

GCE MARKING SCHEME

INTRODUCTION

The marking schemes which follow were those used by WJEC for the SUMMER 2014 examination in GCE CHEMISTRY. They were finalised after detailed discussion at examiners' conferences by all the examiners involved in the assessment. The conferences were held shortly after the papers were taken so that reference could be made to the full range of candidates' responses, with photocopied scripts forming the basis of discussion. The aim of the conferences was to ensure that the marking schemes were interpreted and applied in the same way by all examiners.

It is hoped that this information will be of assistance to centres but it is recognised at the same time that, without the benefit of participation in the examiners' conferences, teachers may have different views on certain matters of detail or interpretation.

WJEC regrets that it cannot enter into any discussion or correspondence about these marking schemes.

	P	age
CH1		1
CH2		7
CH4		13
CH5		21

GCE CHEMISTRY - CH1

SUMMER 2014 MARK SCHEME

SECTION A

Q.1	1s ² 2s ² 2p ⁶ 3s ² 3p ⁶			
Q.2	carbon-12 / ¹² C			
Q.3	any example e.g. iron for Haber process / manufacture of ammonia vanadium(V) oxide in Contact process / manufacture of sulfuric acid platinum / palladium / rhodium in catalytic converters / to remove toxic gases from exhaust fumes nickel in hydrogenation of alkenes / unsaturated oils	[1]		
Q.4		[1] [1]		
Q.5	enthalpy changes = -110	[1]		
Q.6	²³⁴ Th (1) ²³⁴ Pa (1) (award 1 mark for 2 correct symbols)	[2]		
Q.7	Ea ₂ marked, at lower energy than Ea ₁ , and portion to right labelled as molecules the	[1] at [1]		

Section A Total [10]

SECTION B

Q.8	(a)	same number of protons and electrons (1) 0, 1 and 2 neutrons (1)		
	(b)	(i)	3 energy levels between $n=2$ and $n=\infty$ becoming closer together first gap must be < that between $n=1$ and $n=2$	[1]
		(ii)	any arrow pointing upwards (1)	
			from $n = 1$ to $n = \infty$ (1)	[2]
	(c)	(i)	visible	[1]
		(ii)	(not correct because) Balmer series corresponds to energy transition involving $n=2$ (1)	ns
			for ionisation energy need Lyman series / energy transitions involvin $n=1 \ \ (1)$	ng [2]
	(d)	(i)	$Q(g) \rightarrow Q^{+}(g) + e / accept any symbol$	[1]
		(ii)	Group 6	[1]
		(iii)	In T there is more shielding (1)	
			The outer electron is further from the nucleus (1)	
			The increase in shielding outweighs the increase in nuclear charge / there is less effective nuclear charge (1)	[3]
			Legibility of text; accuracy of spelling, punctuation and grammar; clarity of meaning QWC	[1]

Total [14]

Q.9 (a) (i) line drawn that is deflected less by magnetic field [1]

(ii) increase strength of the magnetic field allow decrease charge on charged plates [1]

(b) (i) 1+ (1) ${}^{37}\text{Cl} - {}^{37}\text{Cl}$ (1) ${}^{37}\text{Cl}_2^+$ (2) [2]

(ii) line drawn as m/z 72 (1)

ratio height 6 (1) allow ½ square tolerance [2]

(c) (i) % H = 0.84 (1) C: H: CI = 10.04 / 12: 0.84 / 1.01: 89.12 / 35.5 (1) = 0.84: 0.83: 2.51 = 1: 1: 3 empirical formula = CHCl₃ (1) [3]

(ii) the relative molecular mass / $M_{\rm r}$ / molar mass [1]

(iii) right hand / largest / heaviest m/z peak from mass spectrum [1]

Total [11]

- Q.10 (a) (a reaction in which) the rate of the forward reaction is equal to the rate of the backward reaction [1]
 - (b) goes darker / more brown (1)
 because the (forward) reaction has a +ve ΔH / is endothermic (1)
 goes paler / less brown (1)
 because there are more moles / molecules on RHS (1)
 no change (because catalysts do not affect the position of an equilibrium) (1)
 - (c) (i) moles $N_2H_4 = 14000/32.04 = 437.0$ (1) this produces $437.0 \times 3 = 1311$ moles of gas (1) $volume = 1311 \times 24 = 3.15 \times 10^4 \text{ dm}^3 \text{ (1)} \qquad [minimum 2 \text{ sf}] \qquad [3]$
 - (ii) (large volume of) gas produced [1]
 - (d) (i) an acid is a proton / H⁺ donor [1]
 - (ii) $\rightarrow NO_2^- + H_3O^+$ [1]
 - (iii) sulfuric acid is behaving as the acid / nitric acid is behaving as a base (1)as it donates a proton / as it accepts a proton (1) [2]

Total [14]

[5]

Total [17]

Answer has suitable structure (1)

Q.12 (a) to increase rate of reaction / to increase surface area [1]

(b)
$$MgCO_3 + 2HCI \rightarrow MgCl_2 + CO_2 + H_2O$$
 (ignore state symbols) [1]

(c) rate starts fast and gradually slows (1)

because concentration becomes less so fewer collisions (per unit time) / less frequent collisions / lower probability of collisions (1)

at time =
$$17/18 \text{ min rate} = 0 (1)$$
 [3]

- (d) all the solid would all have disappeared / if more carbonate is added further effervescence is seen [1]
- (e) (i) volume $CO_2 = 200 \text{ cm}^3$ (1) moles $CO_2 = 200 / 24000 = 0.008333 = \text{moles MgCO}_3$ (1) [minimum 2 sf] [2]

(ii) mass MgCO₃ =
$$0.008333 \times 84.3 = 0.702 \text{ g}$$
 (1)
% MgCO₃ = $\frac{0.702}{0.889} \times 100 = 79.0\% / 79\%$ [2]

- (e) carbon dioxide is soluble in water / reacts with water (1)volume collected less therefore % / moles of MgCO₃ less (1)[2]
- (f) use of 40.3 and 84.3 (1) $atom\ economy = 40.3 / 84.3 \times 100 = 47.8\% \ \ (1)$ [2]

Total [14]

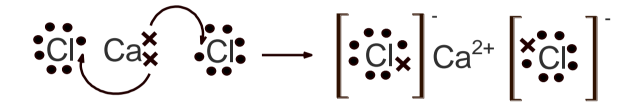
Section B Total [70]

GCE CHEMISTRY - CH2

SUMMER 2014 MARK SCHEME

SECTION A

- Q.1 Van der Waals' forces < Hydrogen bonds < Covalent bonds [1]
- Q.2 2-methylpentan-1-ol allow 2-methyl-1-pentanol [1]
- Q.3 1 mark showing movement of electrons; 1 mark showing dot and cross of CaCl₂ [2]



- **Q.4** (a) δ N—H δ + δ O—Cl δ + both for 1 mark [1]
 - (b) Difference in electronegativity is larger in aluminium oxide (so it is ionic) / the difference is smaller in aluminium chloride (so it is covalent) [1]
- Q.5 Reagent: Bromine (water) (1)
 - Observation(s): hex-2-ene will turn bromine water from orange to colourless, no change for cyclohexane (1) [2]
- Q.6 C (1) and E (1) penalise one mark for each additional incorrect answer [2]

Section A Total [10]

SECTION B

Q.7 (a) (i)

	magnesium	barium	sodium
	nitrate	chloride	hydroxide
potassium	white	white	no visible
carbonate	precipitate	precipitate	change
sodium	WHITE	NO VISIBLE	
hydroxide	PRECIPITATE	CHANGE	
barium chloride	NO VISIBLE CHANGE		,

All three correct for 2 marks, two correct for 1 mark [2]

- (ii) Name of precipitate: Magnesium carbonate (1) Ionic equation: $Mg^{2+} + CO_3^{2-} \rightarrow MgCO_3$ (1) [2]
- (b) (i) Sodium hydroxide solution would turn blue/purple [Ignore references to potassium carbonate] [1]
 - (ii) Potassium carbonate would give a lilac flame
 Sodium hydroxide would give a golden yellow flame
 Barium chloride would give an apple green flame
 (2 for all correct, 1 mark for 2 correct)
 1 max if any reference to white flame for magnesium
 [2]
 - (iii) Barium chloride (1) White precipitate (1) [2]
- (c) Sodium **ions** surrounded by δ on oxygen atoms of water (1) Bromide **ions** surrounded by δ + on hydrogen atoms of water (1) Marks can be obtained from a labelled diagram must show minimum of two oxygen/hydrogen atoms around sodium/bromide ions [2]
 - (ii) Observation with sodium bromide cream precipitate (1)

 Observation with sodium iodide yellow precipitate (1) [2]
 - (iii) Reagent: (dilute) ammonia solution (1)
 Observation with sodium bromide: precipitate dissolves in part
 Observations with sodium iodide: precipitate does not change
 both observations required for (1)
 [If concentrated ammonia (1) used then sodium bromide
 will dissolve completely]
 [2]
 - (iv) $2NaI + Br_2 \rightarrow 2NaBr + I_2$ allow ionic equation [1]

Total [16]

Q.8 (a) Boiling temperatures increase with increasing chain length / number of carbon atoms / relative mass (1)

More carbon atoms leads to greater number of van der Waals' forces between molecules (1)

- (b) (i) Mass of petroleum gases = $1.2\% \times 145$, 000 = 1740g (1) Moles of butane = $1740 \div 58.1 = 30$ mol (1) Volume of butane = $30 \times 24 = 720$ dm³ (1) [3]
 - (ii) I. ultraviolet light [1]
 - II. $Cl_2 \rightarrow 2Cl^{\bullet}$ [1]
 - III. (Propane forms) propyl radicals / C_3H_7 (1) Two C_3H_7 • radicals combine together to make hexane (1) [2]
- (c) Brent crude would be better as it has more naphtha (1)

Naphtha is cracked to produce alkenes (1)

Cracking is caused by heating / zeolites / aluminosilicates / porcelain (1)

Any valid equation that produces ethene e.g. $C_{10}H_{22} \rightarrow C_2H_4 + C_8H_{18}$ (1)

Polymerisation: Many small molecules joining together to make a large molecule (1)

Addition polymerisation (1)

e.g. polystyrene, PVC, PTFE and relevant monomer (1)

QWC: organisation of information clearly and coherently; use of specialist vocabulary where appropriate [1]

Total [16]

[2]

Q.9 (a)
$$1\text{Fe}_2\text{O}_3 + 3\text{CO} \rightarrow 2\text{Fe} + 3\text{CO}_2$$
 [1]

(b) Oxidation state of carbon at start = +2 and at end = +4 so it has been oxidised (1)
Oxidation state of iron at start = +3 and at end = 0 so it has been reduced (1)
Credit 1 mark if all oxidation states are given correctly with incorrect or no
reference to what has been oxidised/reduced [2]

(ii)

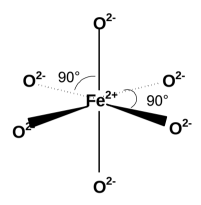


Diagram must be unambiguous, either by showing 3 dimensions, bond angles or through labelling, must identify iron and oxide as ions

[1]

(d) Moles FeO =
$$20,000 \div (55.8 + 16) = 278.6 \text{ mol } (1)$$

Moles Fe = moles FeO = $278.6 \text{ mol } (1)$
Mass Fe = $278.6 \times 55.8/1000 = 15.5 \text{ kg } (1)$ [3]

(e) Pair of shared electrons in both (1)

(f) Lattice / regular arrangement of positive ions (1)
Sea of delocalised electrons (1)
Electrons can move to form an electrical current (1)
Strong forces / bonds between the delocalised electrons and the metal ions require a lot of energy to break / high temperature to overcome (1)

QWC: selection of a form and style of writing appropriate to purpose and to complexity of subject matter

[1]

Total [15]

Q.10	(a)	(i)	Aqueous sodium hydroxide (1) Heat [below 110°C] (1)	[2]
		(ii)	Bromobutane cannot form hydrogen bonds (1) Butan-1-ol can form hydrogen bonds due to its —OH (1) Hydrogen bonds between butan-1-ol and water molecules allow butan-1-ol to dissolve (1)	[3]
	(b)	(i)	Acidified dichromate(VI) / acidified manganate(VII) (1) Heat (1)	[2]
		(ii)	Butanoic acid can form hydrogen bonds between molecules (1) Bromobutane has van der Waals' forces between the molecules Hydrogen bonds are stronger than van der Waals' so require mo energy to break these (1)	` '
				[3]
		(iii)	Fractional distillation	[1]
			Tota	al [11]

Q.11 (a) (i) 1 mark for arrows in first diagram; 1 mark for arrow in second diagram;1 mark for all charges

2 max if incorrect isomer given

[3]

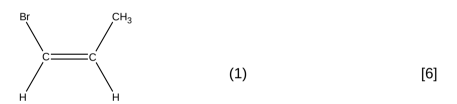
- (ii) 2-bromopropane formed from a secondary carbocation (1) Secondary carbocations are more stable than primary carbocations (1) [2]
- (b) Empirical formula = C_3H_5Br (1) Molecular formula = C_3H_5Br

(must show use of mass spectrum to gain this mark) (1)

Two molecular ion peaks as there are two isotopes of bromine (1)

Peaks at $15 = CH_3^+$ and $41 = C_3H_5^+$ (1)

$$550 \text{ cm}^{-1} = \text{C-Br}$$
 $1630 \text{ cm}^{-1} = \text{C-C}$ $3030 \text{cm}^{-1} = \text{C-H}$ (1) Molecule is:



QWC: legibility of text, accuracy of spelling, punctuation and grammar, clarity of meaning [1]

Total [12]

Section B Total [70]

GCE CHEMISTRY - CH4

SUMMER 2014 MARK SCHEME

SECTION A

Q.1 (a) (i) $CH_3CH_2CH_2CH_3 + Cl_2 \rightarrow CH_3CH_2CH_2CH_2CH_2Cl + HCl$ [1]

(ii) $CH_3CH_2CH_2$ Ĉ HCH_3 [1]

(b) (Anhydrous) aluminium chloride / iron(III) chloride allow AlCl₃ / FeCl₃ [1]

(c) (i) orange / red precipitate [1]

(ii) OCH3

(1) —COCH3 groups in any positions

It must contain a C=O group but it is not an aldehyde as it does not react with Tollens' reagent (1) [2]

- (d) (i) (Alkaline) potassium manganate(VII) (solution) allow $KMnO_4 / MnO_4^-$ [1]
 - (ii) Dilute acid allow HCl / H⁺ [1]
 - (iii) Lithium tetrahydridoaluminate(III) / lithium aluminium hydride allow LiAlH₄ [1]

(iv) CH_2Br [1]

(e) Only the infrared spectrum of benzoic acid would have a peak at 1650–1750 cm⁻¹ (1) This is due to the carbonyl group present in the benzoic acid (1) [2]

Total [12]

Q.2 (a)

[1]

- (b) (i) Acidified potassium dichromate allow H⁺, Cr₂O₇²⁻ [1]
 - (ii) I An equimolar mixture of two enantiomers / optical isomers
 do not accept 'equal mixture' [1]
 - II It has no (apparent) effect on the plane of polarised light [1]
- (c) (i) But-2-enoic acid; this is because each of the carbon atoms of the double bond has two different groups / atoms

 allow reason based on the other isomer [1]
 - (ii) Any TWO from the following for (1) each reagent used / temperature / quantities / time of reaction / catalyst / solvent [2]
- (d) Reagent(s) KOH/I_2 or NaOCl/KI (1) allow names Observation Yellow precipitate (1) [2]
- (e) The NMR spectrum will consist of two peaks, as there are two discrete 'areas' of protons; these will be seen at between 2.0 to 2.5 (CH₃) and between 2.5 to 3.0 (CH₂) (1) The peak area ratio will be 3:2 for the CH₃ and CH₂ protons respectively (1) There will be no splitting of either signal as the protons causing these signals are not bonded directly to other carbon atoms that also have protons (1)

1 max if only one peak described correctly [3]

QWC Legibility of text; accuracy of spelling, punctuation and grammar; clarity of meaning. [1]

Total [13]

Q.3 (a) (i) 2 mol of ethanol gives 1 mol of ethoxyethane (1)

Moles of ethanol = $\frac{69}{46}$ = 1.5

- \therefore Moles of ethoxyethane if theoretical yield = 0.75
- \therefore Moles of ethoxyethane if 45% yield = $0.75 \times 0.45 = 0.34$ (1)

Mass of ethoxyethane = $0.34 \times 74 = 25g$ (1) allow error carried forward [3]

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(ii) Ethene / C_2H_4 [1]

- (1) for correct curly arrows (1) for correct δ^+ and δ^- [2]
- (iv) They need to have an N–H / O–H / F–H bond / a highly electronegative atom bonded to hydrogen [1]
- (b) (i) For example

Accept any polybrominated species Do not accept a monobrominated species

- (ii) Bromine decolorised / orange to colourless / white solid [1]
- (c) Reagent Iron(III) chloride solution / $FeCl_3$ (1)

Observation Purple coloration / solution (1) [2]

(d) (i) $C_{10}H_{12}O_1$ [1]

(ii)
$$H H H$$

 $CH_3 - C - C - C$
 $H Br H$
 $CH_3 - C - C - C$
 $H Br H$
 $CH_3 - C - C - C$
 $H Br H$

(e) Displayed formula, for example

$$HOOC \longrightarrow CH_2-CH_2-CH_3$$
 (1)

Functional group carboxylic acid (1) [2]

Total [15]

SECTION B

- **Q.4** (a) (i) (Fractional) distillation / (preparative) gas chromatography / HPLC / TLC column chromatography / solvent extraction [1]

[1]

- the fragmentation pattern would be different / valid examples given (ii)
- (iii) Ι

$$CH_2NH_2 + CH_3C$$
 CI
 CH_2-N-C
 CH_3
 CH_3
 CH_3
 CH_3
 CH_3
 CH_3

II Heated electrically / by a naked flame with a water bath (1) Add compound **G** to the ethanol until the hot ethanol will (just) not dissolve any more solute (1)

Filter hot (1)

Allow to cool (1)

Filter (1)

Dry in air / window sill / < 60 °C in an oven (1)

[5]

Maximum 4 out of 5 total if second marking point not given Note 5 marks maximum here

QWC Information organised clearly and coherently, using specialist vocabulary where appropriate

[1]

The amine is reacted with sodium nitrite / HCl(aq) or nitrous acid (1) (iv) at a temperature of < 10 °C (1)

[2]

II

$$N = N - CH^3$$

(b) (i) Nucleophilic addition (1)

Accept a mechanism that shows HCN polarisation and nucleophilic addition as a concerted process

polarisation / charges shown (1) curly arrows on first structure (1) regeneration of ${}^{-}C \equiv N$ or capture of H^{+} and curly arrow (1) [4]

(ii) Chromophores (1) The colour will be black (1) as the compound absorbs blue / other colours (1) [3]

Total [20]

Q.5 (a) C 71.3 H 9.6
$$\therefore$$
 O 19.1 (1)
÷ by A_r $\frac{71.3}{12} = 5.94$ $\frac{9.6}{1.0} = 9.6$ $\frac{19.1}{16} = 1.193$
÷ smallest $\frac{5.94}{1.193} = 5$ $\frac{9.6}{1.193} = 8$ $\frac{1.193}{1.193} = 1$ (1)

Only one oxygen atom per molecule

 \therefore Molecular formula is C_5H_8O (1)

Silver mirror produced
$$\therefore -C$$
 present (1)

Ion m/z 29 suggests ethyl group present / CH_3CH_2 (1)

Structure must be

[6]

(b) (i)
$$C_{11}H_{24} \longrightarrow C_6H_{14} + C_2H_4 + C_3H_6$$
 [1]

(ii) Total peak areas 26 + 13 + 46 = 85

% propene =
$$\frac{13 \times 100}{85}$$
 = 15.(3) [1]

(iii) Any THREE points for (1) each [3]

e.g. can it run at a lower temperature (reducing energy costs) is the yield comparable / better than the yield from the propene process is the time taken comparable / better than used in the propene process is there a continued availability of starting materials can the product be easily / better separated from the reaction mixture is relatively more expensive equipment needed is it a batch or continuous process

(iv)
$$CH_2 - CH_2 - CH_2$$

O O

C

C

C

C

C

C

(I)

O CH₃

O CH₃

O CH₃

- (ii) The production of PTT is an example of condensation polymerisation (1)
 The production of poly(propene) is an example of addition polymerisation (1)
 Condensation polymerisation needs bifunctional compounds / COOH,OH etc (1)
 - Addition polymerisation needs a C=C present in the monomer (1)
 - Addition polymerisation has an atom economy of 100% (1)

 Condensation polymerisation has an atom economy of < 100% (as a co-product is formed) (1) [6]
 - QWC Selection of a form and style of writing appropriate to purpose and to complexity of subject matter [1]

Total [20]

GCE CHEMISTRY - CH5

SUMMER 2014 MARK SCHEME

SECTION A

Q.1 (a) (i)
$$NH_4^+(aq) + OH^-(aq) \longrightarrow NH_3(aq) + H_2O(1)$$

Acid 1 Base 2 Base 1 Acid 2 [2]
(1 mark for each pair)

(b) (i)

	$[NH_4^+(aq)]/mol\ dm^{-3}$	$[NO_2^-(aq)]/mol dm^{-3}$	Initial rate/mol dm ⁻³ s ⁻¹
1	0.200	0.010	4.00×10^{-7}
2	0.100	0.010	2.00×10^{-7}
3	0.200	0.030	1.20×10^{-6}
4	0.100	0.020	4.00×10^{-7}

(1 mark for each correct answer)

[3]

(ii)
$$k = 4.00 \times 10^{-7} = 2.0 \times 10^{-4}$$
 (1) 0.200×0.010

Units =
$$\text{mol}^{-1} \text{ dm}^3 \text{ s}^{-1}$$
 (1)

(iv) Increases

If temperature is increased rate increases (1)

and since concentrations do not change the rate constant must increase (or similar) (1) [2]

Total [10]

Q.2 (a)
$$K_w = [H^+][OH^-]$$
 (1) Units = $mol^2 dm^{-6}$ (1) [2]

(b) (i) In pure water
$$[H^+] = [OH^-]$$
 or $[H^+] = \sqrt{1.0 \times 10^{-14}}$ (1)
$$pH = -\log 10^{-7} = 7$$
 (1) [2]

(ii) Final volume of solution is 1000 cm³ so acid has been diluted by a factor of 100 so final concentration of acid is 0.001

or moles acid =
$$\frac{0.1 \times 10}{1000}$$
 = 0.001 (1)
pH = $-\log 0.001 = 3$ (1) [2]

(c)
$$1.78 \times 10^{-5} = \underbrace{[\text{H}^+] \times 0.02}_{0.01}$$
 (1)

$$[H^+] = 8.90 \times 10^{-6} \tag{1}$$

$$pH = 5.05$$
 allow 5 or 5.1 (1)

(Accept correct equations) (1)

When an acid is added, the CH_3COO^- ions react with the H^+ ions, removing them from solution and keeping the pH constant (1) [3]

Total [12]

Total [18]

SECTION B

ΩA	(a)	(i)	Oxidising agent	[1]
Ų.4	(a)	(1)	Oxidising agent	

(ii)
$$A = lead(II) chloride / PbCl_2$$
 (1) $B = chlorine / Cl_2$ (1) [2]

(iii)
$$[Pb(OH)_6]^{4-} / [Pb(OH)_4]^{2-} / Na_4[Pb(OH)_6] etc.$$
 [1]

(v)
$$PbO + 2HNO_3 \longrightarrow Pb(NO_3)_2 + H_2O$$
 [1]

Layers held together by weak intermolecular forces (1)

BN is isoelectronic with C so it forms similar structures (1)

Graphite conducts electricity since electrons are delocalised but in BN, each N has a full unbonded p-orbital and each B has an empty unbonded p-orbital so it does not conduct electricity (1) [4]

(Accept electrons are not delocalised in BN so it does not conduct electricity)

QWC The information is organised clearly and coherently, using specialist vocabulary where appropriate [1]

(ii) Wear-resistant coatings/catalyst support/for mounting high power electronic components / drills in industry / cutting instruments [1]

(c) (i)
$$\Delta G = \Delta H - T \Delta S$$
 ($\Delta G = 0$ for reaction to be spontaneous) (1)

$$T = \frac{1.92}{0.0067} \tag{1}$$

$$T = 286.6 \text{ K}$$
 (1)

(ii) Changes in temperature (above or below 286.6 K) caused the tin to change form making it unstable (and causing it to disintegrate)

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(d) (i) (At the anode) $H_2 \longrightarrow 2H^+ + 2e^-$ (1)

(At the cathode) $O_2 + 4H^+ + 4e^- \longrightarrow 2H_2O$ (1)

(Overall reaction) $2H_2 + O_2 \longrightarrow 2H_2O$ (1)

[3]

(ii) Hydrogen is difficult to store / takes up large volume / too flammable / explosive / produced from fossil fuels which leads to a net energy loss / Pt electrodes very expensive [1]

Total [20]

Q.5 Cold $Cl_2 + 2NaOH$ NaCl + NaClO + H_2O (a) (i) (1) Warm $3Cl_2 + 6NaOH \longrightarrow 5NaCl + NaClO_3 + 3H_2O$ (1) [2] (ii) Disproportionation [1] (b) P can (extend the normal octet of electrons) by using 3d orbitals / P can promote 3s electron to 3d orbital N cannot do this since it is in the second period / 3d orbitals not available (1) [2] (c) The terms involved are: lattice breaking enthalpy which is endothermic (1) and hydration enthalpy which is exothermic (1) ΔH solution = ΔH lattice breaking + ΔH hydration (or similar) (1) If ΔH solution is negative then the ionic solid will be soluble (1) [4] QWC Selection of a form and style of writing appropriate to purpose and to complexity of subject matter [1] (d) (i) Iodide (1) Only one with less positive standard potential than Fe³⁺, Fe²⁺ half-cell (1) [2] (2nd mark can be obtained from calculation value and statement) $Pt(s) | Fe^{2+}(aq), Fe^{3+}(aq) | Ce^{4+}(aq), Ce^{3+}(aq) | Pt(s)$ (ii) (1) EMF = 1.45 - 0.77 = 0.68 V(1) [2] (e) $K_c = [CH_3COOCH_3][H_2O]$ (1) (i) [CH₃COOH][CH₃OH] No units (1) [2] (ii) $moles = \underline{1.25 \times 32.0} = 0.04(0)$ [1] $[CH_3COOH] = 0.04$, therefore 0.06 used in reaction and (iii) $[CH_3COOCH_3] = 0.06$, $[H_2O] = 0.06$ and $[CH_3OH] = 0.083 - 0.06 = 0.023$ (1) $K_c = 0.06 \times 0.06 = 3.91$ (1) [2] 0.04×0.023 Value of K_c decreases since the equilibrium shifts to the left / (iv)

Total [20]

[1]

the forward reaction is exothermic



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