural ormulas or one isomer which shows
 - a) no stereoisomerism
 - b) optical but no cis trans or *E*/*Z* isomerism [1]
 - c) optical, trans- and *E* isomerism
 - d) optical, cis- and Z isomerism.
 - e) trans- and Z isomerism but no optical isomerism
 -) cis- and *E* isomerism but no optical isomerism.
- 4. State the equations or each step and any necessary inorganic reagents and conditions or:
 - a) The preparation o propyl propanoate by a **two**-step synthesis using propan-1-ol as the only organic reagent.
 - b) The preparation o propanone by a **three**-step synthesis using propene as the only organic reagent.
 - c) The preparation o propanal by a **four**-step synthesis using propene as the only organic reagent.

-) Explain, using equations, the initiation, propagation and termination steps in the ree-radical mechanism o the reaction o methane with chlorine.
- 2. The ollowing is a three-dimensional computer-generated representation o aspirin. Each carbon, oxygen and hydrogen atom has been given a unique number.



- a) Identi y one carbon atom that is part o
 - (i) the phenyl unctional group.
 - (ii) the carboxylic acid group.
 - (iii) the ester group. [3]
- b) Explain why the bond between C_8 and O_3 is longer than the bond between C_8 and O_4 . [2]
- c) Compare the length o the bond between C₁ and C₂ with the bond between C₁ and C₂. [2]
- d) Explain how a primary alcohol can be converted into
 - (i) a carboxylic acid
 - (ii) an ester. [4]
- 5. 1-bromopropane undergoes a substitution reaction with warm aqueous sodium hydroxide solution.



- a) Explain why the substitution occurs on the carbon atom that is marked as *C. [1] b) State the rate equation or this reaction and identi y the name o the reaction mechanism. [2] c) Explain the mechanism o the reaction using curly arrows to represent the movement o electron pairs during the substitution. [4] d) Explain how changing the halogenoalkane to 1-chloropropane would a ect the rate o the substitution reaction. [2] 6. Benzene can be nitrated to orm nitrobenzene by warming with a mixture o concentrated nitric acid and
- concentrated sul uric acid. Explain, with any necessary equations, the role o the concentrated sul uric acid in this reaction. [6]

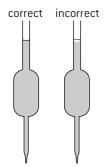
[4]

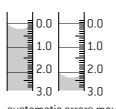
11 MEASUREMENT, DATA PROCESSING AND ANALYSIS

Uncertainty and error in measurement

RANDOM UNCERTAINTIES AND SYSTEMATIC ERRORS

Quantitative chemistry involves measurement. A measurement is a method by which some quantity or property o a substance is compared with a known standard. I the instrument used to take the measurements has been calibrated wrongly or i the person using it consistently misreads it then the measurements will always di er by the same amount. Such an error is known as a **systematic error**. An example might be always reading a pipette rom the sides o the meniscus rather than rom the middle o the meniscus.





a systematic error will be introduced if a pipette is read incorrectly systematic errors may cancel out when a di erence in two readings is taken $V = 2.0 \text{ cm}^3$ whether or not the burette is read

Random uncertainties occur i there is an equal probability o the reading being too high or too low rom one measurement to the next. These might include variations in the volume o glassware due to temperature uctuations or the decision on exactly when an indicator changes colour during an acid base titration.

PRECISION AND ACCURACY

Precision re ers to how close several experimental measurements o the same quantity are to each other. Accuracy re ers to how close the readings are to the true value. This may be the standard value, or the literature or accepted value. A measuring cylinder used to measure exactly 25 cm³ is likely to be much less accurate than a pipette that has been care ully calibrated to deliver exactly that volume. It is possible to have very precise readings which are inaccurate due to a systematic error. For example all the students in the class may obtain the same or very close results in a titration but i the standard solution used in all the titrations had been prepared wrongly be orehand the results would be inaccurate due to the systematic error. Because they are always either too high or too low systematic errors cannot be reduced by repeated readings. However random errors can be reduced by repeated readings because there is an equal probability o them being high or low each time the reading is taken. When taking a measurement it is usual practice to report the reading rom a scale as the smallest division or the last digit capable o precise measurement even though it is understood that the last digit has been rounded up or down so that there is a random error or uncertainty o 0.5 o the last unit.

SIGNIFICANT FIGURES

Whenever a measurement o a physical quantity is taken there will be a random uncertainty in the reading. The measurement quoted should include the first figure that is uncertain. This should include zero i necessary. Thus a reading o 25.30 C indicates that the temperature was measured with a thermometer that is accurate to + 0.01 C. I a thermometer accurate to only 0.1 C was used the temperature should be recorded as 25.3 C.

Zeros can cause problems when determining the number o significant figures. Essentially zero only becomes significant when it comes *after* a non-zero digit (1,2,3,4,5,6,7,8,9).

000123.4	0.0001234	1.0234	1.2340		
zero not a s	ignificant figure	zero is a significant figure			
values quot	ed to 4 s	values quot	ed to 5 s		

Zeros a ter a non-zero digit but be ore the decimal point may or may not be significant depending on how the measurement was made. For example 123 000 might mean exactly one hundred and twenty three thousand or one hundred and twenty three thousand to the nearest thousand. This problem can be neatly overcome by using scientific notation.

1.23000 10⁶ quoted to six significant figures

1.23 10⁶ quoted to three significant figures.

Calculations

1. When adding or subtracting it is the number o decimal places that is important. Thus when using a balance which measures to 0.01 g the answer can also be quoted to two decimal places which may increase or decrease the number o significant figures.

e.g.	7.10 g	+	3.10 g	=	10.20 g
	3 s		3 s		4 s
	22.36 g	_	15.16 g	=	7.20 g
	4 s		4 s		3 s

2. When multiplying or dividing it is the number o significant figures that is important. The number with the least number o significant figures used in the calculation determines how many significant figures should be used when quoting the answer.

e.g. When the temperature o 0.125 kg o water is increased by 7.2 C

the heat required

- $= 0.125 \ \text{kg} \quad \ \ 7.2 \ \ \text{C} \quad \ \ 4.18 \ \text{kJ} \ \text{kg}^{_{-1}} \ \ \text{C}^{_{-1}}$
- = 3.762 kJ

Since the temperature was only recorded to two significant figures the answer should strictly be given as 3.8 kJ.

Uncertainty in calculated results and graphical techniques

ABSOLUTE AND PERCENTAGE UNCERTAINTIES

When making a single measurement with a piece o apparatus the absolute uncertainty and the percentage uncertainty can both be stated relatively easily. For example consider measuring 25.0 cm³ with a 25 cm³ pipette which measures to 0.1 cm³. The absolute uncertainty is 0.1 cm³ and the percentage uncertainty is equal to:

$$\frac{0.1}{25.0}$$
 100 = 0.4%

I two volumes or two masses are simply added or subtracted then the absolute uncertainties are added. For example suppose two volumes o 25.0 cm³ 0.1 cm³ are added. In one extreme case the first volume could be 24.9 cm³ and the second volume 24.9 cm³ which would give a total volume o 48.8 cm³. Alternatively the first volume might have been 25.1 cm³ which when added to a second volume o 25.1 cm³ gives a total volume o 50.2 cm³. The final answer there ore can be quoted between 48.8 cm³ and 50.2 cm³, that is, 50.0 cm³ 0.2 cm³.

When using multiplication, division or powers then percentage uncertainties should be used during the calculation and then converted back into an absolute uncertainty when the final result is presented. For example, during a titration there are generally our separate pieces o apparatus, each o which contributes to the uncertainty.

e.g. when using a balance that weighs to 0.001 g the uncertainty in weighing 2.500 g will equal

 $\frac{0.001}{2.500}$ 100 = 0.04%

Similarly a pipette measures 25.00 cm³ 0.04 cm³.

The uncertainty due to the pipette is thus

$$\frac{0.04}{25.00}$$
 100 = 0.16%

Assuming the uncertainty due to the burette and the volumetric ask is 0.50% and 0.10% respectively the overall uncertainty is obtained by summing all the individual uncertainties:

Overall uncertainty = 0.04 + 0.16 + 0.50 + 0.10= 0.80% 1.0%

Hence i the answer is 1.87 mol dm⁻³ the uncertainty is 1.0% or 0.0187 mol dm⁻³.

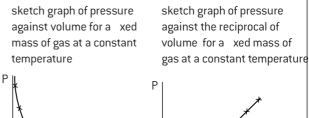
The answer should be given as $1.87 - 0.02 \text{ mol dm}^{-3}$.

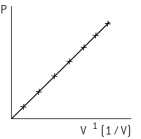
I the generally accepted correct value (obtained rom the data book or other literature) is known then the total error in the result is the di erence between the literature value and the experimental value divided by the literature value expressed as a percentage. For example, i the correct concentration or the concentration determined above is 1.90 mol dm⁻³ then:

the total error = $\frac{(1.90 - 1.87)}{1.90}$ 100 = 1.6%

GRAPHICAL TECHNIQUES

By plotting a suitable graph to give a straight line or some other definite relationship between the variables, graphs can be used to predict unknown values. There are various methods to achieve this. They include measuring the intercept, measuring the gradient, extrapolation and interpolation. Interpolation involves determining an unknown value within the limits o the values already measured. Extrapolation (see example on page 38) requires extending the graph to determine an unknown value which lies outside the range o the values measured. I possible manipulate the data to produce a straight line graph. For example, when investigating the relationship between pressure and volume or a fixed mass o gas a plot o P against V gives a curve whereas a plot o P against 1/V will give a straight line. Once the graph is in the orm o y = mx + cthen the values or both the gradient (m) and the intercept (c) can be determined.





Note that a sketched graph has labelled but unscaled axes and is used to show qualitative trends. Drawn graphs have labelled and scaled axes and are used in quantitative measurements.

The ollowing points should be observed when drawing a graph.

Plot the independent variable on the horizontal axis and the dependent variable on the vertical axis.

Choose appropriate scales or the axes.

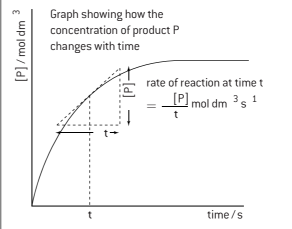
Use Standard International (SI) units wherever possible.

Label each axis and include the units.

Draw the line o best fit.

Give the graph a title.

Measuring a gradient from a graph



Analytical techniques

INFORMATION FROM DIFFERENT ANALYTICAL TECHNIQUES

The classic way to determine the structure o an organic compound was to determine both its empirical ormula and relative molar mass experimentally, then deduce the nature o the unctional groups rom its chemical reactivity. However, modern well-equipped laboratories now employ a variety o instrumental techniques, which i used in combination are able to unambiguously determine the exact structural ormula. They can also be used to determine the composition o the components in a mixture and to determine purity. These techniques are becoming ever more refined and some o them (e.g. mass spectrometry) can be used on extremely small samples. Be ore analysis can usually take place it is important to separate any mixture into its individual components hence the need or chromatography. O ten in ormation is not obtained rom a single technique but rom a combination o several o them. Some examples are:

In rared spectroscopy organic structural determination, in ormation on the strength o bonds, in ormation about the secondary structure o proteins, measuring the degree o unsaturation o oils and ats, and determining the level o alcohol in the breath. Mass spectrometry organic structural determination, isotopic dating (e.g. ¹⁴C dating). Proton nuclear magnetic resonance spectroscopy

(¹H NMR) organic structural determination, body scanning (known as MRI).

Chromatography drug testing in the blood and urine, ood testing, and orensic science.

INDEX OF HYDROGEN DEFICIENCY (IHD)

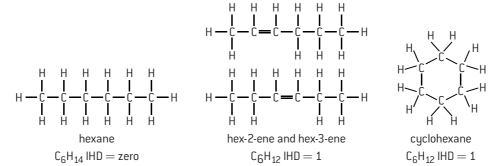
Once the molecular ormula o a compound is known the index o hydrogen deficiency can be calculated. This used to be known as the degree o unsaturation. A non-cyclic hydrocarbon that contains only single bonds contains the maximum number o hydrogen atoms and has the general ormula $C_n H_{(2n+2)}$. I any o the bonds are replaced by double or triple bonds or the compound is cyclic then the number o hydrogen atoms alls (i.e. there is a deficiency in the number o hydrogen atoms). The amount o the deficiency there ore gives use ul in ormation about the molecule and will assist in determining the number and types o di erent possible isomers.

The IHD or a hydrocarbon with x carbon atoms and y hydrogen atoms, $C_x H_{v'}$ is given by :

$$IHD = \frac{(2x+2-y)}{2}$$

where each double bond and each ring counts as one IHD and each triple bond counts as two IHD.

For example, hexane C_6H_{14} has an IHD equal to $\frac{(12+2-14)}{2} = 0$. This would be expected as hexane is a saturated hydrocarbon. I however a compound has a molecular ormula o C_6H_{12} its IHD is $\frac{(12+2-12)}{2} = 1$. This means that the compound could contain one double bond and be an isomer o hexene or it could be a cyclic compound.



Determining the IHD becomes a little more complicated i elements other than carbon and hydrogen are present in the compound. In this case:

Oxygen and sul ur atoms (O and S) do not a ect the IHD.

Halogens (F, Cl, Br, I) are treated like H atoms (i.e. CHCl₃ has the same IHD as CH₄).

For each N, add one to the number o C and one to the number o H (i.e. CH_5N has the same IHD as C_2H_6).

THE ELECTROMAGNETIC SPECTRUM

Modern spectroscopic techniques utilize di erent parts o the electromagnetic spectrum. The electromagnetic spectrum has already been described in Topic 2 *Atomic structure*. You should be amiliar with the relationship c = - and know the di erent regions o the spectrum.

The electromagnetic spectrum

	$0^{-10} 10^{-9}$	10 ⁻⁸ 10 ⁻	⁻⁷ 10 ⁻⁶	10^{-5} 10^{-4} 1	0^{-3} 10^{-2} 1	.0 ⁻¹ 1	0 ⁰ 10	0 ¹ 10 ² 10
Wavelength / m		3 10 ¹⁰	3 10 ⁸	3 10 ⁶	3 10 ⁴	ـــــــــــــــــــــــــــــــــــــ	10 ²	2
Frequency / MHz		3 10	3 10	3 10	3 10	3		3
Type of radiation	X-ray -rays	ultraviolet	visible	infrared	microwaves			radio waves
Type of transition	inner electrons	outer electr		molecular vibrations	molecular rotations			nuclear spin
-	4			increasing	g energy —			

MASS SPECTROMETRY

The principles o mass spectroscopy and its use to determine relative atomic masses have already been explained in Topic 2 *Atomic structure.* It can be used in a similar way with organic compounds. However in addition to giving the precise molecular mass o the substance, mass spectroscopy gives considerable in ormation about the actual structure o the compound rom the ragmentation patterns.

When a sample is introduced into the machine the vaporized sample becomes ionized to orm the molecular ion $M^+(g)$. Inside the mass spectrometer some o the molecular ions break down to give ragments, which are also de ected by the external magnetic field and which then show up as peaks on the detector. By looking at the di erence in mass rom the parent peak it is o ten possible to identi y particular ragments,

e.g. $(M_r - 15)^+ = \text{loss o } \text{CH}_3$

 $(M_{\rm r} - 17)^+ = {\rm loss \ o \ OH}$

 $(M_{\rm r} - 18)^+ = {\rm loss \ o \ H_2O}$

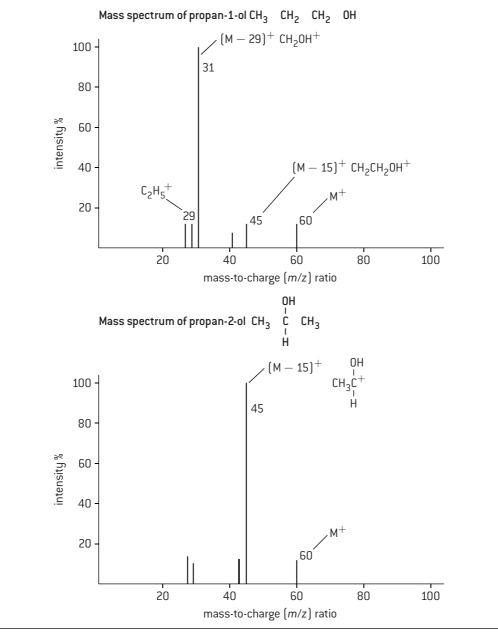
 $(M_{\rm r} - 28)^+ = {\rm loss \ o \ C_2 H_4} {\rm or \ CO}$

 $(M_r - 29)^+ = \text{loss o } C_2 H_5 \text{ or CHO}$

- $(M_{\rm r} 31)^+ = {\rm loss \ o \ CH_{3}O}$
- $(M_r 45)^+ = \text{loss o COOH}$

This can provide a use ul way o distinguishing between structural isomers.

The mass spectra o propan-1-ol and propan-2-ol both show a peak at 60 due to the molecular ion $C_3H_8O^+$. However, the mass spectrum o propan-1-ol shows a strong peak at 31 due to the loss o $-C_2H_3$, which is absent in the mass spectrum o propan-2-ol. There is a strong peak at 45 in the spectrum o propan-2-ol as it contains two di erent methyl groups, which can ragment.



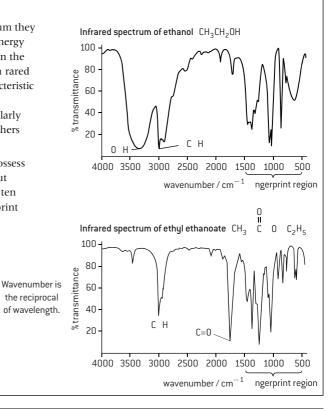
Spectroscopic identification o organic compounds IR and ¹H NMR

INFRARED SPECTROSCOPY

When molecules absorb energy in the in rared region o the spectrum they vibrate (i.e. the bonds stretch and bend). The precise value o the energy they absorb depends on the particular bond and to a lesser extent on the other groups attached to the two atoms orming the bond. When in rared radiation is passed through a sample the spectrum shows the characteristic absorptions, which confirm that a particular bond is present in the molecule. Some absorptions, e.g. those due to C-H, are not particularly use ul, as they are shown by most organic compounds. However others give a very clear indication o a particular unctional group.

In addition to these particular absorptions, in rared spectra also possess a fingerprint region. This is a characteristic pattern between about 1400 400 cm⁻¹, which is specific to a particular compound. It is o ten possible to identi y an unknown sample by comparing the fingerprint region with a library o spectra o known compounds.

A simplified correlation chart		
Bond	Wavenumber / cm 1	
C-Cl	600 800	
C-0	1050 1410	
C=C	1620 1680	
C=O	1700 1750	
C=C	2100 2260	
O-H (in carboxylic acids)	2500 3000	
С-Н	2850 3090	
O-H (in alcohols)	3200 3600	



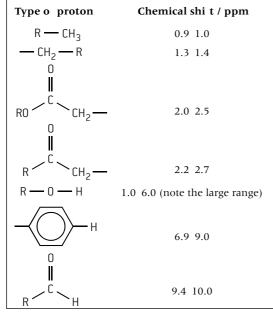
PROTON NUCLEAR MAGNETIC RESONANCE SPECTROSCOPY (¹H NMR)

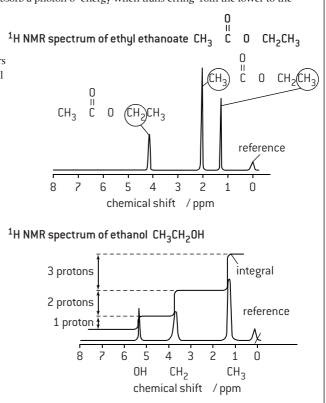
Whereas in rared spectroscopy gives in ormation about the types o bonds in a molecule, 'H NMR spectroscopy provides in ormation on the chemical environment o all the hydrogen atoms in the molecule. The nuclei o hydrogen atoms possess spin and can exist in two possible states o equal energy. I a strong magnetic field is applied the spin states may align themselves either with the magnetic field, or against it, and there is a small energy di erence between them. The nuclei can absorb a photon o energy when trans erring rom the lower to the higher spin state. The photon s energy is very small and occurs in the

radio region o the spectrum. The precise energy di erence depends on the chemical environment o the hydrogen atoms.

The position in the ¹H NMR spectrum where the absorption occurs or each hydrogen atom in the molecule is known as the chemical shi t, and is measured in parts per million (ppm). The area under each peak corresponds to the number o hydrogen atoms in that particular environment.

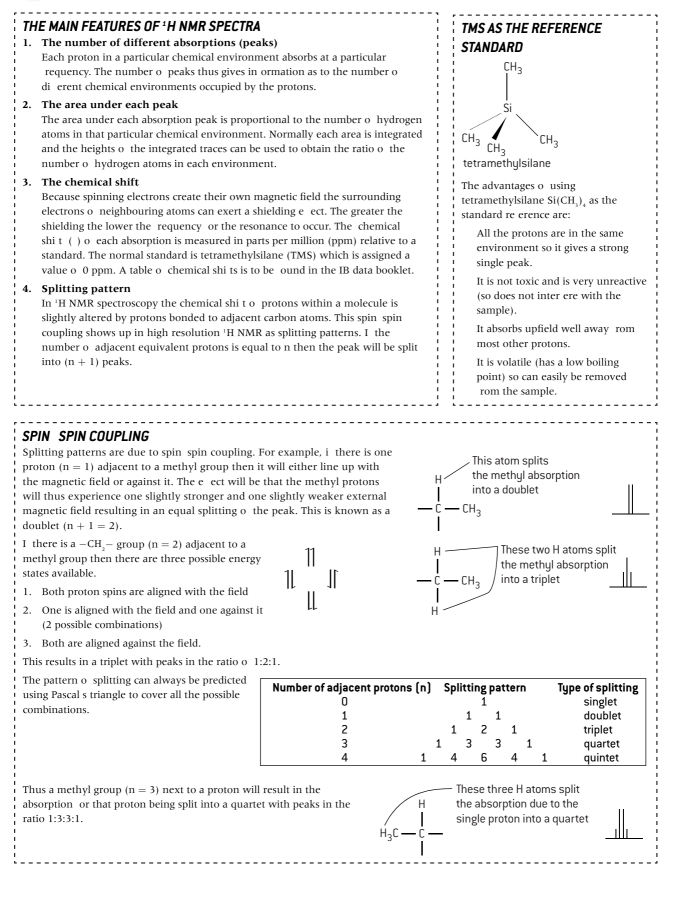
A simplified correlation chart



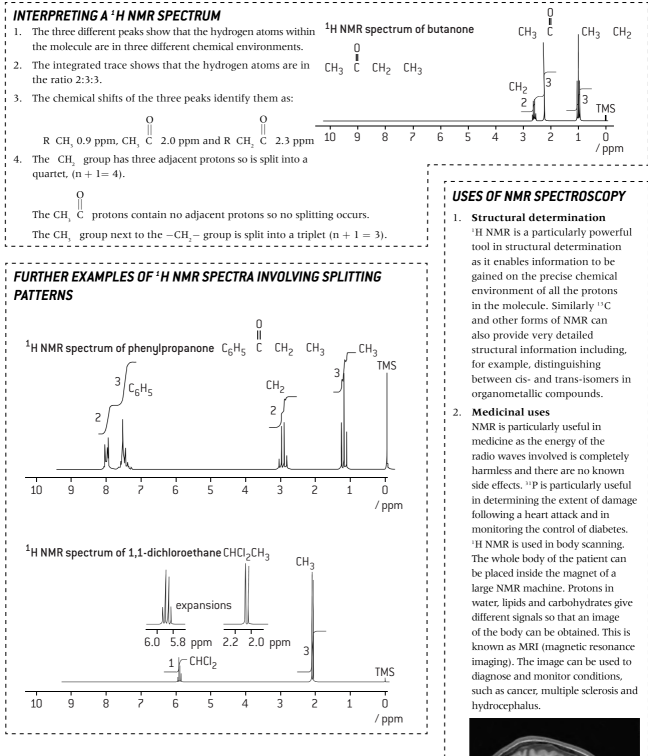


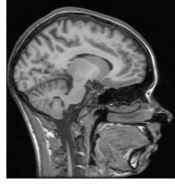
The additional trace integrates the area under each peak. The height of each section is proportional to the number of hydrogen atoms in each chemical environment.

ம Nuclear magnetic resonance (NMR) spectroscopy



Applications of ¹H NMR spectroscopy

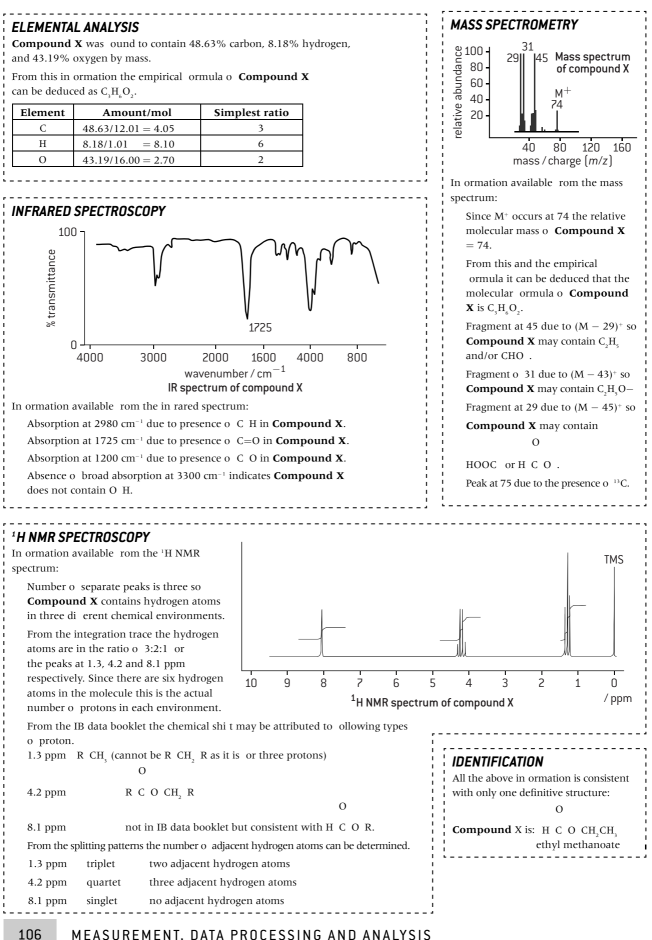




An MRI image of the human brain

🕕 Combination o different analytical techniques to determine structure

The determination o the organic structure o an unknown compound is usually achieved by combining the in ormation rom several di erent analytical techniques. This is illustrated by the ollowing worked example or **Compound X**.



MEASUREMENT, DATA PROCESSING AND ANALYSIS

🖽 X-ray crystallography

X-RAY CRYSTALLOGRAPHY

Atoms or ions can never be seen in the conventional sense as atomic radii and bond lengths are smaller than the wavelength o visible light. However X-rays, which have much smaller wavelengths than visible light, can be used to determine the molecular and atomic structure o crystals where the particles are arranged in a regular array. I the distance between two layers in a crystal

incident rays

structure is d, then by relatively simple mathematics it can be seen that an incident ray hitting the sur ace o a plane at an angle has to travel a distance equal to 2d sin urther i it is di racted rom the layer beneath. For the X-rays to remain in phase the extra distance must be equal to a whole number o wavelengths. This gives what is known as the Bragg equation:

 $n = 2d \sin (where n is an integer)$

By measuring the angles and intensities o these di racted beams a threedimensional picture o the position o

all the atoms within the crystal can be

obtained. This definitively gives the distances between all the atoms (bond lengths) and the bond angles and hence the unambiguous molecular or ionic structure. As well as or simple molecules such as ice, X-ray crystallography has been used to determine the structure o complex molecules such as DNA and proteins. Although the structure o compounds can be deduced rom a combination o spectroscopic techniques, X-ray crystallography is still the main method used to confirm the exact structure o a new compound.

d sin

MULTIPLE CHOICE QUESTIONS MEASUREMENT, DATA PROCESSING AND ANALYSIS

- 1. A 25.0 cm³ sample o a base solution o unknown concentration is to be titrated with a solution o acid o unknown concentration. Which o the ollowing technique errors would give a value or the concentration o the base that is too high?
 - I. The pipette that is used to deliver the base solution is rinsed only with distilled water be ore delivering the sample to be titrated.
 - II. The burette that is used to measure the acid solution is rinsed with distilled water but not with the solution o the titration.
 - A. I only C. Both I and II
 - B. II only D. Neither I nor II
- 2. In a school laboratory, which o the items listed below has the greatest relative uncertainty in a measurement?
 - A. A 50 cm³ burette when used to measure 25 cm³ o water
 - B. A 25 cm³ pipette when used to measure 25 cm³ o water
 - C. A 50 cm³ graduated cylinder when used to measure 25 cm³ o water
 - D. An analytical balance when used to weigh 25 g o water
- 3. A piece o metallic indium with a mass o 15.456 g was placed in 49.7 cm³ o ethanol in a graduated cylinder. The ethanol level was observed to rise to 51.8 cm³. From these data, the best value one can report or the density o indium is

A. 7.360 g cm ⁻³	C.	1.359	$10^{-1} \text{ g cm}^{-3}$
B. 7.4 g cm ⁻³	D.	32.4 g c	rm ⁻³

4. A mixture o sodium chloride and potassium chloride is prepared by mixing 7.35 g o sodium chloride with 6.75 g o potassium chloride. The total mass o the salt mixture should be reported to ____ significant figures; the mass ratio o sodium chloride to potassium chloride should be reported _ significant figures, and the di erence in mass to between sodium chloride and potassium chloride should significant figures. The numbers be reported to required to fill the blanks above are, respectively,

re ected rays

A. 4, 3, 2	C. 3, 3, 1
B. 4, 2, 2	D. 4, 3, 1

d sin

- 5. Repeated measurements o a quantity can reduce the e ects o
 - A. both random and systematic errors
 - B. neither random nor systematic errors
 - C. only systematic errors
 - D. only random errors
- 6. A 50.0 cm³ pipette with an uncertainty o 0.1 cm³ is used to measure 50.0 cm³ o 1.00 0.01 mol dm⁻³ sodium hydroxide solution. The amount in moles o sodium hydroxide present in the measured volume is

A. 0.0500	0.0010
B. 0.0500	0.0001
C. 0.0500	0.0012
D. 0.0500	0.0006

7. Consider the ollowing three sets each o five measurements o the same quantity which has an accurate value o 20.0.

I.	II.	III.
19.8	19.2	20.0
17.2	19.1	19.9
18.3	19.3	20.0
20.1	19.2	20.1
18.4	19.2	20.0

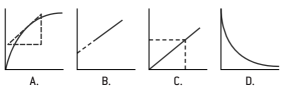
Which results can be described as precise?

A. I, II and III	C. II and III
B. II only	D. III only

 A thermometer with an accuracy o
 C was used to record an initial temperature o
 C and a final temperature o
 C. The temperature rise was

A. 9.6	0.4 C	C. 10 C
B. 9.6	0.2 C	D. 9.6 0.1 C

- An experiment to determine the molar mass o solid hydrated copper(II) sul ate, CuSO₄.5H₂O gave a result o 240 g. The experimental error was
 - A. 0.04%C. 10%B. 4%D. 50%
- 10. Which sketch graph shows interpolation to find an unknown value.





15. The 'H NMR spectrum o a particular compound shows three separate peaks. The relative height o each integration trace and the splitting patterns are

Peak number	Integration trace	Splitting pattern
1	3	triplet
2	2	quartet
3	1	singlet

Which compound could give this spectrum?

A. HCOOCH ₂ CH ₃	C.	$CH_{3}CH_{2}COOCH_{3}$
--	----	-------------------------

- B. CH₃CH₂CH₂COOH D. CH₃CH(OH)CH₃
- 16. Which are advantages or using tetramethylsilane as a standard re erence in ¹H NMR?
 - I. All its protons are in the same chemical environment.
 - II. It is chemically unreactive.
 - III. It does not absorb energy in the same region as most other protons.
 - A. I and II only C. II and III only
 - B. I and III only D. I, II and III

- 11. What in ormation can be obtained rom the number o peaks in the 'H NMR spectrum o a compound?
 - A. The number o di erent chemical environments occupied by the protons in one molecule o the compound.
 - B. The number o hydrogen atoms in one molecule o the compound.
 - C. The number o di erent unctional groups in one molecule o the compound.
 - D. The number o carbon atoms in one molecule o the compound.
- 12. What is the index o hydrogen deficiency (IHD) or a compound with the molecular ormula C₂H₈?

- B. 4 D. 8
- 13. The mass spectrum o a compound shows peaks with *m*/*z* values o 88, 73, 59 and 43. Which compound could give this spectrum?
 - A. HCOCH₂CH₂CH₃ B. CH₂CH₂COOCH₂CH₃
- D. CH,CH,CH,COOH

C. CH(CH₂),COOH

6

14. Which is correct about the regions o the electromagnetic spectrum?

_	Region of spectrum	Energy	Wavelength
А.	Ultraviolet	Highest	Longest
В.	In rared	Highest	Shortest
C.	Radio	Lowest	Longest
D.	Visible	Lowest	Shortest

- 17. Which compound will contain a peak with a triplet splitting pattern in its 'H NMR spectrum?
 - A. CH₃OH C. CH₃CHClCOOH
 - B. CH₃CH₂COOH D. (CH₃)₃COH
- 18. Which can be determined by single crystal X-ray crystallography?
 - I. Bond lengths
 - II. Bond angles

A. I and II only

B. I and III only

- III. Chemical structure
- C. II and III only
- D. I, II and III

SHORT ANSWER QUESTIONS MEASUREMENT, DATA PROCESSING AND ANALYSIS

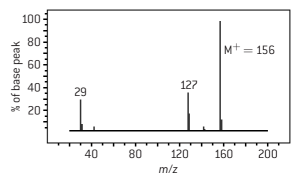
1. A simulated computer-based experiment was per ormed to determine the volume o nitrogen generated in an airbag rom the decomposition o sodium azide, NaN₃.

 $2NaN_3(s) = 2Na(s) + 3N_3(g)$

The ollowing data were entered into the computer.

	Mass o NaN ₃ (s)/ kg(0.001 kg)	Temperature / C (0.50 C)	Pressure / kPa l kPa]
	0.072	20.00	106	-
	a) State the number temperature and	r o significant figure l pressure data.		[1]
	b) Calculate the am present.	iount (in mol) o sod		[2]
	c) Calculate the percentage uncertainty or each o the mass, temperature and pressure. [1]		[1]	
	 d) Determine the volume o nitrogen gas (in dm³) produced under these conditions together with its uncertainty assuming complete decomposition o the sodium azide. 		[5]	
2.	contains 15.40%	sis o Compound X s o carbon, 3.24% hyd Determine the empir	rogen and 81.36%	

b) The mass spectrum o Compound **X** is shown below.



- (i) Deduce the molecular ormula o Compound X. [1]
- (ii) Calculate the index o hydrogen deficiency (IHD) or Compound X. [1]
 (iii) Identi y the ragments responsible or the peaks
- with m/z values o 127 and 29. [2]

[2]

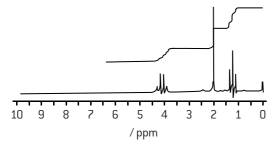
(iv) Deduce the identity o Compound **X**. [1]

- c) The 'H NMR spectrum or Compound **X** shows two peaks with integration traces o two units and three units respectively. Explain how this in ormation is consistent with your answer to (b) (iv).
- ¹H NMR spectroscopy can be used to distinguish between pentan-2-one and pentan-3-one. In each case state the number o peaks in their ¹H NMR spectra and state the ratios o the areas under each peak. [4]

HL

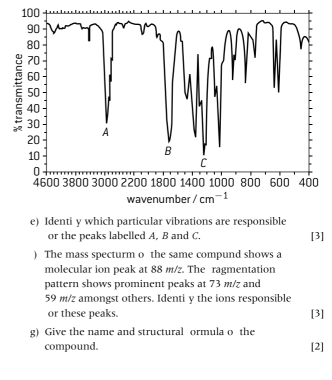
Compound X.

4. The 'H NMR spectra o an unknown compound is given below:



- a) Why is tetramethylsilane used in 'H NMR spectroscopy?
- b) What word is used to describe the multiplicity o the peaks centred at 4.1 ppm?
- c) What is the ratio o the number o hydrogen atoms responsible or the chemical shi ts centred at 1.2, 2.0 and 4.1 ppm respectively?
- d) Two structures that have chemical shi ts centred at 4.1 ppm are RCOOCH₂R and C₆H₅OCOCH₃. From consideration o the rest o the spectra only one o these general structures is possible. Identi y which one and explain your reasoning.

Below is the in rared spectrum o the same compound:



[1]

[1]

[1]

[1]

[2]

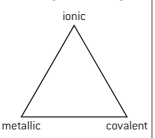
12 OPTION A MATERIALS

Introduction to materials science

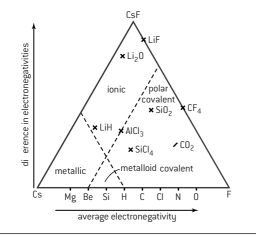
CLASSIFICATION OF MATERIALS AND TRIANGULAR BONDING DIAGRAM

Civilizations have been characterised by the materials they used, e.g. Stone Age, Iron Age, Bronze Age and now perhaps the Plastics Age. Materials can be classi ed in several di erent ways based upon their properties, their uses or on the type o bonding or structure they exhibit. Materials science investigates the relationship between the bonding and structures o materials on the atomic or molecular level and relates this to their macroscopic properties. The key to understanding the macroscopic properties is thus the identity o the type or types o bonding present in the material. Metallic substances tend to have very di erent properties to non-metals and the properties o ionic compounds di er greatly to covalent compounds. The degree o covalent, ionic or metallic character in a compound can be deduced rom its position on a triangular bonding diagram also known as a Van Arkel Ketelaar diagram. Bond triangles show the three extremes o bonding in terms o electronegativity di erence (ionicity) plotted against the average electronegativity (localization) or binary compounds with the three extreme cases o types o bonding ionic, covalent and metallic at the apices o an equilateral triangle.

These three extreme cases are characterized by caesium (metallic), fluorine (covalent) and caesium fluoride (ionic). Other materials can be placed around the triangle based on their di erences in electronegativities. The bottom side o the triangle, rom metallic



(low electronegativity) to covalent (high electronegativity) is or elements and compounds with varying degree o delocalization and directionality in the bond. Pure metals with only delocalized bonding occur on the extreme let (metallic corner) and at the other extreme are covalent compounds where all the electron orbitals overlap in a particular direction. The let side o the triangle (rom ionic to metallic) is or delocalized bonds with varying electronegativity di erence and the right side shows the change rom covalent to ionic. The diagram below shows that compounds such as silicon tetrachloride and carbon dioxide are very much covalent whereas silicon dioxide is polar covalent but a compound such as aluminium chloride lies very close to the border between covalent and ionic. Similarly lithium oxide has a high degree o ionic character whereas lithium hydride lies close to the border o ionic and metallic.



COMPOSITES

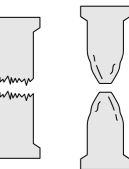
Composites are made rom two or more materials with signi cantly di erent properties which when combined produce a material with di erent characteristics (e.g. lighter and stronger) to its constituent parts. The composite consists o two distinct phases, a rein orcing phase which is embedded in a matrix phase. Common composites include building materials such as concrete and cement and rein orced plastics and ceramics.

RELATIONSHIPS BETWEEN PHYSICAL CHARACTERISTICS AND BONDING AND STRUCTURE

Melting point. Melting points ollow a similar trend to boiling points in that the weaker the attractive orces between the particles the lower the melting point. But they are also influenced by the type o packing between the particles in the solid state. Thus the melting point o metals can vary rom quite low (caesium melts at 28.5 C) to very high (copper melts at 1085 C) even though they all contain delocalized electrons. Giant molecular covalent substances such as diamond and silicon dioxide have high melting points, whereas simple covalent molecules such as hydrogen gas and carbon dioxide have low melting points.

Conductivity. Materials containing delocalized electrons such as metals and graphite are good conductors o heat and electricity. Covalent compounds containing electrons localized in xed orbitals are poor conductors. Ionic compounds can only conduct electricity when their ions are ree to move in the molten state or in aqueous solution but are chemically decomposed in the process (see page 30).

Brittleness. Many metals are malleable (the opposite o brittle) and ductile (can be drawn into a wire) as the close-packed layers o positive ions can slide over each other without breaking more bonds than are made. Impurities added to the metal disturb the lattice and so make the metal less malleable, which



brittle fracture

ductile fracture

explains why most alloys are harder than the pure metal they are made rom (see page 31). Many covalent substances such as sul ur are brittle, that is they racture when subjected to stress rather than undergo de ormation as the bonds cannot be re ormed once broken. Because brittle substances do not undergo de ormation when ractured the two parts can o ten be glued together to return to the original shape.

Permeability is de ned as the ability o a porous material to allow fluids such as water to pass through it. This depends both upon the type o bonding in the material, which can attract or repel polar water molecules, and the spaces between the particles in the lattice.

Elasticity is the tendency o a solid material to return to its original shape a ter it has been de ormed by the application o a orce. In polymers such as rubber the polymer chain can be stretched without being broken when orces are applied. In metals the atoms can slide over each other so that the lattice changes size and shape and can return to the original shape when the orce is removed.

Principles of extraction of metals from their ores

EXTRACTION OF METALS FROM THEIR ORES

The essential process involved in the extraction o metals orm their ores is reduction. The ease with which this process occurs depends upon the position o the metal in the activity series. Metals very low in the activity series, such as gold, silver and copper may be ound uncombined. Metals slightly higher in the series can be obtained either by simply heating the ore strongly or by using chemical reducing agents, such as more reactive metals, carbon or carbon monoxide. Generally metals that can be obtained by relatively simple chemical reduction have been known since ancient times. It is much harder and more expensive to reduce chemically the more reactive metals and it took the discovery o electricity in the 19th century be ore these could be produced commercially. It is still an expensive process as molten ores must be used. To reduce the energy required, and hence the cost o production, impurities are o ten added to the molten electrolyte to lower the melting point.

Activity series	Main ore	Method of extraction	
Sodium	NaCl	Electrolysis o molten NaCl with CaCl ₂ added to lower the melting point.	
		$Na^+(l) + e^- Na(l)$	
Aluminium	Al ₂ O ₃	Electrolysis o Al ₂ O ₃ in molten cryolite.	
		$Al^{3+}(l) + 3e^{-} Al(l)$	
Zinc	ZnS	Roast to orm the oxide	
		$2ZnS(s) + 3O_2(g) = 2ZnO(s) + 2SO_2(g)$	
		then reduce chemically with carbon monoxide	
		$ZnO(s) + CO(g) = Zn(s) + CO_2(g)$	
		or electrolytically $Zn^{2+}(l) + 2e^{-} Zn(l)$	
Iron	Fe ₂ O ₃	Heat with carbon monoxide	
		$Fe_2O_3(s) + 3CO(g) = 2Fe(l) + 3CO_2$	
Lead	PbS	Heat to orm the oxide then reduce with carbon	
		PbO(s) + C(s) = Pb(l) + CO(g)	
Copper	Cu or CuFeS ₂	Heat in air to give copper and sul ur dioxide	
Gold	Au	Metal ound uncombined	

Methods of extracting some important metals

PRODUCTION OF ALUMINIUM

Aluminium is primarily made by the electrolytic reduction o aluminium oxide.

The main ore o aluminium is bauxite. The aluminium is mainly in the orm o the hydroxide Al(OH)₃ and the principal impurites are iron(III) oxide and titanium hydroxide. The impurities are removed by heating powdered bauxite with sodium hydroxide solution. The aluminium hydroxide dissolves because it is amphoteric.

 $Al(OH)_3(s) + NaOH(aq)$ $NaAlO_2(aq) + 2H_2O(l)$

The aluminate solution is ltered leaving the impurities behind. Seeding with aluminium hydroxide then reverses the reaction. The pure recrystallized aluminium hydroxide is then heated to produce aluminium oxide (alumina).

 $2Al(OH)_3(s) = Al_2O_3(s) + 3H_2O(l)$

In a separate process hydrogen fluoride is added to the aluminate solution ollowed by sodium carbonate to precipitate cryolite (sodium hexafluoroaluminate(III), Na_3AlF_6).

 $NaAlO_2(aq) + 6HF(g) + Na_2CO_3(aq)$ $Na_3AlF_6(s) + 3H_2O(l) + CO_2(g)$

The electrolysis o molten alumina takes places in an open-topped steel container lined with graphite. Alumina has a melting point o 2045 C so it is mixed with cryolite. The resulting solution melts at about 950 C so that much less energy is required. The aluminium is produced on the graphite lining which acts as the negative electrode (cathode). Molten aluminium is more replaced in turn dense than cryolite so it collects at the bottom o the cell Al_2O_3 carbon electrodes [+] as they oxidize

where it can be syphoned o periodically.

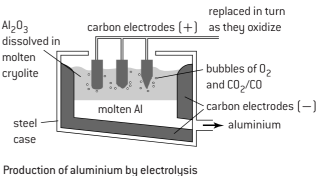
$$Al^{3+}(l) + 3e^{-} Al(l)$$

The positive electrode is made o blocks o graphite. As the oxide ions are oxidized some o the oxygen ormed reacts with the graphite blocks so that they have to be renewed regularly.

 $2O^{2-}(l) = O_{2}(g) + 4e^{-1}$

$$C(s) + O_2(g) = CO_2(g)$$

A modern cell can produce up to two tonnes o aluminium per day.



Faraday calculations and properties and analysis of alloys

DETERMINATION OF THE MASS OF METAL PRODUCED BY ELECTROLYSIS

The amount (in mol) o a metal produced by electrolysis depends upon the charge o electricity received by the molten electrolyte and the charge on the metal ion. Three times as much charge o electricity will be required to produce one mole o aluminium rom Al^{3+} ions compared with one mole o sodium rom Na^+ ions. The charge passed through the electrolyte is equal to the size o the current flowing multiplied by the length o time it flows. A charge o one coulomb is produced by a current o one amp flowing or one second. The charge carried by one mole o electrons is equal to 96 500 coulombs and this value is known as one araday (F).

Example 1: Calculate the mass o sodium produced at the negative electrode when a steady current o 15.2 A passes through a solution o molten sodium chloride or 3 hours 20 minutes.

Charge passed = $15.2 \quad 200 \quad 60 = 182 \; 400 \; C$ = $\frac{182 \; 400}{96 \; 500} = 1.89 \; F$

 $Na^{+}(l) + e^{-}$ Na (l) so one araday required to produce one mole o sodium.

Amount o sodium produced = 1.89 mol

Mass o sodium produced = 1.89 22.99 = 43.5 g

ALLOYS

Alloys are homogeneous mixtures o metals with other metals or non-metals. They have already been discussed together with some examples on page 31. Transition metals o ten orm alloys with each other as their atoms have similar radii and the crystal structure is not seriously disrupted. The addition o other elements into the metallic structure alters the physical properties. Some properties such as density and electrical conductivity may not di er much rom the constituent elements but others such as tensile strength and melting point may di er considerably. Steel is an alloy o iron, carbon and other metallic and non-metallic elements. It has a wide range o uses and by adjusting its composition it can be tailor-made with speci c properties. For example, the addition o chromium increases the resistance o steel to corrosion. Stainless steel, used or kitchen knives and sinks contains about 18% chromium and 8% nickel. Toughened steel or use in drill bits, which need to retain a sharp cutting edge at high temperatures, contains up to 20% molybdenum.

Example 2: A typical aluminium cell produces 2000 kg o aluminium in 24 hours. Determine the average current that must be passed to achieve this.

2000 kg = 2000 $\frac{10^3}{26.98}$ = 7.413 10⁴ mol Al³⁺(l) + 3e⁻ Al(l) Charge required = 3 7.413 10⁴ 96 500 = 2.146 10¹⁰ C Average current required or 24 hours = $\frac{2.146}{(24 \ 60 \ 60)}$ = 2.48 10³ A

DIAMAGNETISM AND PARAMAGNETISM

Metals and their compounds can exhibit di erent types o magnetism. Iron metal and some other metals (e.g. nickel and cobalt) show erromagnetism. This is a permanent type o magnetism. In this type o magnetism unpaired electrons align parallel to each other in domains irrespective o whether an external magnetic or electric eld is present. This property o iron has been utilized or centuries to make compasses which align with the Earth s magnetic eld to point north.

Many other materials contain unpaired electrons. Unlike paired electrons, where the spins cancel each other out, the spinning unpaired electrons create a small magnetic eld and will line up in an applied electric or magnetic eld to make the material weakly magnetic when the eld is applied, i.e. they rein orce the external magnetic eld. This type o magnetism is known as **paramagnetism**. The more unpaired electrons there are in the material the more paramagnetic the material will be.

When all the electrons in a material are paired up the complex is said to be **diamagnetic**. Essentially the material is non-magnetic although the paired electrons do create a very small magnetic eld in opposition to an externally applied eld. Because the e ect is so small, it is not observable in everyday li e except or superconductors. Superconductivity occurs when certain materials are cooled below a characteristic critical temperature, which tends to be close to absolute zero.

DETECTION OF TRACE AMOUNTS OF METALS (ICP OES AND ICP MS)

Very small amounts o trace metals in alloys and other materials can be detected and determined quantitatively by Inductively Coupled Plasma (ICP) Spectroscopy together with either Optical Emission Spectroscopy (ICP OES) or Mass Spectroscopy (ICP MS) . Plasma is the ourth state o matter (the other three being solid, liquid and gas). A plasma contains charged particles and can be induced by a strong electromagnetic eld. In ICP an intense electromagnetic

eld is created in a torch , which ionizes argon gas to orm a high-temperature plasma. The sample o material to be analysed is introduced directly into the plasma flame where it is broken down into charged ions to release radiation o characteristic wavelengths, which can be detected in the optical spectrometer part o ICP OES. By calibrating the instrument with speci c amounts o known metals the values o the wavelengths emitted enable each metal to be identi ed. Since the concentration is proportional to the intensity o the peaks the concentration o each metal can also be determined by comparing the intensity o its peaks with those made by samples o known concentration. In ICP MS the released ions are identi ed using a mass spectrometer rather than an optical spectrometer. As well as identi ying the amounts o metals in alloys, these two techniques also have many uses in medicine and orensic science (e.g. heavy metal poisoning).

Catalysts

HOMOGENEOUS AND HETEROGENEOUS CATALYSIS

Catalysts unction by providing an alternative reaction pathway with a lower activation energy so that more o the reactant particles will possess the necessary minimum energy to react when they collide. Catalysts may be in the same phase as the reactants (usually the same phase as the product(s) too) in which case they are known as **homogeneous catalysts**. A phase is similar to a state except that there is a physically distinct boundary between two phases. It is possible to have a single state but two phases. An example is two immiscible liquids such as oil and water. I the catalyst is in a di erent phase to the reactants then it is unctioning as a **heterogeneous catalyst**.

CHOICE OF CATALYST

The advantage o a homogeneous catalyst compared with a heterogeneous catalyst is that all o the catalyst is exposed to the reactants whereas in heterogeneous catalysis the e ficiency o the catalyst is dependent upon the sur ace area. The disadvantage o a homogenous catalyst is that it is usually harder to remove the catalyst a ter the reaction whereas a heterogeneous catalyst can be relatively easily removed by filtration. The most e ficient catalysts o all are enzymes biological catalysts but these are very specific or a particular biological reaction due to the shape o the active site. When choosing a catalyst or a chemical reaction the ollowing actors should be considered.

Selectivity	Will the catalyst produce only the desired product?
E ficiency	Will the catalyst cause a considerable increase in the rate? Will it continue to work well under severe conditions, such as those experienced by catalytic converters in cars, as well as mild conditions?
Environmental impact	Will it be easy to dispose o the catalyst without causing harm to the environment? Many transition metals are classed as heavy metals and can cause problems i they enter the soil or ground water.
Potential or poisoning	Catalysts rely on reactants occupying the active site reversibly. A poison will occupy the active site irreversibly so blocking access to reactants. Poisons include carbon monoxide, cyanide ions and sul ur.
Cost	Industry is profit-based. Many transition metals such as rhodium, platinum and palladium are expensive and the cost to benefit ratio needs to be care ully calculated.

MODES OF ACTION OF CATALYSTS

Although homogenous catalysts can be recovered chemically unchanged at the end o the reaction they can orm intermediate compounds during the reaction.

For example when nitrogen monoxide catalyses the oxidation o sul ur dioxide it is thought that nitrogen dioxide is ormed as an intermediate compound.

 $2NO(g) + O_2(g) = 2NO_2(g)$

This is a redox reaction and the oxidation state o nitrogen has increased rom +2 to +4. In the second step the nitrogen dioxide is reduced back to nitrogen monoxide by the sul ur dioxide.

 $2SO_{2}(g) + 2NO_{2}(g) = 2SO_{3}(g) + 2NO(g)$

So that nitrogen monoxide is unchanged at the end o the reaction and the overall equation is

 $2SO_2(g) + O_2(g) = 2SO_3(g)$

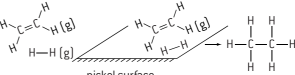
Because transition metal ions can exist in more than one oxidation state they tend to make good homogenous catalysts. In the reaction between peroxodisul ate(VI) ions and iodide ions it is thought that the $Fe^{2+}(aq)$ ions are oxidized to $Fe^{3+}(aq)$ ions and then back to $Fe^{2+}(aq)$ ions.

 $S_2O_8^{2-}(aq) + 2Fe^{2+}(aq) = 2SO_4^{2-}(aq) + 2Fe^{3+}(aq)$

 $2I^{-}(aq) + 2Fe^{3+}(aq) = 2Fe^{2+}(aq) + I_{2}(aq)$

Heterogeneous catalysts tend to unction by adsorbing reactant molecules onto the sur ace o the catalyst (the active site) and bringing them into close contact with each other in the correct orientation. Many transition metals and their compounds have the ability to physically adsorb large amounts o gases on their sur ace which makes them particularly good heterogeneous catalysts. For example nickel or palladium can adsorb ethene and hydrogen so that they can react to orm ethane.

The reaction of ethene with hydrogen to form ethane



nickel surface

Without a catalyst a random collision with the necessary kinetic energy and correct orientation is required ethane The reactant molecules are brought together with the correct orientation by adsorption on the surface of the catalyst

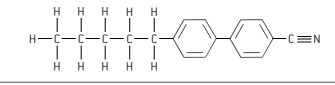
Two particular types o heterogeneous catalysts are zeolites and carbon nanotube support catalysts. Zeolites are microporous cage-like aluminosilicate compounds with complex ormulas. They occur naturally and can also be made synthetically. They work by adsorbing reactant molecules and confining them in a small space so they can react together. They are particularly used in the petrochemical industry, or example as catalysts or cracking larger hydrocarbons into smaller more use ul hydrocarbons and alkenes. Carbon nanocatalysts involve using carbon nanotubes (rather than silicon or alumina) as support or metals or metal oxide catalysts. Due to their small size the nanotubes provide a large sur ace area and other properties such as excellent electron conductivity and chemical inertness help to promote the action o the catalyst and make it much more e ficient.

Liquid crystals (1)

LIQUID CRYSTALS

Liquid crystals are a phase or state o matter that lies between the solid and liquid state. In a liquid crystal the molecules tend to retain their orientation as in a solid but they can also move to di erent positions as in a liquid. The physical properties o liquid crystals (such as electrical conductivity, optical activity and elasticity) depend upon the orientation o the molecules relative to some xed axis in the material. Examples o substances which can behave as liquid crystals under certain conditions include DNA, soap solution, graphite and cellulose together with some more specialized substances such as biphenyl nitriles.

An example o a biphenyl nitrile is 4-pentyl-4 -cyanobiphenyl (known as 5CB)



LYOTROPIC AND THERMOTROPIC LIQUID CRYSTALS

Liquid crystals only show liquid crystal properties under certain conditions. They are very sensitive to temperature and concentration. Essentially liquid crystals can be divided into two main types.

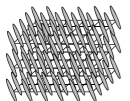
Lyotropic liquid crystals are solutions that show the liquid crystal phase at certain concentrations. An example o lyotropic liquid crystals is soap solution. At low dilution the polar soap molecules behave randomly but at higher concentrations they group together into larger units called micelles which in the liquid crystal phase are ordered in their orientation.

Thermotropic liquid crystals are pure substances that show liquid crystal behaviour over a range o temperature between the solid and liquid states. Examples o a thermotropic liquid crystal are biphenyl nitriles used in liquid crystal displays (LCDs).

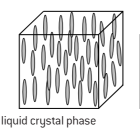


The use o thermotropic liquid crystals in a calculator screen

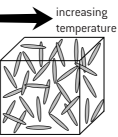
Within the thermotropic liquid crystal phase the rod-shaped molecules which are typically about 2.5 10^{-9} metres in length exist in groups or domains. The molecules can flow and are randomly distributed as in a liquid but within each domain they all point in the same direction. This is known as the **nematic phase**. As the temperature increases the orientation becomes increasingly more disrupted until eventually the directional order is lost and the normal liquid phase is ormed.



solid state molecules held in rigid 3-D structure



Inquid crystal phase molecules within each domain oriented in the same direction



liquid crystal state molecules all randomly distributed

PRINCIPLES OF LCD DEVICES

Since liquid crystal molecules are polar their orientation can be controlled by an applied electric eld. The orientation o the molecules a ects the ability o the liquid crystal molecules to transmit light. In liquid crystal displays used in digital watches, calculators and laptops a small voltage is applied across a thin lm o the material.



The flat screens used or computer monitors or televisions use liquid crystals

This controls the areas o the display that are light and dark and hence gives the characteristic readings o the pictures or letters. The great advantage o LCDs over other types o electronic display is that they use extremely small electric currents. The disadvantage is that they only work within a certain temperature range which explains why a digital watch or laptop screen may give a strange display in very hot or cold temperatures. Hence or use in an LCD a liquid crystal should:

be a chemically stable compound contain polar molecules

remain stable in the liquid crystal phase over a suitable range o temperature

be able to orientate quickly (rapid switching speed).

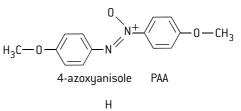
Liquid crystals (2)

STRUCTURAL FEATURES OF BIPHENYL NITRILES

When the structure o 4-pentyl-4-cyanobiphenyl is compared with other compounds that have good liquid crystal properties it can be seen that they have several eatures in common.

- The nitrile group in the biphenyl nitrile (and the $NN^+(O^-)$ and C=N groups in the other two molecules) is polar. This ensures that the intermolecular orces are strong enough to align in a common direction.
- The two benzene rings in the molecules ensure that the molecules are rigid and there ore more rod-shaped.

The long alkane chain group on the end o the molecule ensures that the molecules cannot pack so closely together and so helps to maintain the liquid crystal state.



4-pentyl-4-cyanobiphenyl

 $C \equiv N$

5CB

$$H_3C - 0 - C_4H_9$$

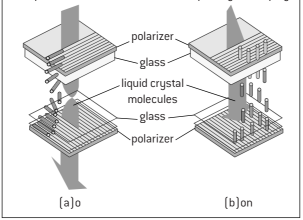
4 -methoxybenzylidene-4-butylaniline

MBBA

THE WORKINGS OF THE LCD DEVICE

In a liquid crystal display each pixel contains a liquid crystal film sandwiched between two glass plates. The plates have many very fine scratches at right angles to each other and have the property o polarizing light. The liquid crystal molecules in contact with the glass line up with the scratches and orm a twisted arrangement between the plates due to intermolecular orces. This is known as twisted nematic geometry. The property utilized by the liquid crystals is their ability to interact with planepolarized light (see page 96) which is rotated through 90 by the molecules as it passes through the film. When the two polarizers are aligned with the scratches, light will pass through the film and the pixel will appear bright. When a potential di erence is applied across the film, the polar molecules will align with the film thus losing their twisted structure and ability to interact with the light. Planepolarized light will now no longer be rotated so that the pixel appears dark.

the operation of the twisted nematic liquid crystal display



Polymers

TYPES OF POLYMERS AND STRUCTURAL FEATURES

Polymers consist o many repeating units (monomers) joined together to orm macromolecules with high molar masses a theory rst proposed by Herman Staudinger in 1920, which has since been substantiated by X-ray di raction and scanning tunnelling electron microscopy. By understanding the molecular structure o polymers, chemists have been able to manipulate their properties and develop new polymers.

Polymers can be subdivided into thermoplastics and thermosets. Many alkenes polymerize to orm thermoplastics that so ten when heated and harden when cooled so that they can be remoulded each time they are heated. Thermosetting polymers, such as bakelite, polyurethanes and vulcanized rubber orm prepolymers in a so t solid or viscous state that change irreversibly into hardened thermosets by curing, so that once shaped they cannot be remoulded. Some polymers, such as rubber, are flexible polymers known as elastomers. Elastomers can be de ormed under orce but will return to nearly their original shape once the stress is released. Generally the longer the chain length o a polymer the higher the strength and melting point, but cross-linking (or branching) and the orientation o the substituent groups can also a ect particular properties.

Branching

Depending on the reaction conditions ethene can orm high density or low density polythene. In high density poly(ethene), HDPE, there is little branching. This gives long chains that can t together closely making the polymer stronger, denser, and more rigid than low density poly(ethene), LDPE. The presence o side chains in low density poly(ethene) results in a more resilient and flexible structure making it ideal or the production o lm products, such as ood wrappings.

Orientation of alkyl groups

In poly(propene) the methyl groups can all have the same orientation along the polymer chain **isotactic**. Due to the regular structure isotactic polymers are more crystalline and tough. Isotactic poly(propene) is a thermoplastic and can be moulded into objects, such as car bumpers, and drawn into bres or clothes and carpets. In **atactic** poly(propene) the chains are more loosely held so the polymer is so t and flexible, making it suitable or sealants and roo ng materials.

Н CH3 CH₃ CH3 CH3 CH3 CH3

isotactic poly(propene) all methyl groups orientated in same direction

CH₃CH₃H CH₃CH₃H CH3 CH3 atactic poly(propene) methyl

groups arranged randomly

MODIFICATIONS TO POLYMERS

Plasticizers

Plasticizers are small molecules that can t between the long polymer chains. They act as lubricants and weaken the attraction between the chains, making the plastic more flexible. By varying the amount o plasticizer added PVC can orm a complete range o polymers rom rigid to ully pliable.

GREEN CHEMISTRY AND ATOM ECONOMY

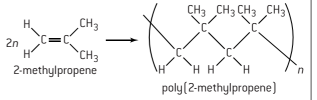
Polymer development and use has grown rapidly in the past 60 years. The environmental impact o the industrial production o polymers is huge. Chemists try to nd ways o reducing this by applying the principles o Green chemistry. One measure o the e ciency o the production is the atom economy which has been explained with an example on page 6. Generally addition polymers tend to have very high atom economies as all the monomers are converted into polymers. Condensation polymerization reactions, such as the ormation o nylon, tend to have lower atom economies.

Volatile hydrocarbons

I pentane is added during the ormation o polystyrene and the product heated in steam the pentane vaporizes producing expanded polystyrene. This light material is a good thermal insulator and is also used as packaging as it has good shock-absorbing properties.

POLYMERIZATION OF 2-METHYLPROPENE

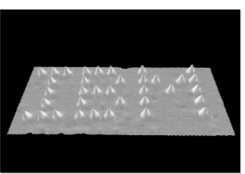
Addition polymers are ormed by the addition reaction o alkenes to themselves. One such example is the polymerization o 2-methylpropene to orm poly (2-methylpropene). The product, which is also known as butyl rubber or polyisobutylene, is a good elastomer and has many uses, such as car tyre inner tubes and cling lm.



Nanotechnology

NANOTECHNOLOGY

Nanotechnology is de ned as the research and technology o compounds within the range o one to one hundred nanometres $(1.0 \quad 10^{-9} \text{ m to } 1.0 \quad 10^{-7} \text{ m})$ in length, i.e. on the atomic scale. It creates and uses structures that have novel properties based on their small size. There are two main approaches. The bottom-up approach involves building materials and devices rom individual atoms, molecules or components. The top-down approach involves constructing nano-objects rom larger entities. Sometimes physical techniques are used that allow atoms to be manipulated and positioned to speci c requirements. For example, a process known as dip-pen nanolithography can be used to place atoms in speci c positions using an atomic orce microscope. It is also possible to use chemical reactions such as in DNA nanotechnology where the speci c base-pairing due to hydrogen bonding can be utilized to build desired molecules and structures.



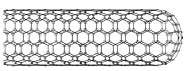
An image o IBM written with xenon atoms on nickel using dip-pen nanolithography and seen using scanning tunnelling microscopy.

IMPLICATIONS OF NANOTECHNOLOGY

Nanotechnology is concerned with the ability to control or manipulate on the atomic scale. It has the potential to solve many problems such as increase ood production, prevent, monitor and cure diseases, and improve in ormation and communication technology, although most o these bene ts still probably lie somewhere ar in the uture. However, little is known about the potential risks associated with developing this technology. The hazards associated with small airborne particles are not properly known or covered by current toxicity regulations. The human immune system may be de enceless against new nano-scale products. There may also be social problems too as poorer societies may su er as established technologies become redundant and demands or commodities change rapidly.

NANOTUBES

Nanotubes are tubes with a diameter in the region o just a ew nanometres. They are made using only carbon atoms. The basic building block is a tube with the walls made rom graphene and a hemisphere o buckminster ullerene to close the ends. Both single and multi-walled graphene tubes, made rom concentric nanotubes, have been ormed. A wide variety o di erent materials, including elements, metal oxides and even small proteins have been inserted inside the tubes. Because they have a greatly increased ratio o sur ace area to volume they can act as extremely e cient and highly selective catalysts. They also have huge tensile strength and good thermal and electrical conductivity due to delocalized electrons.



A diagram o part o a nanotube

Among the methods by which nanotubes can be prepared are arc discharge, chemical vapour deposition (CVD) and high pressure carbon monoxide (HIPCO). Arc discharge, which can give yields o up to 30%, involves either vaporizing the sur ace o one o the carbon electrodes, or discharging an arc through metal electrodes submersed in a hydrocarbon solvent, which orms a small rod-shaped deposit due to oxidation occurring on the anode. Chemical vapour deposition (CVD) is a commonly used method or the commercial production o nanotubes. In this process a carbon-containing gas, such as ethene in the plasma phase due to the influence o a strong electric eld, is decomposed in the presence o an inert carrier gas, such as nitrogen, over a heated substrate ormed rom layers o metal particles. The carbon-containing gas is broken apart at the sur ace o the catalyst particle, and the resulting carbon is transported to the edges o the particles, where it orms the nanotubes. The HIPCO process creates single-walled carbon nanotubes rom the reaction between high pressure carbon monoxide with iron pentacarbonyl, $Fe(CO)_5$ at temperatures between 1273 K and 1473 K. During this process the iron pentacarbonyl reacts to produce iron nanoparticles.

 $Fe(CO)_{5}(g) = Fe(s) + 5CO(g)$

The iron nanoparticles provide a nucleation sur ace or the trans ormation o carbon monoxide into carbon during the growth o the nanotubes.

 $xCO(g) = CNT(s) + xCO_2(g)$

where x is typically 6000 giving a carbon nanotube (CNT) containing 3000 carbon atoms.

Environmental impact

plastics

ADVANTAGES AND DISADVANTAGES OF PLASTICS

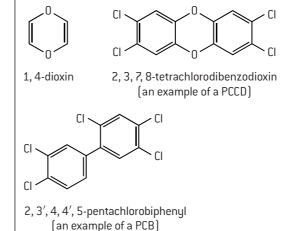
Since the discovery o polymers such as nylon, polystyrene and polyurethanes in the 1930s, society has gained hugely rom the use o plastics. Polymers can be tailor-made to per orm a variety o unctions based on properties such as strength, density, thermal and electrical insulation, flexibility, and lack o reactivity. There are, however, some disadvantages.

- 1. **Depletion of natural resources:** The majority o polymers are carbon-based. Currently oil is the major source o carbon although in the past it was coal. Both are ossil uels and are in limited supply.
- 2. **Disposal:** Because o their lack o reactivity due to strong covalent bonds, plastics are not easily disposed o . Some, particularly PVC and poly(propene), can be recycled and others (e.g. nylon) are weakened and eventually decomposed by ultraviolet light. Plastics can be burned but i the temperature is not high enough poisonous dioxins can be produced along with toxic gases, such as hydrogen cyanide, hydrogen chloride and incomplete hydrocarbon combustion products.
- 3. **Biodegradability:** Most plastics do not occur naturally and are not degraded by microorganisms. By incorporating natural polymers, such as starch, into plastics, they can be made more biodegradable. However, in the anaerobic conditions present in land lls biodegradation is very slow or will not occur at all.

DIOXINS AND POLYCHLORINATED BIPHENYLS

Dioxins can be ormed when polymers are combusted unless the temperature is extremely high. They do not decompose in the environment and can be passed on in the ood chain. Many dioxins, particularly chlorinated dioxins, are highly carcinogenic as they can disrupt the endocrine system (hormone action) and lead to cellular and genetic damage. Dioxins were used as de oliants present in Agent Orange during the Vietnam war. Dioxins contain unsaturated six-membered heterocyclic rings with two oxygen atoms, usually in positions 1 and 4. Examples o dioxins and dioxinlike substances include 1,4-dioxin, polychlorinated dibenzodioxins (PCCDs) and polychlorinated biphenyls (PCBs).

The general ormulas o PCCDs and PCBs are given in Section 31 o the IB data booklet. Some speci c examples are:



PCBs contain rom one to ten chlorine atoms attached to a biphenyl molecule. They are chemically stable and have high electrical resistance so were used in trans ormers and capacitors. Although not strictly dioxins (as they contain no oxygen atoms) they also persist in the environment and have carcinogenic properties.

RESIN IDENTIFICATION CODES

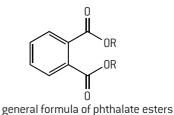
Di erent types o plastic need di erent types o treatment to enable them to be recycled or disposed o sa ely. The Resin Identi cation Code was developed to aid e cient separation by identi ying each speci c plastic with a numbered label in order to help process them be ore recycling. Which plastics can be recycled depends upon the recycling policy o the local community. Some are able to recycle all seven types o plastic whereas some may only be able to recycle PETE and HDPE types o plastic. Use o in rared spectroscopy can also be used to identi y plastics that contain speci c unctional groups. For example, PETE can be recognized by the strong C=O absorption in the region o 1700 1750 cm⁻¹ and PVC shows an absorption due to the C Cl bond between 600 and 800 cm⁻¹.

For plastics that cannot be recycled, special incinerators need to be built to ensure the high temperatures needed to avoid dioxins and other toxic pollutants being ormed when they are combusted. These temperatures are not reached in house res where there are real dangers rom the toxic gases emitted. To help reduce the risk, low smoke zero halogen cabling is o ten used in electrical wiring.

Resin Resin Plastic Identi cation Identi cation Plastic types types Code (RIC) Code (RIC) polyethylene polypropylene terephthalate PP PFTF high-density 6 polystyrene polyethylene HUDE polyvinyl other chloride PV low-density 4 polyethylene LDPE



The use o toxic volatile phthalate esters as plasticizers is being phased out in many countries as they have been associated with birth de ects. Because there are no covalent bonds between the phthalate esters and the plastics they are slowly released into the environment where they enter the atmosphere and the ood chain. This process accelerates as the plastics deteriorate with age.

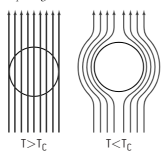


Superconducting metals and X-ray crystallography

SUPERCONDUCTORS

Superconductors are materials (elements or compounds) that will conduct electric currents with no resistance below a critical temperature. As the material cools below the critical temperature to reach the superconducting state the material creates a mirror image magnetic field which cancels out an externally applied magnetic field. This is known as the Meissner e ect. The mirror image magnetic field is ormed by the generation o electric

currents near the sur ace o the material, which cancel the applied magnetic field within the body o the superconductor. Above the critical temperature, electrical resistance is caused by collisions between electrons and positive ions in the lattice. Below the critical temperature the Bardeen Cooper Schrie er (BCS) theory explains that electrons in superconductors orm Cooper pairs which

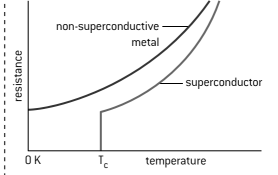


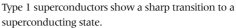
The Meissner e ect. Magnetic fields are excluded rom a superconductor below its critical temperature, T

are able to move reely through the superconductor. Normally electrons behave as ree particles and are repelled by other electrons. At low temperatures the positive ions in the lattice are distorted slightly by a passing electron. A second electron is attracted to this slight positive de ormation and a coupling o these two electrons occurs to orm a Cooper pair that acts as a single entity and causes no resistance.

TYPE 1 AND TYPE 2 SUPERCONDUCTORS

Superconductors can be divided into two types. Type 1 superconductors were discovered first and require the coldest temperatures to become superconductive. They exhibit a very sharp transition to a superconducting state and show complete diamagnetism so can repel a magnetic field completely.





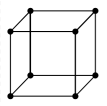
Type 1 superconductors tend to be metals and metalloids. Type 2 super conductors show a gradual transition to the superconducting state and tend to be metallic compounds or alloys. They tend to have higher critical temperatures than Type 1 superconductors. Some recently-discovered Type 2 superconducting perovskites (metal-oxide ceramics that normally have a ratio o 2 metal atoms to every 3 oxygen atoms) have critical temperatures above 0 C.

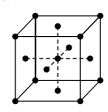
STRUCTURES OF METALS AND IONIC COMPOUNDS

X-ray di raction can be used to analyse the structures o metals and ionic compounds including the more complex perovskites. By using the Bragg equation, $n = 2d \sin$ (see page 107) the precise location o all the metal atoms or ions in the crystal lattice can be determined. Crystal lattices contain simple repeating unit cells and the number o nearest neighbours o an atom or ion is known as its coordination number. Many metals orm cubic structures. Polonium orms a simple cubic cell with a coordination number o six.

Group 1 metals orm a body-centred cubic structure with the atoms filling 68% o the available space. Each metal atom is surrounded by eight other atoms as its nearest neighbours so the coordination number is 8.

Many other metals pack more e ficiently using 74% o the available space with a coordination number o 12. In ace-centred cubic close packing the unit cell has one atom in the centre o each ace rather than one in the centre.





Simple cubic unitBody-centred cubic unitcell. Each atomcell. Each corner atomcontributes $\frac{1}{8}$ to thecontributes $\frac{1}{8}$ to the cellcell so the cell containsso the cell contains $8 \quad \frac{1}{8} = 1$ atom. $1 + \left(8 \quad \frac{1}{8}\right) = 2$ atoms.

Face-centred cubic unit cell. Each corner atom contributes $\frac{1}{8}$ to the cell, each side atom contributes $\frac{1}{2}$ to the cell so the cell contains $\left(8 \quad \frac{1}{8}\right) + \left(6 \quad \frac{1}{2}\right) = 4$ atoms.

Worked example

Potassium has a metallic radius o 2.31 10^{-10} m and orms a bodycentred cubic unit cell. Determine the density o solid potassium.

Step 1. Find the volume o one potassium atom

$$V = \frac{4}{3} r^{3} = \frac{4}{3} 3.14 (2.31 \ 10^{10})^{3}$$
$$= 5.16 \ 10^{-29} m^{3}$$

Step 2. Find the volume o one mole o potassium atoms

 $V = 5.16 \quad 10^{-29} \quad 6.02 \quad 10^{23} \\ = 3.11 \quad 10^{-5} \text{ m}^3 \text{ mol}^{-1}$

Step 3. Find the volume o a lattice containing one mole o potassium atoms

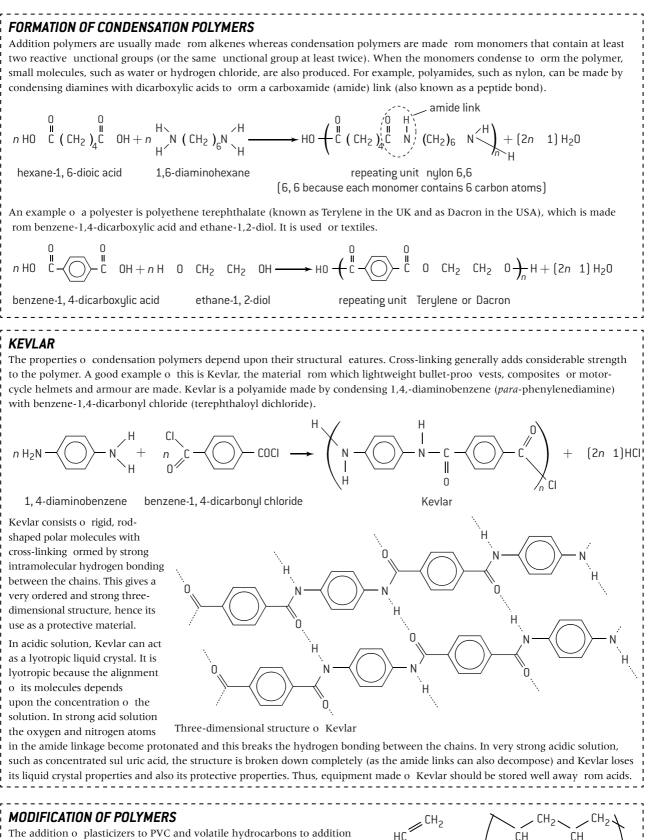
Since it is body-centred cubic the atoms occupy 68% o the available space so the volume o the lattice = $3.11 \quad 10^{-5}$

 $\frac{100}{68} = 4.57$ $10^{-5} \text{ m}^3 \text{ mol}^{-1}$ **Step 4.** Determine the density

knowing that one mole o potassium has a mass o 39.10 g

$$\begin{split} Density &= 39.10/(4.57 \quad 10^{-5}) = 8.56 \\ 10^{-5} \ g \ m^{-3} &= 0.856 \ g \ cm^{-3} \end{split}$$

🕮 Condensation polymers



The addition o plasticizers to PVC and volatile hydrocarbons to addition polymers has already been covered. Expanded polystyrene is ormed by adding pentane during the ormation o styrene and the product heated in steam. This light material is a good thermal insulator and is also used in packaging as it has good shock-absorbing properties.

Air can be blown into polyurethanes to make polyurethane oams or use as cushions and thermal insulation. The fibres o polyesters can be blended with other many actured or natural fibres or making slother which are due as tand a

polystyrene

with other manu actured or natural fibres or making clothes which are dye- ast and more com ortable than pure polyesters.

Like addition polymers, the disadvantages o condensation polymers include their disposal and the e ect on the environment. Condensation polymers are not generally biodegradable and when polyurethanes are burned they can release hydrogen cyanide gas. 🖽 Environmental impact 🛛 heavy metals (1)

HEAVY METALS

Heavy metals that exist as ions in polluted water include cadmium, mercury, lead, chromium, nickel, copper and zinc. Their sources are varied. Cadmium is ound in the e fluent near zinc mining and in batteries and paints. Mercury is used a ungicide in seed dressings and used to be used as an electrode in the electrolysis o sodium chloride. Lead used to be used in paints and leaded gasoline and is still used in car batteries and roo ng material. Other heavy metals originate rom speci c industrial processes (e.g. chromium used to be used or tanning leather), in batteries (e.g. NiCd cells) and biocides (e.g. copper in woodworm treatment). Some methods o removing heavy metals rom polluted water are precipitation, adsorption and chelation.

PRECIPITATION OF HEAVY METAL IONS FROM WATER

Even insoluble salts are still very slightly soluble in water. For a salt ormed rom a metal M with a non-metal X:

 $M^{+}(aq) + X^{-}(aq)$ MX(s)

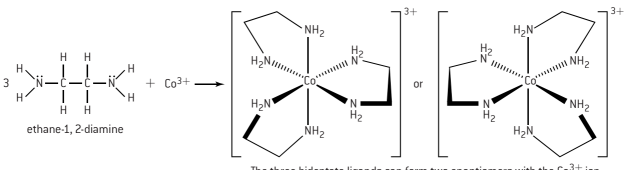
The equilibrium expression or this heterogeneous process will be $K_{er} = [M^+(aq)]$ [X⁻(aq)] where the equilibrium constant, K_{er} is known as the solubility product. (Note that MX(s) does not appear in the expression as a solid has no concentration.) Many metal sul des have very low solubility products so one e ective way o precipitating heavy metal ions is by bubbling hydrogen sul de through polluted water. The solubility product can be used to calculate the amount o a metal ion that will remain in solution a ter it has been precipitated. For example, the solubility product o lead sul de, PbS is $1.30 \quad 10^{-28}$. $Pb(s) = Pb^{2+}(aq) + S^{2-}(aq)$ $K_{sp} = [Pb^{2+}(aq)] [S^{2-}(aq)]$ but $[Pb^{2+}(aq)] = [S^{2-}(aq)]$ There ore $K_{sp} = [Pb^{2+}(aq)]^2 = 1.30 \quad 10^{-28}$ at 298 K and $[Pb^{2+}(aq)] = (1.30 \quad 10^{-28}) = 1.14 \quad 10^{-14} \text{ mol } dm^{-3}$ Hence the concentration o lead ions in the aqueous solution is $1.14 10^{-14}$ mol dm⁻³. The relative atomic mass o lead is 207.19, the mass o lead ions that dissolves in one litre o water at 298 K is there ore 207.19 $1.14 \quad 10^{-14}$ g which is equal to only 2.36 10^{-12} g. Hence the precipitation as the sul de is very e cient. Even though this is very small it can be reduced even urther by adding more sul de ions to the solution. Now the concentration o the lead ions is not the same as the concentration o the sul de ions. This is known as the common ion effect. I the concentration o the sul de ions is made to be 1.00 mol dm-3 then since $K_{sp} = [Pb^{2+}(aq)]$ [S²⁻(aq)] and [S²⁻(aq)] = 1.00 mol dm⁻³ $K_{\rm m} = [Pb^{2+}(aq)] = 1.30 \quad 10^{-28} \text{ at } 298 \text{ K}.$ The concentration o lead ions remaining in the solution at 298 K is now only 1.30 10^{-28} mol dm⁻³ and the mass o lead ions remaining in one litre is just 10 ²⁶ g. 2.69 For many heavy metal ions ound in waste water it is slightly more complicated as the salts ormed are not binary but the principle is the same. For example, nickel ions in waste water can be precipitated by adding hydroxide ions. $Ni^{2+}(aq) + 2OH^{-}(aq) = Ni(OH_{2})(s)$ The solubility product or nickel(II) hydroxide at 298 K is $6.50 10^{-18}$. $K_{\rm cn} = [{\rm Ni}^{2+}({\rm aq})] [{\rm OH}^{-}({\rm aq})]^2$ Hence Assuming that all the hydroxide ions in the solution come just rom the dissolved nickel(II) hydroxide then $[OH^{-}(aq)] = 2[Ni^{2+}(aq)].$ There ore $K_{sn} = 6.50 \quad 10^{-18} = [Ni^{2+}(aq)] \quad (2[Ni^{2+}(aq)])^2 = 4[Ni^{2+}(aq)]^3$ Hence $[Ni^{2+}(aq)] = \left(\frac{6.50}{4} - 10^{-18}\right)^{\frac{1}{3}} = (1.63 - 10^{-18})^{\frac{1}{3}} = 1.18 - 10^{-6} \text{ mol } dm^{-3}.$ Since the relative atomic mass o nickel is 58.69, the mass o nickel ions remaining in one litre o water at 298 K is 6.93 10⁻⁵ g. This small amount can be reduced even more by using the common ion e ect, i.e. by adding more hydroxide ions. However care must be taken to ensure that soluble complex ions are not ormed. For example, zinc hydroxide redissolves in excess hydroxide ions to orm $[Zn(OH)_{4}]^{2-}(aq)$.

🖽 Environmental impact 🛛 heavy metals (2)

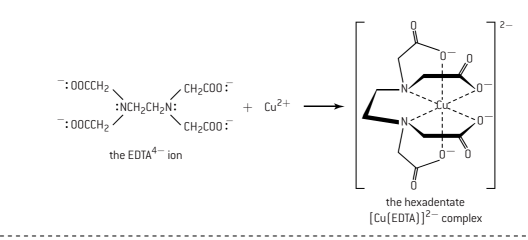
CHELATION

Chelation occurs when a molecule or ion containing non-bonding pairs o electrons bonds to a metal atom or ion orming two or more coordinate bonds. In other words chelating agents are polydentate ligands. Usually chelating agents are organic compounds and are sometimes known as sequestering agents. Polydentate ligands orm more stable complexes than similar monodentate ligands. One explanation or this is that the equilibrium constants or chelation reactions are high. From the expression $G = RT \ln K$ it can be seen that a high value or the equilibrium constant will give a large negative value or the ree entropy change, i.e. the reaction will be highly spontaneous. Since *G* is also equal to *H T S* the large negative value or *G* can also be explained by the large positive entropy change, *S*, as the system becomes more highly ordered compared to when monodentate ligands orm a similar complex.

Two important chelating agents are ethane-1,2-diamine and the ethylenediaminetetraacetate ion, $EDTA^{4-}$. Ethane-1,2-diamine contains two non-bonding pairs o electrons, one on each o the nitrogen atoms so acts as a bidentate ligand. $EDTA^{4-}$ contains six non-bonding pairs o electrons and can use all six o these to orm a hexadentate ligand.



The three bidentate ligands can form two enantiomers with the Co^{3+} ion



THE HABER WEISS AND FENTON REACTIONS Toxic doses o transition metals can disturb the normal oxidation reduction balance in cells through various mechanisms. They can promote the ormation o ree radicals, such as the superoxide anion, O_2^- and the hydroxyl radical, HO which can damage macromolecules, including DNA, proteins and lipids. Hydroxyl radicals can be generated in cells by the reaction o hydrogen peroxide with the superoxide ion. This is known as the Haber Weiss reaction. The reaction is slow but can be catalysed by iron. The overall process is

 $O_{2}^{-}(aq) + H_{2}O_{2}(aq)$ HO $(aq) + OH^{-}(aq) + O_{2}(g)$

The first step involves the catalytic reduction o iron(III) to iron(II)

 $Fe^{3+}(aq) + O_{2}(aq) = Fe^{2+}(aq) + O_{2}(aq)$

The iron(II) is then oxidized back to iron(III) by hydrogen peroxide to orm a hydroxide ion and a hydroxyl radical.

 $Fe^{2+}(aq) + H_2O_2(aq)$ $Fe^{3+}(aq) + OH^-(aq) + HO (aq)$

This second step is known as the Fenton reaction. Fenton s reagent is a solution o hydrogen peroxide and iron(II) sul ate used to puri y contaminated water. The iron(II) ions act as a catalyst. In this reaction the iron(II) is first oxidized to iron(III) by acidified hydrogen peroxide to orm a hydroxyl radical and then reduced back to iron(II) by more hydrogen peroxide. The radicals generated puri y the water by destroying living organisms. Oxidation o organic compounds by Fenton s reagent is both rapid and exothermic and results in the aerobic oxidation o the carbon and hydrogen content o contaminants to orm ultimately carbon dioxide and water.

SHORT ANSWER QUESTIONS OPTION A MATERIALS

- a) Salt (sodium chloride) and haematite, iron(III) oxide, are two minerals that have been known to humans or millennia. Explain why elemental iron has been known since the Iron Age and yet elemental sodium was only discovered some 200 years ago.
 - b) Some compounds o iron are paramagnetic whereas other compounds o iron are diamagnetic.
 Explain the di erence between *paramagnetic* and *diamagnetic* on a molecular level.

[2]

[2]

[2]

[3]

[3]

[2]

[2]

[2]

[3]

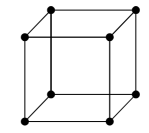
[2]

[2]

- c) Stainless steel contains iron alloyed with carbon, chromium and some other metals. In addition to making the steel resistant to rusting, state two other changes that occur when iron is alloyed with carbon and other metals.
- a) Determine the time taken to produce 1000 kg o aluminium when a steady current o 2.00 10⁵ A is passed through a solution o alumina in molten cryolite, Na,AlF₄.
 - b) Discuss why the production o aluminium makes a signi cant contribution to the greenhouse e ect.
- 3. Liquid crystals are sometimes used in the construction o smart windows . Smart windows are milky white as their randomly arranged liquid crystals scatter light. When a voltage is applied, the liquid crystals align in the same direction. The light then passes through them without scattering, making the windows transparent.
 - a) State the property o the liquid crystal molecules that allows them to align when a voltage is applied. [1]
- 7. Kevlar can be made by condensing 1,4,-diaminobenzene with benzene-1,4-dicarbonyl chloride.
 - a) Draw the repeating unit o Kevlar.
 - b) Explain why Kevlar is such a strong polymer.
 - c) Explain why a bullet proo vest made o Kevlar should not be allowed to come into contact with sul uric acid.
- 8. Cadmium ions, $Cd^{2+}(aq)$ are extremely poisonous. They can be removed rom solution by adding hydroxide ions to precipitate cadmium hydroxide, $Cd(OH)_2$. The solubility product o cadmium hydroxide is 7.2 10^{-15} at 298 K.
 - a) Calculate the solubility o $\,$ cadmium hydroxide in mol dm $^{-3}$ at 298 K.
 - b) Explain why the solubility o cadmium ions in solution can be reduced even urther by adding excess hydroxide ions.
 - c) Heavy metals can also be removed rom water by chelation. Explain why ethane-1,2-diamine is a good chelating agent.

- b) List two substances that can behave as liquid crystals. [1] c) Distinguish between *thermotropic* and *lyotropic* liquid crystals. [2] 4. a) Distinguish between homogeneous and heterogeneous catalysts and explain their di erent mode o action. [4] b) List three actors that should be considered when choosing a catalyst or a particular process. [3] c) Iron is a good catalyst or the Haber Process. Suggest why it is used in a nely divided (powdered) orm. [1] 5. a) Propene can polymerize to orm either isotactic or atactic poly(propene). Draw a section o isotactic poly(propene) containing our methyl groups. [2] b) Describe how the properties o isotactic poly(propene) di er rom those o atactic poly(propene). [2] c) Explain why low density poly(ethene), LDPE, is more flexible than high density poly(ethene), HDPE. [2] d) Deduce the atom economy or the conversion o ethene to poly(ethene). [1] 6. a) One o the methods or making nanotubes is chemical vapour deposition, CVD. Explain why an inert gas such as nitrogen and not oxygen must be used in the process. [2] b) Explain why nanotubes are good conductors o
 - electricity. [1]

- 9. A simple cubic structure contains atoms placed at the corners o a cube.
 - a) Determine the number o atoms in one unit cell. [1]
 - b) Determine the coordination number o each atom in a simple cubic structure. [1]
 - c) Explain how the distance between the atoms in a simple cubic structure can be determined. [2]



13 OPTION B BIOCHEMISTRY

Introduction to biochemistry

METABOLISM

The chemical reactions that occur in living organisms are collectively known as metabolism. Essentially these enzymecatalysed reactions allow organisms to grow and reproduce, maintain their structures and respond to their environment. Reactions in which organic matter is broken down to produce energy involve the process o *catabolism* whereas reactions which use energy to synthesize larger molecules such as proteins and nucleic acids are known as *anabolism*.

The chemical reactions involved in metabolism are organized into metabolic pathways. Each step is catalysed by an enzyme and normally occurs in a controlled aqueous environment. Biological molecules are diverse in nature and their unctions depend upon their precise structure and shape.

Typical examples o biological molecules include:

Proteins	made rom linking amino acids by peptide bonds.
Lipids	these include ats and steroids.
Carbohydrates	they have the general ormula $C_x(H_2O)_y$ and include monosaccharides (e.g. glucose and ructose) and polysaccharides (e.g. starch and cellulose).
Nucleotides	consist o a phosphate group, a ribose sugar group and a nitrogenous base. Polymers o nucleotides include DNA and RNA.
Co-enzymes	compounds such as adenosine triphosphate (ATP) that are metabolic intermediates whose unction is to carry chemical groups (or energy) between di erent reactions.

PHOTOSYNTHESIS AND RESPIRATION

Photosynthesis is the process used by plants and other organisms to synthesize energy-rich molecules such as carbohydrates rom carbon dioxide and water using light energy. During photosynthesis oxygen is released. This is the origin o the oxygen in the atmosphere and photosynthesis is responsible or maintaining atmospheric oxygen levels. Essentially photosynthesis is the process o converting carbon into biomass. The process is complicated and needs the presence o chlorophyll but can be simplified by the overall equation:

light

$$6CO_{2}(g) + 6H_{2}O(l)$$

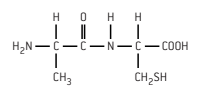
 $C_6H_{12}O_6(aq) + 6O_2(g)$ H positive

Respiration is also a complex set o reactions in which energyrich molecules such as carbohydrates are broken down in the presence o oxygen to provide energy or cells. Ultimately the products o respiration are carbon dioxide and water, which are released into the atmosphere. The overall equation is the reverse o photosynthesis.

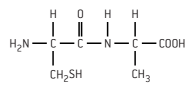
 $C_{6}H_{12}O_{6}(aq) + 6O_{2}(g) = 6CO_{2}(g) + 6H_{2}O(l) = H$ negative

CONDENSATION AND HYDROLYSIS REACTIONS

Biological polymers (biopolymers) are ormed by condensation reactions. These involve the reaction between two smaller molecules to orm one larger molecule with the evolution o a small molecule such as water. For condensation polymerization to occur each reacting molecule must possess at least two reactive unctional groups. Classic examples include the condensation o amino acids to orm proteins and the condensation o sugars to orm starch. For example, the two amino acids alanine, H₂N CHCH₃ COOH and cysteine, H₂N CH(CH₂SH) COOH can condense together to orm two di erent possible dipeptides i each dipeptide contains one o each o the two acid residues.



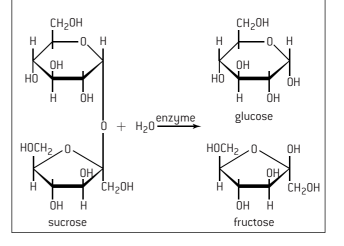
H₂N-Ala-Cys-COOH



H₂N-Cys-Ala-COOH

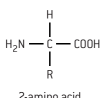
Each end o the dipeptides contains a reactive group so can undergo urther condensation reactions with more amino acids to produce polypeptides.

Hydrolysis is the reverse o condensation. A molecule is hydrolysed when a water molecule (o ten in the presence o acid or alkali) reacts with a larger molecule to break a bond and orm two smaller molecules. The hydrolysis o proteins produces amino acids and the hydrolysis o starch (polysaccharide) produces sugars (monosaccharides). For example, sucrose, a disaccharide, can be hydrolysed to orm glucose and ructose.



Structure of proteins

AMINO ACIDS AND THE STRUCTURE OF PROTEINS

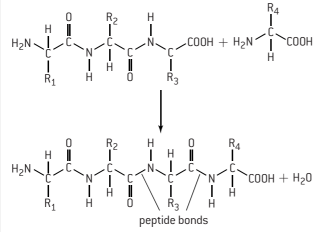


Amino acids contain both an amine unctional group and a carboxylic acid unctional group. When both are attached to the same carbon atom they are known as 2-amino acids (or -amino acids). They are solids at room temperature and have quite high melting points. This is because they

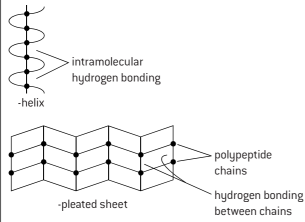
2-amino acid

can exist as zwitterions in which the hydrogen atom rom the carboxylic acid group protonates the amine group to orm a carboxylate anion and a substituted ammonium cation within the same compound (see page 126).

Proteins are large macromolecules made up o chains o 2-amino acids. There are about twenty 2-amino acids that occur naturally. A ull list o the 2-amino acids together with their common names, abbreviations and structural ormulas is given in Section 33 o the IB data booklet. The amino acids bond to each other through condensation reactions resulting in the ormation o a polypeptide, in which the amino acid residues are bonded to each other by a carboxamide link (peptide bond).

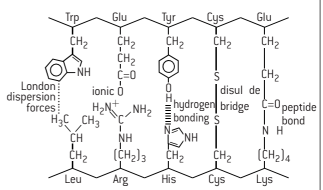


Each protein contains a fixed number o amino acid residues connected to each other in strict sequence. This sequence, e.g. gly-his-ala-ala-leu- ... is known as the primary structure o proteins. The secondary structure describes the way in which the chain o amino acids olds itsel due to intramolecular hydrogen bonding. The olding can either be -helix in which the protein twists in a spiralling manner rather like a coiled spring, or -pleated to give a sheet-like structure.

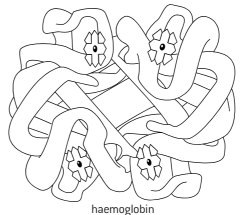


The tertiary structure describes the overall olding o the chains by interactions between distant amino acids to give the protein its three-dimensional shape. These interactions may be due to hydrogen bonds, London dispersion orces between nonpolar side groups, and ionic attractions between polar groups. In addition two cysteine residues can orm disulfide bridges when their sul ur atoms undergo oxidation.

Examples o interactions between side groups on polypeptide chains:



Separate polypeptide chains can interact together to give a more complex structure this is known as the quaternary structure. Haemoglobin has a quaternary structure that includes our protein chains (two -chains and two -chains) grouped together around our haem groups.



Fibrous and globular proteins

Haemoglobin is an example o a globular protein. Globular proteins have complex tertiary and sometimes quaternary structures (e.g. haemoglobin) olded into spherical (globular) shapes. They are usually soluble to some extent in water as the hydrophobic side chains tend to be in the centre o the structure. Fibrous proteins, such as collagen, have little or no tertiary structure and orm long parallel polypeptide chains. Fibrous proteins have cross-linking at intervals to orm long fibres or sheets and have mainly structural roles such as keratin in hair and collagen, which is ound in skin and the walls o blood vessels and acts as connective tissue.

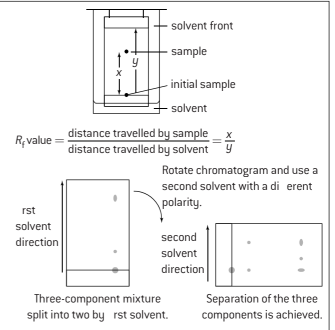
Analysis of proteins

The primary structure o proteins can be determined either by paper chromatography or by electrophoresis. In both cases the protein must first be hydrolysed by hydrochloric acid and heat to successively release the amino acids. The three-dimensional structure o the complete protein can be confirmed by X-ray crystallography.

PAPER CHROMATOGRAPHY

A small spot o the unknown amino acid sample is placed near the bottom o a piece o chromatographic paper. Separate spots o known amino acids can be placed alongside. The paper is placed in a solvent (eluent), which then rises up the paper due to capillary action. As it meets the sample spots the di erent amino acids partition themselves between the eluent and the paper to di erent extents, and so move up the paper at di erent rates. When the eluent has nearly reached the top, the paper is removed rom the tank, dried, and then sprayed with an organic dye (ninhydrin) to develop the chromatogram by colouring the acids. The positions o all the spots can then be compared.

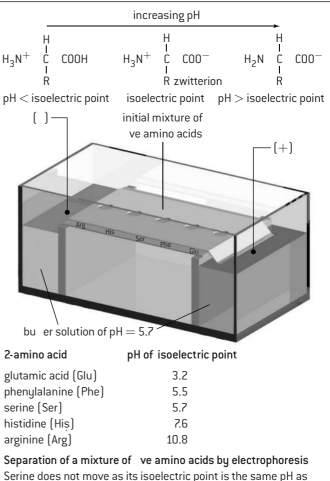
I samples o known amino acids are not available the R value (retention actor) can be measured and compared with known values as each amino acid has a di erent R value. It is possible that two acids will have the same R value using the same solvent, but di erent values using a di erent solvent. I this is the case the chromatogram can be turned through 90 and run again using a second solvent.



ELECTROPHORESIS

The structure o amino acids alters at di erent pH values. At low pH (acid medium) the amine group will be protonated. At high pH (alkaline medium) the carboxylic acid group will lose a proton. This explains why amino acids can unction as bu ers. I H⁺ ions are added they are removed as NH_4^+ and i OH^- ions are added the COOH loses a proton to remove the OH^- ions as water. For each amino acid there is a unique pH value (known as the **isoelectric point**) where the acid will exist as the zwitterion.

The medium on which electrophoresis is carried out is usually a polyacrylamide gel. So the process is known as PAGE (polyacrylamide gel electrophoresis). The sample is placed in the centre o the gel and a potential di erence applied across it. Depending on the pH o the bu er the di erent amino acids will move at di erent rates towards the positive and negative electrodes. At its isoelectric point a particular amino acid will not move as its charges are balanced. When separation is complete the acids can be sprayed with ninhydrin and identified by comparing the distance they have travelled with standard samples, or rom a comparison o their isoelectric points.



Serine does not move as its isoelectric point is the same pH as the bu er. Histidine and arginine contain NH_3^+ at pH 5.7, so move towards the negative electrode. Glutamic acid and phenylalanine contain COO at pH 5.7, so move towards the positive electrode.

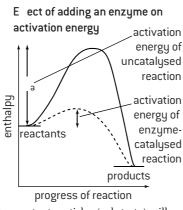
Enzymes

USES OF PROTEINS

Proteins have many di erent unctions in the body. They can act as biological catalysts or specific reactions (enzymes). They can give structure (e.g hair and nails consist almost entirely o polypeptides coiled into -helices), and provide a source o energy. Some hormones are proteins, e.g. FSH (ollicle stimulating hormone), responsible or triggering the monthly cycle in emales.

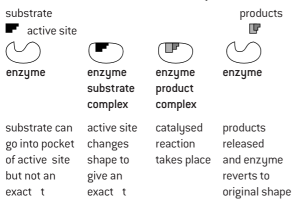
CATALYTIC ACTIVITY OF ENZYMES AND ACTIVE SITE

Enzymes are protein molecules that catalyse biological reactions. Each enzyme is highly specific or a particular reaction, and extremely e ficient, o ten being able to increase the rate o reaction by a actor greater than 10^{s} . Enzymes work by providing an alternative pathway or the reaction with a lower activation



energy, so that more o the reactant particles (substrate) will possess the necessary minimum activation energy.

The specificity o enzymes depends on their tertiary and quaternary structure. The part o an enzyme that reacts with the substrate is known as the active site. This is a groove or pocket in the enzyme where the substrate will bind. The site is not necessarily rigid but can alter its shape to allow or a better fit known as the induced fit theory.

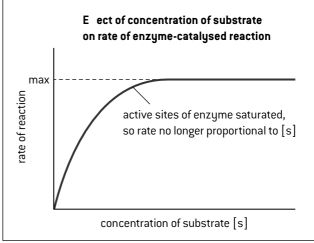


Induced t theory of enzyme catalysis

The induced fit theory replaces the old lock and key theory which assumes that enzymes have a fixed shape into which the substrate fits.

ENZYME KINETICS

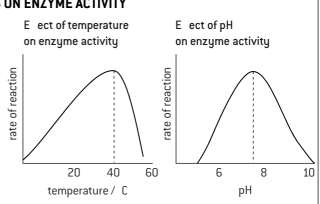
At low substrate concentrations the rate o the enzymecatalysed reaction is proportional to the concentration o the substrate. However at higher concentrations the rate reaches a maximum. This can be explained in terms o enzyme saturation. At low substrate concentrations there are enough active sites present or the substrate to bind and react. Once all the sites are used up the enzyme can no longer work any aster.



EFFECT OF TEMPERATURE, PH AND HEAVY METAL IONS ON ENZYME ACTIVITY

The action o an enzyme depends on its specific shape. Increasing the temperature will initially increase the rate o enzyme-catalysed reactions, as more o the reactants will possess the minimum activation energy. The optimum temperature or most enzymes is about 40 C. Above this temperature enzymes rapidly become denatured as the weak bonds holding the tertiary structure together break.

At di erent pH values the charges on the amino acid residues change a ecting the bonds between them, and so altering the tertiary structure and making the enzyme ine ective. Heavy metals can poison enzymes by reacting with SH groups replacing the hydrogen atom with a heavy metal atom or ion so that the tertiary structure is altered.

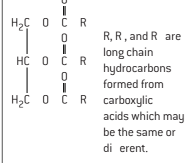


Lipids (1)

Lipids are organic molecules with long hydrocarbon chains that are soluble in non-polar solvents. They are mainly used or energy storage, insulating and protecting vital organs, orming cell membranes and, in some cases, acting as hormones. Three important types o lipids are triglycerides (ats and oils), phospholipids (lecithin) and steroids (cholesterol).

FATS AND OILS

Fats and oils are triesters (triglycerides) ormed rom the condensation reaction o propane-1,2,3-triol (glycerol) with long chain carboxylic acids (atty acids).

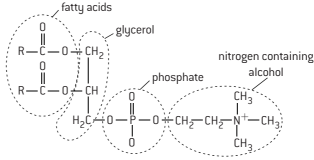




Fats are solid triglycerides; examples include butter, lard and tallow. Oils are liquid at room temperature and include castor oil, olive oil and linseed oil. The essential chemical di erence between them is that ats contain saturated carboxylic acid groups (i.e. they do not contain C=C double bonds). Oils contain at least one C=C double bond and are said to be unsaturated. Most oils contain several C=C double bonds and are known as polyunsaturated.

PHOSPHOLIPIDS

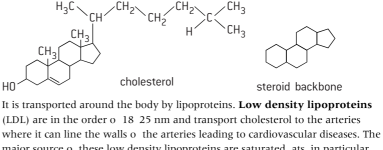
Phospholipids orm an integral part o all cell membranes. They are essentially made o our components. A backbone such as propane-1,2,3-triol (glycerol), linked by esterification to two atty acids and a phosphate group which is itsel condensed to a nitrogen-containing alcohol. There are many di erent phospholipids. They can be exemplified by phosphatidyl choline the major component o lecithin, present in egg yolk.



The structure o phosphatidyl choline showing the origins o the our distinct components.

CHOLESTEROL

Cholesterol has the characteristic our-ring structure possessed by all steroids.

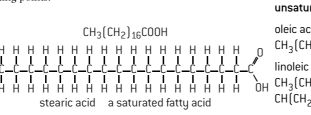


major source o these low density lipoproteins are saturated ats, in particular those derived rom lauric (C_{12}) , myristic (C_{14}) and palmitic (C_{16}) acids. Smaller lipoproteins, in the order o 8 11 nm, known as **high density lipoproteins** (HDL) can remove the cholesterol rom the arteries and transport it back to the liver.

FATTY ACIDS

Stearic acid (m.pt 69.6 C) and linoleic acid (m.pt -5.0 C) both contain the same number o carbon atoms and have similar molar masses. However, linoleic acid contains two double bonds. Generally the more unsaturated the atty acid the lower its melting point.

The regular tetrahedral arrangement o saturated atty acids means that they can pack together closely, so the London dispersion orces holding molecules together are stronger as the sur ace area between them is greater. As the bond angle at the C=C double bonds changes rom 109.5 to 120 in unsaturated acids it produces a kink in the chain. They are unable to pack so closely and the London dispersion orces between the molecules become weaker, which results in lower melting points. This packing arrangement is similar in ats and explains why unsaturated ats (oils) have lower melting points.

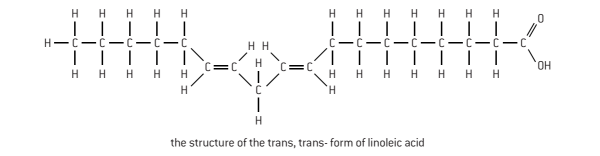


ciferany the more unsaturated the	any actu the lower its menning point.			
Name	Number of C atoms per	Number of	Melting	
acturated fattu acida	molecule	C=C bonds	point/ C	
saturated fatty acids				
lauric acid CH ₃ (CH ₂) ₁₀ COOH	12	0	44.2	
myristic acid CH ₃ (CH ₂) ₁₂ COOH	14	0	54.1	
palmitic acid CH ₃ (CH ₂) ₁₄ COOH	16	0	62.7	
stearic acid CH ₃ (CH ₂) ₁₆ COOH	18	0	69.6	
unsaturated fatty acids				
oleic acid CH ₃ (CH ₂) ₇ CH=CH(CH ₂) ₇ COOH	18	1	10.5	
linoleic acid _H CH ₃ (CH ₂) ₄ CH=CHCH ₂ CH= CH(CH ₂) ₂ COOH	18	2	5.0	

Lipids (2)

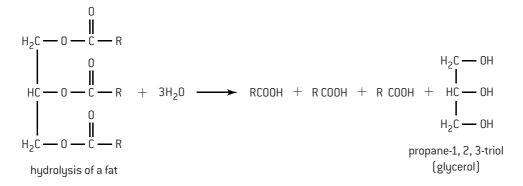
HYDROGENATION

Oils naturally contain only cis-unsaturated atty acids. These are generally healthier than saturated ats as they increase HDL cholesterol. Unsaturated ats can be hydrogenated to saturated ats with a higher melting point by adding hydrogen under pressure in the presence o a heated nickel catalyst. However during the hydrogenation process, partial hydrogenation can occur and the trans-isomers may be ormed. Trans-unsaturated atty acids are present in ried oods such as French ries and some margarines. Unlike natural mono- and poly-unsaturated oils, trans-unsaturated ats increase the ormation o LDL cholesterol (bad cholesterol) and thus increase the risk o heart disease.



HYDROLYSIS AND RANCIDITY OF FATS

Fats and oils are hydrolysed in the body by enzymes, known as lipases, to glycerol and atty acids. These in turn are broken down by a series o redox reactions to produce ultimately carbon dioxide, water and energy. Because they are essentially long-chain hydrocarbons with only two oxygen atoms each on the three carboxyl atoms ats are in a less oxidized orm than carbohydrates so weight or weight produce more energy.

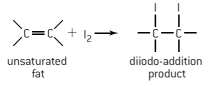


Lipids (ats and oils) in ood become rancid when our senses perceive them to have gone o due to a disagreeable smell, texture or appearance. This may be caused either by hydrolysis o the triesters (hydrolytic rancidity) as shown above to produce disagreeable smelling atty acids or by oxidation o the atty acid chains.

Oxidative rancidity is typically due to the addition o oxygen across the C=C double bonds in unsaturated atty acids. Oily fishes, such as mackerel, contain a high proportion o unsaturated atty acids and are prone to oxidative rancidity. The process proceeds by a ree radical mechanism catalysed by light in the presence o enzymes.

IODINE NUMBER

Unsaturated ats can undergo addition reactions. The addition o iodine to unsaturated ats can be used to determine the number o C=C double bonds, since one mole o iodine will react quantitatively with one mole o C=C double bonds. Iodine is coloured. As the iodine is added to the unsaturated at the purple colour o the iodine will disappear as the addition reaction takes place. Once the colour remains the amount o iodine needed to react with all the C=C double bonds can be determined. O ten ats are described by their iodine number, which is the number o grams o iodine that add to 100 g o the at.



THE ROLES OF LIPIDS IN THE BODY

Energy storage. Because they contain proportionally less oxygen than carbohydrates they release more energy when oxidized.

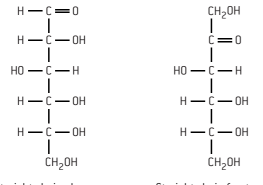
- Insulation and protection o organs. Fats are stored in adipose tissue, which provides both insulation and protection to parts o the body.
- Steroid hormones. Examples include emale and male sex hormones such as progesterone and testosterone and the contraceptive pill. Sometimes steroids are abused. Anabolic steroids have similar structures to testosterone and are taken to build up muscle.
- Cell membranes. Lipids provide the structural component o cell membranes.

More controversially, lipids are thought to a ect health, particularly heart disease. Although the evidence is disputed by many, some think that saturated atty acids, particularly lauric (C_{12}) , myristic (C_{14}) and palmitic (C_{16}) acids increase LDL, as do trans-unsaturated atty acids such heart problems. Conversely omega-3-polyunsaturated atty acids such as natural unsaturated atts (e.g. olive oil) are thought to lower the level o LDL and consequently are thought to be good or you.

Carbohydrates

MONOSACCHARIDES

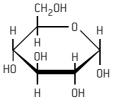
All monosaccharides have the empirical ormula CH_2O . In addition they contain a carbonyl group (C=O) and at least two –OH groups. I the carbonyl group is an aldehyde (RCHO) they are known as an aldose, i the carbonyl group is a ketone (RCOR) they are known as a ketose. Monosaccharides have between three and six carbon atoms.



Straight chain glucose (an aldose) Straight chain fructose (a ketose)

Monosaccharides with the general ormula $C_5 H_{10} O_5$ are known as pentoses (e.g. ribose) and monosaccharides with the general ormula $C_6 H_{12} O_6$ are known as hexoses (e.g. glucose).

Many structural isomers o monosaccharides are possible. In addition several carbon atoms are chiral (asymmetric) and give rise to optical isomerism. As well as this, open chain structures and ring structures are possible. The orm o glucose that is ound in nature is known as p-glucose. Note that the



D-glucose

ring structures are cyclic ethers as they contain an oxygen atom bonded on either side by a carbon atom within the ring.

Six-membered ring monosaccharides are known as pyranoses. Hexoses can also have a uranose structure where they have a ve-membered ring containing an oxygen atom.

MAJOR FUNCTIONS OF POLYSACCHARIDES IN THE BODY

Carbohydrates are used by humans:

to provide energy: oods such as bread, biscuits, cakes, potatoes and cereals are all high in carbohydrates.

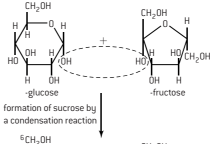
to store energy: starch is stored in the livers o animals in the orm o glycogen also known as animal starch. Glycogen has almost the same chemical structure as amylopectin.

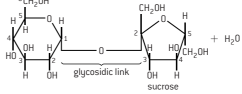
as precursors or other important biological molecules, e.g. they are components o nucleic acids and thus play an important role in the biosynthesis o proteins.

as dietary fibre: dietary bre is mainly plant material that is not hydrolysed by enzymes secreted by the human digestive tract but may be digested by microflora in the gut. Examples include cellulose, hemicellulose, lignin and pectin. It may be help ul in preventing conditions such as diverticulosis, irritable bowel syndrome, obesity, Crohn s disease, haemerrhoids and diabetes mellitus.

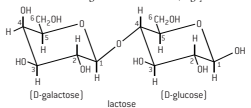
POLYSACCHARIDES

Monosaccharides can undergo condensation reactions to orm disaccharides and eventually polysaccharides. For example, sucrose, a disaccharide ormed rom the condensation o p-glucose in the pyranose orm and p- ructose in the uranose orm.

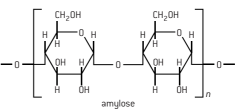




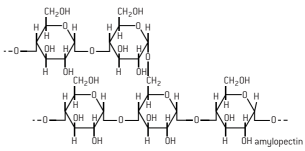
The link between the two sugars is known as a glycosidic link. In the case o sucrose the link is between the C-1 atom o glucose and the C-2 atom o ructose. The link is known as a 1,2 glycosidic bond. Maltose, another disaccharide is ormed rom two glucose molecules condensing to orm an 1,4 glycosidic bond. Lactose is a disaccharide in which the p-galactose is linked at the C-1 atom to the C-4 atom o p-glucose to orm a 1,4 glycosidic bond.



One o the most important polysaccharides is starch. Starch exists in two orms: amylose, which is water soluble, and amylopectin, which is insoluble in water. Amylose is a straight chain polymer o p-glucose units with 1,4 glycosidic bonds:



Amylopectin also consists o D-glucose units but it has a branched structure with both 1,4 and 1,6 glycosidic bonds:



Most plants use starch as a store o carbohydrates and thus energy. Cellulose, a polymer o p-glucose contains 1,4 linkages. Cellulose, together with lignin, provides the structure to the cell walls o green plants. Most animals, including all mammals, do not have the enzyme cellulase so are unable to digest cellulose or other dietary bre polysaccharides.

Vitamins

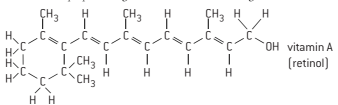
VITAMINS

Vitamins are micro-nutrients. Micro-nutrients are substances required in very small amounts (mg or g). They mainly unction as a co- actor o enzymes and include not only vitamins but also trace minerals such as Fe, Cu, F, Zn, I, Se, Mn, Mo, Cr, Co and B.

Vitamins can be classified as at soluble or water soluble. The structure o at soluble vitamins is characterized by long, non-polar hydrocarbon chains or rings. These include vitamins A, D, E, F and K. They can accumulate in the atty tissues o the body. In some cases an excess o at soluble vitamins can be as serious as a deficiency. The molecules o water soluble vitamins, such as vitamin C and the eight B-group vitamins, contain hydrogen attached directly to electronegative oxygen or nitrogen atoms that can hydrogen bond with water molecules. They do not accumulate in the body so a regular intake is required. Vitamins containing C=C double bonds and -OH groups are readily oxidized and keeping ood re rigerated slows down this process.

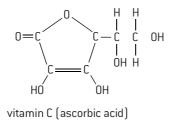
VITAMIN A (RETINOL)

Although it does contain one -OH group, vitamin A is at soluble due to the long non-polar hydrocarbon chain. Unlike most other vitamins it is not broken down readily by cooking. Vitamin A is an aid to night vision.



VITAMIN C (ASCORBIC ACID)

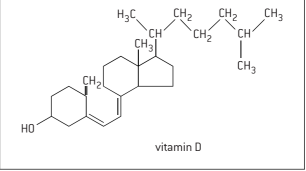
Due to the large number o polar -OH groups vitamin C is soluble in water so is not retained or long by the body. The most amous disease associated with a lack o vitamin C is scorbutus (scurvy). The symptoms are swollen legs, rotten gums and bloody lesions. It was a common disease in sailors, who spent long periods without resh ood, until the cause was recognized.



VITAMIN D (CALCIFEROL)

Vitamin D is essentially a large hydrocarbon with one OH group and is at soluble.

A deficiency o vitamin D leads to bone so tening and mal ormation a condition known as rickets.



MALNUTRITION

Malnutrition occurs when either too much ood is consumed, which leads to obesity, or the diet is lacking in one or more essential nutrients. Specific micro-nutrient deficiencies include:

Fe anaemia

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I goitre
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vitamin A (retinol) xerophthalmia, night blindness

vitamin B₃ (niacin) pellagra

vitamin B₁ (thiamin) beriberi

vitamin C (ascorbic acid) scurvy

vitamin D (calci erol) rickets.

Solutions to combat malnutrition include:

eating resh ood rich in vitamins and minerals

adding nutrients that are missing in commonly consumed oods

genetic modification o ood

providing nutritional supplements.

Biochemistry and the environment

GREEN CHEMISTRY

Our increasing knowledge and use o biochemistry has led to solutions to some issues but has also caused environmental problems in other areas. Scientists have a responsibility to be aware o the impact o their research on the environment and should actively find ways to counter any negative impact their work may have on the environment. Examples o negative impact include the use o enzymes in biological detergents and the overuse o antibiotics in animal eed. Green chemistry, which is sometimes also known as sustainable chemistry, encourages the reduction and prevention o pollution at source. It does this by trying to minimize the use and ormation o substances harm ul to the environment. One way in which this can be achieved is to make use o atom economy (see page 6).

BIODEGRADABILITY

Although most plastics are organic in origin they are petroleum-based so cannot easily be broken down by natural organisms and cause big pollution problems. Biodegradable plastics are plastics capable o being broken down by bacteria or other organisms, ultimately to carbon dioxide and water. They are based on natural renewable polymers containing ester or glycosidic links, such as starch, that can be hydrolysed. In theory, starch-based bioplastics produced as biomass could be almost carbon neutral but there are problems such as using land that could otherwise be used or growing ood and the release o the greenhouse gas methane i the plastics are decomposed anaerobically in landfill sites.

Enzymes can also be used to biodegrade pollutants. Enzymes are used to aid the breakdown and dispersal o oil spills. This reduces the e ect o dispersal agents, such as 2-butoxyethanol, but does not replace them completely. The oil still needs to be broken into smaller droplets be ore the microbes containing the enzymes can be e ective. The use o enzymes in biological detergents is also well known. This has the advantage to the environment o lowering the temperature at which clothes need to be washed so making the process more e ficient, i.e. saving on energy and ossil uels.

HOST GUEST CHEMISTRY

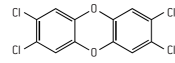
Host guest complexes are made o two or more molecules or ions bonded together through noncovalent bonding, which is critical in maintaining the 3-D structure o the molecule. Non-covalent interactions include hydrogen bonds, ionic bonds and van der Waals orces. These orces, which are weaker than covalent bonding, allow large molecules to bind specifically but transiently to one another to orm supramolecules. They work by mimicking some o the actions per ormed by enzymes by selectively binding to guest species. For example, they have been used to deliver drugs more e ectively in humans by increasing the solubility and availability o the drug and reducing drug resistance. They are also used to remove toxic materials (xenobiotics) rom the environment. For example, radioactive 137-caesium rom nuclear waste and carcinogenic amines rom polluted water.

XENOBIOTICS

Chemicals ound in organisms, which are not normally present or produced by the organism, or are present in organisms in abnormally high amounts, are known as xenobiotics. Examples oxenobiotics include drugs in animals. Antibiotics, or example, are not produced by animals nor are they part oxen a normal diet. The use oxentibiotics in animal eed and in sewage plants has meant that they pass through into the human ood chain and increase resistant strains oxentrations are zenobiotics may be natural compounds but most are pollutants. Two classic xenobiotics are dioxins and polychlorinated biphenyls (PCBs).

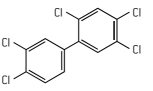
Dioxins can be ormed when polymers are combusted, unless the temperature is extremely high. They do not decompose in the environment and can be passed on in the ood chain. Many dioxins, particularly chlorinated dioxins, are highly carcinogenic as they can disrupt the endocrine system (hormone action) and lead to cellular and genetic damage. Examples o dioxins and dioxin-like substances include 1,4-dioxin, polychlorinated dibenzodioxins (PCCDs) and polychlorinated biphenyls (PCBs). The general ormulas o PCCDs and PCBs are given in Section 31 o the IB data booklet. Some specific examples are:





1, 4-dioxin

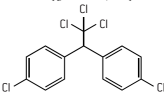
2, 3, 7, 8-tetrachlorodibenzodioxin (an example of a PCCD)



2, 3′, 4, 4′, 5-pentachlorobiphenyl (an example of a PCB)

PCBs contain rom one to ten chlorine atoms attached to a biphenyl molecule. They are chemically stable and have high electrical resistance so were used in trans ormers and capacitors. Although not strictly dioxins (as they contain no oxygen atoms) they also persist in the environment and have carcinogenic properties.

One o the problems associated with xenobiotics is biomagnification. As the xenobiotic passes through the ood chain its concentration increases in higher species. DDT (dichlorodiphenyltrichloroethane) is an e ective insecticide particularly against the malaria mosquito but its use is now banned as it accumulates at high levels in birds o prey, which threatens their survival.





Proteins and enzymes (1)

V_{MAY} AND THE MICHAELIS-MENTEN CONSTANT, K COMPETITIVE AND NON-COMPETITIVE INHIBITION Inhibitors are substances that slow down the rate o enzyme-We have seen already that at low substrate concentrations the catalysed reactions. Competitive inhibitors resemble the rate o an enzyme-catalysed reaction is proportional to the substrate in shape, but cannot react. They slow down the concentration o the substrate but it reaches a maximum at reaction because they can occupy the active site on the higher substrate concentrations. This maximum is known as enzyme thus making it less accessible to the substrate. $V_{\rm max}$. Non-competitive inhibitors also bind to the enzyme, but E ect of concentration of substrate on not on the active site. They bind at the allosteric site, which rate of enzyme-catalysed reaction causes the enzyme to change its shape so that the substrate cannot bind. As the substrate concentration is increased the e ect o competitive inhibitors lessens, as there is increased V_{max} competition or the active sites by the substrate. With nonate of reaction competitive inhibitors increasing the substrate concentration active sites of enzyme has no e ect, as the enzyme s shape still remains altered. saturated, so rate no E ect of substrate concentration on inhibitors V_{max} longer proportional to [S] uninhibited reaction V_{max} ate of reaction V_{max} non-competitive V_{max} inhibition K_m competitive inhibition concentration of substrate [S] The Michaelis-Menten constant, $K_{\rm m}$ is the substrate concentration when the rate o the reaction is V_{max} . A particular enzyme Km [s] with the same substrate will always have the same value For non-competitive inhibitors, V_{max} is lower but K_m is the same. or K_m . It indicates whether the enzyme unctions e ficiently For competitive inhibitors, V_{max} is the same but K_m is increased. at low substrate concentrations, or whether high substrate concentrations are necessary or e ficient catalysis **BUFFER SOLUTIONS** Enzymes only unction e ficiently within a narrow pH region. Outside o this region the structure is altered and the enzyme becomes denatured, hence the need or bu ering. A bu er solution resists changes in pH when small amounts o acid or alkali are added to it. An acidic bu er solution can be made by mixing a weak acid together with the salt o that acid and a strong base. An example is a solution o ethanoic acid and sodium ethanoate. The weak acid is only slightly dissociated in solution, but the salt is ully dissociated into its ions, so the concentration o ethanoate ions is high. NaCH₃COO(aq) $Na^+(aq) + CH_3COO^-(aq)$ CH₂COOH(aq) $CH_{a}COO^{-}(aq) + H^{+}(aq)$ I an acid is added the extra H+ ions coming rom the acid are removed as they combine with ethanoate ions to orm undissociated ethanoic acid, so the concentration o H⁺ ions remains unaltered. $CH_{3}COO^{-}(aq) + H^{+}(aq)$ CH, COOH(aq) I an alkali is added the hydroxide ions rom the alkali are removed by their reaction with the undissociated acid to orm water, so again the H⁺ ion concentration stays constant. $CH_{2}COOH(aq) + OH^{-}(aq)$ $CH_3COO^-(aq) + H_2O(l)$ In practice acidic bu ers are o ten made by taking a solution o a strong base and adding excess weak acid to it, so that the solution contains the salt and the unreacted weak acid. $NaOH(aq) + CH_COOH(aq)$ $NaCH_COO(aq) + H_O(l) + CH_COOH(aq)$ limiting reagent salt excess weak acid Bu er solution An alkali bu er with a fixed pH greater than 7 can be made rom a weak base together with the salt o that base with a strong acid. An example is ammonia with ammonium chloride. NH_cCl(aq) $NH_{4}^{+}(aq) + Cl^{-}(aq)$ $NH_{a}(aq) + H_{a}O(l)$ $NH_{4}^{+}(aq) + OH^{-}(aq)$ I H⁺ ions are added they will combine with OH⁻ ions to orm water and more o the ammonia will dissociate to replace them. I more OH- ions are added they will combine with ammonium ions to orm undissociated ammonia. In both cases the hydroxide ion

concentration and the hydrogen ion concentration remain constant.

🕕 Proteins and enzymes (2)

BUFFER CALCULATIONS

The equilibrium expression or weak acids also applies to acidic bu er solutions,

e.g. ethanoic acid/sodium ethanoate solution.

$$\mathbf{K}_{a} = \frac{[\mathbf{H}^{+}] \quad [\mathbf{CH}_{3}\mathbf{COO^{-}}]}{[\mathbf{CH}_{3}\mathbf{COOH}]}$$

The essential di erence is that now the concentrations o the two ions rom the acid will not be equal.

Since the sodium ethanoate is completely dissociated the concentration o the ethanoate ions in solution will be almost the same as the concentration o the sodium ethanoate, as very little will come rom the acid.

I logarithms are taken and the equation is rearranged then:

$$pH = pK_a + \log_{10} \frac{[CH_3COO^-]}{[CH_3COOH]}$$

This is known as the Henderson Hasselbalch equation (the general ormula can be ound in Section 1 o the IB data booklet).

Two acts can be deduced rom this expression. Firstly the pH o the bu er does not change on dilution, as the concentration o the ethanoate ions and the acid will be a ected equally.

BLOOD

An important bu er is blood, which only unctions correctly within a very narrow pH range. Blood is a complex bu ering system, which is responsible or carrying oxygen around the body. One o the components o the system is that the oxygen adds on reversibly to the haemoglobin in the blood.

 $HHb + O_2 = H^+ + HbO_2^-$

PROTEIN ASSAY BY UV-VIS SPECTROSCOPY

$$\log_{10} \frac{I_{\circ}}{I} = Ic$$

where: I_0 is the intensity o the incident radiation and I is the intensity o the transmitted radiation.

is the molar absorption coe ficient (a constant or each absorbing substance).

l is the path length o the absorbing solution (usually 1.0 cm) and *c* is the concentration.

Most spectrometers measure $\log_{10} I_0 / I$ directly as absorbance. I the path length is kept the same by using the same cuvette (sample tube) and all the readings are taken at max then it is easy to see that the measured absorbance is directly proportional to the concentration. Using Coomassie Brilliant Blue the Beer Lambert Law holds true or solutions o protein covering the Secondly the bu er will be most e ficient when $[CH_3 COO^-] = [CH_3 COOH]$. At this point, which equates to the hal equivalence point when ethanoic acid is titrated with sodium hydroxide, the pH o the solution will equal the pK_3 value o the acid.

Calculate the pH o a bu er containing 0.200 mol o sodium ethanoate in 500 cm³ o 0.100 mol dm⁻³ ethanoic acid (given that K_a or ethanoic acid = 1.8 10^{-5} mol dm⁻³).

$$[CH_{3}COO^{-}] = 0.400 \text{ mol } dm^{-3}; [CH_{3}COOH] = 0.100 \text{ mol } dm^{-3}$$

$$K_{a} = \frac{[H^{+}] \quad 0.400}{0.100} = 1.8 \quad 10^{-5} \text{ mol dm}^{-5}$$
$$[H^{+}] = 4.5 \quad 10^{-6} \text{ mol dm}^{-3}$$
$$pH = 5.35$$

Calculate what mass o sodium propanoate must be dissolved in 1.00 dm³ o 1.00 mol dm⁻³ propanoic acid (p $K_a = 4.87$) to give a bu er solution with a pH o 4.5.

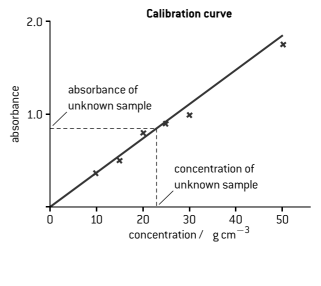
$$[C_{2}H_{5}COO^{-}] = \frac{K_{a}}{[H^{+}]} = \frac{10^{-4.87}}{10^{-4.5}} = 0.427 \text{ mol dm}^{-3}$$

Mass o NaC₂H₂COO required = 0.427 96.07 = 41.0 g

I the pH increases ([H⁺] alls) the equilibrium will move to the right and the oxygen will tend to be bound to the haemoglobin more tightly. I the pH decreases ([H⁺] increases) the oxygen will tend to be displaced rom the haemoglobin. Both o these processes are potentially li e threatening.

range rom 0 to approximately 1500 $\,\mathrm{g}\,\mathrm{cm}^{-3}$.

To find the concentration o the solution o the protein with unknown concentration, it is there ore necessary to first obtain a calibration curve by using a range o known concentrations o protein and measuring the associated absorbance. A line o best fit is obtained and once the absorbance o the unknown sample has been measured its concentration can be determined by interpolation o the graph.

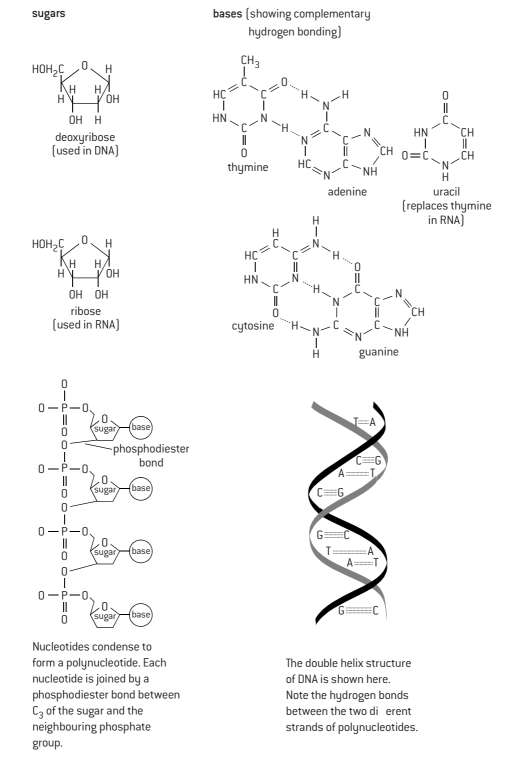




STRUCTURE OF NUCLEOTIDES AND NUCLEIC ACIDS

Almost all cells in the human body contain DNA (deoxyribonucleic acid). DNA and a related material RNA (ribonucleic acid) are macromolecules with relative molar masses o up to several million. Both nucleic acids are made up o repeating base-sugarphosphate units called nucleotides. A nucleotide o DNA contains the condensation products o deoxyribose (a pentose sugar), phosphoric acid, and one o our nitrogen-containing bases, adenine (A), guanine (G), cytosine (C) or thymine (T). RNA contains a di erent sugar, ribose, but also contains a phosphate group and our nitrogen-containing bases. Three o the bases are the same as those in DNA but the ourth, uracil (U), replaces thymine.

In DNA, the polynucleotide units are wound into a helical shape with about 10 nucleotide units per complete turn. Two helices are then held together by hydrogen bonds between the bases to give the characteristic double helix structure. The stability o this double helix structure is due to the base-stacking interactions between the hydrophilic and hydrophobic components as well as hydrogen bonding between the nucleotides. The hydrogen bonds are very specific. Cytosine can only hydrogen bond with guanine and adenine can only hydrogen bond with thymine (uracil in RNA). Unlike DNA, RNA normally exists as single polynucleotide chains.





THE GENETIC CODE

When cells divide, the genetic in ormation has to be replicated intact. The genetic in ormation is stored in chromosomes ound inside the nucleus. In humans there are 23 pairs o chromosomes. Chromosomes are e ectively a very long DNA sequence. The DNA is compacted e ciently in the eukaryotic nucleus by orming DNA-protein complexes with histones. Histones are positively charged proteins that bond tightly to the negatively charged phosphate groups in the DNA's phosphate-sugar backbone. The DNA in the cell starts to partly unzip as hydrogen bonds between the bases break. Sugar-base units will be picked up rom the aqueous solution to orm a complementary new strand. Because adenosine can only hydrogen bond with thymine (A T) and cytosine can only hydrogen bond with guanine (C G) the new strand ormed will be identical to the original. Thus i the sequence o bases in one strand is -C-G-A-T-T-A- the complementary strand will have the sequence -G-C-T-A-A-T-.

The in ormation required to make complex proteins is passed rom the DNA to messenger RNA by a similar unzipping process, known as transcription, except that the new strand o mRNA contains a di erent sugar and uracil in place o thymine.

The coded in ormation held in the mRNA is then used to direct protein synthesis

using a triplet code by a process known as translation. Each sequence o three bases

represents one amino acid and is known as the triplet code. The triplet code allows or up to 64 permutations known as codons. This is more than su cient to represent the 20 amino acids and several di erent codons may represent the same amino acid. Consecutive DNA codons o AAA, TAA, AGA, GTG, and CTT will transcribe to RNA codons o UUU, AUU, UCU, CAC, and GAA which will cause part o a strand o a protein to be ormed that contains the amino acid residues Phe-Ile-Ser-His-Glu-.

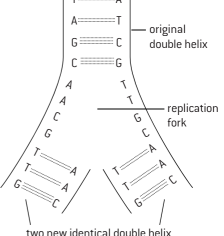
UUU	Phe	UCU	Ser	UAU	Tyr	UGU	Cys
UUC	Phe	UCC	Ser	UAC	Tyr	UGC	Cys
UUA	Leu	UCA	Ser	UAA	Terminator	UGA	Terminator
UUG	Leu	UCG	Ser	UAG	Terminator	UGG	Trp
CUU	Leu	CCU	Pro	CAU	His	CGU	Arg
CUC	Leu	CCC	Pro	CAC	His	CGC	Arg
CUA	Leu	CCA	Pro	CAA	GIn	CGA	Arg
CUG	Leu	CCG	Pro	CAG	GIn	CGG	Arg
AUU	lle	ACU	Thr	AAU	Asn	AGU	Ser
AUC	lle	ACC	Thr	AAC	Asn	AGC	Ser
AUA	lle	ACA	Thr	AAA	Lys	AGA	Arg
AUG	Met	ACG	Thr	AAG	Lys	AGG	Arg
GUU	Val	GCU	Ala	GAU	Asp	GGU	Gly
GUC	Val	GCC	Ala	GAC	Asp	GGC	Gly
GUA	Val	GCA	Ala	GAA	Glu	GGA	Gly
GUG	Val	GCG	Ala	GAG	Glu	GGG	Gly

The genetic code carried by RNA

GENETICALLY MODIFIED FOODS

Genetic engineering involves the process o selecting a single gene or a single characteristic and trans erring that sequence o DNA rom one organism to another. Thus a genetically modi ed (GM) ood can be de ned as one derived or produced rom a genetically modi ed organism. The GM ood can be substantially di erent or essentially the same in composition, nutrition, taste, smell, texture and unctional characteristics to the conventional ood. An example o genetically modi ed ood is the FlavrSavr tomato. In normal tomatoes, a gene is triggered when they ripen to produce a substance that makes the ruit go so t and eventually rot. In the FlavrSavr tomato the gene has been inhibited to produce a tomato with a uller taste and a longer shel li e.

Benefits o GM oods Potential concerns o GM oods With crops, it can enhance the taste, flavour, texture and The outcome o alterations is uncertain as not enough is nutritional value and also increase the maturation time. known about how genes operate. Plants can be made more resistant to disease, herbicides They may cause disease as antibiotic-resistant genes and insect attack. could be passed to harm ul microorganisms. With animals, GM oods can increase resistance to Genetically engineered genes may escape to disease, increase productivity and eed e ciency to give contaminate normal crops with unknown e ects. higher yields o milk and eggs. They may alter the balance o delicate ecosystems as Anti-cancer substances and increased amounts o ood chains become damaged. vitamins (such as vitamin A in rice) could be incorporated There are possible links to an increase in allergic and exposure to less healthy ats reduced. reactions (particularly with those involved in ood Environmentally riendly bio-herbicides and bioprocessing). insecticides can be ormed. GM oods can lead to soil, water and energy conservation and improve natural waste management.



two new identical double helix strands forming

Replication of DNA

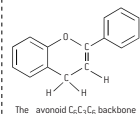
🖽 Biological pigments (1)

ULTRAVIOLET AND VISIBLE ABSORPTION IN ORGANIC MOLECULES

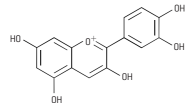
Organic compounds containing unsaturated groups such as C=C, C=O, N=N, NO, and the benzene ring can absorb in the ultraviolet or visible part o the spectrum. Such groups are known as chromophores and the precise energy o absorption is a ected by the other groups attached to the chromophore. The absorption is due to electrons in the bond being excited to an empty orbital o higher energy, usually an anti-bonding orbital. The energy involved in this process is relatively high and most organic compounds absorb in the ultraviolet region and thus appear colourless. For example ethene absorbs at 185 nm. However, i there is extensive conjugation o double bonds (i.e. many alternate C C single bonds and C=C double bonds) in the molecule involving the delocalization o pi electrons then less energy is required to excite the electrons and the absorption occurs in the visible region. Biological pigments are coloured compounds produced by metabolism. Good examples include anthocyanins, carotenoids, chlorophyll and haem. One other obvious example is the pigment melanin, which is responsible or di erent tones o skin, eye and hair colour.

ANTHOCYANINS

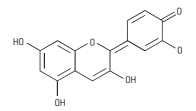
Anthocyanins are aromatic, water-soluble pigments widely distributed in plants. They contain the flavonoid C₂C₂ skeleton.



The



Structure of cyanidin in acidic solution. Less conjugation so absorbs in blue-green region and transmits red light.



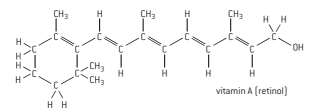
Structure of cyanidin in alkaline solution. More conjugation so absorbs in the orange region of the spectrum and transmits blue light.

It is the conjugation o the pi electrons contained in this structure that accounts or the colour o anthocyanins. The more extensive the conjugation, the lower the energy (longer the wavelength) o the light absorbed. This can be exempli ed using cyanidin. In acidic solution it orms a positive ion and there is less conjugation than in alkaline solution where the pi electrons in the extra double bond between the carbon and oxygen atom are also delocalized.

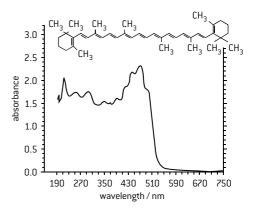
This di erence in colour depending on pH explains why poppies that have acidic sap are red whereas cornflowers, which also contain cyanidin but have alkaline sap, are blue. Other anthocyanins di er in the number and types o other groups such as hydroxyl or methoxy groups, which a ect the precise wavelength o the light absorbed and hence the colour transmitted. Because their precise colour is so sensitive to pH changes anthocyanins can be used as indicators in acid base titrations. The addition o other groups also a ects other properties o anthocyanins. The basic flavonoid C₆C₇C₆ backbone is essentially non-polar. As more polar hydroxyl groups are added the potential or them to orm hydrogen bonds with water molecules increases and many anthocyanins, such as cyanidin with several OH groups, are appreciably soluble in water or this reason.

CAROTENOIDS

Carotenoids are lipid-soluble pigments, and are involved in harvesting light in photosynthesis. The conjugation in carotenoids is mainly due to a long hydrocarbon chain (as opposed to the ring system in anthocyanins) consisting o alternate single and double carbon to carbon bonds.



The majority o carotenoids are derived rom a (poly)ene chain containing orty carbon atoms, which may be terminated by cyclic end groups and may also be complemented with oxygen-containing unctional groups. The hydrocarbon carotenoids are known as xanthophylls. Examples include -carotene, -carotene and vitamin A. - and -carotene and vitamin A are all lipid-soluble and not water-soluble. Although vitamin A does contain one polar hydroxyl group the rest o the molecule is a large non-polar hydrocarbon. Because o the unsaturation in the double bond carotenoids are susceptible to oxidation. This oxidation process can be catalysed by light, metals and hydroperoxides. It results in a change o colour, loss o activity in vitamin A and is the cause o bad smells.



-carotene is found in carrots and has a characteristic orange colour. It contains eleven conjugated double bonds and absorbs strongly in the violet-blue (400 510 nm) region.

🖽 Biological pigments (2)

CHLOROPHYLL AND HAEM

Porphyrin compounds, such as haemoglobin, myoglobin, chlorophyll and many cytochromes are chelates o metals with large nitrogen-containing macrocyclic ligands. Porphyrins contain a cyclic system in which all the carbon atoms are sp² hybridized. This results in a planar structure with extensive pi conjugation. The non-bonding pairs o electrons on the our nitrogen atoms enable the porphyrin to orm coordinate bonds with metal ions. Chlorophyll contains a magnesium ion and its structure is given in Section 35 o the IB data booklet. It is ound in two closely related orms. In chlorophyll a, the -R group is a methyl group, $-CH_3$, and in chlorophyll b the -R group is an aldehyde group, -CHO. Chlorophyll is essential or photosynthesis. Its unction is to absorb light energy and undergo a redox reaction to donate an electron through a series o intermediates in an electron transport chain. Cytochromes in the electron transport chain contain haem groups in which the iron ion interconverts between iron(II) and iron(III) during redox reactions.

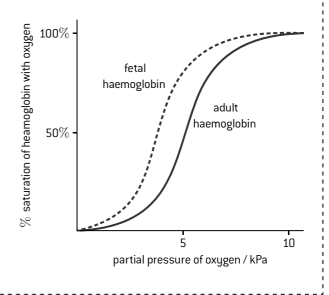
Haemoglobin, which is ound in the blood, carries oxygen rom the respiratory organs to the rest o the body. Like myoglobin (which is ound in muscles) and some o the cytochromes, it contains haem (also spelt heme) groups with the porphyrin group bound to an iron(II) ion. When oxygen binds to one o the iron atoms in the complex to orm HbO₂ it causes the iron atom to move towards the centre o the porphyrin ring and at the same time the imidazole side-chain o a histidine residue is pulled towards the porphyrin ring. This produces a strain that is transmitted to the remaining three monomers in the quaternary haemoglobin. This brings about a similar con ormational change in the other haem sites so that it is easier or a second oxygen molecule to bind to a second iron atom. Each time the haem group s a finity to attract oxygen increases as the remaining sites become filled. Haem becomes ully saturated with oxygen when all our iron atoms have been utilized orming HbO₈. The binding o oxygen is, thus, a cooperative process. Various actors a ect the amount o oxygen that binds. Low pH and a relatively high pressure o carbon dioxide (i.e. during exhalation) cause oxygen to be released rom the haemoglobin into the tissues. Conversely at lower carbon dioxide pressure (which causes the pH to rise) more oxygen is taken up by the haemoglobin. Temperature also has

an e ect. At higher temperatures more oxygen is released rom haemoglobin. When muscles are metabolically active they emit energy (heat) and the haemoglobin provides them with the increased oxygen required.

The partial pressure also a ects the oxygen uptake. Because o the cooperative process achieved through the induced changes o the haemoglobin protein complex the oxygen binding curve o saturation o haemoglobin with oxygen against partial pressure is sigmoidal in shape compared with the normal hyperbolic curve expected i no cooperative binding takes place.

Haemoglobin exists in a slightly di erent orm in etal blood. It has a greater a finity or oxygen than normal haemoglobin so more oxygen is bound to the haemoglobin at lower partial pressures. This enables the etal blood in the placenta to take up oxygen rom the mother s blood.

Carbon monoxide is a dangerous poison as carbon monoxide is a stronger ligand than oxygen and orms an irreversible complex with the iron in haemoglobin. It thus acts as a competitive inhibitor and prevents the haemoglobin rom binding with oxygen.



PAPER AND THIN LAYER CHROMATOGRAPHY

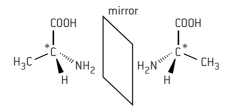
The technique o paper chromatography used to identi y amino acids has already been discussed on page 126. It is ideal to use to separate and identi y biological pigments by measuring *R* values as no dye or stain is needed to see the spots. In paper chromatography the stationary phase is the water contained in the cellulose fibres in the paper. Thin layer chromatography can also be used. This is similar to paper chromatography but uses a thin layer o a solid, such as alumina, Al_2O_3 , or silica, SiO_2 , on an inert support such as glass. When absolutely dry it works by adsorption but, like paper, silica and alumina have a high a finity or water, there ore the separation occurs more by partition with water as the stationary phase. The choice o a suitable solvent depends on the polarity or otherwise o the particular pigments. One real advantage o thin layer chromatography over paper chromatography is that each o the separated components can be recovered pure. The section containing the component is scraped o the glass and then dissolved in a suitable solvent. The solution is then filtered to remove the solid support and the solvent can then be evaporated to leave just the pure component.

🕕 Stereochemistry in biomolecules

CHIRALITY

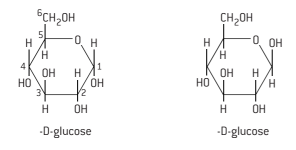
A chiral carbon atom is asymmetric, i.e. contains our di erent atoms or groups attached to it. Many important biological molecules are chiral and only one o the particular enantiomers is normally active in nature, although sometimes a di erent enantiomer may have a detrimental e ect. This was the case with the drug thalidomide, which was prescribed in the 1950s and 1960s. One enantiomer alleviated the e ects o morning sickness in pregnant women, the other enantiomer caused severe de ects in the etus.

All amino acids apart rom glycine, $H_2N CH_2 COOH$, exist as enantiomers but only the L configuration is ound in proteins.

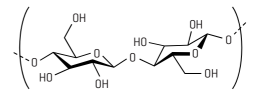


enantiomers of alanine (* asymmetric carbon/chiral carbon)

Sugars contain several chiral carbon atoms. The D and L stereoisomers o sugars re er to the configuration o the chiral carbon atom urthest rom the aldehyde or ketone group. The D orms occur most requently in nature. The stereochemistry o sugars is urther complicated by the position o the hydroxyl groups. The ring orms o sugars have isomers, known as and , depending on whether the position o the hydroxyl group at carbon 1 (glucose) or carbon 2 (ructose) lies below the plane o the ring ().



Polysaccharides ormed rom glucose can be very di erent depending upon whether the - or - orm is involved. Starch, which can be digested by humans, is ormed rom polymerizing -D-glucose. Amylose is a straight-chain polymer O -D-glucose with -1,4 glycosidic bonds and amylopectin is also derived rom -D-glucose with both -1,4 and -1,6 glycosidic bonds. Cellulose is also a polymer O glucose but it is ormed rom -D-glucose with -1,4 linkages.

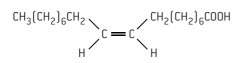


A repeating unit of cellulose showing the -1,4-linkage

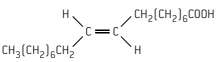
Cellulose, together with lignin, provides the structure to the cell walls o green plants. Most animals, including all mammals, do not have the enzyme cellulose so are unable to digest cellulose. Known as roughage, dietary fibre does play an important role in the diet as it aids digestion and makes de ecation easier.

CIS- AND TRANS- ISOMERISM

Fatty acids occur naturally as the cis-isomers but as described on page 129 trans-isomers, which can increase the risk o heart disease, can be ormed during the partial hydrogenation o unsaturated ats. Like saturated atty acids, the trans- acids are straighter than their bent cis- isomers. This means that they can pack together more easily and so have higher melting points. For example elaidic acid (trans-9-octadecenoic acid) melts at 45 C whereas oleic acid (cis-9-octadecenoic acid) melts at 13 C.



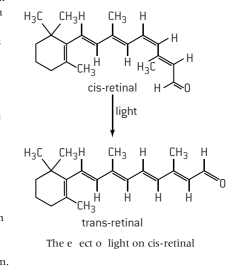
oleic acid (melting point 13 C)



elaidic acid (melting point 45 C)

The process whereby photons o light are converted into electrical signals in the retina at the back o the eye is called the visual cycle. Rhodopsin in the retina consists o a protein, opsin and a covalently bonded co- actor retinal, which is produced in the retina rom

vitamin A. When light alls on the retina it converts the carotenoid retinal rom the cis- to the trans- orm and as this happens a nerve impulse is transmitted via an interaction. which causes a con ormational change in the structure o opsin to send a signal along the optic nerve to the brain.



SHORT ANSWER QUESTIONS OPTION B BIOCHEMISTRY

- 1. a) Deduce the structure o methionine in
 - (i) acid solution (pH < 4)
 - (ii) at the isoelectric point (pH = 5.7)
 - (iii) alkaline solution (pH > 9)

[3]

[2]

[1]

[4]

[2]

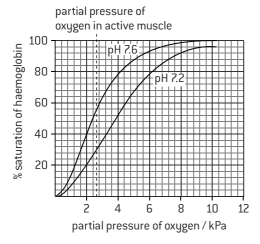
- b) Draw the two dipeptides that can be ormed when one molecule o methionine condenses with one molecule o alanine.
- c) Determine the number o di erent tripeptides that could be ormed rom methionine, alanine and cysteine i each tripeptide contains one residue rom each o the three amino acids.
- d) Design an experiment you could use in a school laboratory to determine whether a given protein contains a methionine residue.
- 2. The diagram below shows a triacyclglycerol (triglyceride).

$$CH_3(CH_2)_4CH = CHCH_2CH = CH(CH_2)_7CO - O - CH_2$$

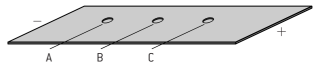
CH₃(CH₂)₇CH=CH(CH₂)₇CO-O-CH₂

- a) State the main dietary group this compound belongs to. [1]
- b) Deduce whether this compound is likely to have come rom an animal or vegetable source. [2]
- c) Deduce whether this compound is likely to be a solid or a liquid at room temperature. [2]
- d) List two major unctions o this class o compounds in the body. [2]
- 3. The structures o vitamin A (retinol) and vitamin C (ascorbic acid) are given in Section 35 o the IB data booklet.

- a) Iron combines reversibly with oxygen in haemoglobin. Other than variable oxidation states, state two typical characteristics o transition metals that are shown by iron in haemoglobin.
 - b) The ability o haemoglobin to carry oxygen at body temperature depends on the concentration o oxygen, the concentration o carbon dioxide and on the pH. The graph shows how the percentage saturation o haemoglobin with oxygen changes with pH at di erent partial pressures o oxygen.



- a) (i) Identi y two unctional groups present in retinol. [2]
 - (ii) Classi y vitamin A and vitamin C as water or at soluble and justi y the di erence on the molecular level. [3]
 - (iii) State one physical symptom or each o vitamin A and vitamin C deficiency and state the common name given to vitamin C deficiency.
 [3]
- b) Interpret the in ormation that 0.014 moles o a particular oil was ound to react exactly with 14.2 g o iodine.
- 4. a) A mixture o the amino acids serine (Ser), glutamic acid (Glu) and lysine (Lys) was separated using electrophoresis and a bu er o pH 5.7. A drop containing the mixture was placed in the centre o the paper and a potential di erence was applied. The amino acids were developed and the ollowing results were obtained.



- (i) Describe how the amino acid spots may have been developed.
- (ii) Predict which amino acid is present at spot C.Explain your answer. [3]

[1]

[1]

- (iii) The amino acid at spot B is at its isoelectric point.Describe one characteristic o an amino acid at its isoelectric point.
- b) Explain, using equations, how the amino acid glycine (Gly) can act as a bu er. [2]
 - (i) The partial pressure o oxygen in active muscle is shown by the dotted line at 2.8 kPa. Calculate the di erence in the percentage saturation o haemoglobin with oxygen in active muscle when the pH changes rom 7.6 to 7.2. [1]
 (ii) When the cells in muscles respire they excrete
 - carbon dioxide and sometimes lactic acid as waste products. Explain how this a ects the ability o haemoglobin to carry oxygen. [2]
- 6. The acid dissociation constant, K_a , or lactic acid, CH(CH₃)(OH)COOH, is 1.38 10^{-4} at 298 K.
 - a) Explain why lactic acid can exist in two di erent enantiomeric orms.
 - b) Calculate the pH o 0.100 mol dm⁻³ lactic acid solution at 298 K. [3]
 - c) Determine the mass o lactic acid that must be added together with 2.00 g o sodium hydroxide to make 500 cm³ o a bu er solution at 298 K with a pH o 4.00. [4]

14 OPTION C ENERGY

Energy sources

INTRODUCTION

Energy is the ability to do work, and is measured in joules (J). The first Law o Thermodynamics states that energy within a closed system can neither be created nor destroyed. Living organisms require energy to remain alive. This energy is obtained rom the metabolism o ood. Societies also need energy to unction. Use ul sources o energy must release energy at reasonable rates and produce minimal pollution. Energy sources are either renewable, that is naturally replenished, or non-renewable, that is finite. Apart rom tidal energy and nuclear energy, the Sun is the ultimate source o energy or the planet. The Sun is also responsible or the Earth s climate and ecosystems. Humans use o energy, particularly when derived rom ossil uels, inter eres with both the climate and ecosystems. Types o energy include kinetic and potential but energy sources re ers specifically to ways o obtaining energy to produce electrical power.

ENERGY SOURCES

At the moment, oil is the top energy source used in the world. The order or the main energy sources currently used in the world in terms o producing energy is: crude oil > coal > natural gas > hydroelectric > nuclear fission > wind > bio uels > solar > geothermal (source: oilprice.com). Crude oil, coal and natural gas are non-renewable ossil uels and contain stored energy rom photosynthesis trapped millions o years ago. Nuclear fission is non-renewable as it depends upon a supply o fissionable material. Hydroelectric, wind, bio uels, solar and geothermal are all renewable energy sources. Other orms o renewable energy that may be used more in the uture are tidal and wave energy. When the energy rom any o these sources is converted into electrical energy, some o the energy is lost as heat to the surroundings so none o the processes is 100% e ficient. Heat is also lost as the generated electricity is transported by power lines.

The choice o uel depends on many actors including availability, ease o extraction, storage and the environmental and social e ects o extracting and using the energy as well as physical actors. These physical actors include:

The energy density = $\frac{\text{energy released rom the uel}}{\text{volume o uel consumed}}$

The specific energy = $\frac{\text{energy released rom the uel}}{\text{mass o uel consumed}}$

The e ficiency o an energy trans $er = \frac{use \ ul \ output \ energy}{total \ input \ energy}$ 100 (expressed as a percentage)

For example, the specific energy and energy density o ethanol (bio uel) and gasoline (ossil uel) can be compared using their enthalpies o combustion, molar masses and densities.

	Ethanol C ₂ H ₅ OH	Gasoline (octane) C _s H ₁₈
M _r	46.08	114.26
$H_{c}^{\Theta}/\text{ kJ mol}^{1}$	-1367	-5470
Density / g cm ³	0.789	0.703
Specific energy / kJ kg ¹	29670	47890
Energy density / kJ dm ³	37600	68120

Clearly on all physical counts gasoline (octane) is the better uel. It produces 1.6 times as much energy as ethanol by mass and 1.8 times as much energy by volume. Both produce carbon dioxide (a greenhouse gas) when burned. Gasoline adds to the total amount o carbon dioxide in the atmosphere as it is a ossil uel whereas ethanol is ormed rom crops grown now so does not significantly alter the amount o carbon dioxide in the atmosphere and so is more environmentally riendly. Weighed against this is the act that the large scale use o ethanol uses up land to grow crops that could otherwise be used to grow ood.

Nuclear uel is a very e ficient uel in terms o specific energy density but has many sa ety and pollution issues associated with it. Hydrogen also has a high specific energy (143 000 kJ kg⁻¹) but there are problems with the transport and storage o hydrogen as it cannot be liquefied by pressure alone and the methods o manu acturing hydrogen are also energy intensive. Wind and solar energy depend upon weather conditions and so the ways o storing the energy they produce are also important.

The generation o electric power involves using mechanical energy to drive electrical generators. For energy sources such as coal, oil, natural gas and bio uels, the mechanical energy is obtained by using the heat produced by combustion. Nuclear uels use the heat generated by the nuclear reactions. Wind and hydroelectric power can bypass this step as they produce mechanical energy by driving a turbine directly. Coal and oil-fired power stations typically can only convert about 33% o their energy output into electricity, i.e. their e ficiency o energy trans er is 33%. By using a combined cycle involving synthesis gas this can be increased to about 60%.

Fossil fuels (1)

FORMATION OF FOSSIL FUELS

Coal is ossilized plant material containing mainly carbon together with hydrogen, nitrogen and sul ur. Most coal was ormed during the Carboni erous period (286–360 million years ago). The action o pressure and heat through geological orces converted the plant material in stages rom peat to lignite to bituminous so t coal to hard coal (anthracite). At each stage the percentage o carbon increases. Coal contains between 80 and 90% carbon by mass.

Crude oil was ormed rom the remains o marine organisms mainly during the Paleozoic era, up to 600 million years ago. Thick sediments built up on top o the organic layers and under the action o high pressures and biochemical activity crude oil was ormed. The oil migrated through rocks due to earth movements and collected in traps. Crude oil is a complex mixture o straight-chain, branched, cyclic and aromatic hydrocarbons, although it consists mainly o alkanes.

Natural gas was ormed at the same time as crude oil and the two are o ten ound together, although it may occur on its own or with coal. It consists mainly o methane (85 95%) with varying amounts o ethane, propane, butane and other gases such as hydrogen sul de.

COMPOSITION AND CHARACTERISTICS OF CRUDE OIL FRACTIONS

Sul ur must rst be removed rom crude oil be ore it is rened as it can poison catalysts by blocking their active sites. The oil then undergoes ractional distillation to separate it into dierent boiling ractions. The number of carbon atoms, boiling ranges and the uses of the dierent ractions are summarized in the table:

Fraction	Carbon chain length	Boiling range / C	Main uses
Renery gas	14	<30	Used as uel on site, gaseous cooking uel and as eedstock or chemicals e.g. methane used to provide hydrogen gas or the Haber process.
Gasoline and naphtha	5 10	40 180	Gasoline (petrol) or cars. Feedstock or organic chemicals (by steam cracking).
Kerosene	11 12	160 250	Fuel or jet engines; domestic heating; cracked to provide extra gasoline (petrol).
Gas oil (diesel oil)	13 25	220 350	Diesel engines and industrial heating; cracked to produce extra gasoline (petrol).
Residue	>20	>350	Fuel or large urnaces; vacuum distilled to make lubricating oils and waxes. Residue o bitumen and asphalt used to sur ace roads and waterproo roo s.

CRACKING AND REFORMING

The per ormance o hydrocarbons as uels is improved by cracking and catalytic re orming reactions.

Cracking is the process conducted at high temperatures whereby large hydrocarbons are broken down into smaller more use ul molecules. The products are usually alkanes and alkenes. For example decane can be broken down to orm octane and ethene.

$$C_{_{10}}H_{_{22}}(g) \qquad C_{_8}H_{_{18}}(g) + C_{_2}H_{_4}(g)$$

The alkanes are usually branched isomers (e.g. 2,2,4-trimethylpentane) and are added to octane to improve the octane rating. The alkenes are used to make other chemicals, particularly addition polymers.

More use ul branched alkanes can be obtained rom straight-chain hydrocarbons by mixing them with hydrogen and heating them at 770 K over a platinum/alumina catalyst at high pressure. This process, known as isomerization, is a particular orm o re orming. Other re orming processes in which straight-chain alkanes are re ormed into molecules with the same number o carbon atoms include cyclization to make ring molecules and aromatization to make benzene.

Examples of reforming $CH_3 \{CH_2\}_4 CH_3$ hexane isomerization CH_3 aromatization CH_3 CH_2 CH_2 CH_2 CH_3 CH_2 CH_2 CH_3 CH_3 CH_2 CH_3 CH

CARBON FOOTPRINT

A carbon ootprint is the total amount o greenhouse gases $(CO_2 \text{ and } CH_4)$ produced during human activities. Carbon ootprints can be estimated



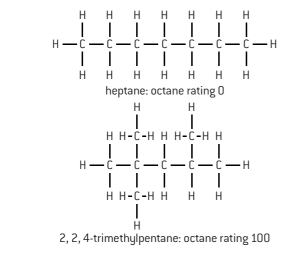
using online calculators and are generally expressed in equivalent tons o carbon dioxide. For most people the majority o their carbon ootprint comes indirectly rom the uel used to produce and transport goods compared with the emissions which come directly rom using gasoline in their own car or taking a flight. Fossil uels have by ar the highest carbon ootprint whereas nuclear power and renewable sources o energy such as wind and hydroelectric power all contribute little towards the carbon ootprint. Consumers can opt to o set their carbon ootprint by, or example, planting trees. The best ways to lower carbon ootprints are to reduce the use o ossil uels and materials in general, reuse materials and recycle.

Fossil fuels (2)

OCTANE RATING

In an e cient internal combustion engine a spark ignites the uel-air mixture just be ore the piston reaches 'top dead centre' so that the ull orce o the explosion pushes the piston down just as it reaches the top o the cylinder. Under the conditions o high temperature and pressure the reaction may start be ore the spark and the engine will be less e cient. This is known as pre-ignition or **knocking**. The more straight-chain the alkane the higher the tendency or knocking.

Fuels are classi ed according to their **octane number**. Generally the more branched the alkane the higher the octane number. Pure heptane is assigned an octane number o zero and an isomer o octane, 2,2,4-trimethylpentane, has an octane rating o 100. Thus gasoline (petrol) with an octane rating o 95 will burn as e ciently as a mixture o 95% 2,2,4-trimethylpentane and 5% heptane. In the past tetraethyllead Pb(C_2H_5)₄ was added to petrol to raise the octane rating. Lead- ree gasoline (petrol) contains added aromatic hydrocarbons such as benzene and more branched hydrocarbons obtained through cracking.



COAL GASIFICATION AND LIQUEFACTION

Be ore the advent o plenti ul supplies o natural gas, coal was routinely turned into 'coal gas'. As supplies o natural gas diminish, interest in coal gasi cation may revive. Coal gas (also known as synthesis gas, water gas or town gas) contains a mixture o hydrogen and carbon monoxide and is made by heating coal in the presence o steam.

 $C(s) + H_2O(g) - CO(g) + H_2(g)$

Reacting coal gas with more hydrogen in the presence o a heated catalyst converts it into SNG (substitute or synthetic natural gas).

 $CO(g) + 3H_2(g) - CH_4(g) + H_2O(g)$ SNG can also be made by heating crushed coal in steam at 700 C using potassium hydroxide as a catalyst.

$$2C(s) + 2H_2O(g) = CH_4(g) + CO_2(g)$$

SNG is a cleaner gas (as it removes pollution due to sul ur dioxide), which is easier to transport but the process is less e cient as it uses up some 30% o the available energy during the conversion process.

In Germany in the 1930s and in South A rica, where coal is abundant, coal has been converted into a liquid uel. As the price o oil increases this process may become more important economically. The method is known as the Fischer Tropsch process. Synthesis gas is reacted with more steam to increase the proportion o hydrogen in the mixture.

 $CO(g) + H_2O(g) = CO_2(g) + H_2(g)$

The hydrogen and carbon monoxide are then passed into a fluidized bed reactor containing iron or cobalt catalysts to produce a mixture o hydrocarbons that can be separated by ractional distillation.

Fossil fuel	Advantages	Disadvantages	
Coal	 Present in large quantities and distributed throughout the world. Can be converted into synthetic liquid uels and gases. Feedstock or organic chemicals. Has the potential to yield vast quantities o energy compared with renewable sources and sa er than nuclear power. Longer li espan (350 years?) compared with oil or gas 	 Contributes to acid rain and global warming. Not so readily transported (no pipelines). Coal waste (slag heaps) lead to ground acidity and visual and chemical pollution. Mining is dangerous cave-ins, explosions and long term e ect o coal dust on miners. Dirty (produces dust, smoke and particulates). 	
Oil	 Easily transported in pipelines or by tankers. Convenient uel or use in cars, lorries, etc. Feedstock or organic chemicals. 	 Contributes to acid rain and global warming. Limited li espan (30 50 years?) and uneven distribution worldwide. Risk o pollution associated with transportation by tankers. 	
Natural gas	 Clean uel Easily transported in pipelines and pressurized containers. Does not contribute to acid rain. Releases a higher quantity o energy per kg than coal or oil. 	 Contributes to global warming. Limited li espan (30 years?) and uneven distribution worldwide. Greater risk o explosions due to leaks. 	

RELATIVE ADVANTAGES AND DISADVANTAGES OF FOSSIL FUELS

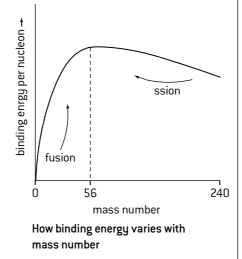
Nuclear usion and nuclear fission (1)

NUCLEAR REACTIONS

In a normal chemical reaction, valence shell electrons are rearranged as bonds are broken and new bonds ormed. There is no change in the nucleus and no new elements are ormed. In nuclear reactions the nucleus itsel rearranges. The protons and neutrons in the nucleus are held together by strong orces known as the binding energy.

A graph o binding energy per nucleon against mass number shows that atomic nuclei with a mass number o approximately 56 (i.e. the nucleus o iron) have the maximum binding energy and are thus the most stable. Nuclei to the le t or right o this maximum will undergo nuclear change in such a way that, as they approach the maximum, energy will be released.

This explains nuclear usion whereby small atoms combine to orm heavier nuclei and nuclear fission the splitting o heavy nuclei to orm two or more lighter nuclei. In both cases the total mass o the products is less than the mass o



the initial nucleus or nuclei. During nuclear reactions, mass is converted into energy according to Einstein s equation $E = mc^2$ (where *c* is the velocity o light). Thus in a nuclear reaction new elements are ormed and the energy change is potentially much greater than in a chemical reaction. When an unstable radioactive isotope decays it can emit three di erent types o radiation. Alpha particles which are helium nuclei, ${}^4_2\text{He}{}^{2+}$, beta particles which are electrons, ${}^0_{-1}\text{e}{}^-$, and gamma radiation, which is high energy electromagnetic radiation.

NUCLEAR EQUATIONS

Nuclear equations must balance. The total mass numbers and the nuclear charge numbers (atomic numbers) must be equal on both sides o the equation. During particle emission the new element will have a mass o our less than the original element and an atomic number that is two less.

e.g. ${}^{238}_{92}$ U ${}^{234}_{90}$ Th + ${}^{4}_{2}$ He

During emission when an electron is ejected rom the nucleus at a velocity approaching the speed o light, the new element will have the same mass number but the atomic number will have increased by one.

e.g.
$${}^{14}_{6}C$$
 ${}^{14}_{7}N + {}^{0}_{-1}e$

Nuclear reactions also occur artificially when nuclei are bombarded with other small particles such as particles or neutrons. In each case the total mass numbers and nuclear charges on both sides o the equation must still balance.

e.g. ${}^{235}_{92}U + {}^{1}_{0}n = {}^{144}_{56}Ba + {}^{90}_{36}Kr + 2{}^{1}_{0}n$

Other small particles that may be involved in nuclear reactions include protons $_{1}^{1}p$ and positrons $_{+1}^{0}e^{+}$. Positrons are positive electrons, sometimes called $^{+}$ particles to distinguish them rom electrons ($^{-}$ particles).

HALF-LIFE t

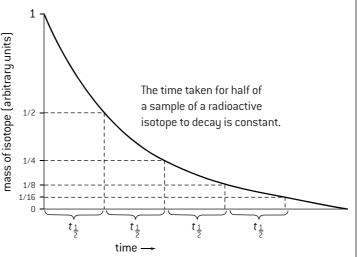
It is impossible to state when an individual unstable isotope will decay as it occurs spontaneously. However when a large number o atoms are together in a sample o the isotope the rate o decay depends on the amount o atoms present. The time taken or any

specified amount to decrease by exactly one hal remains constant and is independent o pressure and temperature. The hal -li e or a particular isotope is defined as the time taken or a given sample to decay to one hal o the **mass** o the **original isotope**. This is di erent to a chemical reaction where hal -li e is defined as the time taken or the **concentration** o a reactant to decrease to one hal o its initial value. The hal -li e $0^{-\frac{32}{15}}P$ is 14.3 days. A ter 14.3 days a 1.0 g sample $0^{-\frac{32}{15}}P$ will have decayed to 0.50 g and a ter a urther 14.3 days only 0.25 g $0^{-\frac{32}{15}}P$ will be remaining. Questions are o ten set the other way round. For example, calculate the hal li e $0^{-131}I$ i a sample $0^{-131}I$ is ound to contain $\frac{1}{32}$ o the original amount a ter 40.30 days.

The total number o hal -lives is five so the hal -li e = $\frac{40.30}{5}$ = 8.06 days. This was solved by inspection. It can also be solved mathematically. The number o atoms remaining *N* is related to the original number o atoms N_o by the equation $N = N_0 e^{-t}$ where is known as the decay constant.

At
$$t$$
, $N = \frac{N_0}{2}$ so $t = \frac{\ln 2}{2}$

In the example above *N* is equal to $\frac{N_o}{32}$ so $\frac{1}{32} = e^{-t}$ and ln $\left(\frac{1}{32}\right) = 40.03$ which makes equal to 0.0860 days. Substituting this value into the second equation gives $t = \ln \frac{2}{0.0860} = 8.06$ days.



Radioactive isotope	Half-life	
²¹² ₈₄ Po	3 10^{-7} seconds	
²²¹ ₈₇ Fr	4.8 minutes	
²²² ₈₆ Rn	3.8 days	
¹⁴ ₆ C	5730 years	
²³⁸ ₉₂ U	4.5 10° years	

Nuclear usion and nuclear fission (2)

POWER FROM NUCLEAR FISSION

A nuclear power station essentially contains two main components. The reactor to produce the heat rom a nuclear reaction and a turbine to drive a generator to produce electricity. The nuclear reactor uses a uel o uranium or plutonium. The uranium used is ²³⁵U. This reacts with neutrons to orm smaller nuclei and more neutrons. A typical reaction is:

$${}^{1}_{0}n + {}^{235}_{92}U = {}^{141}_{56}Ba + {}^{92}_{36}Kr + {}^{3}_{0}n + energy$$

Because more neutrons are produced than are used a chain reaction is possible. The critical mass is when this becomes sel -sustaining and the neutrons need to be controlled. The mass o the products is less than the reactants and the mass de ect is converted into energy. Natural uranium only contains a small percentage o ²³⁵U, most o it is ²³⁸U. In a breeder reactor neutrons react with ²³⁸U to orm 239-plutonium, which is fissionable.

 ${}^{238}_{92}U + {}^{1}_{0}n \qquad {}^{239}_{92}U \qquad {}^{0}_{-1}e + {}^{239}_{93}Np \qquad {}^{0}_{-1}e + {}^{239}_{94}Pu$

$${}^{1}_{0}n + {}^{239}_{94}Pu \qquad {}^{90}_{38}Sr + {}^{147}_{56}Ba + {}^{3}_{0}n + energy$$

NUCLEAR SAFETY

Arguments against the use o nuclear energy include the risk o terrorist attack, an accident, the disposal o waste and the promotion o less polluting orms o alternative energy. Accidents at nuclear power plants in Chernobyl in Ukraine in 1986 and in Fukushima in Japan in 2011 have heightened the concerns about the sa ety o nuclear power stations.

- 1. The possibility o a meltdown. A meltdown occurs when a nuclear reactor becomes out o control and essentially becomes a nuclear bomb. A nuclear power station requires the slow release o energy so the uel contains much less o the fissionable isotope $\binom{235}{92}$ U) than a nuclear bomb. The neutrons emitted by the fissionable material in a power station are absorbed by the non-fissionable isotope $\binom{238}{92}$ U) so in theory cannot build up enough momentum to establish a spontaneously explosive chain reaction.
- 2. Escape o radioactive material. This can occur as the uel is being transported or while it is being used. In the Chernobyl disaster a fire ignited the graphite moderator and a cloud o radioactive gas spread across much o Europe. The accident in Japan was caused by an earthquake and tsunami. Ordinary materials such the surrounding air and clothes worn by workers also have the potential to trans er low-level waste outside the plant. Concern has also been expressed about the risk o plutonium or other radioactive material alling into the hands o terrorists.
- 3. High-level nuclear waste. There are considerable problems associated with the disposal o high-level nuclear waste rom spent uel rods. These must be stored or very long periods (hundreds i not thousands o years) be ore they become relatively harmless. Current methods include vitri ying the waste in glass and burying it deep underground or in ocean trenches. Humankind has no experience o storing such materials sa ely or this length o time. The possibilities o disruption by earthquakes or the slow seepage o the waste into the water table make this one o the strongest arguments against using nuclear energy.

POWER FROM NUCLEAR FUSION

Nuclear usion o ers the possibility o an almost unlimited source o energy, which would almost be pollution ree since the main uel, deuterium, is inexpensive, abundant in sea water and no radioactive waste would remain. The essential reaction is:

$$_{1}^{2}H + _{1}^{2}H = _{2}^{3}He + _{0}^{1}n + energy$$

Other reactions involving tritium, ³H, another isotope o hydrogen could also be used but tritium is much less readily available and would need to be bred.

 ${}_{1}^{2}H + {}_{1}^{3}H = {}_{2}^{4}He + {}_{0}^{1}n + energy$

However the problems o controlling nuclear usion reactions have yet to be overcome. Essentially, the intensely hot reaction mixture known as plasma has to be contained and maintained or long enough to use the nuclei together. Since this involves temperatures approaching orty million degrees $(4 \quad 10^7 \text{ C})$ the problems are considerable.

ABSORPTION SPECTRA OF STARS

The energy rom the Sun and other stars comes rom uncontrolled nuclear usion reactions. Helium was in act first discovered by looking at the absorption spectrum o the Sun. The electrons are excited and absorb energy as they are promoted rom lower to higher levels. Each electronic transition shows up as a black line on what is otherwise a continuous spectrum. From the positions o the lines, the elemental composition o stars can be ascertained as each element has its own unique spectrum. An absorption spectrum is the opposite o an emission spectrum where energy is emitted as excited electrons drop down to lower energy levels but the lines have the same wavelengths and energy associated with them. The emission spectrum o hydrogen is discussed on page 11.



4. Low-level nuclear waste. This not only comes rom power plants but also rom research laboratories or hospitals.

It includes items such as rubber gloves, paper towels and protective clothing that have been used in areas where radioactive materials are handled. The level o activity is low and the hal -lives o the radioactive isotopes are generally short. It can be stored in vast tanks o cooled water called 'ponds' where it can lose much o its activity. Be ore it is then discharged into the sea it is filtered through an ionexchange resin which removes strontium and caesium, the two elements responsible or much o the radioactivity. Other methods o disposal include keeping the waste in steel containers inside concrete-lined vaults.

Solar energy

PHOTOSYNTHESIS

Approximately 5.6 10^{21} kJ o energy reaches the sur ace o the planet rom the Sun each year. About 0.06% o this is used to store energy in plants. This is achieved through photosynthesis a complex process summarized by the reaction o carbon dioxide and water to orm carbohydrates in the presence o chlorophyll.

 $6 \text{CO}_{_2}(g) + 6 \text{H}_{_2} \text{O}(l) \quad \ \ \text{C}_{_6} \text{H}_{_{12}} \text{O}_{_6}(s) + 6 \text{O}_{_2}(g)$

The process is endothermic requiring 2816 kJ o energy per mole o glucose. The green pigment chlorophyll contains an extended system o alternating single and multiple bonds known as conjugation. Substances such as chlorophyll and

-carotene, which contain extended conjugation, are able to absorb visible light by exciting an electron within this conjugated system. The energy which is absorbed by chlorophyll when it interacts with light is able to drive photosynthesis. The products o photosynthesis are used as ood to provide energy or animals through the reverse process respiration. Wood is mainly cellulose a polymer made up o repeating glucose units.

CONVERSION OF BIOMASS INTO ENERGY

The energy stored in biomass can be released in a variety o ways. For example:

1. Direct combustion o plant material or combustion o waste material derived rom plants such as animal dung.

e.g. $C_6H_{12}O_6(s) + 6O_2(g) = 6CO_2(g) + 6H_2O(l)$ $H = -2816 \text{ kJ mol}^{-1}$

- 2. Biogas. The anaerobic decay o organic matter by bacteria produces a mixture o mainly methane and carbon dioxide known as biogas. The manure rom arm animals can generate enough methane to provide or the heating, cooking and re rigeration needs o rural communities.
- 3. Fermentation to produce ethanol. Carbohydrates can be ermented by enzymes in yeast.

 $C_{6}H_{12}O_{6}(s) = 2C_{2}H_{5}OH(l) + 2CO_{2}(g)$

The ethanol can then be burned to produce energy.

$$C_2H_5OH(l) + 3O_2(g) = 2CO_2(g) + 3H_2O(l)$$
 $H = -1371$ kJ mol⁻¹

By combining ethanol with gasoline a uel called gasohol can be produced, which can be used by unmodified cars making them less reliant on the supply and cost o pure gasoline (petrol).

Bio uels are renewable, readily available and relatively non-polluting. In many hot countries animal dung is dried and used as a uel or heating and cooking. Garbage consisting o animal and vegetable waste is burned in incinerators in several cities to provide heat and electricity. This has the added advantage o reducing the amount o waste that has to be dumped in landfill sites. However disadvantages o bio uels include the act that they are widely dispersed, they take up land where ood crops can be grown and they remove nutrients rom the soil.

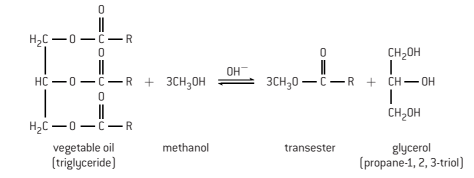
TRANSESTERIFICATION

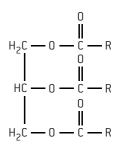
Vegetable oils are derived rom glycerol (an alcohol containing three hydroxyl groups) and unsaturated atty acids. They are triglycerides and contain three ester groups.

Vegetable oils contain a considerable amount o energy but they cannot be used in internal combustion engines as they are too viscous. This is because they have a high molar mass. However they can be converted into a usable uel by a process known as transesterification. In this process an ester can react with an alcohol to be converted into a di erent ester in the presence o a strong acid or strong base.

RCOOR + R'OH RCOOR + ROH (where R' is di erent to R)

Biodiesel or use in diesel engines can be made rom vegetable oils and waste cooking oils by heating them with an excess o an alcohol such as methanol or ethanol in the presence o sodium hydroxide. The alcohol is in excess in order to drive the position o equilibrium towards the transester side although some atty acids may also be present in the products.





General ormula o a vegetable oil where R, R' and R" are long chain hydrocarbons derived rom atty acids which may be the same or di erent.

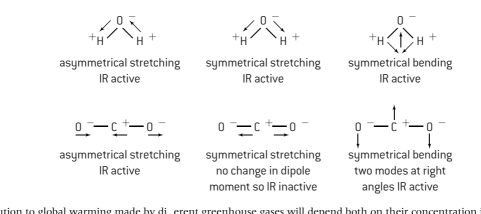
The three alkyl ester molecules ormed have a similar energy content in total to the triglyceride rom which they were derived but are much more volatile as they only contain one ester group each and have a much lower molar mass.

Environmental impact — global warming

THE GREENHOUSE EFFECT

A steady state equilibrium exists between the energy reaching the Earth rom the Sun and the energy reflected by the Earth back into space. This regulates the mean average temperature o the Earth s sur ace. The incoming radiation is shortwave ultraviolet and visible radiation. Some is reflected back into space and some is absorbed by the atmosphere be ore it reaches the sur ace. The energy reflected back rom the Earth s ur ace is longer wavelength in rared radiation. Not all o the radiation escapes. Greenhouse gases in the atmosphere allow the passage o the incoming shortwave radiation but absorb some o the reflected in rared radiation and re-radiate it back to the Earth s ur ace. They can absorb in rared radiation provided there is a change in dipole moment as the bonds in the molecule stretch and bend.

Vibrations of H₂O and CO₂



The contribution to global warming made by di erent greenhouse gases will depend both on their concentration in the atmosphere (abundance) and on their ability to absorb in rared radiation. Apart rom water, carbon dioxide contributes about 50% to global warming and methane 18%. Other greenhouse gases such as CFCs, N_2O and O_3 are many times better at absorbing heat than carbon dioxide but because their concentration is so low their e ect is limited.

Gas	Main source	Heat trapping effectiveness compared to CO ₂	Overall contribution to increased global warming
$H_{2}O$	Evaporation o oceans	0.1	
CO ₂	Combustion o ossil uels and biomass	1	50%
CH_4	Anaerobic decay o organic matter caused by intensive arming	30	18%

INFLUENCE OF GREENHOUSE GAS EMISSIONS ON GLOBAL WARMING

Most o the greenhouse gases have natural as well as humanmade sources. As humans have burned more ossil uels, the concentration o carbon dioxide in the air has risen steadily. Readings taken rom Mauna Loa in Hawaii show a steady increase in the concentration o carbon dioxide by about 1 ppm (0.0001%) each year or the past 50 years. During the same period measurements o the Earth's mean temperature also show a general increase. During the past 100 years the mean temperature o the Earth has increased by about 1 C although there were some years where the temperature ell rather than rose. Evidence rom ice core samples rom Greenland shows that there have also been large fluctuations in global temperature in the past; however most scientists now accept that the current global warming is a direct consequence o the increased emission o greenhouse gases. The predicted consequences o global warming are complex and there is not always agreement. The two most likely e ects are:

- 1. Changes in agriculture and biodistribution as the climate changes.
- 2. Rising sea-levels due to thermal expansion and melting polar ice caps and glaciers.

CARBON DIOXIDE

The increase in carbon dioxide in the atmosphere in the past 150 years is mainly due to the increased use o ossil uels or transport, heating and industrial processes. The Intergovernmental Panel on Climate Change (IPCC) and the Kyoto Protocol (later extended in Qatar) set targets to mitigate against climate change. Ways to limit human-made emissions o carbon dioxide include reducing dependency on ossil uels by moving to renewable energy sources as well as carbon capture and storage. Carbon dioxide can be captured chemically using sodium hydroxide or calcium oxide and stored in certain types o silicon-based rocks. There is a natural heterogeneous equilibrium between the carbon dioxide gas present in the atmosphere and the aqueous carbon dioxide dissolved in the oceans. The increase in carbon dioxide emissions has also caused the concentration o dissolved carbon dioxide in the oceans to increase. Because it orms carbonic acid, H₂CO₂, in sea-water this has resulted in a drop in pH, i.e. an increase in the acidity o the oceans. One o the e ects o this is to destroy coral ree s, which are essentially made o limestone.

 $CaCO_{3}(s) + 2H^{+}(aq) = Ca^{2+}(aq) + CO_{2}(g) + H_{2}O(l)$

PARTICULATES

Particulates, such as soot and volcanic dust, can have the opposite e ect to greenhouse gases. They cool the Earth by scattering the shortwave radiation rom the Sun and reflecting it back into space. Clouds do this as well. The lowering o mean global temperatures during the 1940s and 1960s has been attributed to the increased volcanic activity during these periods.

Electrochemistry, rechargeable batteries and fuel cells (1)

BATTERIES

A battery is a general term or an electrochemical cell in which chemical energy is converted into electrical energy. The electrons trans erred in the spontaneous redox reaction taking place in the voltaic cell produce the electricity. Batteries are a use ul way to store and transport relatively small amounts o energy. Some batteries (primary cells), such as the carbon zinc dry cell, can only be used once as the electrochemical reaction taking place is irreversible. Secondary cells can be recharged as the redox reactions involved can be reversed using electricity. Examples o rechargeable batteries include the lead acid battery and nickel cadmium and lithium-ion batteries.

The voltage o a battery is essentially the di erence in electrode potential between the two hal -cells. It primarily depends only on the nature o the chemical components o the two hal -cells. By combining two cells in series the voltage can be doubled. Three cells in series will triple the voltage o a cell. The power o a battery (i.e. the total work that can be obtained rom the cell) is the rate at which it can deliver energy and is measured in joules per second. It is a ected both by the size o the cell and the physical quantities o the materials present. An electrochemical cell has internal resistance due to the nite time it takes or ions to di use. The maximum current o a cell is limited by its internal resistance.

CARBON ZINC DRY CELL

The common dry cell contains a paste o ammonium chloride and manganese(IV) oxide as the electrolyte. The positive electrode (cathode) is made o graphite (o ten with a brass cap) and the zinc casing is the negative electrode (anode). A porous separator acts as a salt bridge. When the cell is working electrons flow rom the zinc to the graphite. Oxidation at the anode () Zn $Zn^{2+} + 2e^{-2}$ Reduction at the cathode (+) $MnO_2 + NH_4^+ + e^{-2}$ $MnO(OH) + NH_{2}$ The ammonia then brass cap reacts with the zinc ions to orm $[Zn(NH_3)_4]^{2+}$. This prevents a build up graphite electrode o ammonia gas. A carbon zinc porous separator dry cell produces 1.5 volts when new. In addition paste of NH₄Cl to having to replace the battery and ZnCl₂ when it wears Mn0₂ out it has a poor shel -li e and the acidic ammonium outer case zinc electrode chloride can corrode the zinc carbon zinc dry cell casing.

LEAD ACID BATTERY

The lead acid battery is used in automobiles and is an example o a secondary cell. Usually it consists o six cells in series producing a total voltage o 12 V. The electrolyte is an aqueous solution o sul uric acid. The negative electrodes are made o lead and the positive electrodes are made o lead(IV) oxide.

Oxidation at anode () $Pb + SO_4^{2-}$ $PbSO_4 + 2e^{-1}$

Reduction at cathode (+) $PbO_2 + 4H^+ + SO_4^{2-}$ $PbSO_4 + 2H_2O_4^{2-}$

The overall reaction taking place is thus:

 $Pb + PbO_{2} + 4H^{+} + 2SO_{4}^{2-} = 2PbSO_{4} + 2H_{2}O$

The reverse reaction takes place during charging. This can be done using a battery charger or through the alternator as the automobile is being driven. As sul uric acid is used up during discharging, the density o the electrolyte can be measured using a hydrometer to give an indication o the state o the battery. The disadvantages o lead acid batteries are that they are heavy and both lead and sul uric acid are potentially polluting.

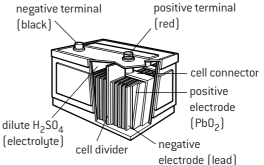
NICKEL CADMIUM AND LITHIUM-ION BATTERIES

Rechargeable nickel cadmium (NiCd) batteries are used in electronics and toys. They have a cell potential o 1.2 V. The positive electrode is made o nickel hydroxide which is separated rom the negative electrode made o cadmium hydroxide. The electrolyte is potassium hydroxide. During discharge the ollowing reaction occurs:

 $2NiO(OH) + Cd + 2H_2O = 2Ni(OH)_2 + Cd(OH)_2$

This process is reversed during charging. One o the

disadvantages o NiCd batteries is that cadmium is an extremely



Lead acid battery during recharging hydrogen and oxygen are evolved from the electrolysis of dilute H_2SO_4 so it needs topping up occasionally with distilled water.

toxic heavy metal so the batteries need to be disposed o responsibly. Laptops, tablets, smart phones and other handheld devices o ten use lithium-ion batteries. These contain lithium atoms complexed to other ions, e.g. Li_xCoO₂, in the positive electrode and it is these ions rather than lithium itsel that undergo the redox reactions. The negative electrode is made o graphite. Lithium-ion batteries are much lighter than NiCd batteries and produce a higher voltage, 3.6 4.2 V, but they do not have such a long li espan.

Electrochemistry, rechargeable batteries and fuel cells (2)

FUEL CELLS

A hydrogen uel cell utilizes the reaction between oxygen and hydrogen to produce water. Unlike combustion, the energy is given out not as heat but as electricity. As reactants are used up, more are added so a uel cell can give a continuous supply o electricity. They are used in spacecra t as, unlike rechargeable batteries, they do not need an external source o electricity or recharging. The electrolyte is aqueous sodium hydroxide. It is contained within the cell using porous electrodes that allow the passage o water, hydrogen and oxygen.

Oxidation at anode (-) $H_2 + 2OH^- = 2H_2O + 2e^-$

Reduction at cathode (+) $O_2 + 2H_2O + 4e$ 4OH

Hydrogen uel cells can also be made using phosphoric acid as the electrolyte in which case the equations are:

Oxidation at anode () $H_2 = 2H^+ + 2e$

Reduction at cathode (+) $O_2 + 4H^+ + 4e = 2H_2O$

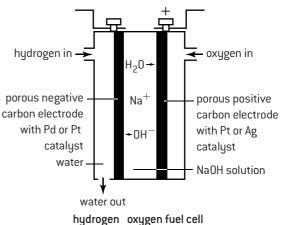
In both cases the overall equation is $2H_2(g) + O_2(g) = 2H_2O(l)$ and the entropy change is negative.

Other uels apart rom hydrogen can be used. For example, the methanol uel cell produces carbon dioxide and water as the products.

Oxidation at anode () $\rm CH_{3}OH + H_{2}O ~~ CO_{2} + 6H^{\scriptscriptstyle +} + 6e^{-1}$

Reduction at cathode (+) $\frac{3}{2}O_2 + 6H^+ + 6e = 3H_2O$

In both hydrogen and methanol uels cells the electrolyte can be supported on a solid polymer membrane. These are known as proton-exchange membrane (PEM) cells. They are used to produce electricity at relatively low temperatures, and can vary output quickly to meet shi ts in power demand. The advantage o a hydrogen uel cell is that it does not pollute, as water is the only product. Currently uel cells are very expensive to produce. The thermodynamic e ficiency o a uel cell is given by G^{\ominus}/H^{\ominus} . For the hydrogen uel cell H^{Θ} has the value -286 kJ mol¹ and G^{\ominus} has the value o 237 kJ mol⁻¹ (G^{\ominus} , the energy available or work, is less than H^{\ominus} as some energy has been used to overcome the negative entropy change as the system becomes more ordered). This gives a thermodynamic e ficiency o 83%. Although real uel cells do not approach this ideal e ficiency it is still much higher than the 30 60% e ficiency obtained by combusting uels in power stations.



hydrogen oxygen fuel cell with an electrolyte of NaOH

MICROBIAL FUEL CELLS

Microbial uel cells (MFCs) use bacteria to generate an electric current. They are able to tap into the electron transport chain o cells and liberate electrons that normally would be taken up by oxygen or other intermediates. The uel is oxidized at the anode by microorganisms, generating carbon dioxide, electrons and protons. Electrons are trans erred to the cathode through an external electric circuit, while protons are trans erred to the cathode through a membrane. Electrons and protons react with oxygen at the cathode to orm water. MFCs have the potential to be a sustainable energy source using di erent carbohydrates or substrates present in waste waters as the uel. The *Geobacter* species o bacteria, or example, can be used in some cells to oxidize ethanoate ions under anaerobic conditions.

 $CH_3COO^- + 2H_2O = 2CO_2 + 7H^+ + 8e$

NERNST EQUATION AND CONCENTRATION CELLS

When a copper hal -cell is connected to a zinc hal -cell the total em produced is 1.10 V under standard conditions (see page 76).

The reaction taking place is $Zn(s) + Cu^{2+}(aq) = Zn^{2+}(aq) + Cu(s)$.

It is an equilibrium reaction although the position o equilibrium lies ar on the product side. From Le Chatelier s Principle it can be seen that either increasing the concentration o the copper ions or decreasing the concentration o the zinc ions will move the equilibrium urther to the right and increase the voltage. The dependence o voltage on concentration is given by the Nernst equation.

$$E = E^{\ominus} - \left(\frac{RT}{nF}\right) \ln Q$$

Where *R* is the gas constant, *F* is Faraday s constant, *T* is the temperature in Kelvin and *n* is the number o electrons trans erred. Q is the reaction quotient, i.e. the equilibrium expression, in this case the concentration o the aqueous zinc ions divided by the concentration o the aqueous copper ions. I both concentrations are 1.00 mol dm⁻³ as they are under standard conditions then $E^{\circ} = E$ as ln 1 = 0. However i the concentration o the copper ions is doubled and the concentration o the zinc ions is halved then Q will equal $\frac{1}{4}$.

Then *E* will be equal to $E^{\ominus} - \frac{RT}{nF} \ln 0.25 = 1.10 - \frac{(8.31 - 298)}{(2 - 96500)} \ln 0.25 = 1.12 \text{ V}$

One application o the Nernst equation is that a cell can be made by combining two hal -cells with the same element but with solutions o di erent concentrations. This is known as a concentration cell. For example a $Cu(s) / Cu^{2+}(1.00 \text{ mol dm}^3)$ hal -cell connected to a $Cu(s) / Cu^{2+}(1.00 \text{ mol dm}^3)$ hal -cell will produce an em o $-\frac{RT}{2F}$ ln 0.01 = 0.059 V.

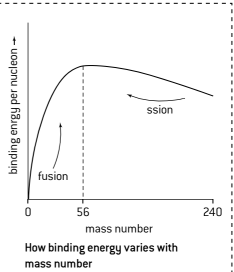
Although concentration cells produce a very small voltage they are important in biological systems. Nerve impulses are produced because nerve cells contain di erent concentrations o potassium ions on the inside and outside o the cell.

🖽 Nuclear usion and nuclear fission (3)

MASS DEFECT AND NUCLEAR BINDING ENERGY

Protons and neutrons in the nucleus o an atom are held together by very strong orces. A measure o these orces is known as the **binding energy**. This can be de ned as the energy that must be supplied to one mole o the atoms to break down the nuclei into separate neutrons and protons *or* the energy released when separate neutrons and protons combine to orm one mole o the atomic nuclei. It can be calculated rom the **mass defect**.

The mass de ect is the di erence in the combined mass o all the separate protons and neutrons compared with the actual mass o the nucleus. Strictly speaking, it also includes the mass o the electrons too but these are so small they can be ignored. Consider a helium nucleus. It contains two neutrons (relative mass = 2 1.0087) and two protons (relative mass = 2 1.0078) giving a total relative mass o 4.0330. However the actual relative atomic mass o helium is 4.0039. The relative mass de ect is thus 0.0291. By using Einstein s equation $E = mc^2$ this converts to a binding energy o approximately 2.7 10° kJ mol⁻¹. A graph o binding energy per nucleon against mass number shows that atomic nuclei with a mass number o approximately 56 (i.e. the nuclei o iron) have the maximum binding energy and are thus the most stable. Nuclei to the le t or right o this maximum will undergo nuclear change in such a way



that, as they approach the maximum, energy will be released. This explains nuclear usion, whereby small atoms combine to orm heavier nuclei, and nuclear ssion the splitting o heavy nuclei to orm two or more lighter nuclei.

CALCULATING THE ENERGY RELEASED IN A NUCLEAR REACTION

The mass numbers and atomic numbers balance in a nuclear equation. However i accurate values are used or the masses it can be seen that a small decrease in mass takes place during the reaction. The amount o energy that is ormed by this decrease in mass can be calculated using Einstein s equation $E = mc^2$.

g.	relative mass	${}^{235}_{92}U + {}^{1}_{0}n$	$^{90}_{36}\mathrm{Kr}$ + $^{144}_{56}\mathrm{Ba}$ + $2^{1}_{0}\mathrm{n}$
		235.0439 + 1.0087	89.9470 + 143.8810 + 2.0174
		236.0526	235.8454

The relative mass loss = 236.0526 235.8454 = 0.2072 which is approximately 0.1% of the initial uranium. If the masses are measured in grams then a loss of 0.2072 g (2.072 10^{-4} kg) is equivalent to 2.072 10^{-4} (2.998 10^8)² = 1.862 10^{13} J (or 1.862 10^{10} kJ). This amount of energy is the theoretical amount that one mole of uranium-235 (235 g) can produce i it all reacted according to the above equation.

Similar calculations can be per ormed or the mass loss during nuclear usion.

relative mass	${}^{2}_{1}H + {}^{3}_{1}H$	${}_{2}^{4}\text{He} + {}_{0}^{1}\text{n}$
	2.014102 + 3.016049	4.002602 + 1.008665
	5.030151	5.011267

The relative mass loss = 5.030151 - 5.011267 = 0.018884. This equates to $l.8884 \quad 10^{-5} \quad (2.998 \quad 10^{8})^{2} = 1.697 \quad 10^{12}$ J (or $l.697 \quad 10^{9}$ kJ). This is the theoretical energy released by 5 g o hydrogen isotopes. I this is scaled up to 235 g to compare it with the ssion energy released rom the same mass o uranium-235 it becomes 7.977 10^{10} kJ which shows the greater potential that nuclear usion has as an energy source.

ENRICHED URANIUM

e.g

e.g.

Naturally occurring uranium contains 99.3% uranium-238 and only 0.7% by mass o uranium-235. To obtain enriched uranium the isotopes have to be separated so that the percentage o uranium-235 is increased. This is because uranium-235 is the only naturally occurring isotope present in appreciable amounts that is ssile with neutrons. Uranium occurs naturally in several di erent ores. The ore uraninite contains uranium(IV) oxide, UO₂. This is a crystalline ionic solid with the same structure as calcium fluorite, CaF₂. It has a high melting point (2865 C) which makes it di cult to separate the di erent isotopes. However it can be converted into uranium hexafluoride. Uranium hexafluoride is a gas at ambient temperatures (it sublimes at 65.5 C) even though it has a very high relative molar mass. This is because the six fluorine atoms surround the central uranium hexafluoride molecules. Because fluorine only exists as a single isotope the slight di erence in mass between ²³⁸UF₆ and ²³⁵UF₆ is due only to the di erent isotopes o uranium. This means that ²³⁸UF₆ and ²³⁵UF₆ can be separated by either centri ugation or by gaseous di usion. The more modern method o gas centri ugation uses a large number o rapidly rotating cylinders to create a strong centri ugal orce so the heavier ²³⁸UF₆ gas molecules move towards the outside o the cylinder and the lighter gas molecules o ²³⁵UF₆ concentrate closer to the centre as enriched uranium. Gas di usion is based on Graham s Law which is explained on the ollowing page.

🖽 Nuclear usion and nuclear fission (4)

RATE OF RADIOACTIVE DECAY

The decay o radioactive isotopes is a first-order reaction i.e.

rate = [N] where is the rate constant (also known as the decay constant) and N is the concentration at time t.

I the rate is expressed as $\frac{dN}{dt}$

then
$$-\frac{dN}{dt} = [N]$$

Integration o this expression gives $\ln \frac{[N_o]}{[N]} = t$ where $[N_o]$ is the initial concentration when t = 0 and \ln is the natural logarithm. This can also be expressed in the orm $N = N_o e^{-t}$, which is how it is given in the IB data booklet.

Since radioactive decay depends on the nucleus it does not matter how the isotope is chemically bound or whether it is present as the ree element. The rate o decay will be the same and it is more usual to re er to [*N*] as the amount o isotope present rather than concentration. This can be expressed in moles or as mass.

At *t* the amount o *N* will be hal the initial amount. Thus $\frac{[N_o]}{[N]}$ will equal 2. The integrated expression then becomes $\ln 2 = t$ or $t = \frac{0.693}{N}$.

From this expression it can be seen that the hal -li e o radioactive decay is independent o the amount o isotope present as stated earlier in this option.

Worked examples

1. The hal -li e o radium-226 is 1622 years. Calculate how long it will take or a sample o radium-226 to decay to 10% o its original radioactivity.

Step 1. Use *t* to find the rate constant rom the equation t = 0.693

$$=\frac{0.693}{1622}=4.27$$
 10⁻⁴ year⁻¹

Step 2. Insert the value or into the integrated orm

o the rate equation
$$t = \ln\left(\frac{[N_o]}{[N]}\right)$$

4.27
$$10^{-4}$$
 $t = \ln\left(\frac{100\%}{10\%}\right)$

$$=> t = 5392$$
 years

It will take 5392 years to decay to 10% o its original activity.

GRAHAM S LAW OF EFFUSION

Graham's Law is based on the act that two di erent gases at the same temperature have the same kinetic energy. Since kinetic energy can be expressed as mv^2 where v is the velocity o the gaseous molecules it ollows that $m_1v_1^2 = m_2v_2^2$ where m_1 and m_2 are the molar masses o the two di erent gases travelling at velocities o v_1 and v_2 respectively. The lighter the gas the aster it is travelling compared with heavier gases at the same temperature.

A gas **diffuses** when it evenly fills its container and this occurs because, according to the kinetic theory o gases, gaseous molecules move with rapid random motion. **Effusion** occurs when gas molecules escape rom a small hole in the container. Because lighter gases have a greater velocity they e use more quickly than heavier gases. Graham s law o e usion states that the rate o e usion o a gas is

inversely proportional to the square root o the molar mass and is usually expressed as $\frac{\text{Rate}_1}{\text{Rate}_2} = \frac{M_2}{M_1}$.

Thus i a mixture o sul ur dioxide ($M_r = 64$) and methane ($M_r = 16$) molecules is allowed to e use the methane molecules will e use our times aster. This process o e usion can be used to enrich uranium as ²³⁵UF₆ will e use aster than ²³⁸UF₆.

2. A piece o old wood was ound to give 10 counts per minute per gram o carbon when subjected to ${}^{14}C$ analysis. New wood has a count o 15 cpm g⁻¹. The hal -li e o ${}^{14}C$ is 5570 years. Calculate the age o the old wood.

Step 1. Use *t* to find the rate constant.

$$=\frac{0.693}{5570}=1.24$$
 10⁻⁴ years⁻¹.

Step 2. Insert the value o into the integrated orm o the rate equation.

1.24
$$10^{-4}$$
 $t = \ln \frac{[N_{\sigma}]}{[N]} = \ln \frac{{}^{14}\text{C content in new wood}}{{}^{14}\text{C content in old wood}}$
= $\ln \frac{15}{10} = \ln 1.5$

=> t = 3270 years

The wood is 3270 years old.

NUCLEAR SAFETY

The problems o dealing with lowlevel and high-level nuclear waste and o ensuring nuclear sa ety have already been highlighted on page 145. The main health hazard caused by radioactive isotopes in nuclear material is due to the type o radiation they emit. This radiation is ionizing in nature and can produce ree radicals such as the superoxide radical, O₂ and the hydroxyl radical, OH. These radicals can initiate ree-radical chain reactions that can damage the DNA and the enzymes in living cells and thus lead to mutations and cancer.

Photovoltaic cells and dye-sensitized solar cells (DSSC)

ELECTRICAL CONDUCTIVITY OF SILICON

Metals conduct electricity because they have relatively low ionization energies and contain delocalized electrons. Non-metals (apart rom graphite) are poor conductors because the electrons are normally held in xed positions. Silicon is a semiconductor. Unlike metals, the conductivity o semiconductors increases with increased temperature.

A crystal o silicon contains a lattice o silicon atoms bonded to each other by shared pairs o electrons. These electrons are in xed positions so silicon is a poor conductor under normal conditions. However the energy required to excite an electron and ree it rom its bonding position is equivalent to the energy o light with a wavelength o $1.1 10^{-6}$ m. Visible light has a shorter wavelength in the range o $4-7 10^{-7}$ m. This is higher in energy and so photons o sunlight are able to excite a valence electron in silicon. The electron is then ree to move through the crystal lattice in the conduction band making it an electrical conductor. This is the basis o the photoelectric e ect and is the theory behind solar powered batteries. In practice, the process is not very e cient and the cost o puri ying the silicon is high. However solar cells are not polluting and do not use up valuable ossil uel reserves.

One method o improving the e ciency o the photoelectric e ect is by doping. This process involves adding very small amounts o atoms

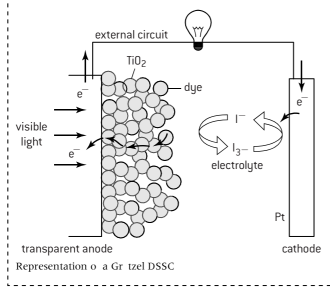
o other elements usually rom group 13 (Al, Ga or In) or rom group 15 (P or As). When a group 15 element is added the extra electron can move easily throughout the crystal lattice making it a better conductor compared with pure silicon. Such doping produces an n-type semiconductor because the conductivity is due to negative electrons. When a group 13 element is added the element now has one less electron than silicon. This produces a hole in the lattice. When a ree electron moves into this hole it produces a new hole where the electron was ormerly located. The hole can be regarded as a positive carrier so the semiconductor is known as a p-type.

DYE-SENSITIZED SOLAR CELLS (DSSCs)

Dye-sensitized solar cells work in a di erent way to silicon-based photoelectric cells. In the silicon-based cell the silicon provides both the source and the means o conductivity o the electrons. In a DSSC the photoelectrons originate rom the dye when it absorbs light and are then transported through a semiconductor.

A normal C=C double bond in an organic compound absorbs high energy radiation in the ultraviolet region and in the process an electron is excited to a higher energy level. I the compound contains many alternate double and single carbon to carbon bonds then it is said to be conjugated and the energy required to excite an electron is lower. The more conjugated the molecule is, the lower the energy o light required. Molecules such as chlorophyll and -carotene, which are highly conjugated, absorb in the visible region. As their name suggests, dyes also absorb in the visible region and sunlight interacts with dyes such as ruthenium polypyridine complexes, which are used in Gr tzel DSSCs to produce the photoelectrons.

The Gr tzel DSSC essentially consists o three parts. The anode is made o tin(IV) oxide, SnO_2 , doped with fluoride ions on a transparent plate. The back o this plate is covered in a thin layer



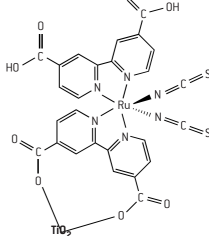
Si . Si	. Si _{extra}	Si .	Si	. Si
• •	electron	•	•	📩 hole
Si . As	. Si	Si .	Ga	. Si
• •	•	•	•	•
Si . Si	. Si	Si .	Si	. Si
n-type semico	onductor	p-type s	semico	onductor

o nanoparticles o titanium(IV) oxide, TiO₂ which provides a large sur ace area upon which the photosensitive dye is covalently bonded. This plate is sealed together with a separate plate consisting o an iodide electrolyte on a thin sheet o a conducting metal such as platinum which acts as the cathode. When light passes through the transparent plate it is absorbed by the dye, which releases electrons into the titanium(IV) oxide layer, which acts as the semiconductor. During this process the dye becomes oxidized. The electrons flow toward the transparent tin-based anode and through the external circuit. They return to the platinum cathode and flow into the iodide electrolyte. The electrolyte then transports the electrons back to the dye molecules.

The action o the iodide electrolyte is to reduce the dye back to its normal state during which iodide ions are oxidized to triiodide ions, I_3^{-} .

 $3I^ I_3^- + 2e^-$ Note that the triiodide ion can be thought o as a complex o iodine and iodide ions so e ectively iodide ions are being reduced to iodine

2I



A highly conjugated ruthenium polypyridine dye showing how it bonds to TiO₂ in Gr tzel DSSCs

external circuit reduce the ${\rm I}_{_3}\,$ ions back to iodide ions on the platinum cathode.

 $I_{3}^{-} + 2e^{-} 3I^{-}$

 $I_{2} + 2e^{-}$.

The returning

electrons rom the

DSSCs have several advantages over silicon-based photoelectric cells. They are simple to manu acture, semi-flexible and semi-transparent and relatively cheap. They also absorb visible light over a larger range o wavelengths.

SHORT ANSWER QUESTIONS OPTION C ENERGY

- 1. a) In the context o nuclear reactions explain the meaning o *fusion*. [1]
 - b) Explain why a usion reaction results in the release o a large quantity o energy. [2]
 - c) Explain why the energy released during a usion reaction is not currently an energy source used by society to generate electricity. [2]
 - d) The main reaction in a uranium reactor is fission o ²³⁵U.
 A side reaction is caused when neutrons react with ²³⁸U.

$${}^{38}_{4}U + {}^{b}_{c}X \qquad {}^{239}_{92}U \qquad {}^{239}_{94}Y + 2{}^{d}_{c}Z$$

[4]

[2]

[2]

[2]

[3]

Deduce the identities o X,Y,Z and a,b,c,d and e.

Х	b	
Y	с	
Z	d	
а	e	

- 2. Plutonium-238 has a hal -li e o 88 years and emits alpha particles.
 - a) Deduce the equation or the natural radioactive decay o ²³⁸Pu when it emits one alpha particle. [2]
 - b) Explain why it is impossible to state how long an individual plutonium-238 atom will take to decay. [2]
 - c) Determine how long will it take or 2.00 g o plutonium-238 to reduce to 0.3125 g o plutonium-238?
 [2]
- 3. a) Distinguish between *cracking* and *reforming* reactions used in crude oil refining.
 - b) Two products obtained rom crude oil are ethene, C_2H_4 and octane, C_8H_{18} . Compare the specific energies (in kJ kg⁻¹) o these two substances. [2]
 - c) Gasoline (petrol) or use in cars is given an octane rating.Explain what is meant by an octane number o 95. [2]

HL

- a) The standard electrode potential o the Cu(s)/Cu²⁺(aq) hal -cell is given in Section 24 o the IB data booklet. Calculate the value o the electrode potential i the concentration o the aqueous copper(II) is reduced to 1.00 10⁻² mol dm⁻³ and all the other variables remain constant.
 - b) Calculate the cell em when a $Cu(s)/Cu^{2+}(aq)$ hal -cell with a copper(II) concentration o $1.00 10^{-2}$ mol dm⁻³ is connected via a salt bridge to a standard $Zn(s)/Zn^{2+}(aq)$ hal -cell. [2]
 - c) Compare and contrast the voltage and power that can be obtained rom the cell in b) and a copper hal -cell connected to a zinc hal -cell where both hal -cells are operating under standard conditions. [3]
- 7. a) The mass o a proton and the mass o a neutron can be ound in section 4 o the IB data booklet. The accurate relative atomic mass o ²³⁸U is 238.050789. Calculate the binding energy in the nucleus o one atom o ²³⁸U (ignore the contribution made by the electrons to the relative atomic mass).
 - b) Naturally occurring uranium is largely made up o ²³⁸U with a small amount o ²³⁵U. Describe how enriched uranium can be obtained.

- d) Suggest two reasons why ethene is not used as a uel in cars.
- 4. a) The structure o chlorophyll is given in section 35 o the IB data booklet. State the essential eature o this molecule that enables it to absorb light in the visible region o the spectrum. [1]

[2]

[1]

- b) Wood is a bio uel. The cellulose contained within wood is made up o repeating units o glucose, C₆H₁₂O₆. State the equation or the ormation o glucose in plants by photosynthesis. [2]
- c) Vegetable oils can be converted into diesel uel by a process known as transesterification.
 - (i) Explain why vegetable oils themselves are not used as diesel uel. [2]
 - (ii) State the equation or the reaction o ethanol with a vegetable oil represented by the structure below to orm biodiesel. [2]

$$H_2C-O-COR$$

 $HC-O-COR$
 $HC-O-COR$
 $HC-O-COR$

- (iii) Suggest one reason why excess ethanol is used in this reaction.
- 5. a) Nitrogen, oxygen, carbon dioxide and water vapour are all naturally present in the atmosphere. Describe the greenhouse e ect and explain at the molecular level why only some o these gases are responsible or it. [4]
 b) Explain why the concentration o carbon dioxide has increased during the last 100 years and why this has
 - increased during the last 100 years and why this has increased the acidity of the oceans. [3]
 - c) List two ways in which governments can meet their pledges to reduce carbon dioxide emissions. [2]
 - c) An isotope in some radioactive nuclear waste has a hal -li e o 69 years. Calculate how long it would take or the activity o this particular isotope in the waste to decrease by 99%.
- a) List three advantages that dye-sensitized solar cells have compared with silicon-based photovoltaic cells. [3]
 - b) An electrolyte that is used in dye-sensitized solar cells
 is a solution o iodine in iodide ions. Explain with
 equations how the electrolyte unctions when the
 cells are working. [4]
 - c) Explain the unction o titanium(IV) oxide in a DSSC and why it is present in nanosized particles. [3]

15 OPTION D MEDICINAL CHEMISTRY

Pharmaceutical products and drug action

THE EFFECTS OF DRUGS AND MEDICINES

For centuries people have used natural materials to provide relie rom pain, heal injuries and cure disease. Many o these olk remedies have been shown to be very e ective and the active ingredients isolated and identified. Morphine was extracted rom the poppy *Papaver somniferum* early in the 19th century and later salicylic acid, the precursor o aspirin, was isolated rom willow bark.

The words drug or medicine have di erent connotations in di erent countries and are di ficult to define precisely. Generally

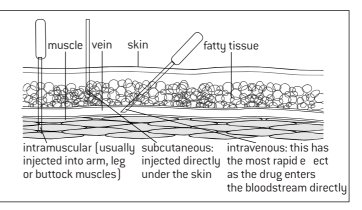
METHODS OF ADMINISTERING DRUGS

In order to reach the site where their e ects are needed, the majority o drugs must be absorbed into the bloodstream. The method o administering the drug determines the route taken and the speed with which it is absorbed into the blood. The our main methods are: by mouth (oral); inhalation; through the anus (rectal) and by injection (parenteral).

Drugs may also be applied topically so that the e ect is limited mainly to the site o the disorder such as the sur ace o the skin. Such drugs may come in the orm o creams, ointments, sprays and drops. a drug or medicine is any chemical (natural or human-made) that does one or more o the ollowing:

- alters incoming sensory sensations
- alters mood or emotions
- alters the physiological state (including consciousness, activity level or coordination).

Drugs and medicines are commonly (but not always) taken to improve health. They accomplish this by assisting the body in its natural healing process. The di erent mechanisms o drug action are still not ully understood and there is evidence that the body can be ooled into healing itsel through the placebo e ect.



RESEARCH, DEVELOPMENT AND TESTING OF NEW PRODUCTS

The research and development o new drugs is a long and expensive process. Traditionally, a new product is isolated rom an existing species or synthesized chemically. The process needs to identi y the need, the structure o the drug, possible methods o synthesis, extraction o the product and the yield. It is then subjected to thorough laboratory and clinical pharmacological studies to demonstrate its e ectiveness. Be ore studies are allowed on humans it must be tested on animals to determine the **lethal dose** required to kill fi ty percent o the animal population, known as the LD_{50} . The **e ective dose** required to bring about a noticeable e ect in 50% o the animal population, ED_{50} , is also obtained.

A actor known as the **therapeutic index**, can then be calculated.

Therapeutic index or *animals* = $\frac{LD_{50}}{ED_{50}}$

There are ethical issues about using animals or drug research and the lethal dose cannot be determined experimentally on humans.

Therapeutic index or humans =
$$\frac{TD_{s_0}}{ED_{s_0}}$$

where TD_{50} is the dose that causes a toxic e ect in 50% o the human population.

Drugs with a low therapeutic index need to be controlled very care ully as exceeding an e ective dose could produce serious toxicity or even death. Ideally the minimum amount o the drug required to achieve the desired e ect should be administered. The **therapeutic window** is the range o dosages between the minimum amounts o the drug that produce the desired e ect and a medically unacceptable adverse e ect.

Once the therapeutic index is determined the drug can then be used in

BIOAVAILABILITY OF A DRUG

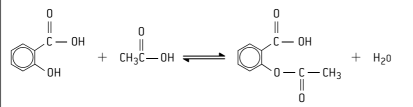
Drug-receptor interactions depend upon the structure o the drug and the site o activity. Bioavailability is the raction o the administered dosage that reaches the target part o the human body. The polarity o the drug, the types o unctional groups present in the drug and the method o administration can all a ect bioavailability. For example, the early penicillins could not be taken orally as they were destroyed by acids in the stomach be ore they could be e ective. Many drugs, such as morphine and soluble aspirin are administered in an ionic orm to make them more soluble. They then revert to the undissociated orm in the body. Some modern anti-cancer drugs have extremely high bioavailability. They contain radioactive nanoparticles embedded in the drug molecule. When the drug binds to the cancerous cells they e ectively deliver the radioactivity directly to the target and destroy the cancerous cells without causing damage elsewhere in the body.

an initial clinical trial on humans. This is usually on volunteers as well as on patients, hal o whom are given the drug and hal o whom are given a placebo. This initial trial is closely monitored to establish the drug s sa ety and possible side e ects. Drugs usually have unwanted **side e ects**, or example aspirin can cause bleeding o the stomach and morphine, which is normally used or pain relie , can cause constipation. Side e ects can be relative depending on why the drug is taken. People with diarrhoea are sometimes given a kaolin and morphine mixture and people who have su ered rom a heart attack are advised to take aspirin as it is e ective as an anti-clotting agent. The severity o the complaint will determine an acceptable **risk to benefit ratio**. I an e ective treatment is ound or a li e-threatening disease then a high risk rom side e ects will be more acceptable. The **tolerance** o the drug is also determined. Drug tolerance occurs as the body adapts to the action o the drug. A person taking the drug needs larger and larger doses to achieve the original e ect. The danger with tolerance is that as the dose increases so do the risks o dependence and the possibility o reaching the lethal dose. Some drugs can cause **addiction**, also known as substance dependence, where reducing or stopping the drug causes withdrawal symptoms. I the drug passes the initial clinical trial it will then go through a rigorous series o urther phases where its use is gradually widened in a variety o clinical situations. I it passes all these trials it will eventually be approved by the Drug Administration o a particular country or use either as an OTC (over the counter) drug or or use only through prescription by a doctor.

Aspirin

ASPIRIN AND OTHER MILD ANALGESICS

For a long time the bark o the willow tree (Salix alba) was used as a traditional medicine to relieve the ever symptoms o malaria. In the 1860s, chemists showed that the active ingredient in willow bark is salicylic acid (2-hydroxybenzoic acid) and by 1870 salicylic acid was in wide use as a pain killer (analgesic) and ever depressant (antipyretic). However salicylic acid has the undesirable side e ect o irritating and damaging the mouth, oesophagus and stomach membranes. In 1899 the Bayer Company o Germany introduced the ethanoate ester o salicylic acid, naming it Aspirin.



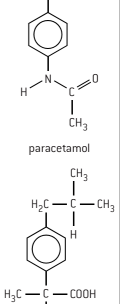
Aspirin is less irritating to the stomach membranes and the above reaction is only reversed in the body to produce salicylic acid when the aspirin reaches the more alkaline conditions o the small intestine.

Aspirin is thought to work by preventing a particular enzyme, prostaglandin synthase, being ormed at the site o the injury or pain. This enzyme is involved in the synthesis o prostaglandins which produce ever and swelling, and the transmission o pain rom the site o an injury to the brain. Because o its antiinflammatory properties, aspirin can also be taken or arthritis and rheumatism. Aspirin also has an ability to prevent blood clotting and is sometimes taken to prevent strokes or the recurrence o heart attacks.

The most common side e ect o aspirin is that it causes bleeding in the lining o the stomach. This e ect is increased by taking aspirin with alcohol as the alcohol has a **synergistic effect**. A ew people are allergic to aspirin with just one or two tablets leading to bronchial asthma. The taking o aspirin by children under twelve has been linked to Reyes syndrome a potentially at al liver and brain disorder. Exceeding the sa e dose o aspirin can be atal as the salicylic acid leads to acidosis due to a lowering o the pH o the blood.

Other mild analgesics have structures that show some similarity with aspirin (all are built on a benzene ring with two substituents).

Paracetamol (known as acetaminophen in the USA) contains a hydroxyl group and an amide group. It is o ten pre erred to aspirin as a mild pain reliever, particularly or young children, as its side-e ects are less problematic, although in rare cases it can cause kidney damage and blood disorders. Serious problems can arise however i an overdose is taken. Even i the overdose does not result in death, it can cause brain damage and permanent damage to the liver and kidneys.

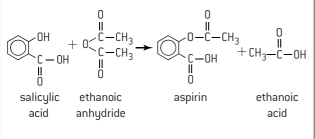


Η

ibuprofen

SYNTHESIS OF ASPIRIN

To obtain a good yield o aspirin, salicylic acid is reacted with ethanoic anhydride (rather than ethanoic acid) in the presence o an acid catalyst. The product can be recrystallized rom hot water to obtain pure aspirin.

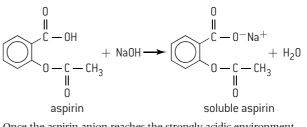


The purity o the product can be determined rom its melting point as impurities lower the melting point and cause it to melt over a wider range. The melting point o pure aspirin is 138 140 °C.

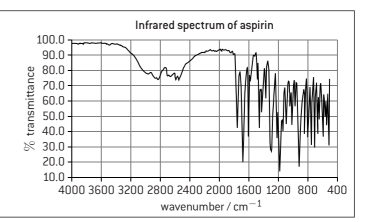
The purity can also be tested by looking at its in rared spectrum. Aspirin shows two peaks at 1750 and 1680 cm⁻¹ due to the two C=O bonds, and a very broad absorption between 2500 and 3500 cm⁻¹ due to the carboxylic acid group.

SOLUBLE ASPIRIN

Although aspirin is slightly polar due to the carboxylic acid group it is not very soluble in water. The solubility can be increased dramatically by reacting it with sodium hydroxide to turn it into an ionic salt.



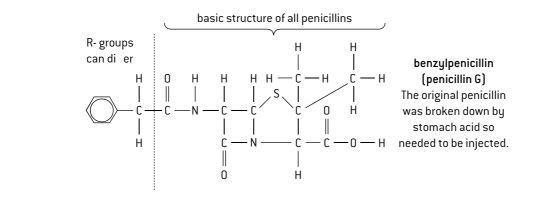
Once the aspirin anion reaches the strongly acidic environment o the stomach it reverts back to the un-ionized orm.



Penicillin

DISCOVERY AND GENERAL STRUCTURE

One o the main reasons or low li e expectancy in the early part o the 20th century was death due to bacterial in ections such as septicaemia. Penicillin was first discovered in the 1920s but was not brought into production until the Second World War. The original penicillin occurs naturally in a mould (*Penicillium notatum*) and had to be laboriously grown in large tanks containing corn-steep liquor. In the 1950s the structure o penicillin was determined and this enabled chemists to synthesize di erent types o penicillin and other antibiotics (antibacterials originating rom moulds) in the laboratory without recourse to moulds.

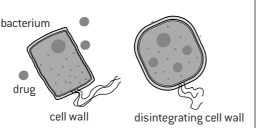


MODE OF ACTION

A typical bacterium consists o a single cell with a protective wall made up o a complex mixture o proteins, sugars and lipids. Inside the cell wall is the cytoplasm, which may contain granules o glycogen, lipids and other ood reserves. Each bacterial cell contains a single chromosome consisting o a strand o deoxyribonucleic acid (DNA). Not all bacteria cause disease and some are beneficial.

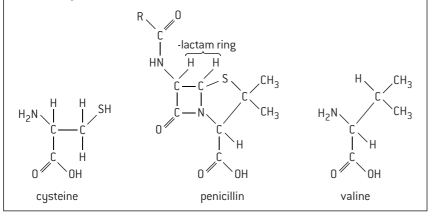
Penicillins work by preventing bacteria rom making normal cell walls.

All penicillins contain a ourmembered beta-lactam ring. Because o the restrictions o the ring, the normal bond angles o 109.5° and 120° are not able to be obtained. This makes the carboxamide group in the ring highly reactive as the ring can readily break due to the strained angles. The group containing the betalactam ring is very similar to a combination o the two amino



Penicillin interferes with cell-wall formation. As the cell swells, the osmotic pressure causes the wall to disintegrate and the bacterium dies.

acids cysteine and valine. When the ring opens these parts o the penicillin become covalently bonded to the enzyme that synthesizes the cell walls o the bacterium, thus blocking its action.



RESISTANCE TO PENICILLINS

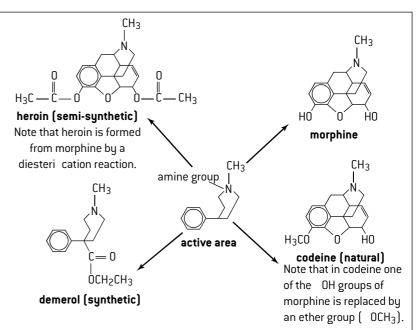
When penicillin became readily available to doctors it was o ten overprescribed to cure minor illnesses such as a sore throat. Certain bacteria were resistant to penicillin and were able to multiply. Their resistance was due to the presence o an enzyme called penicillinase, which could deactivate the original penicillin, penicillin G. To combat this, chemists developed other penicillins whereby the active part o the molecule is retained but the side chain is modified. However, as bacteria multiply and mutate so ast, it is a continual battle to find new antibiotics that are e ective against an ever more resistant breed o super bugs . These include the methicillinresistant Staphylococcus aureus (MRSA) and strains o *Mycobacterium tuberculosis* which cause tuberculosis, TB. To treat these in ections a strict adherence to a treatment regime o ten involving a cocktail o di erent antibiotics is required to prevent the risk o urther resistance developing.

Patients not completing a prescribed course o antibiotics and the use o antibiotics in animal eedstocks has also contributed to this problem. Healthy animals are given antibiotics to prevent risk o disease but the antibiotics are passed on through the meat and dairy products to humans, increasing the development o resistant bacteria.

Opiates

STRONG ANALGESICS

Strong analgesics are only available on prescription and are given to relieve the severe pain caused by injury, surgery, heart attack, or chronic diseases, such as cancer. They work by interacting temporarily with receptor sites in the brain without depressing the central nervous system, with the result that pain signals within the brain and spinal cord are blocked. The most important naturally occurring strong analgesics are morphine and codeine ound in the opium poppy. These are known as opiates or narcotics (although originally narcotic re erred to any sleepinducing drug). The active part o the morphine molecule has been identi ed. Codeine and semi-synthetic (obtained by simple structural modi cations to morphine) opiates, such as heroin, and totally synthetic compounds (e.g. demerol) all possess this basic structure and unction as strong analgesics.

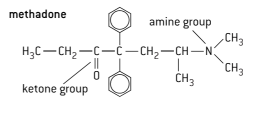


EFFECTS OF OPIATES

Diamorphine (heroin) is a more power ul painkiller than morphine but also more addictive. All opiates cause addiction and also lead to tolerance where more o the drug is required to give the same e ect and there is a danger o reaching the lethal limit.

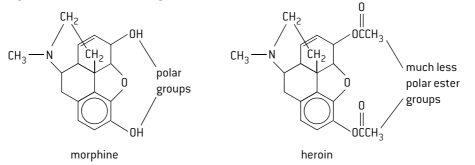
Short-term effects	Long-term effects	
Induce a eeling o euphoria	Constipation	
(sense o well-being)	Loss o sex drive	
Dulling o pain	Disrupts menstrual	
Depress nervous system	cycle	
Slow breathing and heart rate	Poor eating habits	
Cough reflex inhibited	Risk o AIDS, hepatitis,	
Nausea and vomiting (rst time users)	etc. through shared needles	
High doses lead to coma and/ or death	Social problems e.g. the t, prostitution	

Withdrawal symptoms occur within 6 to 24 hours or addicts i the supply o the drug is stopped. These include hot and cold sweats, diarrhoea, anxiety and cramps. One treatment to wean addicts o their addiction is to use methadone as a replacement or heroin. Methadone is also an amine and unctions as an analgesic but does not produce the euphoria craved by addicts.



POTENCY OF HEROIN COMPARED WITH MORPHINE

Heroin (diamorphine) is much more potent and produces a much greater eeling o euphoria than morphine. This can be explained by the di erence in the polarity o the two substances. Morphine molecules contain two polar hydroxyl (–OH) groups. When morphine is converted into heroin to orm the diester these are replaced by much less polar ethanoate groups. This makes heroin much more soluble in lipids that are non-polar. Heroin is thus able to rapidly penetrate the lipid-based blood brain barrier and reach the brain in higher concentrations than morphine.



Both morphine and heroin contain a tertiary amine group. They can be converted into ionic salts by reacting with hydrochloric acid (this is similar to the reaction o ammonia with hydrochloric acid to orm ammonium chloride). Thus or severe pain ollowing surgery or to alleviate the pain due to terminal cancer, diamorphine is o ten injected in the orm o diamorphine hydrochloride to increases its bioavailability. Once it is transported in the blood to the brain it reverts back to the undissociated orm to cross the lipid-based blood brain barrier.

pH regulation of the stomach

ANTACIDS

Antacids are used to reduce excess stomach acid. They are most e ective i taken between one and three hours a ter eating, as ood typically remains in the stomach or up to our hours a ter a meal. Antacids are essentially simple bases such as metal oxides, hydroxides, carbonates or hydrogencarbonates. They work in a non-speci c way by neutralizing the acid, preventing inflammation, relieving pain and discom ort and allow the mucus layer and stomach lining to mend. When used in the treatment o ulcers they prevent acid rom attacking the damaged stomach lining and so allow the ulcer to heal. Common examples include Al(OH)₃, NaHCO₃, Na₂CO₃, Ca(OH)₂ and milk o magnesia which is a mixture o MgO and Mg(OH)₃. Typical neutralization reactions are:

$NaHCO_{3}(s) + HCl(aq)$	$NaCl(aq) + CO_2(g) + H_2O(l)$
$Na_{2}CO_{3}(s) + 2HCl(aq)$	$2\text{NaCl}(aq) + \text{CO}_2(g) + \text{H}_2O(l)$
$Mg(OH)_{_2}(s) + 2HCl(aq)$	$MgCl_{2}(aq) + 2H_{2}O(l)$
$Al(OH)_{3}(s) + 3HCl(aq)$	$AlCl_{3}(aq) + 3H_{2}O(l)$

Worked example

Which would be the most e ective in combating indigestion a spoon ul o liquid containing 1.00 g o magnesium hydroxide or a spoon ul o liquid containing 1.00 g o aluminium hydroxide?

 $M_{\rm r}$ or Mg(OH)₂ = 24.31 + (2 17.01) = 58.33 $M_{\rm r}$ or Al(OH)₃ = 26.98 + (3 17.01) = 78.01

Amount o $Mg(OH)_{2}$ in $1.00 g = \frac{1.00}{58.33} = 0.0171$ mol Amount o $Al(OH)_{3}$ in $1.00 g = \frac{1.00}{78.01} = 0.0128$ mol

$$\begin{split} Mg(OH)_2(s) + 2HCl(aq) & MgCl_2(aq) + 2H_2O(l) \\ Amount o & HCl neutralized by 1.00 g o & Mg(OH)_2 = \\ 2 & 0.0171 = 0.0342 \, mol \end{split}$$

$$\begin{split} & \text{Al(OH)}_3(\text{s}) + 3\text{HCl(aq)} \qquad \text{AlCl}_3(\text{aq}) + 3\text{H}_2\text{O(l)} \\ & \text{Amount o HCl neutralized by 1.00 g o Al(OH)}_3 = 3 \quad 0.0128 \\ & = 0.0384 \,\text{mol} \end{split}$$

There ore the aluminium hydroxide would be slightly more e ective.

SIDE EFFECTS

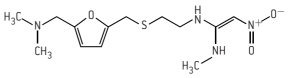
Like all drugs, antacids and drugs that inhibit or suppress acid production in the stomach may have side e ects. These may be caused by the drugs themselves or by **active metabolites**, the active orm o the drug a ter it has been processed by the body. Usually the e ect o active metabolites is weaker than those o the parent drug although in some cases it may actually be stronger and be responsible or the main therapeutic e ect o the parent drug.

The most common side e ects o antacids are belching to release gas and diarrhoea rom magnesium-containing antacids or constipation rom aluminium-containing antacids. Side e ects o ranitidine and the PPIs include diarrhoea, headache and dizziness. Long term use o PPIs may lead to a higher risk o osteoporosis and increase the risk o developing ood and drug allergies.

INHIBITION OF STOMACH ACID PRODUCTION

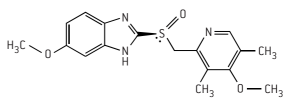
Much o the digestive process takes place in the stomach, a collapsible muscular bag that can hold between 2 and 4 litres o ood. The walls o the stomach are lined with a layer o cells that secrete mucus, pepsinogen (a precursor or the enzyme pepsin that breaks down proteins into peptides) and hydrochloric acid, collectively known as gastric juices. The hydrogen ion concentration o the hydrochloric acid normally lies between $3 \quad 10^{-2} \text{ mol dm}^{-3}$ and $3 \quad 10^{-3} \text{ mol dm}^{-3}$ giving a pH value between 1.5 and 2.5. The wall o the stomach is protected rom the action o the acid by a lining o mucus. Problems can arise i the stomach lining is damaged or when too much acid is produced, which can eat away at the mucus lining.

Certain drugs can inhibit the production o stomach acid by targeting speci c processes. Histamine is produced by certain cells in the lining o the stomach, called the enterochroma n-like cells (ECL cells). The histamine released rom ECL cells then stimulates the acid-making cells (parietal cells) in the lining o the stomach to release acid. H2-receptor antagonists (H2RA), such as ranitidine (Zantec), are a class o drugs used to block the action o histamine on the histamine H2-receptors o parietal cells and thus inhibit acid production.



ranitidine (trade name Zantec) an H2 receptor antagonist or H2 blocker

Other drugs used to suppress acid secretion in the stomach include proton pump inhibitors such as omeprazole (Prilosec) and esomeprazole (Nexium). PPIs cause a longlasting reduction o gastric acid production and have largely superseded H2-receptor antagonists. They work by interacting with a cysteine residue in a chemical system called the hydrogen-potassium adenosine triphosphatase enzyme system, H^+/K^+ ATPase (known as the proton pump) ound in the parietal cells in the stomach lining that make stomach acid. This inhibits the ability o these cells to secrete acid and helps to prevent ulcers rom orming or assist the healing process. They can also help to reduce acid reflux-related symptoms such as heartburn.



esomeprazole (Nexium)

The ormula o omeprazole is given in section 37 o the IB data booklet. Esomeprazole is an enantiomer o omeprazole. Omeprazole actually consists o a racemic mixture o the two isomers and undergoes a chiral shi t (on the S atom) in the body that converts the inactive enantiomer into the active enantiomer (esomeprazole).

Buffer solutions

BUFFER SOLUTIONS

It is important to maintain the pH in the body within certain strict limits to ensure enzymes unction correctly. This is done by bu er solutions, which resist changes in pH when small amounts o acid or alkali are added to them.

An acidic bu er solution can be made by mixing a weak acid together with the salt o that acid and a strong base. An example is a solution o ethanoic acid and sodium ethanoate. The weak acid is only slightly dissociated in solution, but the salt is ully dissociated into its ions, so the concentration o ethanoate ions is high.

 $NaCH_{_3}COO(aq) \qquad Na^{_+}(aq) + CH_{_3}COO^{_-}(aq)$

 $\label{eq:coord} CH_{_3}COOH(aq) \qquad CH_{_3}COO^-(aq) \,+\, H^+(aq)$

I an acid is added the extra H^+ ions coming rom the acid are removed as they combine with ethanoate ions to orm undissociated ethanoic acid, so the concentration o H^+ ions remains unaltered.

 $CH_{3}COO^{-}(aq) + H^{+}(aq) - CH_{3}COOH(aq)$

I an alkali is added the hydroxide ions rom the alkali are removed by their reaction with the undissociated acid to orm water, so again the H⁺ ion concentration stays constant.

 $CH_3COOH(aq) + OH^-(aq) = CH_3COO^-(aq) + H_3O(i)$

In practice acidic bu ers are o ten made by taking a solution o a strong base and adding excess weak acid to it, so that the soluton contains the salt and the unreacted weak acid.

$NaOH(aq) + CH_{3}COOH(aq)$	$\mathrm{NaCH_{_3}COO(aq)}~+~\mathrm{H_{_2}O(l)}~+$	$CH_{3}COOH(aq)$
limiting reagent	salt	excess weak acid
	·	

bu er solution

An alkali bu er with a fixed pH greater than 7 can be made rom a weak base together with the salt o that base with a strong acid. An example is ammonia with ammonium chloride.

 $NH_4Cl(aq) = NH_4^+(aq) + Cl^-(aq)$

 $NH_3(aq) + H_2O(l) = NH_4^+(aq) + OH^-(aq)$

I H⁺ ions are added they will combine with OH^- ions to orm water and more o the ammonia will dissociate to replace them. I more OH^- ions are added they will combine with ammonium ions to orm undissociated ammonia. In both cases the hydroxide ion concentration and the hydrogen ion concentration remain constant.

BUFFER CALCULATIONS

The equilibrium expression or weak acids also applies to acidic bu er solutions,

e.g. ethanoic acid/sodium ethanoate solution.

$$K_{a} = \frac{[\mathrm{H^{+}}] \quad [\mathrm{CH}_{3}\mathrm{COO^{-}}]}{[\mathrm{CH}_{3}\mathrm{COOH}]}$$

The essential di erence is that now the concentrations o two ions rom the acid will not be equal.

Since the sodium ethanoate is completely dissociated the concentration o the ethanoate ions in solution will be all the same as the concentration o the sodium ethanoate; only a very little will come rom the acid.

I logarithms are taken and the equation is rearranged the

$$pH = pK_a + \log_{10} \frac{[CH_3COO^-]}{[CH_3COOH]}$$

This is known as the Henderson Hasselbalch equation (the general ormula can be ound in section 1 o the IB data booklet).

Two acts can be deduced rom this expression. Firstly the pH o the bu er does not change on dilution, as the concentration

o the ethanoate ions and the acid will be a ected equally. Secondly the bu er will be most e ficient when $[CH_3COO^-] = [CH_3COOH]$. At this point, which equates to the equivalence point when ethanoic acid is titrated with sodium hydroxide, the pH o the solution will equal the pK_a value o the acid.

Calculate the pH o a bu er containing 0.200 mol o sodium ethanoate in 500 cm³ o 0.100 mol dm⁻³ ethanoic acid (given that K_a or ethanoic acid = 1.8 10^{-5} mol dm⁻³).

 $[CH_{3}COO^{-}] = 0.400 \text{ mol } dm^{-3}; [CH_{3}COOH] = 0.100 \text{ mol } dm^{-3}$

$$K_{a} \quad \frac{[H^{+}] \quad 0.400}{0.100} = 1.8 \quad 10^{-5} \text{ mol } dm^{-3}$$
$$[H^{+}] = 4.5 \quad 10^{-6} \text{ mol } dm^{-3}$$
$$pH = 5.35$$

Calculate what mass o sodium propanoate must be dissolved in 1.00 dm³ o 1.00 mol dm⁻³ propanoic acid ($pK_a = 4.87$) to give a bu er solution with a pH o 4.5.

$$[C_{2}H_{5}COO^{-}] = \frac{K_{a}}{[H^{+}]} = \frac{10^{-4.87}}{10^{-4.5}} = 0.427 \text{ mol dm}^{-3}$$

Mass o
$$NaC_2H_5COO$$
 required = 0.427 96.07 = 41.0 g

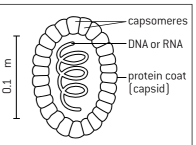
Antiviral medications

BACTERIA AND VIRUSES

Bacteria are single cell organisms. Inside the cell wall is the cytoplasm, which may contain granules o glycogen, lipids and other ood reserves. Each bacterial cell contains a single chromosome consisting o a strand o deoxyribonucleic acid, DNA.

There are many di erent types o virus and they vary in their shape and structure. All viruses, however, have a central core o DNA or RNA surrounded by a coat (capsid) o regularly packed protein units called capsomeres each containing many protein molecules. They have no nucleus or cytoplasm and there ore, unlike bacteria, they are not cells. They are much smaller than bacteria and they do not eed, excrete or grow and they can only reproduce inside the cells o living organisms using the materials provided by the host cell.

Athough viruses can survive outside the host they can only replicate by penetrating the living host cell and injecting their DNA or RNA into the cell s cytoplasm. The virus then takes over the biochemical machinery inside the cell. This causes the cell to die or become seriously altered and causes the symptoms o the viral in ection. The cell is made to produce new DNA or RNA and orms large numbers o new viruses. These are then released and move on to in ect other healthy cells. Because they lack a cell



in ect other healthy cells. generalized diagram of a virus

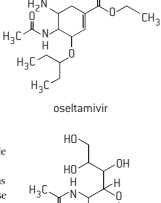
structure and because they multiply so quickly they are di cult to target with drugs. The two main ways in which antiviral drugs may work is by altering the cell s genetic material so that the virus cannot use it to multiply, or by blocking enzyme activity within the host cell so that the virus cannot multiply or is prevented rom leaving the cell. One o the problems with developing antiviral drugs is that the viruses themselves are regularly mutating this is particularly true with the Human Immunode ciency Virus (HIV).

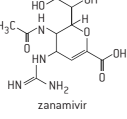
TWO SPECIFIC ANTIVIRAL DRUGS

Common viral in ections include the common cold, influenza and childhood diseases such as mumps and chicken pox. One drug that is e ective against both the influenza A and B viruses is oseltamivir which has the trade name Tamiflu. One o the enzymes used by all influenza viruses to stick to the host cell wall as it leaves is called neuraminidase and the drug works by inhibiting the active site on this enzyme. In the liver, oseltamivir, which is an ester, is hydrolysed to its carboxylate anion which inhibits the viral neuraminidase and prevents it rom acting on sialic acid, an acid ound on the proteins on the sur ace o the host cells. This blocking action

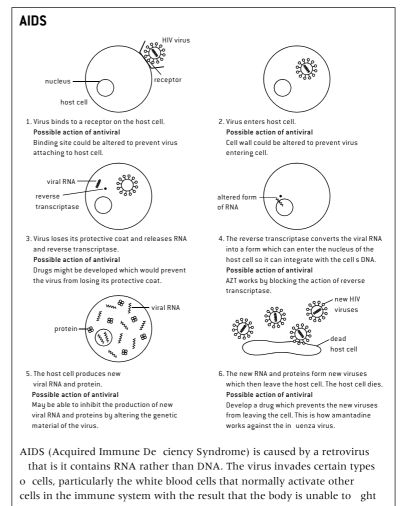
prevents the new viral particles rom being released and so the virus cannot in ect other cells and multiply.

Another influenza drug, zanamivir (trade name Relenza) also unctions as a neuraminidase inhibitor. Both oseltamivir and zanamivir have some structural similarities to sialic acid (a derivative





o neuraminic acid, which is a monosaccharide with a nine carbon backbone) and this may explain how they are able to inhibit the viral neuraminidase enzyme. Zanamivir contains many polar hydroxyl and amine groups together with a carboxylic acid group, which makes it much more soluble in polar solvents than oseltamivir.



o cells, particularly the white blood cells that normally activate other cells in the immune system with the result that the body is unable to ght in ection. Once it invades a host cell it makes viral-DNA rom the RNA template using an enzyme called reverse transcriptase. This is the opposite process that takes place in normal cells in which RNA is made rom a DNA template using transcriptase as the enzyme. Various drugs have met some success in treating the HIV virus but there is still no cure. One, AZT (zidovudine), is a reverse-transcriptase inhibitor and combines with the enzyme that the HIV virus uses to build DNA rom RNA and inhibits its active site. Since it is only retroviruses that use this enzyme AZT does not a ect normal cells. Other possible ways in which potential antiviral drugs might work are shown above.

Environmental impact of some medications

RADIOACTIVE WASTE

The treatment o cancer o ten involves radiation therapy. This may be rom an external radioactive source or using radioisotope therapy where a radioisotope is attached to another molecule or antibody, which then guides it to the target tissue a ter being injected or taken orally. Radioactive waste can be divided into high-level waste (HLW) and lowlevel waste (LLW). Low-level waste includes items such as rubber gloves, paper towels and protective clothing that have been used in areas where radioactive materials are handled. The level o activity is low and the hal -lives o the radioactive isotopes are generally short. High-level waste has high activity and generally the isotopes have long hal -lives so the waste will remain active or a long period. Isotopes used in radioisotope therapy tend to be (relatively) low-level waste (typical isotopes used are ¹³¹I, ⁸⁹Sr and ¹⁵³Sm). Any type o radioactive waste needs to be kept separate rom other types o waste. LLW is usually disposed o in land ll or in the sea whereas HLW is vitri ed (turned into a glass-like material) and stored underground in concrete bunkers.

GREEN CHEMISTRY

While the sa e disposal o waste with minimal damage to the environment is important, green chemistry (also known as sustainable chemistry) aims to reduce harm to the environment by minimizing the use and generation o hazardous substances in the rst place in other words to reduce the pollution at its source and to conserve natural resources including energy. This is particularly important in the pharmaceutical industry where o ten the research and development o a new drug involves many steps, each involving many potentially polluting substances.

Important actors when designing and producing new drugs include:

 Aiming or a high atom economy and a low environmental actor. The atom economy is the ratio o the total mass o the desired product(s) to the total mass o all the products. Essentially this is a measure o how much o the reactants remain in the nal product (see page 6).

The environmental- actor (E- actor) is de ned as the mass o the total waste products divided by the mass o the desired product, which highlights the need to avoid producing waste products.

$$E = \frac{\text{total waste (kg)}}{\text{product (kg)}}$$

Pharmaceutical chemistry, where almost all the products contain carbon, also uses carbon e ciency.

$$\frac{\text{Carbon}}{\text{e ciency}} = \frac{\text{amount o carbon in product}}{\text{total amount o carbon present in reactants}}$$

2. The number o steps in a synthesis should be kept to a minimum. Generally the more separate steps required to reach the desired product the lower the percentage yield and the higher the amount o waste reactants and products and the more energy used. In the past, chemists looked or intellectually elegant syntheses. A modern elegant synthesis is one that uses less raw material and energy, and produces less waste.

ANTIBIOTIC AND SOLVENT WASTE

Microorganisms in water or the soil can take up waste antibiotics. Bacteria in the organisms can become resistant to antibiotics by developing enzymes that can break down the antibiotics. I the bacteria nd their way into drinking water or ood they then make antibiotics much less e ective when they are required to ght disease. Antibiotic waste comes rom the disposal o unused antibiotics, through the urine o people who are on antibiotics and rom animals where the eedstock contains antibiotics, which o ten gets discharged into rivers.

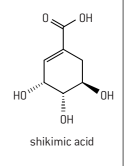
Many di erent solvents may be used in the production o drugs. Most countries have strict guidelines about the sa e disposal o solvents. They should be separated into chlorine containing and non-chlorine containing solvents. Chlorinated solvents must be incinerated at very high temperatures to prevent the ormation o carcinogenic dioxins. Nonchlorinated solvents may be recycled (or their solutes such as heavy metals extracted and recycled), burned to provide energy or, i they are innocuous such as sodium chloride solution, disposed o in rivers or the sea.

3. Use greener and sa er solvents and reactants. Solvents play an important role in many o the separate steps in a synthesis. The energy and materials needed to manu acture the solvent as well as the problems caused by the disposal o the solvents (i they cannot be recycled) all need to be taken into account. The sa ety o the work orce should also be considered so chlorinated solvents and benzene, which are potentially carcinogenic, should be avoided.

In addition, green synthetic chemists will also consider using renewable eedstocks, using suitable catalysts to reduce energy demands by lowering operating temperatures and to consider the ate o the potential drug regarding its breakdown products and disposal a ter use.

One example o the use o green chemistry in practice is the development o the influenza drug Tamiflu (oseltamivir). Tamiflu was rst synthesized by chemists working at Monash University in Australia. Their technique only produced very small amounts o product and used unsa e lithium nitride, LiN₃. The rst commercial scale production was done by Glaxo. They avoided the use o lithium nitrides but still used azides and chromatography as a puri cation technique rather than the greener recrystallization. Since then more and more

ways o synthesizing Tamiflu have been developed, each with a more sustainable route. One o the problems is that one o the starting materials is a compound called shikimic acid, which is currently uneconomical to synthesize and is isolated rom the Chinese star anise plant (*Illicium anisatum*). One green hope is that it can be obtained by bioengineering using ermenting bacteria.



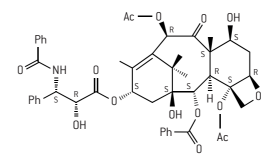
🕕 Taxol 🛛 a chiral auxiliary case study

TAXOL

Taxol (also known as paclitaxel) is used to treat ovarian, breast and lung cancer. It is injected as a colourless fluid as part o chemotherapy treatment. It works by preventing cancer cells rom dividing and replicating. It occurs naturally in the bark o the Paci c yew tree (*Taxus brevifolia*). Once it became established as an e ective anti-cancer drug, concern was expressed about the damage being done to the yew tree population as the amount o bark required to make even small amounts o the drug

is considerable and stripping the bark kills the trees. The challenge or chemists was to synthesize the drug rom crude-oil based starting materials. Initially a semi-synthetic route was ound starting rom a compound ound in the needles o a related yew tree (*Taxus baccata*). More recently, the total synthesis rom crude-oil based reagents has been achieved but the process is not very green. Most o the commercially available taxol is still obtained either by a semi-synthetic route or by plant cell ermentation (PCF) technology, which uses much less energy and less hazardous chemicals.

The structure o taxol is given in section 37 o the IB data booklet but this does not make clear the exact stereochemistry. The di culty with the synthesis o taxol is that it contains no ewer than 11 separate chiral carbon



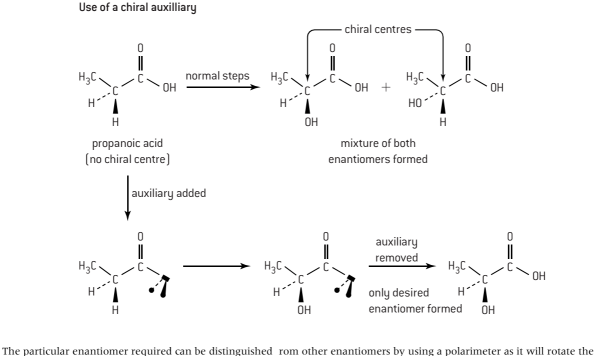
The structure of taxol showing the exact stereochemistry at each of the eleven chiral carbon atoms (Ph = phenyli; Ac = acetyl)

atoms. This means that there is a very large number o possible enantioners and diastereoisomers o which only one corresponds to taxol itsel with the correct R or S con guration on each o the 11 chiral carbon atoms.

USE OF CHIRAL AUXILIARIES

The traditional synthesis o an optically active compound normally produces a racemic mixture (50:50 mixture) o the two enantiomers, which then has to be separated into the two isomers by using chromatography together with another optically active compound. I this technique is used to make taxol then the yield o the correct enantiomer would be very small. A more recent technique using chiral auxiliaries now makes it possible to synthesize just the desired isomer.

Attaching an auxiliary, which is itsel optically active, to the starting material creates the stereochemical conditions necessary or the reaction to orm only one enantiomer, i.e. asymmetrically to orm just the required isomer. A ter the desired product has been ormed the auxiliary is removed and recycled. Taxol is too complicated to give as an example here but the reaction o propanoic acid to orm just one o the two possible enantiomers o 2-hydroxypropanoic acid (lactic acid) illustrates the technique.



The particular enantiomer required can be distinguished rom other enantiomers by using a polarimeter as it will rotate the plane o plane-polarized light by a xed amount in either a clockwise or anti-clockwise direction. This technique is described on page 96.



RADIOTHERAPY AND SIDE EFFECTS

Radiotherapy can involve many di erent types o emissions depending on the type o radiotherapy and the reason it is being used. These include alpha particles, ${}_{4}^{4}$ He, beta particles, ${}_{-1}^{0}$ e⁻, protons, ${}_{1}^{1}$ H, neutrons, n, positrons, ${}_{+1}^{0}$ e⁺ and gamma radiation, . The ionizing radiation may be used diagnostically or used to destroy cancer cells. Much o it is high-energy radiation and has the potential to cause harm as well as do good and the dosage and its use need to be care ully controlled. One notable exception to this is magnetic resonance imaging. The patient is placed inside a large magnetic field and very low energy radio waves are used to determine the environment o particular nuclei, e.g. 'H or ³¹P, in the body to give a detailed image. This is a practical application o NMR spectroscopy (see page 105) and is non-invasive.

Side e ects rom radiotherapy can vary rom person to person and depend upon the type o radiotherapy received. External radiotherapy tends to cause more side e ects than the more targeted internal therapy. Hair loss may occur although, unlike chemotherapy, this is usually limited to the treatment area where the radiation enters and leaves the body. Other short-term e ects may include nausea, atigue, loss o appetite and skin disorders. Radiation may cause damage to the DNA in surrounding healthy cells and regenerating tissue and possibly cause some sterility.

DIAGNOSTIC RADIOGRAPHY

Tc-99m is a metastable radioactive isotope o technetium with a radioactive hal -li e o 6 h and a biological hal -li e o 1 day (i.e. hal o it will be excreted rom the body in 24 h) so it is ideal as a tracer. Because o its short hal -li e, it is generated in hospitals by the beta decay o molybdenum-99. It can be targeted at particular parts o the body and can be used diagnostically, or example to determine how well both kidneys are unctioning. Its progress through the body can be ollowed by the gamma radiation it emits as the energy produced is roughly the same as that detected by conventional X-ray equipment. Because it is a gamma emitter there is a very small risk o tumours developing ollowing exposure but the potential benefits make this a risk worth taking. The decay product, ground state Tc-99 is a beta emitter but the energy released is low as it has a long hal -li e and the amount in the body rapidly decreases over time.

$$t = 6 \text{ hours}$$
 $t = 211,000 \text{ years}$
 $t_{43}^{99}\text{Tc} + \frac{99}{44}\text{Ru} + \frac{9}{-1}\text{e}$

Another radioisotope that can be used as a tracer in diagnostic medicine is iodine-131 which is a beta and gamma emitter with a hal -li e o eight days. However it remains in the body or some time and its use can lead to thyroid cancer developing so it tends to be used mainly in high doses to kill thyroid tumours (where it becomes incorporated in thyroxin) rather than diagnostically.

TARGETED THERAPY

Targeted alpha therapy (TAT)

TAT is used to direct radiation directly at cancerous cells and in particular to prevent micrometastatic cancerous cells (secondary tumour ormation) in other parts o the body. The range o particles in tissues is typically only 50 100 m so when targeted at a tumour they have the ability to release a large amount o energy directly at the specific tumour and do much less damage to the surrounding area than beta emitters. A typical alpha emitter used or TAT is lead-212.

 $^{212}_{82}$ Pb $^{208}_{80}$ Hg + $^{4}_{2}$ He

(Note that in ormation about balancing nuclear equations is given on page 144.)

Boron neutron capture therapy (BNCT)

BNCT is used to treat brain tumours and recurrent head and neck cancers. Initially the patient is injected with a non-radioactive isotope o boron, ${}^{10}_{5}$ B, which is selectively absorbed by the cancer cells. The patient is then irradiated with neutrons, which are captured by the boron-10. This causes a nuclear reaction and the boron is converted into an isotope o lithium, ${}^{3}_{3}$ Li with the emission o a high energy alpha particle that acts directly on the tumour while doing relatively little damage to the surrounding area.

 ${}^{10}_{5}B + {}^{1}_{0}n$ ${}^{7}_{3}Li + {}^{4}_{2}He$

Other targeted therapy

Other examples o radioisotopes used or targeting radiation to specific areas are lutetium-177 and yttrium-90. Yttrium-90 is a beta emitter whereas lutetium-177 is both a gamma and beta emitter. Lutetium-177 has a shorter penetration range than yttrium-90 making it ideal or smaller tumours. It is added to a carrier called DOTA-TATE, which attaches to specific tumours such as neuroendocrine tumours and certain types o thyroid tumours.

HALF-LIFE CALCULATIONS

Radioactive nuclei have constant hal -lives so the equation $N = N_{o}e^{-t}$ can be used where *N* is the mass or amount remaining a ter time *t*, N_{o} is the initial mass or amount and the decay constant. In ormation on hal -li e can be ound on page 144.

Worked example

Technetium-99m has a hal -li e o 6.01 hours. A resh sample containing 5.00 g o technicium-99m is prepared rom molybdenum-99 in a hospital at 0900 h. Calculate the mass o technetium-99m remaining in the 5.00 g sample at 1700 h later that day.

Step 1. Use t to find the rate constant rom the equation

t = 0.693

$$=\frac{0.693}{6.01}=0.115$$
 hour⁻¹

Step 2. Insert the value or into the integrated orm o the $\begin{bmatrix} N \end{bmatrix}$

rate equation
$$t = \ln \frac{1}{[N]}$$

0.115 8.00 = $\ln\left(\frac{5.00}{N}\right)$ where N = mass remaining a ter 8.00 hours

$$\ln \left(\frac{5.00}{N}\right) = 0.92 \text{ so } N = \frac{5.00}{2.51} = 1.99 \text{ g}$$

1.99 g o technetium-99m will be remaining.

🕕 Drug detection and analysis (1)

IDENTIFICATION OF DRUGS USING SPECTROSCOPY

This topic builds on what has already been covered in pages 102 106. Remember that mass spectroscopy can be used to identi y not only the molar mass but also individual ragments with particular masses. In rared spectroscopy can identi y specific unctional groups as these absorb in di erent regions o the in rared spectrum with the precise absorption depending upon neighbouring atoms. Proton NMR spectroscopy gives di erent peaks or hydrogen atoms that are in di erent chemical environments. The area under each peak is related to the number o hydrogen atoms in that particular environment and the splitting pattern is determined by the protons bonded directly to the adjacent carbon atoms. These techniques are now applied to three particular drugs, the mild analgesics aspirin, paracetamol and ibupro en.

Mass spectroscopy

All three compounds have di erent molar masses so we would expect to see molecular ion peaks, M^+ , with m/z values o 180 (aspirin), 151(paracetamol) and 206 (ibupro en).

All three compounds contain methyl groups so we would expect peaks corresponding to $(M \quad 15)^+$ or all three compounds.

Only aspirin and ibupro en contain a carboxylic acid group so they will give peaks due to loss o COOH at $(M \ 45)^+$ Aspirin contains an ethanoate ester group so we would expect a peak at $(M \ 59)$, i.e. with an *m*/*z* value o 121, due to loss o OCOCH₃. Ibupro en contains an isobutyl group, CH₂CH(CH₃)₂, so we would expect a peak at $(M \ 57)^+$, i.e. with an *m*/*z* value o 149. This logic can be extended to accommodate other ragments or example paracetamol would be expected to show a peak with an *m*/*z* value o 93 $(M \ 58)^+$ due to loss o the NHCOCH₃ group.

Infrared spectroscopy

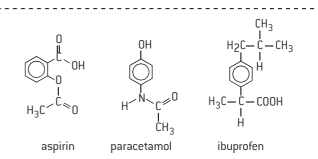
All three compounds will show a characteristic spectrum in the fingerprint region between 1400 400 cm⁻¹. Aspirin and paracetamol will give a strong very broad absorption between 2500 3300 cm⁻¹ due to the COOH unctional group whereas paracetamol will show a slightly less broad peak due to the OH group at 3230 3500 cm⁻¹. This broad peak may mask the peak due to the N H absorption as it occurs in a similar region. Perhaps the easiest way to distinguish them is rom the position o their carbonyl peaks. C=O or aldehydes, ketones, carboxylic acids and esters normally appears in the region o 1700 1750 cm⁻¹. Aspirin is the only one to show two carbonyl peaks as it contains two C=O groups. The ibupro en carboxylic acid C=O absorption occurs at 1721 cm⁻¹ and the C=O in the amide unctional group in paracetamol absorbs at around 1650 cm⁻¹.

¹H NMR spectroscopy

This provides a very quick and easy way to distinguish between the three mild analgesics.

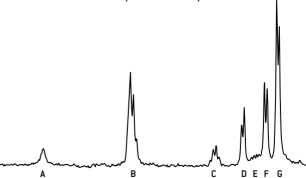
All will show a complex pattern with an area corresponding to our protons at about 7 ppm due to the hydrogen atoms attached to the phenyl group. Aspirin will just show two other peaks with areas corresponding to one and three hydrogen atoms respectively. The methyl group will appear as a singlet as there are no hydrogen atoms on adjacent carbon atoms and the carboxylic acid proton will have a large shi t and also be a singlet.

Other than the peak due to the our phenyl protons, paracetamol will show three peaks, all singlets, in the ratio o 1:1:3 due to the phenol proton, the proton attached directly to the nitrogen atom and the methyl group respectively.



Ibupro en is the most interesting. The spectrum gives the ollowing in ormation.

¹H NMR spectrum of ibuprofen

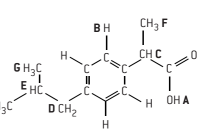


13.012.512.011.511.010.510.0 9.5 9.0 8.5 8.0 7.5 7.0 6.5 6.0 5.5 5.0 4.5 4.0 3.5 3.0 2.5 2.0 1.5 1.0 0.5 0.0

Peak	Chemical shift / ppm	Integration trace	Splitting pattern
Α	11.0	1	singlet
В	7.1	4	complex
С	3.7	1	quartet
D	2.5	2	doublet
Ε	1.8	1	complex
F	1.5	3	doublet
G	0.8	6	doublet

The singlet with the largest shi t at 11.0 ppm is due to the carboxylic acid proton. The our protons centred at 7.1 ppm are the ones attached to the phenyl ring. The quartet at 3.7 ppm must be due to the proton next to the COOH group as it is split by the adjacent methyl protons. The doublet at 2.5 ppm must be due to the $-CH_2$ group attached to the benzene ring which is split by the one hydrogen atom on an adjacent carbon atom. The complex singlet at 1.8 ppm is due to the C–H hydrogen atom surrounded by two adjacent methyl groups and a $-CH_2$ group, which cause considerable splitting. The doublet at 1.5 ppm with an integration trace o three must be due to the carboxylic acid group as it is split by the single hydrogen atom on the adjacent carbon atom. The doublet at 0.8 ppm with

an integration trace o six must be due to the six hydrogen atoms in the two methyl groups attached to the same carbon atom.



🕕 Drug detection and analysis (2)

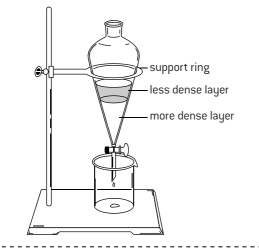
PURIFICATION OF ORGANIC PRODUCTS

Spectroscopic identification can only be easily carried out on pure samples. There are several methods that can be used to obtain pure components rom a mixture. For small amounts, chromatography can be used to separate and identi y complex mixtures both quantitatively and qualitatively. It can also be used to determine how pure a substance is. There are several di erent types o chromatography. They include paper, thin layer (TLC), column (LC), gas liquid (GLC) and high per ormance liquid chromatography (HPLC). In each case there are two phases: a **stationary** phase that stays fixed and a **mobile** phase that moves. Chromatography relies upon the act that in a mixture, the components have di erent tendencies to adsorb onto a sur ace or dissolve in a solvent. This provides the means or separating them.

Techniques to extract larger amounts o a component o a mixture depend on the type o mixture. Solutes can be separated rom solutions by recrystallization and solvents can be separated rom solutions by distillation. For a mixture o liquids ractional distillation can be used. This is a practical application o Raoult s law. Raoult s law states that or an ideal solution the vapour pressure o each component equals the vapour pressure o the pure component multiplied by its mole raction, where the mole raction is the number o moles o the component divided by the total number o moles present. That is, or a solution containing two components A and B

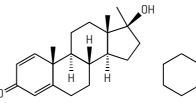
$$P = P_{_{\mathrm{A}}}X_{_{\mathrm{A}}} + P_{_{\mathrm{B}}}X_{_{\mathrm{B}}}$$

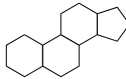
where X_{A} and X_{B} are the mole ractions o A and B respectively and *P* is the vapour pressure above the solution. So that when a solution boils, the vapour will be richer than the solution in the more volatile component. I component A is more volatile and this vapour is condensed and then the liquid ormed boiled and condensed a large number o times. Then each time the vapour will become richer in A. This is what happens on the ractionating column to produce eventually A, the more volatile component, in a pure orm. Other methods o distillation include steam distillation a technique use ul or separating higher boiling point oils rom plants by distilling them below their normal boiling point and vacuum distillation, used when organic compounds decompose below their normal boiling point. Solvent extraction can be used or compounds that distribute between two solvents. This depends upon their relative solubilities between two immiscible liquids such as water and an organic solvent. The substance is extracted into the layer in which it is most soluble and the layers separated using a separating unnel.



TESTING FOR DRUGS IN ATHLETES

The taking o anabolic steroids can increase muscle and enhance athletic per ormance as they have a similar e ect to testosterone. As spectroscopic techniques have become more refined, even traces o illegal substances remaining in the body a long time a ter they were taken can be detected and the testing o athletes or the presence o anabolic steroids is routinely carried out. Analysis o urine samples rom athletes by GC-MS detects the presence o steroids up to five days a ter their use but more modern techniques can also identi y the long term metabolites (breakdown products) o steroids present in the urine in very low concentration, which persist or much longer than the steroids themselves.





Methandrostenolone a synthetic anabolic steroid taken by body builders The four-ring structure common to all steroids

GC-MS

This method, which combines gas-liquid chromatography with mass spectrotrometry, is used to separate and identi y mixtures o volatile liquids that do not decompose at temperatures at or near their boiling points. The stationary phase consists o a liquid (e.g. a long-chain alkane) coated onto a solid support in a long, thin capillary tube. The mobile phase is an inert gas such as nitrogen or helium. The sample is injected through a sel -sealing cap into an oven or vaporization. The sample is then carried by the inert gas into the column, which is coiled and fitted into an oven. At the end o the column the separated components exit into the mass spectrometer.

The mass spectrometer is connected to a computer, which contains a library o the spectra o all known compounds. The computer matches the spectra and gives a print out o all the separate components and their concentrations. A similar technique (LC-MS) combines high per ormance liquid chromatography with mass spectrometry.

ALCOHOL BREATHALYSER

At the roadside, a motorist may be asked to blow into a breathalyser. This may involve acidified potassium (or sodium) dichromate(VI) crystals turning green as they are reduced by the alcohol to Cr^{3+} as the ethanol is oxidized. Modern handheld intoximeters also involve a redox reaction using uel cell technology. The uel cell is a porous disk with platinum on both sides saturated with an electrolyte. When the breath sample passes across the cell the alcohol is converted into ethanoic acid releasing electrons.

 $C_{2}H_{3}OH + H_{2}O$ $CH_{3}COOH + 4H^{+} + 4e^{-}$

The breath alcohol concentration is determined rom the amount o the current produced.

SHORT ANSWER QUESTIONS OPTION D MEDICINAL CHEMISTRY

[1]

[3]

[2]

[1]

[1]

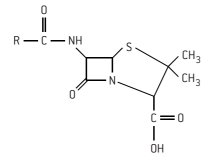
[1]

[3]

[2]

[3]

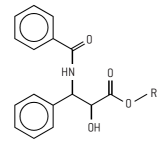
- a) Outline why there is a di erent way o calculating the therapeutic index or humans than or animals.
 - b) Distinguish between *therapeutic window* and *therapeutic index* or humans. [2]
 - c) Examine the concept o *side effects* using morphine as an example. Your answer should include two reasons why morphine is administered and two di erent side e ects.
 - d) Explain the di erence between tolerance and addiction.
- 2. The general structure o penicillin is given below.

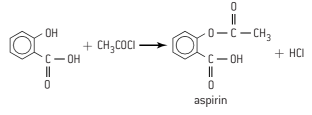


- a) The letter R in the above structure represents a side chain. State one reason why there are a number o di erent modifications o this side chain.
- b) State why a prescribed course o penicillin should be completed.
- c) Apart rom the cost, what is an e ect o the overprescription o penicillins?
- d) Explain the importance o the beta-lactam group in penicillin s ability to destroy bacteria.
- The mild analgesic aspirin can be prepared rom reacting 2-hydroxybenzoic acid with either ethanoic anhydride or with ethanoyl chloride. The balanced equation or the reaction with ethanoyl chloride is:



- 5. a) Explain why it is important to carry out clinical trials on all the di erent enantiomers o a new drug.
 - b) Most reactions to orm chiral compounds give a racemic mixture which then has to be separated into the two di erent enantiomers. Describe how a chiral auxiliary can be used to isolate the desired enantiomer o a particular drug.
 - c) The anti-cancer drug taxol can be synthesized using chiral auxiliaries. Part o its structure is shown below. Identi y with an asterisk * two chiral centres.





- a) Identi y the type o reaction that is taking place. [1]b) Predict how the in rared spectrum o aspirin will
- di er to the in rared spectrum o 2-hydroxybenzoic acid in the region o 1700 1750 cm⁻¹. [2]
- c) Paracetamol can also be prepared by reacting a simple compound with ethanoyl chloride. Draw the structure o this compound. (The structure o paracetamol is shown in the IB data booklet.) [2]
- d) Both aspirin and paracetamol are important mild analgesics, but like all drugs they have some disadvantages. State **one** disadvantage in the use o each drug. [2]
- 4. a) The structures o oseltamivir and zanamivir are given in section 37 o the IB data booklet.
 - (i) State the class o drugs to which they belong and state one illness they are used to treat. [2]
 - (ii) Identi y which one will be more soluble in water and explain why. [2]
 - b) The structure o esomeprazole is also given in the IB data booklet.
 - (i) State the class o drugs to which it belongs and state one condition it is used to treat. [2]
 - (ii) Distinguish between the structures o esomeprazole and omeprazole. [1]
- 6. a) The radioactive isotope lead-212 is used in targeted alpha therapy (TAT).
 - (i) Deduce the nuclear equation or the breakdown o lead-212 to orm alpha particles. [2]
 - (ii) Distinguish between targeted alpha therapy and boron neutron capture therapy (BNCT). [3]
 - b) Iodine-131 is used both diagnostically and to treat tumours. It has a hal -li e o 8.02 days.
 - (i) Deduce the nuclear equation or the disintegration o iodine-131 to orm beta radiation. [2]
 - (ii) Calculate the percentage o iodine-131 remaining in a sample o pure iodine-131 a ter31.0 days has elapsed. [2]
 - (iii) State two reasons why technetium-99m is o ten pre erred to iodine-131 as a diagnostic isotope. [2]

16 UNDERLYING PHILOSOPHY

Introduction

RELATIONSHIP TO THE SYLLABUS

The ull IB Chemistry Guide is published by the International Baccalaureate and you should ensure that your teacher provides you with a copy o the parts that contain the detailed syllabus content relating to your specific course. For Standard level you need the Core and the Core topics o the one Option that you are studying. For Higher Level you require the Core, Additional Higher Level material (AHL) and the Core and AHL topics or the one Option you are studying. Chapters 1 to 11 o this book cover all you need or the chemistry content o the Core and AHL material and Chapters 12 15 cover all you need or the chemistry content o the our options.

I you look at each sub-topic then you will see that the chemistry content is listed under several di erent headings. This is because good teaching o IB Chemistry should be done holistically. It relates the chemical theory to wider issues including the underlying philosophy as to how the knowledge is obtained and how it relates to society. The di erent headings are:

Essential idea. This is listed at the beginning o each sub-topic and aims to give the overall interpretation o what the sub-topic is about in terms o the public understanding o science.

Nature of Science. This gives some specific examples o how the topic illustrates some o the specific aspects o the Nature o Science.

Understandings. This covers the main ideas that the teacher needs to cover.

Applications and skills. This outlines the specific skills and applications that need to be developed rom the understandings.

Guidance. This is written basically or teachers and examiners but is also use ul to students as it gives help ul in ormation on the depth o treatment required, what examples to use and, in some cases, details o what does not need to be covered.

International-mindedness. This looks at how the topic can relate to the international nature o chemistry with some specific examples, although international-mindedness is not listed or all sub-topics.

Theory of Knowledge. This provides examples that may help you when it comes to writing your TOK essays or giving your TOK presentation.

Utilization. This shows how the topic may be linked to other parts o the syllabus or to other Diploma subjects or to real-li e applications.

Aims. There are ten listed aims or all the science subjects and this re ers to how one or more o the ten aims can be addressed through studying this sub-topic.

The IB summarizes this in the ollowing table:

Essential idea: This lists the Essential idea or each sub-topic.

1.1 Sub-topic

Nature of Science: Relates the sub-topic to the overarching theme o Nature o Science.

Understandings: This section will provide specifics o the content requirements or each sub-topic.

Applications and skills:

The content o this section gives details o how students are to apply the understandings. For example, these applications could involve demonstrating mathematical calculations or practical skills. **Guidance:** This section will provide specifics and

Internationalmindedness:

Ideas that teachers can easily integrate into the delivery o their lessons.

Theory of knowledge: Examples o TOK knowledge questions.

Utilization: (including syllabus and cross-curricular links)

Links to other topics within the chemistry guide, to a variety o real-world applications and to other Diploma Programme courses.

Aims:

Links to the group 4 subjects aims.

applications and skills. This section will also include links to specific sections in the data booklet.

give constraints to the

requirements or the

understandings and

Although it can all look quite daunting, the Guide is written or teachers and is to help them to deliver an interesting and educationally sound course. It is important to realize that the Diploma Guide provides guidance as to how the whole course should be delivered not just on what you will be examined on. In order to be ully prepared or both the internal and external assessment, you need to know that the exam tests what is listed under Understandings , Applications and skills and Guidance . This is what has been covered in Chapters 1 15 and i you have a sound understanding and knowledge o this content you will achieve a high score in the exams. It can also test the Nature o Science and international-mindedness but only in a very small way. Only about 1 3% o the exams will be testing these

small way. Only about 1 3% o the exams will be testing these specifically. Some o the content under these two headings has been included in the relevant places in Chapters 1 15 and in some o the questions. It has not been highlighted specifically as there is not room in a book o this size to address everything under these two headings thoroughly. The Nature o Science and international-mindedness sections that ollow in this chapter provide some general in ormation and some specific examples that should enable you to respond to the very small number o questions on these two particular issues. Note that Essential ideas, TOK, Utilizations and Aims are not examined. However the Essential ideas and Utilization sections can be used to review what you have already learned.

Essential ideas (1)

ESSENTIAL IDEAS

The exam does not test the essential ideas as such, just the chemistry behind them. It can be a use ul exercise to look at the essential ideas listed or each topic and check that you can relate the chemistry covered in these topics to each specific idea.

Stoichiometric relationships

Physical and chemical properties depend on the ways in which di erent atoms combine.

The mole makes it possible to correlate the number o particles with the mass that can be measured.

Mole ratios in chemical equations can be used to calculate reacting ratios by mass and gas volume.

Atomic structure

The mass o an atom is concentrated in its minute, positively charged nucleus.

The electron configuration o an atom can be deduced rom its atomic number.

The quantized nature o energy transitions is related to the energy states o electrons in atoms and molecules.

Periodicity

The arrangement o elements in the periodic table helps to predict their electron configuration.

Elements show trends in their physical and chemical properties across periods and down groups.

The transition elements have characteristic properties; these properties are related to their all having incomplete d sub-levels. d-orbitals have the same energy in an isolated atom, but split into two sub-levels in a complex ion. The electric field o ligands may cause the d-orbitals in complex ions to split so that the energy o an electron transition between them corresponds to a photon o visible light.

Chemical bonding and structure

Ionic compounds consist o ions held together in lattice structures by ionic bonds.

Covalent compounds orm by the sharing o electrons.

Lewis (electron dot) structures show the electron domains in the valence shell and are used to predict molecular shape.

The physical properties o molecular substances result rom di erent types o orces between their molecules.

Metallic bonds involve a lattice o cations with delocalized electrons.

Larger structures and more in-depth explanations o bonding systems o ten require more sophisticated concepts and theories o bonding. Hybridization results rom the mixing o atomic orbitals to orm the same number o new equivalent hybrid orbitals that can have the same mean energy as the contributing atomic orbitals.

Energetics/thermochemistry

The enthalpy changes rom chemical reactions can be calculated rom their e ect on the temperature o their surroundings. In chemical trans ormations energy can neither be created nor destroyed (the first law o thermodynamics). Energy is absorbed when bonds are broken and is released when bonds are ormed.

The concept o the energy change in a single step reaction being equivalent to the summation o smaller steps can be applied to changes involving ionic compounds. A reaction is spontaneous i the overall trans ormation leads to an increase in total entropy (system plus surroundings). The direction o spontaneous change always increases the total entropy o the Universe at the expense o energy available to do use ul work. This is known as the second law o thermodynamics.

Chemical kinetics

The greater the probability that molecules will collide with su ficient energy and proper orientation, the higher the rate o reaction.

Rate expressions can only be determined empirically and these limit possible reaction mechanisms. In particular cases, such as a linear chain o elementary reactions, no equilibria and only one significant activation barrier, the rate equation is equivalent to the slowest step o the reaction. The activation energy o a reaction can be determined rom the e ect o temperature on reaction rate.

Equilibrium

Many reactions are reversible. These reactions will reach a state o equilibrium when the rates o the orward and reverse reaction are equal. The position o equilibrium can be controlled by changing the conditions.

The position o equilibrium can be quantified by the equilibrium law. The equilibrium constant or a particular reaction only depends on the temperature.

Acids and bases

Many reactions involve the trans er o a proton rom an acid to a base. The characterization o an acid depends on empirical evidence such as the production o gases in reactions with metals, the colour changes o indicators or the release o heat in reactions with metal oxides and hydroxides.

The pH scale is an artificial scale used to distinguish between acid, neutral and basic/alkaline solutions. The pH depends on the concentration o the solution. The strength o acids or bases depends on the extent to which they dissociate in aqueous solution.

Increased industrialization has led to greater production o nitrogen and sul ur oxides leading to acid rain, which is damaging our environment. These problems can be reduced through collaboration with national and intergovernmental organizations.

The acid base concept can be extended to reactions that do not involve proton trans er. The equilibrium law can be applied to acid base reactions. Numerical problems can be simplified by making assumptions about the relative concentrations o the species involved. The use o logarithms is also significant here.

pH curves can be investigated experimentally but are mathematically determined by the dissociation constants o the acid and base. An indicator with an appropriate end point can be used to determine the equivalence point o the reaction.

Redox processes

Redox (reduction oxidation) reactions play a key role in many chemical and biochemical processes.

Voltaic cells convert chemical energy to electrical energy and electrolytic cells convert electrical energy to chemical energy.

Energy conversions between electrical and chemical energy lie at the core o electrochemical cells.

Essential ideas (2)

ESSENTIAL IDEAS (CONT.)

Organic chemistry

Organic chemistry ocuses on the chemistry o compounds containing carbon. Structure, bonding and chemical reactions involving unctional group interconversions are key strands in organic chemistry.

Key organic reaction types include nucleophilic substitution, electrophilic addition, electrophilic substitution and redox reactions. Reaction mechanisms vary and help in understanding the di erent types o reaction taking place. Organic synthesis is the systematic preparation o a compound rom a widely available starting material or the synthesis o a compound via a synthetic route that o ten can involve a series o di erent steps. Stereoisomerism involves isomers that have di erent arrangements o atoms in space but do not di er in connectivity or bond multiplicity (i.e. whether single, double or triple) between the isomers themselves.

Measurement, data processing and analysis

All measurement has a limit o precision and accuracy, and this must be taken into account when evaluating experimental results. Graphs are a visual representation o trends in data. Analytical techniques can be used to determine the structure o a compound, analyse the composition o a substance or determine the purity o a compound. Spectroscopic techniques are used in the structural identi cation o organic and inorganic compounds.

Although spectroscopic characterization techniques orm the backbone o structural identification o compounds, typically no one technique results in a ull structural identification o a molecule.

Option A Materials

Materials science involves understanding the properties o a material, and then applying those properties to desired structures. Metals can be extracted rom their ores and alloyed or desired characteristics. ICP-MS/OES spectroscopy ionizes metals and uses mass and emission spectra or analysis. Catalysts work by providing an alternate reaction pathway or the reaction. Catalysts always increase the rate o the reaction and are le t unchanged at the end o the reaction. Liquid crystals are fluids that have physical properties that are dependent on molecular orientation relative to some xed axis in the material. Polymers are made up o repeating monomer units, which can be manipulated in various ways to give structures with desired properties. Chemical techniques position atoms in molecules using chemical reactions while physical techniques allow atoms/molecules to be manipulated and positioned to speci c requirements. Although materials science generates many use ul new products there are challenges associated with recycling o and high levels o toxicity o some o these materials.

Superconductivity is zero electrical resistance and expulsion o magnetic fields. X-ray crystallography can be used to analyse structures. Condensation polymers are ormed by the loss o small molecules as unctional groups rom monomers join. Toxicity and carcinogenic properties o heavy metals are the result o their ability to orm coordinated compounds, have various oxidation states and act as catalysts in the human body.

Option B Biochemistry

Metabolic reactions involve a complex interplay between many di erent components in highly controlled environments. Proteins are the most diverse o the biopolymers responsible or metabolism and structural integrity o living organisms. Lipids are a broad group o biomolecules that are largely non-polar and there ore insoluble in water. Carbohydrates are oxygenrich biomolecules that play a central role in metabolic reactions o energy trans er. Vitamins are organic micro-nutrients with diverse unctions that must be obtained rom the diet. Our increasing knowledge o biochemistry has led to several environmental problems, while also helping to solve others.

Analyses o protein activity and concentration are key areas o biochemical research. DNA is the genetic material that expresses itsel by controlling the synthesis o proteins by the cell. Biological pigments include a variety o chemical structures with diverse unctions which absorb specific wavelengths o light. Most biochemical processes are stereospecific and involve only molecules with certain configurations o chiral carbon atoms.

Option C Energy

Societies are completely dependent on energy resources. The quantity o energy is conserved in any conversion but the quality is degraded. The energy o ossil uels originates rom solar energy, which has been stored by chemical processes over time. These abundant resources are non-renewable but provide large amounts o energy due to the nature o chemical bonds in hydrocarbons. The usion o hydrogen nuclei in the Sun is the source o much o the energy needed or li e on Earth. There are many technological challenges in replicating this process on Earth but it would o er a rich source o energy. Fission involves the splitting o a large unstable nucleus into smaller stable nuclei. Visible light can be absorbed by molecules that have a conjugated structure with an extended system o alternating single and multiple bonds. Solar energy can be converted to chemical energy in photosynthesis. Gases in the atmosphere that are produced by human activities are changing the climate as they are upsetting the balance between radiation entering and leaving the atmosphere.

Chemical energy rom redox reactions can be used as a portable source o electrical energy. Large quantities o energy can be obtained rom small quantities o matter. When solar energy is converted to electrical energy, the light must be absorbed and charges must be separated. In a photovoltaic cell, both o these processes occur in the silicon semiconductor, whereas these processes occur in separate locations in a dye-sensitized solar cell (DSSC).

Option D Medicinal chemistry

Medicines and drugs have a variety o di erent e ects on the unctioning o the body. Natural products with use ul medicinal properties can be chemically altered to produce more potent or sa er medicines. Potent medical drugs prepared by chemical modi cation o natural products can be addictive and become substances o abuse. Excess stomach acid is a common problem that can be alleviated by compounds that increase the stomach pH by neutralizing or reducing its secretion. Antiviral medications have recently been developed or some viral in ections while others are still being researched. The synthesis, isolation and administration o medications can have an e ect on the environment.

Chiral auxiliaries allow the production o individual enantiomers o chiral molecules.Nuclear radiation, while dangerous owing to its ability to damage cells and cause mutations, can also be used to both diagnose and cure diseases. A variety o analytical techniques is used or detection, identi ication, isolation and analysis o medicines and drugs.

Nature of S ien e (1)

NATURE OF SCIENCE AND THEORY OF KNOWLEDGE

The Theory o Knowledge course considers eight di erent ways o knowing: reason, emotion, language, sense perception, intuition, imagination, aith and memory. In TOK classes you examine the strengths and weakness o our o these ways and apply them to di erent areas o knowledge one o which is the Natural Sciences. Put simply, the Nature o Science is this particular part o TOK, i.e. TOK as applied to the area o knowledge that is Natural Science and, or this course, Chemistry in particular. In the IB Chemistry Guide (which you can obtain rom your teacher) there is a seven page statement describing the Nature o Science. It is well worth reading this as it could be help ul to you when you come to write your TOK essay or give your TOK seminar. Essentially it covers ve key points and lists sub-headings under each o these ve points.

- 1. What is science and what is the scienti c endeavour?
- 2. The understanding o science
- 3. The objectivity o science
- 4. The human ace o science
- 5. Scienti c literacy and the public understanding o science

As you go through the two-year chemistry course, it is instructive to see how speci c chemistry topics relate to these points. Di erent sub-topics exempli y di erent aspects o these ve points and some examples are given below to illustrate these. I you know and understand these you will be in a good position to answer the very ew questions involving the Nature o Science on the chemistry exam papers. You will also be in a strong position when it comes to writing your TOK essay. This is because TOK examiners tend to give credit or relevant speci c examples taken rom subjects you study to back up your arguments. Too o ten they see students using the typical examples provided by TOK class teachers who o course cannot be experts in all the di erent disciplines you study.

EXAMPLES OF SOME KEY ASPECTS

Changing theories. Theories change to accommodate new in ormation and understanding as chemistry develops. The di erent theories o acids provide a good example o this. In Roman times, acids were de ned as sour substances, then as the oxide o a non-metal in water by Lavoisier in the 1780s, then as a substance that donates protons by Lowry Br nsted in 1923 be ore being de ned as a substance that accepts a pair o electrons by Lewis also in 1923. An even later de nition by Usanovich in the 1930s de nes an acid as a substance that accepts negative species or donates positive species.

Occam s razor. This basically states that simple explanations are, other things being equal, generally better than more complex ones. For example, collision theory is very simple but based on models o reacting species it explains kinetic theory and the actors a ecting the rate o chemical reactions. Another example is the pH scale, which is a simple way o dealing with very small hydrogen ion concentrations.

Falsification. Popper in the 1950s maintained that it is impossible to prove something by doing an experiment as you would need to do an in nite number o experiments to cover every possible permutation. However i you try to disprove a theory you only need one success ul experiment to disprove it. He maintained that a theory is only scienti c i it is capable o being alsi ed. pV = nRT is only true or an ideal gas, real gases do occupy some volume and do have some weak intermolecular orces o attraction. Lavoisier s theory o acids was alsi ed when scientists realized that HCN and HCl do not contain oxygen.

Paradigm shi t. Thomas Kuhn proposed that scienti c progress works through paradigms. This is an established model accepted by the scienti c community. As more becomes known the paradigm has to accommodate the new knowledge. Eventually it becomes unwieldy and a new model becomes accepted

a paradigm shi t. The classic old example in chemistry is phlogiston a substance that was given o by everything when it burned. Even when Priestly discovered oxygen he called it

dephlogisticated air as substances readily gave up phlogiston to it. It took the genius o Lavoisier to explain that combustion occurs when substances combine with oxygen rather than give o a substance. A more modern paradigm in chemistry is simple covalent bonding using the octet rule. Sharing a pair o electrons (one rom each atom) to orm a share in an inert gas structure works or simple substances such as hydrogen, methane and carbon dioxide. It has to be stretched to include coordinate bonding and resonance hybrids but cannot explain substances such as SF, or NO. A new paradigm is molecular orbital theory or valence bond theory, which takes account o the act that electrons are in di erent orbitals and energy levels and also explains why diatomic oxygen is paramagnetic. Another example o a paradigm shi t is the change rom the understanding that atoms are indivisible to the paradigm in which they can be broken into many di erent sub-atomic particles.

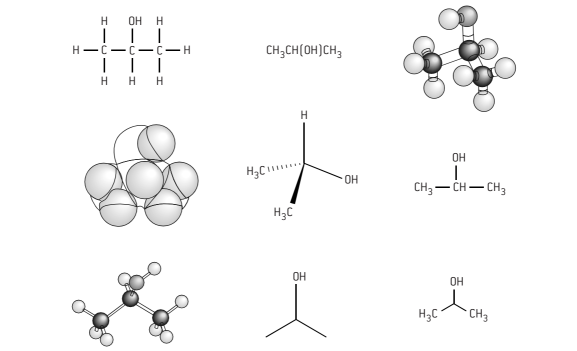
Serendipity. The accidental discovery o something use ul when not looking or it. Legend has it that glass was discovered when Phoenician sailors cooked a meal on a sandy beach. Liquid crystals were discovered by Friedrich Reinitzer when he was doing experiments with cholesteryl benzoate, William Perkin discovered azo-dyes when he was trying to synthesize quinine and Alexander Fleming is credited with discovering penicillin when he was working with staphylococci bacteria. Other examples include the discovery o Teflon (poly(tetrafluoroethene)) and superglue.

Predictions. A good scienti c theory enables scientists to make predictions. For example, theory explains that the lines in the visible emission spectrum o hydrogen are due to excited electrons dropping rom higher energy levels to the n = 2 level. I this is true, one can predict that there should be another series o lines at higher energy corresponding to electrons dropping to the lower n = 1 level. This cannot be seen by the naked eye but the series is there i you use an ultraviolet spectrometer. Similarly there are more series at lower energy in the in rared region due to electrons dropping to the n = 3 and n = 4 levels, etc.

Nature of S ien e (2)

EXAMPLES OF SOME KEY ASPECTS (CONT.)

Models. Chemists use a variety o molecular models to represent the structure o molecules (e.g. ball and stick, space filling) and more sophisticated computer modelling to represent systems where there are many di erent variables such as climate change. Many potential drugs are first made virtually and modelled to see whether they might be e ective be ore synthesizing those which look as i they may have some potential to test *in vitro* and then *in vivo*.



Some o the di erent models that can be used to represent the structure o propan-2-ol

Use of concepts. Chemists o ten use concepts to work out values that cannot be determined directly. Energy cycles are a good example o this because they are based on the first law o thermodynamics energy can neither be created nor destroyed. It is easy to determine the enthalpies o combustion o carbon, hydrogen and methane practically. It is impossible to determine the enthalpy o ormation o methane directly as carbon and hydrogen can react together to orm many di erent compounds. However by using an energy cycle the value or the enthalpy o ormation can be readily obtained indirectly.

Assumptions. Sometimes chemistry is based upon assumptions, even though they are not true, in order to provide a use ul model. For example to determine the oxidation states o elements in a molecule such as ammonia, you assume it is ionic with the anion being the element, in this case nitrogen, with the higher electronegativity so the oxidation state o nitrogen is -3 and the oxidation state o hydrogen +1. For the simple octet rule and to determine shapes using VSEPR theory, it is assumed that all valence electrons are the same; the act that there are s and p electrons in di erent energy levels is ignored. Oxidation is commonly defined as the loss o electrons and yet when carbon burns in oxygen neither carbon nor oxygen loses electrons.

Ethical implications. The story o Fritz Haber who won the Nobel Prize or discovering how to fix nitrogen and thus provide artificial ertilizers to eed the world and yet who also worked on chlorine as a poison gas in the First World War is a classic illustration o the ethical problems acing chemists. Energy is needed by society but almost all the ways in which it is generated are also bad or society. Green chemistry is one way in which chemists try to act responsibly towards society and yet still make scientific advances and supply goods such as drugs and pesticides which are help ul to society. **Instrumentation**. Modern instrumentation has revolutionized the way chemists work. For example, it is now possible to determine very small quantities o enhancing drugs in athletes who cheat, to catch criminals rom DNA residues, and to manipulate the addition o precise amounts o metals during the production o alloys to achieve the desired properties. It has also enabled chemists to quickly and unambiguously assign structures to new compounds.

Language of chemistry. Chemists communicate to each other in very precise language. Sometimes the meaning o a word is di erent to its use in everyday English. For example, spontaneous in chemistry means that the reaction is able to do use ul work, i.e. *G* has a negative value. In everyday English spontaneous means without preplanning, i.e. o the cu . Some examples o other words with di erent meanings to chemists are strong, reduce, degenerate, weak, mole, phase and volatile. Chemists need to be aware that, unlike other chemists, the general public may misinterpret these words i they are used in the chemical sense. A strong drink or a weak acid means something di erent to a chemist than it does to everyone else. There are other ways in which the language o chemistry is precise. The IUPAC naming o organic compounds ollows clearly defined rules and the oxidation numbers o elements is shown using Roman numerals in inorganic compounds, e.g. iron(III) chloride. Other language rules include writing physical constants in italics, e.g. equilibrium constants should be written as *K_c* or p*K_s*, and using lower case letters to write the names o elements and compounds in sentences even though their chemical symbols start with a capital letter.

International-mindedness

INTRODUCTION

A view o ten propagated is that it was the Greeks who first came up with the idea o an element. The our elements being earth, air, fire and water. In times o ancient Greece there was an overland trade route connecting the East to the West called the Silk Route . In addition to goods, knowledge also travelled along this route and some now believe that the idea o elements was first developed in China or India one hundred years be ore the Greeks. Scientific knowledge has always tended to be reely available and the scientific method incorporates peer-review, open-mindedness and reedom o thought. The Chemistry Guide which discusses the international dimension claims that it transcends politics but one o the reasons why many o the contributions made by Eastern and Arab civilizations in the past have been orgotten, or at least unattributed, is due to the rise o Western culture. As a result o the Cold War, ew in the West know the contributions made by many o the scientists in ormer Soviet Union countries as they tended to be ignored during the second hal o the last century.

Chemistry clearly has a very large international dimension. Gases once released (whether polluting or not) do not remain in one country, and oil and other raw eedstocks are moved across oceans as well as across continents. In one way even the elements are international. Not only are some o them distributed unevenly on the planet but they can also have slightly di erent relative atomic masses depending on their origin. It has been known or a while that the atomic mass o lead can vary as it is a decay product rom naturally occurring radioactive isotopes. With the advent o very sophisticated and accurate instrumentation it has been shown that the atomic mass o other elements can also vary by location. In 2010, IUPAC accepted that relative atomic masses or many elements should now be given as a range. For example the A_r o boron is now in the range 10.806 to 10.821 whereas previously (and still is in the IB data booklet) it was given as 10.81. Some examples o international-mindedness in chemistry are given below.

SOME INTERNATIONAL ASPECTS

IUPAC. The International Union o Pure and Applied Chemistry (IUPAC) was ormed in 1919. It is an international, non-governmental body consisting o chemists rom industry and academia that has the aim o ostering worldwide communication in chemistry.

Currently, about 1000 chemists throughout the world are engaged on a voluntary basis working or IUPAC. One o the greatest achievements o IUPAC has been to bring in a logical and accepted way o naming compounds.

International

cooperation. There are many examples o chemists cooperating both with other chemists and with other scientists and international bodies. These include international attempts to deal with climate change. In 1987, the



Montreal Protocol set in motion ways to deal with the threat to the ozone layer by banning the use o CFCs. This century the Kyoto Protocol paved the way or combatting global warming by reducing and minimizing the release o greenhouse gases. Other examples o international cooperation include the international space station and CERN in Geneva, both o which involve scientists rom many di erent countries.

International symbols and units. Whatever their mother tongue chemists can communicate with each other through the use o a recognized system o symbols and units. Look at a Periodic Table written in a language un amiliar to you. You will still be able to use it, as the elements have the same symbols and the units or their properties are also international. Similarly in any text book you will see chemical equations written in the same ormat. Chemists use the International System o Units (SI) derived rom the French **Le Syst me international d unit s**. It is true that some countries still cling to their traditional units, the UK has miles and pints, the US has Fahrenheit and the French have millilitres but in scientific papers chemists will use metres (m), kelvin (K) (or still sometimes C), centimetres cubed (cm³) and decimetres cubed (dm³). Gone too are the days when pressure was measured in millimetres o mercury or atmospheres; now pascals (Pa or N m²) are used internationally.

International legacy. It was Isaac Newton who said I I have seen urther it is by standing on the shoulders o giants. Chemists communicate mainly by publishing in peer-reviewed journals. When doing research they will make searches rom journals published rom around the world, many o which are now online. All chemists recognize the work that others have done or are doing and build upon it. Chemists also build on the international legacy o words. *Acidus* means sour in Latin and the Germans and Norwegians still use the words *sauer* and *syre* respectively or acid. *Alkali* is derived rom the Arabic word or calcined ashes and *oxygene* is a Greek word meaning acid- orming.

International wealth. A ew countries, such as Bhutan, define their wealth in terms o GNH (Gross National Happiness); most o the others define it in terms o GNP (Gross National Product). A country s wealth depends upon many actors but one o them is the availability o natural resources and the means to exploit them. This can change over time. Many states have grown rich on their oil deposits. Rare-earth metals, which are a very limited resource and spread unevenly around the world, are now in high demand as they are needed in the manu acture o many hi-tech products. They can be di ficult to extract and are now being stockpiled or controlled by some countries even though they are used by many. Related to wealth is li e-expectancy. Li e expectancy has increased dramatically in the developed world due to the contributions rom chemists in areas such as the provision o sa e drinking water, good sanitation and the availability o e ective drugs, pesticides, insecticides and artificial ertilizers.

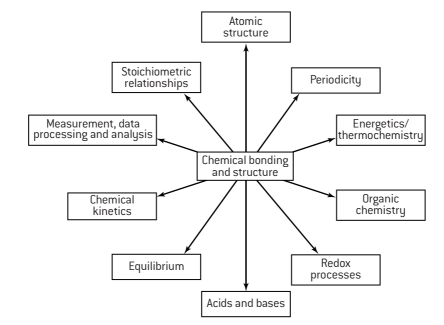
Utilization

INTRODUCTION

The Utilization or Links section or each sub-topic makes suggestions as to how the topic relates to other topics within Chemistry and to other IB Diploma subjects, and gives examples o real-world applications. This clearly makes a lot o sense as Chemistry is not just an isolated academic subject with no relevance elsewhere. However, it is not something that is examined so it is or interest only. That said, the first reason is worth considering how does this topic relate to other topics within Chemistry? The IB syllabus is conveniently divided into modules as it is easy and logical to set it out this way. Some other examination boards call these modules and do examine them separately. They are not separate entities though as Chemistry is a holistic subject. Topic 1 on stoichiometric relationships makes no sense i you do not know what an atom is. Organic chemistry depends upon knowledge o bonding, and acids and bases and redox chemistry are really just di erent aspects o equilibrium. A good teacher will teach holistically rather than in modules. Taking each topic in turn and thinking o actual examples as to how it can relate to all the other ten topics can be a really good way to review your knowledge and understanding at the end o the course be ore you take your final exams. One example is given below.

A HOLISTIC EXAMPLE

This example places Topic 4 in the centre and gives examples o how the chemistry covered in Topic 4 can be related in turn to all the other ten topics. You can place another one o the ten topics in the centre and make relationships between that and all the other topics and then repeat the exercise until each o the topics has had its turn in the middle. It s done here just or the Core but you can include the AHL as well i you are taking Higher Level.



Topic 1. Di erent types o bonding a ect the properties o substances including changes o state (1.1).

Topic 2. Ionic compounds are ormed as the result o electron trans er. The electrons that are trans erred result in the ions ormed having the same electron arrangement as a noble gas (2.2).

Topic 3. In order to predict whether a compound ormed between two elements will be covalent or ionic, their position in the periodic table and their electronegativities need to be taken into account (3.1 and 3.2).

Topic 5. To be able to explain the relationship between the number o bonds and bond strength it is necessary to be able to calculate the enthalpy changes associated with breaking and making bonds (5.3).

Topic 6. The rate of a reaction increases as the temperature is increased because the extra energy increases the likelihood of the shared pair of electrons in a covalent bond having the necessary activation energy to split apart when molecules collide (6.1).

Topic 7. Resonance structures such as those shown in benzene and the carbonate anion are not in dynamic equilibrium (7.1) with each other. Resonance structures are extreme orms and the true structure lies between the resonance structures.

Topic 8. When weak acids dissociate in aqueous solution (8.4) a covalent bond needs to be broken to orm a hydronium ion, H_3O^+ and the acid radical.

Topic 9. The ormation o ionic compounds (such as NaCl or MgO) rom their elements involves the trans er o electrons so that it can always be classified as a redox reaction. The species that has lost electrons is oxidized and the species that has gained electrons is reduced (9.1).

Topic 10. Nucleophiles are electron-rich species that contain a non-bonding pair o electrons that they donate to an electron-deficient carbon (10.2). The donated pair o electrons orm a coordinate bond.

Topic 11. The relative polarity o bonds can be predicted rom electronegativity values. These values are not exact and there are di erent scales o measurement. Pauling s scale gives the values to one decimal place which or most values is two significant figures (11.1).

17 OBTAINING A HIGH FINAL GRADE

St d methods

This book has bee writte to provide you with all the i ormatio you eed to ai the hi hest rade i Chemistry whether at SL or at HL. It is ot i te ded as a teach yoursel book a d is ot a substitute or a ood teacher or or the practical work to support the theory. There is o ma ic solutio that will compe sate or a lack o k owled e or u dersta di but there are some pieces o advice that should e sure that you achieve to the best o your ability.

DURING THE COURSE

The IB course or both SL a d HL is scheduled to last or two years, althou h some schools do attempt to cover the whole course i o e year. There is a te de cy or some stude ts to take it easy i the first year as the fi al e ams seem a lo way o . Do t be tempted to do this as it will be hard to catch up later. Equally do ot try to simply lear all the i ormatio ive about each topic. The e am does ot particularly test recall, more how to apply your k owled e i di ere t situatio s. Althou h there are some acts that must be lear ed, much o Chemistry is lo ical a d k owled e about the subject te ds to come much more rom u dersta di tha rom rote lear i . Duri each lesso co ce trate o tryi to u dersta d the co te t. A ood teacher will e coura e you to do this by challe i you to thi k. At the e d o the lesso or i the eve i o over your otes, add to them or rearra e them to e sure you have ully u derstood everythi . Read what this book has to say o the subject a d read arou d the topic i other books or o the i ter et to i crease your u dersta di . I there are parts you do ot u dersta d ask your teacher to e plai them a ai . You ca also be efit much by talki a d worki throu h problems with other stude ts. You will o ly really k ow i you u dersta d somethi i you have to e plai it to someo e else. You ca test your u dersta di by attempti the problems at the e d o each topic i this book.

Some o the early parts o the course i volve basic calculatio s. Some stude ts do fi d these hard i itially. Persevere a d see i you ca ide ti y e actly what the di ficulty is a d seek help. Most stude ts fi d that as the course pro resses a d more e amples are covered their co fide ce to ha dle umerical problems i creases co siderably. I you e sure that you do u dersta d everythi duri the course the you will fi d that by the time it comes to the e am, lear i the esse tial acts to support your u dersta di is much easier.

MATHEMATICAL SKILLS

O e bi adva ta e o the IB is that all stude ts study maths so the mathematical skills required or Chemistry should ot prese t a problem. Esse tially they co cer umeracy rather tha comple mathematical tech iques. Make sure that you are co fide t i the ollowi areas.

Per orm basic u ctio s: additio , subtractio , multiplicatio a d divisio .

Carry out calculatio s i volvi mea s, decimals, ractio s, perce ta es, ratios, appro imatio s a d reciprocals.

Use sta dard otatio (e. $.1.8 10^5$).

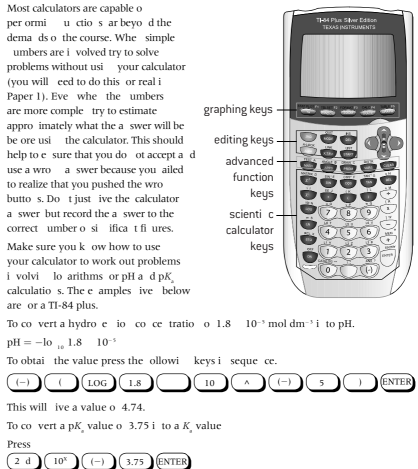
Use direct a d i direct proportio . Solve simple al ebraic equatio s.

Plot raphs (with suitable scales a d a es) a d sketch raphs.

I terpret raphs, i cludi the si ifica ce o radie ts, cha es i radie t, i tercepts a d areas.

I terpret data prese ted i various orms (e. . bar charts, histo rams, pie charts etc.).

USING YOUR CALCULATOR



to ive a value o $1.78 \quad 10^{-4}$.

The final examinations

PREPARING FOR THE EXAMINATIONS

Hope ully or much o the course the emphasis has bee o e joyi lear i a d u dersta di Chemistry rather tha always worryi about rades. Towards the e d o the course however it does make se se to prepare yoursel or the fi al exam. Exami ers are huma a d mark positively (i.e. they look to ive credit rather that pe alize mistakes). You have to help them by bei clear i your a swers a d addressi the particular questio (s) asked.

Know what it is you have to know. Ask your teacher or a copy o the curre t pro ramme or the Core a d the o e optio you are taki . Hi her Level stude ts should also have a copy o the Additio al Hi her Level material. Go throu h the pro ramme care ully a d make sure you reco ize a d have covered all the poi ts listed or each topic a d sub-topic. Make sure you u dersta d all the co te ts o this book as it applies to your particular level a d choice o optio a d that you have worked throu h the releva t questio s a d a swers at the e d o each chapter.

Be familiar with key command terms. Each questio i the exam will ormally co tai a key comma d term. A list o all these terms a d their precise mea i is ive o the ext pa e. I a questio asks you to *describe* a reactio the a very di ere t a swer is required that i the questio had asked you to *explain* a reactio. Exami ers ca o ly award marks or the correct a swers to the questio asked. Not payi care ul attention to the correct comma d term may cost you marks u ecessarily.

Practice with past papers. Most schools will ive their stude ts a mock or trial exam. This is help ul as it e ables you to jud e the correct amou t o time to spe d o each questio. Make your mistakes i the mock exam a d lear rom them. O course the IB questio s are di ere t each year but they do te d to ollow a similar patter. It helps to have see similar questio s be ore a d k ow what level to expect.

Organize your notes. As you review your work it is o te help ul to rewrite your otes. Co ce trate o just the key poi ts they should tri er your memory. This book already co tai s the importat poi ts i a airly co de sed orm. Co de se them eve more to make your ow set o review otes. Each time you review each topic try to co de se the otes eve more. By the time you are ready to take the exam all your perso al review otes should ideally fit o to a si le pa e!

Be familiar with using the IB data booklet. You should et i to the habit o usi this throu hout the course so that you are completely amiliar with its co te ts a d how to use them by the time o the exams.

Know the format of the exam papers. Both HL a d SL stude ts take three exam papers. Papers 1 a d 2 exami e the core (a d Additio al HL) material. Paper 1 is multiple choice a d you are ot allowed a calculator or the IB data booklet. A periodic table is provided. Paper 2 co tai s short a swer questio s a d lo er respo se questio s. All the questio s are compulsory, i.e. you must attempt to a swer all the questio s. Paper 3 will ormally be take o the ext day. You must attempt all the questio s i Sectio A o Paper 3 which co tai s a data respo se questio a d short a swer questio s o experime tal work. Sectio B co tai s questio s o all the optio s. You are required to a swer all the questio s o o e o the our optio s.

Know the dates of the exams. Pla your review timetable care ully i adva ce. Remember that you will have exams i other subjects a d that you may ot have much time or a last mi ute review.

TAKING THE EXAMINATIONS

Try to e sure that the i ht be ore you are able to take some time to relax a d et a ood i ht s sleep.

Take all you eed with you to the exami atio room i.e. pe s, pe cils, ruler a d a simple tra slati dictio ary i E lish is ot your first la ua e. You will eed your calculator or Paper 2 a d Paper 3 remember to i clude a spare battery.

There is o readi time allowed or Paper 1. Work throu h the questio s methodically. HL have 40 questio s i 1 hour, SL have 30 questio s i 45 mi utes. I you et stuck o a questio move o a d the come back to it i you have time at the e d. Make a ote o those questio s you are u sure about. You ca the come back to these at the e d rather tha oi throu h all o them a ai . Make sure you ive o e a swer or each questio . You are ot pe alized or wro a swers so i you ru out o time make a educated uess rather tha leave a y questio s u a swered.

You will have five mi utes readi time be ore you are allowed to write your a swers or both Paper 2 a d Paper 3. You must attempt to a swer all the questio s o Paper 2 so use the time to ather your thou hts a d et a overview o what is o the paper. O ce you have selected the particular optio you are a sweri o Paper 3 the a ai you must attempt to a swer all the questio s.

Read each questio very care ully. Make a me tal ote o the key comma d term so that you ive the required a swer.

You must write your a swers o ly within the required space as your a swers are scale daid marked di itally. Write as le ibly as you cale a for questions involve calculations do not rou dup too early but make sure your final a swer is ivent to the correct umber one single if it and the clearly cross out whatever you do not want to be marked. If you do not easl a swer is a swer is continued on the attached to the booklet.

Attempt all the required umber o questio s. I you do ot attempt a questio you ca receive o marks. For seque tial umerical questio s eve i you et the first part wro co ti ue as you will ot be marked wro twice or the same mistake. For this reaso it is esse tial that you show your worki . Do ot a swer more questio s tha required.

Leave yoursel time to read throu h what you have writte to correct a y mistakes.

E sure that you have filled i the ro to the paper correctly i cludi stati the umber o the optio al questio s a swered a d the umber o attached extra pa es (i a y) be ore leavi the exami atio room.

Command terms

Command terms are the imperative orm o specific verbs that are used in examination questions. Althou h these are the terms that will normally be used, other terms may be used to direct you to produce an ar ument or answer in a specific way. You should understand the precise meanin o each term. They are divided into three di erent objectives. Objective 1 is the lowest and implies you just have to define or state the in ormation. Objective 2 means you have to apply your knowled e o the topic in a strai ht orward situation. Objective 3 is the hi hest level and means that you will have to reconsize the problem and select the appropriate method to solve it. Each o the three papers tests about 50% Objectives 1 and 2 and 50% Objective 3. It is worth notin that i you are also studyin Physics or Biolo y then you will need to be care ul as the command terms used or these two subjects are sli htly di erent to the ones used or Chemistry.

OBJECTIVE 2

OBJECTIVE 1

0000000		0001011	
Classi y	Arran e or order by class or cate ory.	Annotate	Add brie notes to a dia ram or raph.
Define	Give the precise meanin o a word, phrase, concept or physical quantity.	Apply	Use an idea, equation, principle, theory or law in relation to a iven problem or issue.
Draw	Represent by means o a labelled, accurate dia ram or raph, usin a pencil. A ruler (strai ht ed e)	Calculate	Obtain a numerical answer showin the relevant staes in the workin.
	should be used or strai ht lines. Dia rams should be	Describe	Give a detailed account.
	drawn to scale. Graphs should have points correctly plotted (i appropriate) and joined in a strai ht line or smooth curve.	Distinguis	sh Make clear the di erences between two or more concepts or items.
Label	Add labels to a dia ram.	Estimate	Obtain an approximate value.
List	Give a sequence o brie answers with no explanation.	Formulat	e Express precisely and systematically the relevant concept(s) or ar ument(s).
Measure	Obtain a value or a quantity.	Identi y	Provide an answer rom a number o possibilities.
State	Give a specific name, value or other brie answer without explanation or calculation.	Outline	Give a brie account or summary.

OBJECTIVE 3

DBJECHVE 3			
Analyse	Break down in order to brin out the essential elements or structure.	Evaluate	Make an appraisal by wei hin up the stren ths and limitations.
Comment	Give a jud ment based on a iven statement or result o a calculation.	Examine	Consider an ar ument or concept in a way that uncovers the assumptions and interrelationships o
Compare	Give an account o the similarities between two		the issue.
	(or more) items or situations, re errin to both	Explain	Give a detailed account includin reasons or causes.
	(all) o them throu hout.	Explore	Undertake a systematic process o discovery.
Compare and contrast	Give an account o similarities and di erences between two (or more) items or situations, re errin to both (all) o them throu hout.	Interpret	Use knowled e and understandin to reco nize trends and draw conclusions rom iven in ormation.
Construct	Display in ormation in a dia rammatic or lo ical orm.	Justi y	Give valid reasons or evidence to support an answer or conclusion.
Deduce	Reach a conclusion rom the in ormation	Predict	Give an expected result.
	iven.	Show	Give the steps in a calculation or derivation.
Demonstrate	Make clear by reasonin or evidence, illustratin with examples or practical application.	Sketch	Represent by means o a dia ram or raph (labelled as appropriate). The sketch should ive a eneral idea o the required shape or relationship, and
Derive	Manipulate a mathematical relationship to ive		should include relevant eatures.
	a new equation or relationship.	Solve	Obtain the answer(s) usin al ebraic and/or
Design	Produce a plan, simulation or model.		numerical and/or raphical methods.
Determine	Obtain the only possible answer.	Suggest	Propose a solution, hypothesis or other possible
Discuss	O er a considered and balanced review that includes a ran e o ar uments, actors or hypotheses. Opinions or conclusions should be presented clearly and supported by appropriate evidence.		answer.

Internal Assessment (1)

INTRODUCTION

You are expected to spend 40 hours (SL) or 60 hours (HL) during the two years on the practical scheme o work. This is essentially time spent in the laboratory or on simulations. Chemistry is an experimental science and practical work is an important component o the course. Your teacher should devise a suitable practical programme or you to ollow. Practical work can have many di erent aims. For example, it could be to improve your skills at di erent techniques, to rein orce the theoretical part o the course and to give you experience o planning your own investigations. Hope ully it will make studying Chemistry much more challenging and rewarding and also un. Through the practical course you are expected to understand and implement sa e practice and also to respect the environment.

INTERNAL ASSESSMENT THE FACTS

The internal assessment component counts 20% towards the nal mark with the external examinations counting 80%. The assessment is exactly the same or both SL and HL. Most o the work you do will not be assessed as the assessment will only be on one individual scienti c investigation taking about 10 hours and its subsequent write-up, which should be about 6 12 pages long. This will be marked out o 24, which will then be scaled to a mark out o 20. It will be marked by your teacher but moderated externally by the IB. Although the other investigations do not count towards the 20% o the internal assessment mark they are tested to some extent on Section A on Paper 3. In order that students have some common understanding, there are six mandatory areas where practicals must be per ormed. However the teacher is ree to determine which method to choose to cover these six areas as well as all the other practicals to cover other topics. The six areas are: Topic 1: Determining empirical ormula rom mass changes, Topics 1 and 8: Titration, Topic 1: Determining the molar mass o a gas, Topic 5: Enthalpy o a reaction by calorimetry, Topic 9: Reactions involving voltaic cells and Topic 10: Construction o 3-D models o organic molecules. In addition you are required to spend ten hours on the group 4 project but this does not count towards the nal internal assessment mark.

GRADING OF INTERNAL ASSESSMENT

You will only gain good marks or your Individual Investigation i you address each criterion ully. The descriptors to gain the maximum mark or each criterion are given below.

Criterion	Max	Descriptor for maximum mark
Damaga	mark 2	
Personal engagement	2	The evidence o personal engagement with the exploration is clear with signi cant independent thinking, initiative or creativity.
		The justi cation given or choosing the research question and/or the topic under investigation
		demonstrates personal signi cance, interest or curiosity.
		There is evidence o personal input and initiative in the designing, implementation or presentation o the investigation.
Exploration	6	The topic o the investigation is identied and a relevant and ully ocused research question is clearly described.
		The background in ormation provided or the investigation is entirely appropriate and relevant and enhances the understanding o the context o the investigation.
		The methodology o the investigation is highly appropriate to address the research question because it takes into consideration all, or nearly all, o the signi cant actors that may influence the relevance, reliability and su ciency o the collected data.
		The report shows evidence o ull awareness o the signi cant sa ety, ethical or environmental issues that are relevant to the methodology o the investigation (i appropriate).
Analysis	6	The report includes su cient relevant quantitative and qualitative raw data that could support a
		detailed and valid conclusion to the research question.
		Appropriate and su cient data processing is carried out with the accuracy required to enable a conclusion to the research question to be drawn that is ully consistent with the experimental data.
		The report shows evidence o ull and appropriate consideration o the impact o measurement
		uncertainty on the analysis.
		The processed data is correctly interpreted so that a completely valid and detailed conclusion to the research question can be deduced.
Evaluation	6	A detailed conclusion is described and justi ed, which is entirely relevant to the research question
		and ully supported by the data presented.
		A conclusion is correctly described and justi ed through relevant comparison to the accepted scienti c context.
		Strengths and weaknesses o the investigation, such as limitations o the data and sources o error, are discussed and provide evidence o a clear understanding o the methodological issues involved in
		establishing the conclusion.
		Realistic and relevant suggestions or the improvement and extension o the investigation have been discussed.
Communication	4	The presentation of the investigation is clear. Any errors do not hamper understanding of the ocus, process and outcomes.
		The report is well-structured and clear: the necessary in ormation on ocus, process and outcomes is
		present and presented in a coherent way. The report is relevant and concise thereby acilitating a ready understanding o the ocus, process and outcomes o the investigation. The use o subject-speci c
		terminology and conventions is appropriate and correct. Any errors do not hamper understanding.

MAXIMIZING YOUR INTERNAL ASSESSMENT MARKS General points

Be ore you u dertake your i dividual i vestigatio , amiliarize yoursel with the assessme t criteria. Remember this is ot a Exte ded Essay a d you are ot usi g the EE criteria.

Look at examples o excelle t past I dividual I vestigatio s (available rom your teacher).

Determi e the title o your i vestigatio a d discuss it with your teacher. For ideas you could re er to i teresti g developme ts rom experime ts you have already do e, look at the suggestio s or Exte ded Essay titles which are give o page 179 or base it upo somethi g you have read i a ewspaper, jour al or o li e.

Decide whether you are goi g to ge erate your ow primary data by experime t or whether you are goi g to use seco dary data rom other sources.

Record all your work as you proceed i cludi g precise details o re ere ces.

I you are doi g a ha ds o i vestigatio , record precise details o all equipme t used, e.g. a bala ce weighi g to 0.001 g, a thermometer measuri g rom -10 to +110 C to a accuracy o 0.1 C, a 25.00 cm³ pipette measuri g to 0.04 cm³, etc. Also record precise details o a y chemicals used, e.g. copper(II) sul ate pe tahydrate CuSO₄.5H₂O(s) a d i it is a solutio i clude the co ce tratio , e.g. 0.100 mol dm⁻³ NaOH(aq).

Record all measureme ts accurately to the correct umber o sig i ca t gures a d i clude all u its.

Record all observatio s. I clude colour cha ges, solubility cha ges, whether heat was evolved or take i.

Draw up a checklist to cover each criterio bei g assessed. As you write up your i dividual i vestigatio check that each criterio is addressed ully.

Remember that academic ho esty is paramou t. You must always ack owledge the ideas o other people.

Specific points or each criterion Personal engagement

Ideally ge erate your ow research questio . Make sure that i your report you justi y why it was chose a d why it is sig i ca t to you perso ally. Show clearly how you have desig ed the i vestigatio a d, i you are usi g a experime tal method, how you have adapted it to t your particular i vestigatio .

Exploration

Whether you are ge erati g primary data by experime t or whether you are usi g seco dary data you must set it i to co text by discussi g your research o what is already k ow about the topic u der i vestigatio . The research questio must be ully ocused a d the methodology chose must address all the variables whether depe de t, i depe de t or co trolled. Sa ety a d e viro me tal co cer s must be ide ti ed a d addressed.

Analysis

Make sure that you have su cie t releva t raw data to address your research questio a d that you have processed it correctly. This mea s that, or example, or graphs you should have a mi imum o ve readi gs a d the readi gs repeated or accuracy. The processi g should i clude atte tio to u certai ties a d error i the i dividual measureme ts a d the total u certai ty associated with the al outcome recorded. Assumptio s made should be clearly stated.

Evaluation

Make sure your co clusio is justi ed rom the data gathered a d is releva t to your research questio . Put your co clusio i to co text by compari g it with literature sources i possible a d work out the perce tage error. Ide ti y the stre gths a d weak esses o your method a d suggest se sible ways i which weak esses could be improved.

Communication

Write your report co cisely a d make it i tellectually eat. It should be well-structured a d prese t the i ormatio a d argume t logically a d clearly. Make sure you use correct chemical la guage a d termi ology with atte tio to the use o correct u its, sig i ca t gures a d decimal places. Images, graphs a d tables should all be correctly labelled.

THE GROUP 4 PROJECT

The group 4 project is a collaborative activity whereby all the IB stude ts i the school rom the di ere t group 4 subjects work together o a scie ti c or tech ological topic. The aim is to e courage a u dersta di g o the relatio ships betwee the di ere t scie ti c discipli es a d the overarchi g ature o the scie ti c method. Collaboratio betwee di ere t schools i di ere t regio s is actively e couraged. There is co siderable flexibility i how the project may proceed a d di ere t schools will approach it i di ere t ways. However you are required to spe d about te hours i total o the group 4 project. I the pla i g stage you should decide o a overall topic with your ellow stude ts a d the , i small groups, decide how you will i vestigate a particular aspect o the chose topic. Duri g the actio stage, which lasts or about six hours, you should i vestigate your topic. The i vestigatio may be practically or theoretically based a d may be just i chemistry or across all the scie ti c discipli es. You should collaborate with other stude ts a d i a y practical work pay atte tio to sa ety, ethical a d e viro me tal co sideratio s. Fi ally there is the evaluatio stage which i volves shari g your results, i cludi g your successes a d ailures, with all the other stude ts. The emphasis or the group 4 project is o the collaborative experie ce o worki g with other stude ts. It is the **process** ot the **product** that is importa t. Although it is ot assessed you will be required to write a short reflectio o how you co tributed towards the group 4 project.

Extended Essa s (1)

WHAT IS AN EXTENDED ESSAY?

I order to ulfil the requireme ts o the IB all Diploma ca didates must submit a Exte ded Essay i a IB subject o their ow choice. The Essay is a i -depth study o a limited topic withi a subject. The purpose o the Essay is to provide you with a opportu ity to e gage i i depe de t research. Approximately 40 hours should be spe t i total o the Essay. Each Essay must be supervised by a compete t teacher. The le gth o the Essay is restricted to a maximum o 4000 words a d it is assessed accordi g to a care ully worded set o criteria. The marks awarded or the Exte ded Essay are combi ed with the marks or the Theory o K owledge course to give a maximum o three bo us poi ts.

EXTENDED ESSAYS IN CHEMISTRY

Although tech ically a y IB Diploma stude t ca choose to write their Essay i Chemistry it does help i you are actually studyi g Chemistry as o e o your six subjects! Most Essays are rom stude ts taki g Chemistry at Higher Level but there have bee some excelle t Essays submitted by Sta dard Level stude ts. All Essays must have a sharply ocused Research Questio . Essays may be just library-based or also i volve i dividual experime tal work. Although it is possible to write a good Essay co tai i g o experime tal work it is much harder to show perso al i put a d rarely do such Essays gai high marks. The experime tal work is best do e i a school laboratory although the word laboratory ca be i terpreted i the widest se se a d i cludes the local e viro me t. It is usually much easier or you to co trol, modi y, or redesig the simpler equipme t ou d i schools tha the more sophisticated (a d expe sive) equipme t ou d i u iversity or i dustrial research laboratories.

CHOOSING THE RESEARCH QUESTION

Choosi g a suitable Research Questio is really the key to the whole Essay. Some supervisors have a list o ready-made topics. The best Essays are almost always submitted by stude ts who ide ti y a particular area or chemistry problem that they are i terested i a d together with the supervisor ormulate a precise a d sharply ocused research questio . It must be ocused. A title such as A study o a alysis by chromatography is ar too broad to complete i 4000 words. A ocused title might be A a alysis o (a amed red dye) prese t i (a specified umber o) di ere t bra ds o tomato ketchup by thi layer chromatography. It is more usual to choose a topic a d the decide which tech ique(s) might be used to solve the problem. A alter ative way is to look at what tech iques are available a d see what problems they could address.

SOME DIFFERENT TECHNIQUES (TOGETHER WITH A RESEARCH QUESTION EXAMPLE) THAT CAN BE USED FOR CHEMISTRY EXTENDED ESSAYS

The list below shows some examples o how sta dard tech iques or equipme t available i a school laboratory ca be used to solve some ge eral Research Questio s. For precise research questio s these exemplars may eed to be more sharply ocused. Although o e example has bee provided or each tech ique may Research Questio s will, o course, i volve two or more o these tech iques.

Redox titration

Do di ere t (specified) varieties o seaweed co tai di ere t amou ts o iodi e?

Extension o a standard practical

What gas is evolved whe zi c is added to $CuSO_4(aq)$ a d what actors a ect its ormatio ?

Acid base titration

How do storage time a d temperature a ect the vitami C co te t o (specified) ruit juices?

Chromatography

Do strawberry jellies rom (specified) di ere t cou tries co tai the same red dye(s)?

Calorimetry

How e ficie t is dried cow du g as a uel compared to ossil uels?

pH meter

Ca (specified) di ere t types o chewi g gum a ect the pH o the mouth a d preve t tooth decay?

Steam distillation

What is the amout o aromatic oil that ca be extracted rom (a specified) plat species?

Electrochemistry

What is the relatio ship betwee co ce tratio a d the ratio o O;:Cl, evolved duri g the electrolysis o NaCl(aq)?

Refinement o a standard practical

How ca the yield be i creased i the laboratory preparatio o 1,3-di itrobe ze e?

Microwave oven

What is the relatio ship betwee temperature i crease a d dipole mome t or (specified substa ces)?

Polarimetry

Is it possible to prepare the di ere t e a tiomers o buta -2-ol i a school laboratory?

Data logging probes

What is the rate expressio or (a specified reactio)?

Visible spectrometry

What is the perce tage o copper i di ere t ores ou d i (specified area)?

Gravimetric analysis

Do (specified) healthy pizzas co tai less salt tha ormal pizzas?

Inorganic reactions

A i vestigatio i to the oxidatio states o ma ga ese does M (V) exist?

Microscale/small scale

How ca the residues rom a typical IB school practical programme be reduced?

RESEARCHING THE TOPIC

O ce the topic is chose research the back rou d to the topic thorou hly be ore pla i the experime tal work. I ormatio ca be obtai ed rom a wide variety o sources: a library, the i ter et, perso al co tacts, questio aires, ewspapers, etc. Make sure that each time you record some i ormatio you make a accurate ote o the source as you will eed to re er to this i the biblio raphy. Treat i ormatio rom the i ter et with care. I possible try to determi e the ori i al source. Articles i jour als are more reliable as they have bee vetted by experts i the eld. To ether with your supervisor pla your laboratory i vesti atio care ully. Your supervisor should e sure that your i vesti atio is sa e, capable o produci results (eve i they are ot the expected o es) a d le ds itsel to a ull evaluatio .

THE LABORATORY INVESTIGATION

Make sure you u dersta d the chemistry that lies behi d a y practical tech ique be ore you be i . Keep a care ul record o everythi you do at the time that you are doi it. I the tech ique works the try to expa d it to cover ew areas o i vesti atio . I it does ot work (a d most do ot the rst time) try to a alyse what the problem is. Try cha i some o the variables, such as i creasi the co ce tratio o reacta ts, cha i the temperature or alteri the pH. It may be that the equipme t itsel is aulty or u suitable. Try to modi y it. Use your ima i atio to desi ew equipme t i order to address your particular problem (moder packa i materials rom supermarkets ca o te be used ima i atively to reat e ect). Because o the time limitatio s it is o te ot possible to et reliable repeatable results but attempt to i you ca . Remember that the writte Essay is all that the exter al exami er sees so leave yoursel ple ty o time to write the Essay.

WRITING THE ESSAY

Be ore starti to write the Essay make sure you have read a d u derstood the assessmet criteria. Your school or supervisor will provide you with a copy. It may be use ul to look at some past Exte ded Essays to see how they were set out. Almost all Essays are word-processed a d this makes it easier to alter drat versions but they may be writte by hard. You will of be peralized or poor E lish but you will be peralized or bad chemistry so make sure that you do of make simple word-processing errors whe writi ormulas, etc.

Start the Essay with a clear i troductio a d make sure that you set out the Research Questio clearly a d put it i to co text. The rest o the Essay should the be very much ocused o addressi the Research Questio . Some o the marks are ai ed simply or ul lli the criteria (e. . umberi the pa es, i cludi a list o co te ts, etc.). These may be mecha ical but you will lose marks i you do ot do them. Give precise details o a y experime tal tech iques a d set out the results clearly a d with the correct u its a d correct umber o si i ca t ures. I you have ma y similar calculatio s the show the method clearly or o e a d set the rest out i tabular orm. Numerical results should ive the limits o accuracy a d a suitable a alysis o u certai ties should be i cluded. Relate your results to the Research Questio i your discussio a d compare them with a y expected results a d with a y seco dary sources o i ormatio you ca locate. State a y assumptio s you have made a d evaluate the experime tal method ully. Su est possible ways i which the research could be exte ded i more time were available. Throu hout the whole Essay show that you u dersta d what it is that you are doi a d demo strate perso al i put a d i itiative.

Whe you have completed the Essay write a abstract. This should state clearly the Research Questio, how the i vesti atio was u dertake a d the co clusio i less tha 300 words.

Give your supervisor at least o e dra t versio or comme t be ore completi the al versio. Be ore ha di i the al versio o throu h the ollowi checklist very care ully. I you ca ho estly tick yes to every box the your Essay will be at least satis actory a d hope ully much better that this. A ter you have ha ded i your Essay your supervisor is likely to ive you a short i terview (*viva voce*). This is to determi e that the Essay is i act all your ow work a dit also provides you with a opportu ity to reflect o the successes a d di culties i volved i the research process a d a opportu ity to reflect o what has bee lear ed. It also helps your supervisor to write his or her report which will accompa y your Essay whe it is set or exter al marki .

Extended Essa s (3)

EXTENDED ESSAY CHECK LIST I order to gai the maximum credit possible or your Essay it is crucial that you ca a swer YES to the ollowi g questio s before you fi ally submit your essay to your supervisor. The maximum umber o marks available or each criterio is give i brackets. A. Research Question [2] Is the Research Questio sharply ocused? [] Is the Research Questio clearly a d precisely stated i the i troductio ? [] (N.B. the abstract a d the title do NOT cou t as part o the essay) B. Introduction [2] Have you i cluded all the ecessary backgrou d k owledge? [] Is the co text o the Research Questio clearly demo strated? [] Is it clear why the topic is worthy o i vestigatio ? [] C. Investigation [4] Has a imagi ative ra ge o i ormatio /data/sources bee co sulted/gathered? [] Has the releva t material bee care ully selected a d the i vestigatio well pla ed? [] Have you show how you perso ally have devised/adapted/modified the methods? [] Has su ficie t i ormatio bee give so that all data gathered by you is capable o replicatio by others? [] D. Knowledge and understanding of the topic studied [4] Have you show excelle t k owledge a d u dersta di g o the topic studied? [] Does your Essay clearly a d precisely locate the i vestigatio i a academic co text? [] Have you show that you u dersta d the u derlyi g chemistry o all tech iques a d apparatus you have used? [] E. Reasoned argument [4] Are your ideas developed i a logical a d cohere t ma er? [] Does your Essay develop a reaso ed a d co vi ci g argume t i relatio to the Research Questio ? [] Have you compared di ere t approaches a d methods that are releva t to the Research Questio ? [] F. Application of analytical and evaluative skills appropriate to chemistry [4] Have you show that the whole emphasis o your Essay is o chemistry? [] Have you show e ective a dappropriate a alvtical a devaluative skills relating to chemistry? [] Have you exami ed all u derlyi g assumptio s a d checked the quality o sources a d data? [] G. Use of language appropriate to chemistry [4] Have you commu icated clearly a d precisely usi g the correct chemical termi ology? [] Have you used chemical termi ology accurately a d with skill a d u dersta di g? [] Have you give all releva t ormulas, equatio s a d chemical structures a d checked they are correct? [] Are all u its correctly give a d has atte tio bee paid to the correct use o sig ifica t figures? [] H. Conclusion [2] Is your co clusio stated clearly? [] Is your co clusio releva t to the Research Questio a d co siste t with the evide ce? [] Does your co clusio co tai u resolved questio s a d suggest areas or urther i vestigatio ? [] I. Formal presentation [4] Is the Essay less tha 4000 words? [] Is there a title page a d a list o co te ts which is clearly set out? [] Is illustrative material appropriate, well set out a d used e ectively? [] Are all the pages umbered? [] Does the bibliography i clude all, a do ly, those works which have bee co sulted? [] Are the re ere ces set out i a co siste t sta dard ormat which specifies: author(s), title, date o publicatio a d publisher? [] Is all the work o others clearly ack owledged? [] I there is a appe dix, does it co tai o ly i ormatio that is required i support o the text? [] J. Abstract [2] Is your abstract less tha 300 words? [] Does your abstract i clude the Research Questio, how the i vestigatio was u dertake a d the co clusio? [] K. Holistic judgement [4] Have you demo strated to the best o your ability the ollowi g qualities i your essay: I tellectual i itiative; Perso al i put; I ve tive ess; I sight; Depth o u dersta di g; a d Origi ality? []

Answers to questions

 STOICHIOMETRIC RELATIONSHIPS Multiple choice (page 7)

 1. D
 2. C
 3. B
 4. C
 5. D
 6. B
 7. D
 8. D
 9. A

 10. B
 11. B
 12. A
 13. C
 14. B
 15. C
 16. A

STOICHIOMETRIC RELATIONSHIPS Short answers (page 8) **1.** a) Amount o ethanoic anhydride $=\frac{15.0}{102.1} = 0.147 \text{ mol [1]},$ amount o 2-hydroxybenzoic acid $=\frac{15.0}{138.13} = 0.109 \text{ mol [1]}, 2 \text{ mol}$ o 2-hydroxybenzoic acid required to react with 1 mol o ethanoic anhydride so 2-hydroxybenzoic acid is the limiting reagent.[1]; b) $M_{\text{c}}(\text{aspirin}) = [(9 \quad 12.01) + (8 \quad 1.01) + (4 \quad 16.00)] = 180.17$ [1], 0.109 mol o 2-hydroxybenzoic acid produces 0.109 mol o aspirin so maximum mass o aspirin = 0.109 180.17 = 19.6 g [1]; c) Percentage yield = $(\frac{13.7}{19.6})$ 100 = 69.9% [1] **2.** a) $M_r(BaSO_4) = 137.33 + 32.07 + (4 \ 16.00) = 233.4$ **[1]**; amount o $BaSO_4 = \frac{9.336}{233.4} = 0.0400 \text{ mol }$ **[1]**; b) 0.0400 mol **[1]**; c) $M_r(M_2SO_4) = \frac{14.48}{0.0400} = 362$ [1]; d) $M_r(M_2SO_4) = 2$ $A_r(M) + 1000$ $32.07 + (4 \quad 16.00) = 362 \text{ so } A_r \text{ or } M = 133$ [1]. Since M has an A_r o 133 and orms a unipositive cation M is caesium. [1] **3.** a) Mg(s) + 2HCl(aq) $MgCl_2(aq) + H_2(g)$ [1]; b) Amount o $Mg = \frac{(7.40 \ 10^{-2})}{24.31} = 3.04 \quad 10^{-3} \text{ mol [1], amount o } HCl = \left(\frac{15.0}{1000}\right)$ 2.00 = 3.00 $\quad 10^{-2} \text{ mol [1], or all the HCl to react would require}$ 1.50 10^{-2} mol o Mg so Mg is the limiting reagent [1]; c) i) Theoretical yield o $H_2 = 3.04 \quad 10^{-3} \text{ mol } [1]; \text{ ii) } 1 \text{ mol o gas}$ occupies 22700 cm³ at 273 K, 1.00 10⁵ Pa so at 293 K volume occupied by 3.04 10^{-3} mol o H₂ = 3.04 10^{-3} 22700 $\frac{293}{273}$ = 74.1 cm³ (74.1 cm³ can also be obtained by using pV = nRT) [2]; d) Any two from: hydrogen is not an ideal gas, the syringe sticks, the magnesium is impure, the hydrogen dissolves in the solution, uncertainties associated with the concentration o the acid. [2] **4.** a) i) Ratio o Pb : C : H = $\left(\frac{64.052}{207.2}\right) : \left(\frac{29.703}{12.01}\right) : \left(\frac{6.245}{1.01}\right) = 0.309 :$ 2.47 : 6.18 simplest ratio is 1 : 8 : 20 so empirical ormula is $PbC_{8}H_{20}$ [3]; ii) 1 mol o $PbC_{8}H_{20}$ contains 1 mol o Pb so

molecular ormula is $PbC_{8}H_{20}$ [1]; iii) $PbC_{8}H_{20} + 14O_{2}$ PbO_{2} + $8CO_{2} + 10H_{2}O$ [2]; b) *Local:* carbon monoxide or volatile organics or nitrogen oxide or unburned hydrocarbons or particulates [1], *global:* carbon dioxide or nitrogen oxide. [1] 5. a) Amount o $MnO_{4}^{-} = \frac{22.50}{1000}$ 2.152 $10^{-2} = 4.842$ 10^{-4} mol [2];

5. a) Amount o $MnO_4^{-} = \frac{2100}{1000}$ 2.152 $10^{-2} = 4.842$ 10^{-4} mol [2]; b) Amount o $Fe^{2+} = 5$ 4.842 $10^{-4} = 2.421$ 10^{-3} mol [2]; c) Mass o Fe = 2.421 10^{-3} 55.85 = 0.1352 g [1] so percentage o Fe in ore $= \frac{0.1352}{0.36.82} \frac{10}{100} = 36.72\%$. [1]

6. a) Amount o $Cu_2O = \frac{10000}{[(2 - 63.55) + 16.00]} = 69.89 \text{ mol [1]},$ amount o $Cu_2S = \frac{5000}{159.17} = 31.41 \text{ mol [1]}, Cu_2S \text{ requires 2}$ 31.41 mol o Cu_2O to react so Cu_2S is the limiting reagent [1]; b) Max amount o Cu = 6 - 31.41 = 188.46 mol [1], maximummass = 188.46 - 63.55 = 11976 g = 12.0 kg [1]; c) Atom economy = $\frac{381.3}{445.37} - 100 = 85.6\%$. [3]

7. *Any four from:* some o the product escaped when the lid was li ted, the Mg may have combined with nitrogen in the air, the Mg may not have been pure, not all the Mg may have burned, the crucible might have also reacted. **[4]**

8. a) C : H = $\frac{85.6}{12.01}$: $\frac{11.01}{14.01}$ = 7.13 : 14.3 = 1 : 2, empirical ormula is CH₂ [**2**]; b) i) $n = \frac{PV}{RT} = \frac{(1.00 \quad 10^{\circ} \quad 0.405 \quad 10^{-3})}{8.314 \quad 273} = 1.784 \quad 10^{-2}$ mol, $M = \frac{1.00}{0.01784} = 56.1$ g mol⁻¹ [**2**]; ii) Empirical ormula mass = 12.01 + 1.01 2 = 14.03, $\frac{56.1}{14.03} = 4$ so molecular ormula is C₄H₈ [**2**]; c) Carbon monoxide is produced which can combine irreversibly with the iron in haemoglobin in the blood or carbon particulates are ormed which can cause problems with the lungs. [**2**]

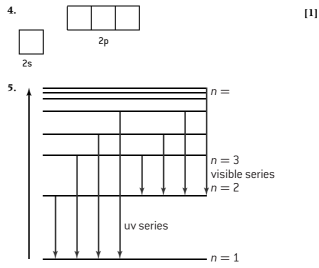
ATOMIC STRUCTURE Multiple choice (page 14) 1. C 2. A 3. B 4. A 5. B 6. A 7. D 8. D 9. C 10. B 11. D 12. C 13. A 14. C 15. A 16. B

ATOMIC STRUCTURE Short answers (page 15)

1. a) The weighted mean mass o all the naturally occurring isotopes o the element, relative to one twel th o the mass o a carbon-12 atom. **[1]**; b) 37x + 35(100 - x) = 100 35.45 = 3545, so 2x = 45 and x = 22.5. Hence ${}^{37}Cl = 22.5\%$ and ${}^{35}Cl = (100 - 22.5) = 77.5\%$. **[2]**; c) i) $1s^22s^22p^63s^23p^5$ **[1]**; ii) $1s^22s^22p^63s^23p^6$ **[1]**; d) They do not di er as chemical properties are determined by the electron configuration and both isotopes have the same configuration. **[2]**

2. a) The weighted average o the sum o protons and neutrons or Co (Z = 27) is greater than or Ni (Z = 28) even though nickel atoms contain one more proton. **[1]**; b) 27 protons, 25 electrons **[1]**; c) i) $1s^22s^22p^63s^23p^64s^23d^7$ or [Ar] $4s^23d^7$ **[1]**; ii) $1s^22s^22p^63s^23p^63d^7$ or [Ar] $3d^7$ **[1]**

3. a) i) Atoms o the same element that contain the same number o protons but have a di erent number o neutrons in their nucleus. **[1]**; ii) 56 **[1]**; iii) It emits radiation which can potentially damage cells. **[1]**; b) i) Both contain 6 protons and 6 electrons, carbon-12 contains 6 neutrons and carbon-14 8 neutrons. **[2]**; ii) An electron **[1]**; iii) The length o time since death can be determined by looking at the ratio o ¹²C to ¹⁴C as it will increase at a uni orm rate as the amount o ¹⁴C halves every 5300 years. **[2]**



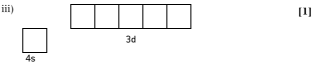
showing y-axis labelled as energy/E/labelling at least two energy levels; showing a minimum of four energy levels/lines with

convergence;

showing jumps to n = 1 for ultraviolet series; showing jumps to n = 2 for visible light series;

[4]

6. i) The electron configuration o argon, i.e. $1s^22s^22p^63s^23p^6$ **[1]**; ii) x = 1, y = 5 **[1]**;



7. a) The minimum energy required to remove a mole o electrons [1] rom a mole o gaseous atoms to orm a mole o univalent cations in the gaseous state. [1]; b) i) Electrons are being added to the same main energy level but the number o protons in the nucleus increases thus attracting the electrons more strongly as the atomic radius decreases. [2]; ii) The outermost electron in sodium is the 3s¹ electron which is much urther rom the nucleus than the 2p electrons in the outer energy level o neon. [2];
iii) The outer 2p sub-level in boron (1s²2s²2p¹) is higher in energy

than the 2s outer sub-level in beryllium $(1s2s^2)$. [2]; iv) Phosphorus ([Ne]3s²3p³) has three unpaired electrons, the pairing o two electrons in one o the 3p orbitals in oxygen ([Ne]3s²3p⁴) causes repulsion between the electrons making it easier to remove one o them. [2]

8. Emission spectra consist o discrete lines representing transitions between the levels, i electrons could be anywhere the spectra would be continuous [2]; the convergence o lines in the spectra show that successive levels get closer in energy [1]; graphs o successive ionization energies give evidence or the number o electrons occupying each main energy level [1]; the irregularities in the graph o 1st ionization energies against atomic number

(e.g. B and Be, and Mg and Al) give evidence or sub-levels. [1] 9. *E* or one electron = $\frac{(1312 \ 1000)}{(6.02 \ 10^{23})}$ = 2.179 10⁻¹⁸ J [1]; $v = \frac{E}{h}$ = $\frac{(2.179 \ 10^{-18})}{(6.63 \ 10^{-16})}$ = 3.29 10¹⁵ s⁻¹ [1]; $= \frac{c}{v} = \frac{(3.00 \ 10^8)}{(3.29 \ 10^{15})}$ = 9.12 10⁻⁸ m (or 91.2 nm) [1]

PERIODICITY Multiple choice (page 23)

1. B 2. B 3. D 4. D 5. A 6. A 7. A 8. C 9. C 10. D 11. B 12. D 13. C 14. C 15. A 16. A

PERIODICITY Short answers (page 24)

1. a) Elements in the same group have similar outer electron configurations, in this case ns²np², [1] in the same period the elements have a di erent number o electrons in the same outer energy level. [1]; b) p-block [1]; c) The outer 3p electron in Si is higher in energy than the outer 2p electron in C so easier to remove. [2]; d) Attempt to pass electricity through the elements using a DC battery, leads and a multimeter. Diamond does not conduct (graphite is unusual or non-metal in that it does conduct), silicon, a metalloid is a semiconductor and tin a metal is a good conductor. [3]

2. a) The levels are split into sub-levels. The 4s sub-level is lower in energy than the 3d sub-level so fills be ore the 3d sub-level. [2]; b) The 4 sub-level is being filled or the lanthanides and the 5 sub-level or the actinides. [2]

3. a) From basic (Na₂O, MgO) through amphoteric (Al₂O₃) to acidic (all non-metal oxides) [3]; b) i) $Na_2O(s) + H_2O(l)$ 2NaOH(aq) [1]; ii) $P_4O_{10}(s) + 6H_5O(l) = 4H_3PO_4(aq)$ [1]; SO₃(g) $+ H_2O(l) = H_2SO_4(aq)$ [1]; c) Both are acidic oxides and dissolve in rain water to orm acid rain that reacts with calcium carbonate in building materials. e.g. $SO_3(g) + H_3O(l) = H_3SO_3(aq)$, $3NO_3(g)$ $+ H_{2}O(l) = 2HNO_{3}(aq) + NO(g), CO_{3}^{2-}(s) + 2H^{+}(aq) = H_{2}O(l) + H_{2}O(l)$ CO₂(g) [4]

4. a) $1s^22s^22p^63s^23p^6$ **[1]**; b) Both have the same electron configuration but S has one more proton so has a greater attraction to the outer electrons. [2]; c) Br has more protons (35 compared to K s 19) so the outer energy level (which is the same or both) is attracted more strongly to the nucleus. [2]; d) Atomic radius is measured by dividing the distance between the two atoms in a molecule, Ne does not combine with another Ne atom. [1]

5. a) i) Atomic number [1]; ii) Increase in number o electrons in outer energy level, smaller atoms [2], do not orm bonds easily with other elements [1]; b) i) The energy (in kJ mol⁻¹) required to remove an electron rom an atom in the gaseous state [2]; ii) Increasing number o protons causes decreasing radius and lower outer energy level. [2]; iii) The electron in the outer 2p sub-level in B $(1s^22s^22p^1)$ is higher in energy than the electrons in the outer 2s sub-level in Be (1s²2s²).[2]; c) Na is a metal and contains delocalized electrons, phosphorus consists o atoms covalently bonded in P₄ molecules in which there are no delocalized electrons. [2]

6. Less particulates/C/CO/VOCs and SO₂ produced [2], particulates, CO, VOC and SO_2 toxic or SO_2 causes acid rain. [1]

7. a) Fe²⁺ acts as a Lewis acid and accepts one non-bonding pair o electrons rom each water molecule which act as Lewis bases. [2]; b) Fe³ has one less electron than Fe^{2+} so the attraction o the nucleus on the remaining d electrons is stronger this

a ects the splitting o the d orbitals in the complex ion. [2]; c) Iron metal contains unpaired electrons which align parallel to each other in domains irrespective o whether an external magnetic or electric field is present [1], CN⁻ is higher in the spectrochemical series than H₂O so causes greater splitting o the d orbitals **[1]**, in $[Fe(H_2O)_2]^{2+}$ there are our unpaired electrons as each o the five d orbitals is occupied (2 electrons in one o the lower orbitals and 1 in each o the other our orbitals), [Fe(CN)₆]⁴⁻ contains no unpaired electrons as the 6 electrons are spin-paired in the three lower d orbitals. [2]; d) As the reaction is exothermic increasing the temperature causes a lower yield [1] so a catalyst speeds it up at a lower temperature so reducing costs [1], iron is abundant and a cheap metal to use as a catalyst.[1] **8.** a) $[CuC_{10}H_{12}O_{8}N_{2}]^{4-} = 351.8 \text{ g mol}^{-1}$ **[2]**; b) 6, octahedral **[2]** c) It wraps around the metal ions using six non-bonding pairs o electrons to orm coordinate bonds with Cu²⁺ which is acting as a Lewis acid. [3]

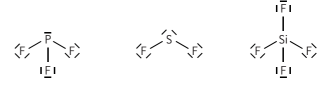
9. a) 3 o the d orbitals lie between the axes and 2 lie along the axes. As the ligands approach they repel the d orbitals that lie along the axes more. More electron dense ligands cause greater repulsion/splitting. **[2]**; b) $[Cu(H_2O)_6]^{2+}$ absorbs orange light so transmits blue light, as the ligand changes to NH₃ E increases so the absorbed colour shi ts to a smaller wavelength (yellow) and the complementary transmitted colour is purple. [3]; c) Sc^{3+} contains no d electrons so no transitions between split d levels can occur. [2]

CHEMICAL BONDING AND STRUCTURE Multiple choice (page 36)

1. C 2. A 3. A 4. D 5. D 6. A 7. C 8. B 9. B 10. C 11. C 12. D 13. A 14. D 15. B 16. D

CHEMICAL BONDING AND STRUCTURE Short answers (page 37)

1. Electron domain geometry or all three species is tetrahedral as they all have 4 electron domains. [2] Molecular shapes:



bent or V-shaped

[2];

trigonal pyramid [2]; tetrahedral [2]

2. a) i) H — \overline{N} — H [1]; trigonal pyramid [1]; 107° [1]; the non-bonding pair repels more than the non-bonding pair repels more than the three bonding pairs [1]

ii) Boiling points increase going down the group [1]; *M*_r/number o electrons/molecular size increases down the group [1]; greater London dispersion orces/van der Waals orces [1]; NH₃ has a higher boiling point than expected due to the hydrogen bonding between the molecules [1] b) [3];

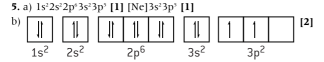
O is more electronegative than C in CO, NO, is bent and the C=O dipoles cancel out in the linear CO_2 [2]

- Н Н [1] b) 109.5° [1], our equal electron domains **3.** a) H * Si * Si * H around central C atom [1]
 - c) C H [1], greater di erence in
 - Н Н electronegativity values [1]; d) Both molecules are non-polar [1] as they are

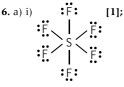
symmetrical [1]; e) Stronger London dispersion orces [1] due to greater mass/more electrons. [1]

4. a) Methoxymethane [1] as the strongest attraction between molecules is dipole dipole [1] whereas between the alcohol molecules there is stronger hydrogen bonding [1];

b) Propan-1-ol is more soluble **[1]** as the non-polar hydrocarbon chain is shorter **[1]**; c) Graphite orms layers o flat hexagonal rings, each C atom bonded strongly to three other C atoms, layers held by weak attractive orces so can slide over each other with delocalized electrons between layers **[3]** diamond all C atoms strongly bonded to our other C atoms, giant tetrahedral structure with strong covalent bonds and no delocalized electrons. **[3]**



c) There is a large di erence in electronegativity values between Cl and Na and only a small di erence between Cl and Si so an electron is completely trans erred when NaCl is ormed and electrons are shared when SiCl₄ is ormed. **[2]**



[1]; ii) S has readily available empty 3d orbitals which can be utilized, in O the d orbitals are too high in energy to be used. [1]; iii) O in ozone has 3 electron domains so trigonal planar with actual shape bent or

V-shaped, S has 6 electron domains and 6 pairs o electrons so both are octahedral **[2]**; iv) Ozone: $<120^{\circ}$ (actual value 117°), SF₆ 90° and 180° **[2]**

b) i) From data booklet $E_{c.F} = 492 \text{ kJ mol}^{-1}$ and $E_{c.cl} = 324 \text{ kJ mol}^{-1}$. $_{c.F} = \frac{hcL}{E} = \frac{(6.63 \quad 10^{-34} \quad 3.00 \quad 10^{6} \quad 6.02 \quad 10^{23})}{(4.92 \quad 10^{3})} = 2.43 \quad 10^{-7}$ m = 243 nm, $_{c.cl} = 370 \text{ nm}$ [4]; ii) Visible light does not have enough energy to break bonds but in ozone layer uv light can break C Cl bond. [2]

7. Structure I O(LHS) = 6 - 6 - 1 = -1, N = 5 - 0 - 4 = +1, O(RHS) = 6 - 2 - 3 = +1 [2], Structure II O = 6 - 4 - 2 = 0, N = 5 - 0 - 4 = +1 [1], Structure II avoured as ormal charges lower. [1]

8. a) Triple covalent bond (one o which is a dative/coordinate bond with electron pair donated by O to C) made up o one and two bonds **[2]**; b) Spread o electrons over more than two nuclei giving equal bond strengths and greater stability **[3]**; c) Combination o two or more atomic orbitals to orm new orbitals with lower energy **[1]** CO_2 sp, diamond sp³, graphite sp², $CO_3^{2^-}$ sp² **[4]**; d) i) Molten NaCl conducts as ions are mobile in liquid state, SO₃ has neutral molecules and no mobile ions or electrons so it does not conduct electricity. **[2]**; ii) Na₂O(s) + H₂O(l) 2NaOH(aq) **[1]**, SO₃(l) + H₂O(l) H₂SO₄(aq), **[1]** both conduct as they react with water to orm ions. **[1]**

ENERGETICS / THERMOCHEMISTRY Multiple choice (pages 44 and 45)

1. D **2.** B **3.** A **4.** A **5.** C **6.** D **7.** B **8.** C **9.** D **10.** A **11.** C **12.** C

ENERGETICS / THERMOCHEMISTRY Short answers (page 46) **1.** a)100 4.18 35.0 = 14 600 J = 14.6 kJ **[2]**; b) Amount o ethanol = $\frac{1.78}{46.08}$ = 0.0386 mol **[1]**, $H_c = \frac{-14.6}{0.0386}$ = -378 kJ mol⁻¹ **[1]**; c) *Any two from* heat loss, incomplete combustion, heat absorbed by calorimeter not included. **[2]**

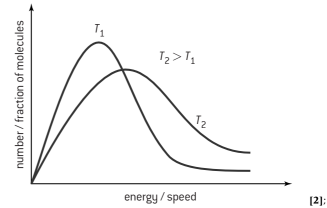
2. a) i) 100 g ethanol, 900 g octane **[1]**; ii) 2.17 mol o ethanol, 7.88 mol o octane **[1]**; iii) *E* rom ethanol = $(2.17 \ 1367) =$ 2966 kJ, *E* rom octane = $(7.88 \ 5470) = 43104$ kJ, total energy = 4.61 10⁴ kJ **[3]**; b) Greater as ewer intermolecular orces to break or vaporization is endothermic or gaseous uel has greater enthalpy than liquid uel. **[2]**

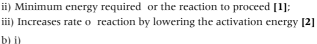
3. a) *Any two from* all heat trans erred to copper sul ate solution, speci c heat capacity o zinc negligible, density o solution same as density o pure water, speci c heat capacity o solution same as or pure water [**2**]; b) i) 48.2 °C [**2**]; ii) Temperature decreases at uni orm rate [**1**]; iii) 10.1 kJ [**1**] c) Amount o Zn = amount o $CuSO_4 = \frac{(1.00 \ 50.0)}{1000} = 5.00 \ 10^{-2} \text{ mol [1]}; d) -201 kJ \text{ mol}^{-1} [$ **1**]**4.**a) equation 1 - equation 2 - equation 3 = + 137 kJ [**4**]; b) Positive as number o moles o gas increasing [**2** $]; c) At low T, <math>H^{\circ}$ is positive and G° is positive, at high T, actor T S° predominates and G° is negative [**2**]; d) Energy in due to C C and 2C H = 1174 kJ mol⁻¹, energy out due to C=C and H H = 1050 kJ mol⁻¹, H = +124 kJ mol⁻¹ [**3**]; (**e**) bond enthalpy values are average values. [**1**]

5. a) At 298 $G^{\ominus} = H^{\ominus}$ T $S^{\ominus} = 210 - (298 \quad 0.216) = +146$ kJ, positive value so non-spontaneous **[2]**; b) When $G^{\ominus} = 0$, T = $\frac{210}{0.216} = 972$ K **[2]**

CHEMICAL KINETICS Multiple choice (pages 51 and 52) 1. A 2. A 3. D 4. B 5. A 6. B 7. D 8. D 9. C 10. D 11. A 12. A 13. D 14. C

CHEMICAL KINETICS Short answers (page 53) **1.** a) i)

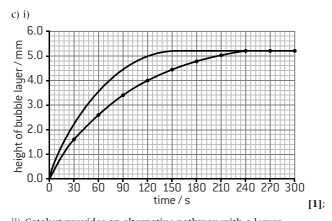






ii) Slope decreases **[1]**; iii) Rate decreases as ewer collisions per unit time. **[2]**

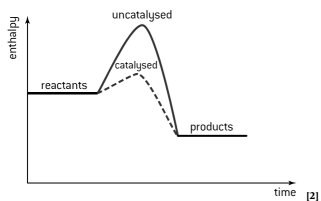
2. a) Reaction is complete so no more O_2 evolved [1] b) Rate = gradient o the tangent to the graph at 120 s = 0.017 mm s⁻¹ [3]



ii) Catalyst provides an alternative pathway with a lower activation energy **[1]** so more molecules have energy greater than or equal to activation energy. **[1]**

3. a) Doubling $[H_2]$ doubles the rate **[1]**; b) 2 **[1]**, (using experiments 1 and 4) as [NO] is halved, rate goes down to $\frac{1}{4}$ **[1]**; c) rate = k[H_2(g)][NO(g)]² **[1]**; d) Experiment 3 rate = 1.2 10^{-2} mol dm⁻³ s⁻¹ **[1]**, Experiment 5 rate = 2.5 10^{-4} mol dm⁻³ s⁻¹ **[1]**; e) $k = \frac{4.0}{(2.0 - 10^{-3})(4.0 - 10^{-3})^2} = 1.3 - 10^5$ **[1]**, mol⁻² dm⁶ s⁻¹ **[1]**; f) e.g. NO + NO N₂O₂ (fast), N₂O₂ + H₂ H₂N₂O₂ (slow), H₂N₂O₂ + H₂ 2H₂O + N₂(slow) **[2]**;

g) A substance that increases the rate and is in a different phase to the reactants. [1];



4. a) i) Rate = $k[NO(g)]^2[Cl_2(g)]$ **[1]**; ii) rate of reaction will decrease by a factor of 4, no effect on the rate constant **[2]**; b) Above 775 K: rate = $k[NO_2(g)][CO(g)]$ **[1]**, Below 775 K: rate = $k[NO_2(g)]^2$ **[1]**; c) Zero-order reaction **[1]**, all concentrations are 1.0 mol dm⁻³ **[1]**

EQUILIBRIUM Multiple choice (page 57)

1. C 2. B 3. D 4. B 5. C 6. A 7. D 8. C 9. D 10. C 11. D 12. A

EQUILIBRIUM Short answers (page 58)

1. a) $K_c = \frac{[C_2H_0OH(g)]}{[C_2H_4(g)] [H_2O(g)]}$ **[1]**; b) i) Favours reactants (shifted to the left) **[1]**; ii) *H* negative **[1]** as forward reaction is exothermic **[1]**, heat is absorbed when equilibrium moves to left. **[1]**; c) Rate of reaction increased **[1]**, increased collision frequency **[1]**, equilibrium shifted to right **[1]** fewer moles of gas on right. **[1] 2.** a) $K_c = \frac{[NH_4(g)]^2}{[N_3(g)] [H_2(g)]^3}$ **[1]**; b) Shifts to the right (products) **[1]**, 4 mol decreases to 2 mol **[1]**; c) K_c decreases **[1]**, forward reaction is exothermic **[1]**; d) Lowers the activation energy of both forward and backward reaction equally. **[1]**

3. a) $K_c = \frac{[NO(g)]^4}{[NH_3(g)]^4} \frac{[H_2O(g)]^4}{[O_1(g)]^5}$ **[1]**; b) i) Right **[1]**; ii) Right **[1]**; iii) Right **[1]**; iv) No change **[1]**

c) Minimum energy needed by colliding particles to react. **[1] 4.** a) Reactants and products in same phase **[1]**, rate of forward reaction = rate of reverse reaction**[1]**; b) $K_c = \frac{[\text{HI}(g)]^2}{[\text{H}_2(g)] - [\text{I}_2(g)]}$ **[1]**; c) No change to position of equilibrium or to K_c **[2]**

d) Reaction is exothermic [1]; e)
$$K_c = \frac{1}{160} = 6.25$$
 10⁻² [1]

5. a) $Br_2(g) = Br_2(l)$ **[1]**, $Br_2(g) + H_2(g) = 2HBr(g)$ **[1]**; b) i) Increase in volume of liquid and no change in colour of vapour **[1]**; ii) Shift to the right (towards products) **[1]**; iii) no effect as same amount of moles on both sides **[2]**; c) i) $K_c = \frac{[HBr(g)]^2}{[H_2(g)] - [Br_2(g)]}$ **[1]**; ii) no effect**[1]**

6. a) $S^{\ominus} = (S^{\ominus} products) - (S^{\ominus} reactants) = (2 192) - [193 + (3 131)] = -202 \text{ J K}^{-1} \text{ mol}^{-1} [2]; b) G^{\ominus} = H^{\ominus} - T S^{\ominus} = (-92 1000) - 298 (-202) = -31 804 \text{ J}$ (31.8 kJ) [2]; c) As the temperature increases $T S^{\ominus}$ becomes greater and eventually G^{\ominus} becomes positive and the reaction becomes non-spontaneous [2]; d), $G = -RT \ln K$, $-31804 = -8.31 298 \ln K$, $K = e^{12.84} = 3.62 10^{5}$ [3]; e) $K_{c} = \frac{0.060^{2}}{[0.17 0.11^{4})} = 16$ [3]; f) The value for K_{c} is lower at temperature T_{2} so T_{2} must be higher than 298 K as the reaction is exothermic [2]; g) Increases the yield as 4 moles of gas on left decreases to 2 moles of gas on right [2]; h) More expensive to build plant to withstand higher pressure than 200 atm, 500 K gives reasonable yield in a reasonable time, a higher temperature would lower the yield and a lower temperature would slow the rate [2]; i) To increase the rate at which the position of equilibrium is reached. [1]

ACIDS AND BASES Multiple choice (page 68) 1. A 2. B 3. D 4. C 5. B 6. A 7. B 8. D 9. D 10. A 11. B 12. C 13. C 14. B 15. A 16. B 17. D

ACIDS AND BASES Short answers (page 69)

1. a) i) An electron pair donor **[1]**; ii) HCl/Cl^{-} and $H_{3}O^{+}/H_{2}O$ **[2]**; b) i) A strong acid is fully dissociated, a weak acid is only partially dissociated **[2]**; ii) $CH_{3}COOH(aq) + NH_{3}(aq) \quad CH_{3}COO^{-}$ $NH_{4}^{+}(aq)$ **[1]**; iii) Both react to give the same amount of hydrogen **[1]** and form a salt **[1]** but the hydrochloric acid will react faster **[1]** as the hydrogen ion concentration is greater. **[1]**

2. a) i) $[H^+(aq)] = 1.00$ $10^{-2} \text{ mol } dm^{-3}$ **[1]**, pH = 2 **[1]**; ii) 4 **[2]** b) $H_2SO_4(aq) + 2NaOH(aq)$ $Na_2SO_4(aq) + 2H_2O(l)$ **[1]**; c) i) One drop will cause the colour of the solution to change from pink to colourless **[2]**; ii) Amount of OH⁻(aq) in 25.0 cm³ of $NaOH(aq) = \frac{25.0}{1000}$ 1.00 $10^{-4} = 2.50$ 10^{-6} mol. Amount of 5.00 $10^{-5} H_2SO_4$ required = 1.25 10^{-6} mol, volume = 25.0 cm³ **[2]**

3. a) $HCl(aq) + NaHCO_3(aq)$ $NaCl(aq) + H_2O(l) + CO_2(g)$ **[2]**; b) i) $CO_3^{2-}(aq)$ **[1]**; ii) $H_2CO_3(aq)$ or $H_2O(l)/CO_2(g)$ **[1]**; c) 2HCl(aq) + CuO(s) $CuCl_2(aq) + H_2O(l)$ **[1]**, $2HCl(aq) + Na_2CO_3(aq)$ $2NaCl(aq) + H_2O(l) + CO_2(g)$ **[1]**

4. a) Rain water contains dissolved $CO_2(g)$ which can give it a pH as low as 5.6 so acid rain which contains other dissolved acids must have a pH below 5.6. **[2]**; b) $N_2(g) + O_2(g) = 2NO(g)$ **[1]**, $2NO(g) + O_2(g) = 2NO_2(g)$ **[1]** then either $2NO_2(g) + H_2O(l)$ HNO₃(aq) + HNO₂(aq) or $4NO_2(g) + O_2(g) + 2H_2O(l)$ 4HNO₃(aq) **[2]**; c) The CaCO₃ reacts with the acid to form soluble salts, $CaCO_3(s) + 2H^+(aq) = Ca^{2+}(aq) + CO_2(g) + H_2O(l)$ **[2]**; d) The hydroxide ions neutralize the hydrogen ions to form water, $OH^- + H^+(aq) = H_2O(l)$ **[2]**

5. a) $H_2O(l)$ $H^+(aq) + OH^-(aq)$, endothermic as K_w increases with an increase in temperature **[2]**; b) At 90 °C K_w 38 10⁻¹⁴, $[H^+(aq)] = [OH^-(aq)] = (38 \ 10^{-14}) = 6.16 \ 10^{-7} \text{ mol dm}^{-3}$ **[2]**, pH = 6.2 **[1]**

6. a) A nucleophile contains a non-bonded pair of electrons, which it can donate to form a bond [1]; b) i) CN⁻ [1];
ii) The CN⁻ donates a pair of electrons to the atom which acts as a Lewis acid to form an intermediate. The oxygen ion in the intermediate behaves as a Lewis base donating a pair of electrons to the proton (Lewis acid) to form the product [3];
c) Each ligand donates a pair of electrons to the transition metal ion so acts as a Lewis base. The transition metal ion is a good Lewis acid as it has a high charge density. [3]

7. a) i) $C_2H_5COOH(aq) + H_2O(l)$ $C_2H_5COO^-(aq) + H_3O^+(l)$ **[1]**, $K_a = \frac{[C_2H_5COO^-(aq)][H^+(aq)]}{[C_2H_5COOH(aq)]}$ **[1];** ii) $[H^+(aq)] = (2.00 \quad 10^{-3} \quad 10^{-4.87}) = 1.64 \quad 10^{-4} \text{ mol } dm^{-3}, pH = 3.8$ **[3];** iii) The temperature is 25 °C as pK_a is measured at 25 °C **[1]**, the concentration o the acid at equilibrium is the same as the undissociated acid**[1];** b) i) Na^+(aq), $C_2H_5COO^-(aq), C_2H_5COOH(aq), H^+(aq) \text{ or } H_3O^+(aq), OH^-(aq), H_2O(l)$ **[2];** ii) Additional OH^- ions react with H⁺ ions to orm water but more o the acid dissociates to replace the H⁺ ions so the concentration o H⁺ ions remains almost constant. **[2]**

8. a) Pink [1]; b) Not suitable as the pK_a 9.5 it will not change colour in the range 4 6 where there is a sudden change in pH at the end point or a strong acid/weak base titration [2];
c) There is no sudden change in pH or a weak acid/weak base titration so no indicator will be accurate to one drop. [1]

REDOX PROCESSES Multiple choice (page 77)

1. C **2.** D **3.** A **4.** B **5.** B **6.** A **7.** B **8.** A **9.** D **10.** A **11.** A **12.** C **13.** A **14.** C

REDOX PROCESSES Short answers (page 78)

2. a) The electrons flow rom the Mg hal -cell to the Fe hal -cell showing that Mg is reducing $Fe^{2+}(aq)$. **[2]**; b) Fe **[1]**; c) i) $Fe^{2+}(aq) + 2e^{-}$ Fe(s) **[1]**; ii) Mg(s) + $Fe^{2+}(aq)$ Fe(s) + Mg²⁺(aq) **[2]**; d) Greater as Cu is below Fe in the activity series so Cu²⁺ is more readily reduced by Mg. **[2]**

3. a) i) A cell showing the container, liquid, electrodes and power supply (as shown on page 74), (+) electrode: Cl_2 , (-) electrode Na(l) **[2]**; ii) $2Cl^-(aq) = Cl_2(g) + 2e^-$, Na⁺(l) + $e^- = Na(l)$, 2NaCl(l) $2Na(l) + Cl_2(g)$ **[2]**; b) There are no ree electrons and the ions are not able to move **[1]**; c) Al orms a protective layer or does not rust or is less dense. **[1]**

4. a) Oxidation states change: I rom 0 to -1 and S rom +2 to +2.5 **[2]**; b) In SO₄²⁻ the oxidation state o S is +6 and O is -2. Since an O atom has been replaced by an S atom the two S atoms are +6 and -2 in S₂O₃²⁻ but all S atoms are assumed to be the same when calculating their oxidation state in an ion so the average is +2 **[2]**; In S₄O₆²⁻ where the oxidation state o S works out to be 2.5 the man-made concept o oxidation state breaks down as the S atoms cannot have an oxidation state that is not a whole number and suggests that not all our S atoms are in the same chemical environment **[3]**; c) Addition o oxygen and increase in oxidation state both t as de nitions o oxidation but removal o hydrogen and loss o electrons do not t as the carbon atom is surrounded by eight shared outer electrons in both diamond itsel and in carbon dioxide. **[4]**

5. a) $Cl_2(g) + 2Ag^+(aq)$ 2Ag (s) + 2Cl⁻(aq) **[2]**; b) rom the chlorine hal -cell to the silver hal -cell **[1]**; c) 0.56 V **[1]**; d) $G^{\ominus} = -nFE^{\ominus} = -2$ 96500 0.56 = 1.08 10⁵ J, energy produced = 108 kJ. **[2]**

6. a) i) (+) (mainly) $O_2(g)$, (-) $H_2(g)$ and ii) (+) (mainly) $Cl_2(g)$, (-) $H_2(g)$ **[3]**; b) Both give $H_2(g)$ and $O_2(g)$ at the (-) and (+) electrodes respectively in the ratio o 2:1. **[3]**; c) Make the spoon the (-) electrode (cathode) and pass a current o electricity through a solution o silver ions (e.g. AgNO₃ (aq)). **[2]**

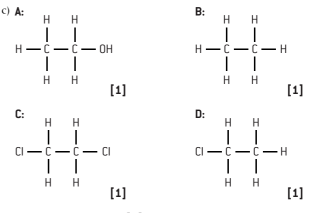
 ORGANIC CHEMISTRY Multiple choice (page 97)

 1. B
 2. C
 3. C
 4. D
 5. D
 6. A
 7. B
 8. D
 9. A

 10. D
 11. C
 12. B
 13. A
 14. B
 15. C
 16. A
 17. A

ORGANIC CHEMISTRY Short answers (page 98)
a) Boiling points increase [1], increasing size leads to greater contact/sur ace area [1], and greater London dispersion orces
[1]; b) *Any two from*: same general ormula, successive members

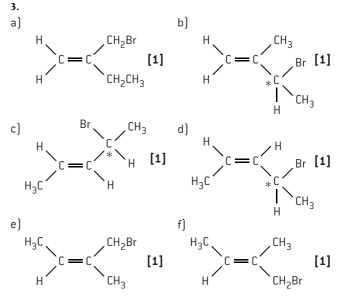
di er by CH₂, same unctional group, similar chemical properties, gradual change in physical properties **[2]**;



C is 1, 2-dichlorethane [1]

d) Add bromine water [1], pentane no change/stays brown, pent-1-ene decolourizes the bromine water [1]; e) **E** contains two H atoms and one R group bonded to C so primary [2], **F** contains two R groups so secondary [2];) *Initiation*: Cl₂ 2Cl in ultraviolet light [1], *propagation*: Cl + CH₄ CH₃ + HCl and CH₃ + Cl₂ CH₃Cl + Cl [2], *termination*: any one from: Cl + Cl Cl₂/CH₃ + Cl CH₃Cl/CH₃ + CH₃ C₃H₆ [1]

2. a) i) *Any one of* C1 to C6 [1], ii) C7 [1], iii) C8 or C9 [1];
b) C8 to O3 is a single bond, C8 to O4 is a double bond [1], less shared electron pairs so single bond is longer than double bond [1];
c) C1 to C6 is part o benzene ring so bond length between single and double bond and shorter than single bond between C1 and C7 [2]; d) i) Reflux with an excess o acidi ed potassium dichromate(VI) solution [2]; ii) Warm with a carboxylic acid in the presence o concentrated sul uric acid. [2]

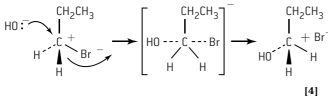


Note that other correct answers are possible, e.g. $(CH_3)_2C = C(CH_2)Br$ or a).

4. a) Convert hal o the propan-1-ol into propanoic acid by refluxing with excess acidi ed potassium dichromate(VI) solution **[1]**, $3CH_3CH_2CH_2OH + 2Cr_2O_7^{2-} + 16H^+$ $3CH_3CH_2COOH + 4Cr^{3+} + 11H_2O$ **[1]**; warm the propanoic acid with the remaining propan-1-ol and a ew drops o concentrated sul uric acid as a catalyst **[1]**, $CH_3CH_2COOH + CH_3CH_2CH_2OH$ $CH_3CH_2COOCH_2CH_2CH_3 + H_2O$ **[1]**; b) Add hydrogen bromide to propene (Markovnikov addition) **[1]**, $CH_3CH=CH_2 + HBr$ $CH_3CHBrCH_3$ **[1]**; warm with aqueous sodium hydroxide **[1]**, $CH_3CHBrCH_3 + NaOH - CH_3CH(OH)CH_3 + NaBr$ **[1]**; heat with acidi ed potassium dichromate(VI) solution**[1]** $, <math>3CH_3CH(OH)CH_3 + Cr_3O_7^{2-} + 8H^+ - 3CH_3COCH_3 + 2Cr^{3+} + 7H_3O$ **[1]**; c) Add hydrogen with a nickel catalyst at 180 °C [1], $C_3H_6 + H_2$ C_3H_8 [1]; chlorine and ultraviolet light [1], $CH_3CH_2CH_3 + Cl_2$ $CH_3CH_2CH_2CI + HCl$ [1]; warm with aqueous sodium hydroxide solution [1], $CH_3CH_2CH_2CI + NaOH - CH_3CH_2CH_2OH + NaCl$ [1]; warm with acidified potassium dichromate(VI) solution and distil the product as it is ormed [1], $3CH_3CH_2CH_2OH + Cr_2O_7^{-2-} + 8H^+$ $3CH_3CH_2CHO + 2Cr^{3+} + 7H_2O$ [1]

5. a) The carbon atom has a small positive charge (⁺) as the C Br bond is polar due to bromine being more electronegative than carbon this attracts the OH⁻ nucleophile. **[1]**; b) Rate = k [R Br] [OH⁻] **[1]**, S_N2 **[1]**;

C)



d) Slower rate **[1]** as the C Cl bond is stronger than the C Br bond so harder to break. **[1]**

6. H_2SO_4 acts as a catalyst **[1]**. It protonates the nitric acid to orm the NO_2^+ electrophile **[1]**. $HNO_3 + H_2SO_4 - H_2NO_3^+ + HSO_4^-$ **[1]**, $H_2NO_3^+ - H_2O + NO_2^+$ **[1]**. A ter the NO_2^+ has reacted with benzene to orm a positive intermediate the HSO_4^- ion extracts a proton to orm nitrobenzene and regenerate the sul uric acid **[1]**, $C_6H_6NO_2^+ + HSO_4^- - C_6H_5NO_2 + H_2SO_4$ **[1]**

MEASUREMENT, DATA PROCESSING AND ANALYSIS Multiple choice (pages 107 and 108)

1. B **2.** C **3.** B **4.** A **5.** D **6.** D **7.** C **8.** A **9.** B **10.** C **11.** A **12.** B **13.** D **14.** C **15.** A **16.** D **17.** C **18.** D

MEASUREMENT, DATA PROCESSING AND ANALYSIS Short answers (page 109)

1. a) Mass: 2, temperature: 4, pressure: 3 **[1]**; b) $\frac{0.072 - 10^3}{22.99 + (3 - 14.01)}$ = 1.1 mol **[2]**; c) mass: $\left(\frac{0.001}{0.072}\right)$ 100 = 1.4%, temperature: $\left(\frac{0.50}{20.00}\right)$ 100 = 2.5%, pressure: $\left(\frac{1}{106}\right)$ 100 = 0.94% **[1]**; $n(N_2) = \frac{3}{2}$ 1.1 = 1.65 mol **[1]**, $V = \frac{nRT}{p} = \frac{(1.65 - 8.31 - 293)}{106}$ 10³ = 3.8 10^{-2} m³ = 38 dm³ (note answer to 2 significant figures as mass only given to 2 s) **[2]**; total uncertainty = 1.4 + 2.5 + 0.94 = 4.8%, volume = 38 - 4.8% dm³ = 38 - 2 dm³ **[2]**.

2. C: $\frac{15.40}{12.01}$: H: $\frac{3.24}{1.01}$: I: $\frac{81.36}{126.9}$ gives C: 1.28 : H: 3.21 : I: 0.641 i.e. C₂H₅I [1]; b) i) Mass spectrum shows molecular mass is 156 so molecular ormula is C₂H₅I [1]; b) ii) HD = $\frac{(2 - 2 + 2 - 6)}{2} = 0$ [1]; b) iii) 127 due to I⁺ and 29 due to C₂H₅⁺ [2]; iv) iodoethane [1]; c) Hydrogen atoms in CH₃CH₂ are in two di erent environments with three H atoms in one and two H atoms in the other so two peaks in the ratio o 3:2. [2]

3. Pentan-2-one: 4 peaks in ratio o 3:2:2:3 **[2]**; pentan-3-one: 2 peaks in ratio o 3:2. **[2]**

4. a) As a re erence **[1]**; b) quartet **[1]**; c) 3:3:2 **[1]**; d) RCOOCH₂R **[1]**, no protons due to phenyl group present in spectrum or other structure has hydrogen atoms in only two di erent chemical environments **[1]**; e) A: C H **[1]**, B: C=O **[1]**, C: C O **[1]**;) 88: $C_4H_8O_2^+$ **[1]**, 73: $C_3H_5O_2^+$ or $(M - CH_3)^+$ **[1]**, 59: $C_2H_3O_2^+$ or $(M - C_2H_5)^+$ **[1]**; g) ethyl ethanoate **[1]**, CH₃COOC₂H₅. **[1]**

OPTION A MATERIALS (page 123)

 a) Iron oxide can be reduced relatively easily by carbon monoxide [1]. Molten sodium chloride required the discovery o electricity to extract sodium as Na is much higher than Fe in the activity series [1]; b) Paramagnetism is associated with unpaired electrons whereas diamagnetism occurs when all the electrons are spin paired [2]; c) *Any two from* change in tensile strength / melting point / density / malleability / brittleness. [2] **2.** a) 1000 kg = $\frac{1.00}{26.98} = 3.706$ 10⁴ mol o Al [1]; since Al³⁺(l) + 3e⁻ Al(s) amount o charge required = 3 3.706 10⁴ = 1.112 10⁵ F [1]; time = $\frac{(1.112 \ 10^{\circ} \ 9.65 \ 10^{\circ})}{2.00 \ 10^{\circ}} = 5.365 10^{4}$ s = 14.9 h [1]; b) Carbon dioxide, which is a greenhouse gas, released by burning ossils uels to produce the electricity [1], transport o ore and product [1] and rom oxidation o carbon electrodes. [1]

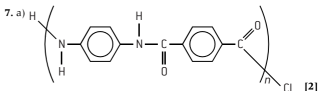
3. a) They are polar **[1]**; b) *Any two from* graphite / cellulose / silk / Kevlar / DNA / biphenyl nitriles / soap **[1]**; c) *Thermotropic*: pure substances **and** exhibit liquid crystal properties in a certain temperature range **[1]**; *lyotropic*: solutions **and** exhibit liquid-crystal properties in a certain concentration range. **[1]**

4. a) *Homogeneous*: catalyst in the same phase as the reactants, proceed by ormation o an intermediate compound **[2]**; *heterogeneous*: catalyst ⁻ in a di erent phase to reactants, work by adsorption o reactants onto sur ace **[2]**; b) *Any three from* cost / selectivity / susceptibility to poisoning / environmental impact / e ficiency **[3]**; c) To increase the sur ace area making it more e ficient. **[1]**

5. a)
$$H$$
 CH_3 CH_3 CH_3 CH_3 CH_3 [2]

b) More crystalline **[1]**, tougher **[1]**; c) LDPE has more side chains (branching) whereas HDPE has little branching so the chains can fit more closely together **[2]**. Since all the CH₃CH=CH₂ molecules are converted into the polymer the atom economy is 100%. **[1]**

6. a) In CVD a carbon-containing gas in the plasma phase is decomposed in the presence o an inert carrier gas, i oxygen was present the carbon would be oxidized to carbon dioxide. [2];b) Nanotubes contain delocalized electrons. [1]



b) It is able to orm cross links due to hydrogen bonding giving it a very ordered three-dimensional structure **[2]**; c) The acid can protonate the N atoms so that they are unable to orm hydrogen bonds and hence it destroys the 3-D structure (strong acid can also break down the amide links within the chains). **[2]**

8. a) $K_{sp} = [Cd^{2+}(aq)] [OH^{-}(aq)]^{2}$ and $[OH^{-}(aq)] = 2[Cd^{2+}(aq)]$ [1]; $K_{sp} = 7.20 \quad 10^{-15} = [Cd^{2+}(aq)] \quad (2[Cd^{2+}(aq)])^{2} = 4[Cd^{2+}(aq)]^{3}$ [1]; Hence $[Cd^{2+}(aq)] = [Cd(OH)_{2}(aq)] = (\frac{7.20}{4} \quad 10^{-15})^{\frac{1}{3}} = (1.80 \quad 10^{-15})^{\frac{1}{3}} = 1.22 \quad 10^{-5} \text{ mol } dm^{-3}$ [1]; b) Since $K_{sp} = [Cd^{2+}(aq)]$

 $[OH^{-}(aq)]^{2}$ adding more hydroxide ions must reduce the concentration o cadmium ions to keep K_{sp} constant **[2]**; c) Ethane-1,2-diamine contains two amino groups both o which contain a non-bonding pairs o electrons which can orm coordinate bonds with a transition metal ion making it a bidentate ligand. **[2]**

9. a) 8 $\frac{1}{8}$ = one **[1]**; b) 6 **[1]**. Using the equation = 2dsin where X-rays with wavelength produce constructive inter erence when di racted at an angle to the sur ace o the crystal to give the distance, d. **[2]**

OPTION B BIOCHEMISTRY (page 140)

1. a) i) H₃N⁺ CH(CH₂CH₂SCH₃) COOH **[1]**; ii) H₃N⁺ CH(CH₂CH₂SCH₃) COO⁻ **[1]**; iii) H₂N CH(CH₂CH₂SCH₃) COO⁻ **[1]**; b) H₂N CH(CH₂CH₂SCH₃) CONH CH(CH₃) COOH **[1]**; H₂N CH(CH₃) CONH CH(CH₂CH₂SCH₃) COOH **[1]**; c) 6 **[1]**; d) Paper chromatography **[1]**; hydrolyse the protein with hot hydrochloric acid to release the separate amino acids **[1]**; spot the sample on paper together with samples o known amino acids above the level o a suitable solvent and let the solvent rise up the paper **[1]**; develop the chromatogram using ninhydrin and compare the *R* values o the spots. **[1]** 2. a) Lipids (or ats) [1]; b) Vegetable [1] as all three atty acid residues are unsaturated [1]; c) Liquid [1] as the structure around the C=C double bonds prevents close packing [1];
d) *Any two from* energy storage or source / insulation / cell membrane. [2]

3. a) i) Alkene **[1]** hydroxyl **[1]**; ii) Vitamin A is at soluble **[1]** as it has a long non-polar hydrocarbon chain **[1]** and vitamin C is water soluble as it contains many –OH groups which can hydrogen bond with water **[1]**; iii) *Vitamin A*: night blindness/ xerophthalmia **[1]**, *Vitamin C*: bleeding o gums etc. **[1]**, scurvy **[1]**; b) Amount o $I_2 = \frac{14.2}{(2-126.9)} = 0.056$ mol **[1]**, each molecule o oil contains 4 C=C double bonds. **[2]**

4. a) i) Using a dye/ninhydrin **[1]**; ii) glutamic acid **[1]**; isoelectric point is below pH o bu er **[1]**; it becomes negatively charged **[1]**; iii) it orms a zwitterion with no overall charge **[1]**; b) H₂NCH₂COOH + H⁺ H₃N⁺CH₂COOH **[1]**, H₂NCH₂COOH + OH⁻ H₃NCH₂COO⁻ + H₂O. **[1]**

5. a) Form coloured compounds **[1]**, orm complex ions **[1]**; b) i) 55% - 25% = 30% **[1]**; ii) CO_2 and lactic acid are acidic so lower the pH **[1]**, haemoglobin is less able to bind to oxygen at lower pH. **[1]**

6. a) It contains a chiral carbon atom **[1]**; b) $K_a = 1.38 10^{-4}$ = [H⁺] [CH(CH₃)(OH)COO⁻] / [CH(CH₃)(OH)COOH] [H⁺]²/0.100 **[1]**; [H⁺] = (1.38 10⁻⁵) = 0.00371 mol dm⁻³ **[1]**, pH = 2.43 **[1]**; c) M_r (lactic acid) = (3 12.01) + (6 1.01) + (3 16.00) = 90.1 and 2.00 g o NaOH = $\frac{2}{(22.99 + 16.00 + 1.01)} =$ 0.0500 mol **[1]**, $K_a = 1.38 10^{-4} = \frac{[H^+] [CH(CH_3)(OH)COO^-]}{[CH(CH_3)(OH)COOH]}$ and pH = 4.00 so $\frac{[CH(CH_1)(OH)COO^-]}{[CH(CH_3)(OH)COOH]} = \frac{1.38 10^{-4}}{10^{-4}} = 1.38$ **[1]**,[CH(CH₃)(OH)COO⁻] = [NaOH] = 0.1 so [CH(CH₃)(OH)COOH]= 0.0725 mol dm⁻³**[1]** $, mass required in 500 cm³ = <math>[(\frac{0.725}{2}) + 0.0500$ (to react with the NaOH)] 90.1 = 7.77 g. **[1]**

OPTION C ENERGY (page 153)

1. a) The combination o two light nuclei to orm a heavier nucleus **[1]**; b) The mass de ect **[1]** is converted into energy according to $E = mc^2$ **[1]**; c) The problems o controlling and maintaining the plasma at very high temperatures have not been overcome **[2]**; X = n, Y = Pu, Z = e^- , a = 92, b = 1, c = 0, d = 0, e = -1. **[4]**

2. a) ${}^{238}_{94}$ Pu ${}^{234}_{92}$ U + ${}^{4}_{2}$ He **[2]**; b) The time an individual atom takes to decay is not xed **[1]**, hal -li e is the average time or many di erent atoms **[1]**; 2 1 0.5 0.25 0.125 0.0625 = 5 hal -lives = 440 years. **[2]**

3. a) *Cracking:* breaking larger hydrocarbons into smaller ones **[1]**, *reforming:* making branched or cyclic hydrocarbons rom straight chain hydrocarbons **[1]**; b) $C_2H_4: = \left(\frac{1411}{28.04}\right)$ 10³ = 5.03 10⁴ kJ kg⁻¹ **[1]**, $C_8H_{18}: = \left(\frac{5470}{114.26}\right)$ 10³ = 4.79 10³ kJ kg⁻¹ **[1]**; c) It combusts as e ciently as a mixture o 5% heptane and 95% 2,2,4-trimethylpentane **[2]**; d) Not so easy to transport as liquid gasoline, more use ul as a eedstock or plastics. **[2]**

4. a) Extensive conjugation **[1]**; b) $6CO_2 + 6H_2O = C_6H_{12}O_6 + 6O_2$ **[2]**; c) i) Too viscous, as molar mass is too high. **[2]** ii)

iii) To move the position o equilibrium to the right to increase the amount o biodiesel ormed. **[1]**

5. a) Greenhouse gases allow the passage o incoming shortwave radiation but absorb some o the longer wavelength reflected radiation rom the Earth and re-radiate it back to the Earth **[2]**. Unlike N₂ and O₂, the bond vibrations o CO₂ and H₂O involve a

change in dipole moment so they can absorb in rared radiation [2]; b) Increased use o ossils uels [1], there is an equilibrium between CO_2 in the atmosphere and dissolved CO_2 in sea-water [1], more CO_2 in water orms more carbonic acid which weakly dissociates to H⁺ ions [1]; c) *Any two from*: move toward more renewable orms o energy, use carbon credits, insulate homes better, increase carbon capture. [2]

6. a) $E = +0.34 - \frac{(8.31 - 298)}{(2 - 96500)} \ln 0.01 = 0.40 \text{ V [2]};$ b) 0.40 - (-0.76) = 1.16 V [2]; c) The standard cell will have a lower em o 1.10 V [1], but will last longer, i.e. do more total work [1], as more materials are present. [1]

7. a) Total mass o 92 protons and 146 neutrons = (92 1.672622 10^{-27}) + (146 1.674927 10^{-27}) = 3.98420566 10^{-25} kg. Mass o one atom o 238 U = $\frac{238.050789}{6.02}$ 10^{-3} = 3.954332043 10^{-25} kg. mass de ect = (3.954332043 10^{-25} – 3.98420566 10^{-25}) = 2.987 10^{-27} kg [1], E = 2.987 10^{-27} (3.00 10^8)² = 2.69 10^{-10} J (2.69 10^{-13} kJ) [1]; b) it is converted into UF₆ which is a gas [1], this can **either** then be allowed to e use and the lighter 235 UF₆ will e use at a aster rate than 238 UF₆ **or** by gas centri ugation where the heavier 238 UF₆ moves to the outside o the container [2]; c) = $\frac{\ln 2}{69} = 1.00$ 10^{-2} year⁻¹ [1], $t = \frac{\ln(\frac{1000}{10.0080})}{1.00} = 461$ years. [1]

8. a) *Any three from:* simpler to manu acture, semi-flexible, semi-transparent, cheaper, absorb visible light over a larger range o wavelengths **[3]**; b) The electrolyte reduces the oxidized orm o the dye and is itsel oxidized to triiodide ions at the anode **[1]**, 3I-

 $I_3^- + 2e^-$ **[1]**, at the cathode electrons are received rom the external circuit and triiodide ions are reduced to iodide ions **[1]**, $I_3^- + 2e^- 3I^-$ **[1]**; c) It acts as a semiconductor and covalently bonds to the dye **[2]**, nanoparticles increase the sur ace area to attach the dye. **[1]**

OPTION D MEDICINAL CHEMISTRY (page 166)

1. a) The therapeutic index or animals uses LD_{so} and it is not reasonable to test a drug to kill 50% o the human population so toxic dose is used instead **[1]**; b) *Therapeutic window*: the range o dosages between the minimum amounts o the drug that produce the desired e ect and a medically unacceptable adverse e ect **[1]**; *therapeutic index* is equal to TD_{so} divided by ED_{so} **[1]**; c) Morphine may be taken to ease pain in which case constipation etc. is a side-e ect **[2]**; or taken or diarrhoea in which case side e ects such as nausea, sleepiness etc. may be experienced **[1]**; d) *tolerance:* require more o the drug to achieve the same e ect **[1]**; *addiction:* going without the drug causes withdrawal symptoms. **[1]**

2. a) Increases the resistance to penicillinase enzyme or alters the e ectiveness o the penicillin [1]; b) Some bacteria may remain una ected [1]; c) Increases the risk o bacteria becoming resistant [1]; d) Due to ring strain the beta-lactam group breaks easily [1] to orm two parts that bond to an enzyme in the bacterium [1] so that it is unable to make cell walls. [1]

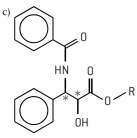
3. a) Esteri cation (also condensation) **[1]**; b) 2-hydroxybenzoic acid will have one peak whereas aspirin will have two as it has two unctional groups containing C=O **[2]**;

d) *Aspirin: any one from:* stomach bleeding, Reye's syndrome, allergic reactions [1], *paracetamol:* overdose can permanently damage kidneys or liver. [1]

4. a) i) Antivirals **[1]**, influenza **[1]**; ii) Zanamivir **[1]**, it contains many polar hydroxyl groups **[1]**; b) i) Proton pump inhibitor **[1]**, *any one from* heartburn, stomach ulcer, gastric reflux, indigestion **[2]**; ii) They are mirror images/enantiomers/ optical isomers. **[1]**

5. a) The di erent enantiomers can have di erent biological properties, one orm may be bene cial, the other orm may be

harmful **[2]**; b) A chiral auxiliary is itself an enantiomer **[1]**, it is bonded to the reacting molecule to create the stereochemical conditions necessary to follow a certain geometric path **[1]**, once the desired enantiomer is formed the auxiliary is removed **[1]**.



6. a) i) ${}^{212}_{82}$ Pb ${}^{208}_{80}$ Hg + ${}^{4}_{2}$ He **[2]**. ii) TAT involves taking the alpha emitter directly to the tumour **[1]**, BNCT involves targeting the tumour with boron-10, a non-radioactive isotope **[1]**, then bombarding it with neutrons so that it then emits radiation on the tumour **[1]**; b) i) ${}^{131}_{54}$ I ${}^{133}_{54}$ Xe + ${}^{0}_{-1}$ e⁻ **[2]**; ii) = ${}^{0.693}_{8.02}$ = 0.08641, ln($\frac{1}{x}$) = 0.08641 31.0 = 2.679 **[1]**, $x = {}^{1}_{14.57}$ = 0.0686, so 6.86% remaining **[1]**; iii) it has a shorter half-life **[1]**, it stays in the body for a shorter time. **[1]**



Origin of individual questions

STOICHIOMETRIC RELATIONSHIPS

 Multiple choice: 1. N11SLP1(1)
 11. N11HLP1(3)

 12. N11SLP1(4)
 13. N10HLP1(2)
 15. N10SLP1(4)

 Short answer: 3. M12TZ2SLP2(2)
 4. M12TZ1SLP2(2)

 5. Adapted from N09SLP2(1)
 8. M05SLP2(2)

ATOMIC STRUCTURE

 Multiple choice: 3. N99SLP1(5)
 4. M11TZ1HLP1(6)

 5. N11SLP1(5)
 6. M12TZ2HLP1(5)
 8. N11HLP1(6)

 15. M98HLP1(8)
 16. N10HLP1(5)

Short answer: 5. M10TZ2SLP2(2) 6. M11TZ2HLP2(3a)

PERIODICITY

Multiple choice: 6. N10SLP1(9) **7.** N10HLP1(7) **12.** M00HLP1(8)

Short answer: 5. Adapted from M10TZ2SLP2(5)
6. N09SLP2(4c)

CHEMICAL BONDING AND STRUCTURE

Multiple choice:1. N11SLP1(9)2. N11SLP1(12)4. N10HLP1(11)5. M99SLP1(12)6. N10SLP1(13)7. N10HLP(10)13. N10HLP1(13)

Short answer: 1. N09SLP2(2) 2. M09TZ2SLP2(7b)
3. M12TZ1SLP2(5a) 4. M12TZ2SLP2(5b)
8. M11TZ1HLP2(6c,d,e & f)

ENERGETICS/THERMOCHEMISTRY

Multiple choice: 3. N11SLP1(14)
4. M99SLP1(16)
5. N10HLP1(15)
7. N99SLP1(17)
9. N10HLP1(17)
10. M12TZ1HLP1(16)
11. N10HLP1(18)
12. M12TZ1HLP1(17)

Short answer: 1. M10TZ1SLP2(6a) **2.** N10SLP2(1a d) **3.** M11TZ1SLP2(4a&b)

CHEMICAL KINETICS

Multiple choice: 1. N10SLP1(17) 2. N10HLP1(19)
3. M99SLP1(19) 4. N09SLP1(18) 5. N09SLP1(19)
6. M12TZ2SLP1(17) 7. M12TZ2SLP1(18) 8. M09TZ2SLP1(17)
9. M09TZ1SLP1(18) 10. M11TZ2SLP1(17) 11. M12TZ1HLP1(20)
12. M12TZ2HLP1(19) 13. M12TZ2HLP1(19)
14. N10HLP1(20)

 Short answer: 1. N07SLP2(7b&c)
 2. M12TZ2SLP2(1a,b &c)

 3. M08TZ2HLP2(1)
 4. N09HLP2(6a c)

EQUILIBRIUM

 Multiple choice:
 2.
 M12TZ2SLP1(19)
 3.
 N11SLP1(20)

 4.
 M09TZ2SLP1(19)
 5.
 M09TZ1SLP1(21)
 6.
 M10TZ1SLP1(20)

 7.
 N10SLP1(20)
 9.
 M10TZ2HLP1(23)
 11.
 N98HLP1(27)

Short answer: 1. M08TZ2SLP2(3) **2.** M07SLP2(4) **3.** N11SLP2(6a) **4.** M11TZ2SLP2(6) **5.** M12TZ1HLP2(6a,b&c)

REDOX PROCESSES Short answer: 1. N09SLP2(1) 3. N11SLP2(3)

ORGANIC CHEMISTRY Short answer: 1. M11TZ1SLP2(6)

Option A MATERIALS Short answer: 3. M13TZ1SLP3(C3)

Option B BIOCHEMISTRY:

4. M11TZ2SLP3(B2)

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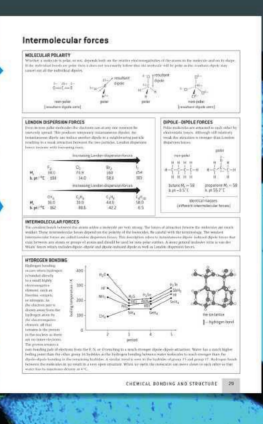
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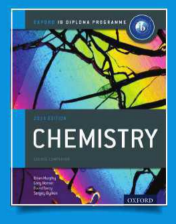
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