

Cambridge International Examinations Cambridge International Advanced Subsidiary and Advanced Level

CHEMISTRY

Paper 4 A Level Structured Questions SPECIMEN MARK SCHEME 9701/04 For Examination from 2016

2 hours

MAXIMUM MARK: 100

This document consists of 8 printed pages.



Mark scheme abbreviations

- ; separates marking points
- *I* alternative answers for the same point
- R reject
- A accept (for answers correctly cued by the question, or by extra guidance)
- **AW** alternative wording (where responses vary more than usual)
- **<u>underline</u>** actual word given must be used by candidate (grammatical variants excepted)
- max indicates the maximum number of marks that can be given
- ora or reverse argument
- mp marking point (with relevant number)
- ecf error carried forward
- I ignore
- **AVP** Alternative valid point (examples given as guidance)

- 1 (a) (i) carbonates become more stable down the Group/higher decomposition temperature (1) cation/ M^{2+} radius/size increases down the group/ M^{2+} charge density decreases (1) anion/carbonate ion/ CO_3^{2-} suffers less polarisation/distortion (1) [3]
 - (ii) ionic radii quoted: $Ca^{2+}: 0.099 \text{ nm}, Zn^{2+}: 0.074 \text{ nm}, Pb^{2+}: 0.120 \text{ nm} (1)$ thus we expect $ZnCO_3$ to be less stable, but $PbCO_3$ to be more stable (1) [2]

if candidate states $PbCO_3$ is more stable than $ZnCO_3$ (or converse) with no reference to $CaCO_3$ (1)

(b) hydroxides become more soluble down the group (1) both lattice energy and hydration decrease (1) but lattice energy decreases more than hydration energy so enthalpy of solution become less endothermic (1)

[4]

[Total: 9]

2 (a)

(a)							
		[CH ₃ CHO] /mol dm ⁻³	[CH ₃ OH] /mol dm ⁻³	[H*] /mol dm ⁻³	[acetal A] /mol dm ⁻³	[H ₂ O] /mol dm ⁻³	
	at start	0.20	0.10	0.05	0.00	0.00	
	atequilibrium	(0.20 - x)	(0.10 – 2x)	0.05	x	x	
	atequilibrium	0.175	0.05	0.05	0.025	0.025	
	(i) 3 values in	second row 3	× (1)				[3]
(ii) 4 values in	third row 4 \times	(1)				[4]
(i	ii) K _c = {[ace units = mo	tal A][H ₂ O]}/{[(ol ⁻¹ dm ³ (1)	CH₃CHO][CH₃	OH] ² } (1)			[2]
(i	v) $K_{\rm c} = 0.025$	²/(0.175 × 0.0	5 ²) = 1.4(3) (mo	ol ^{–1} dm ³)			[1]
(b)	(i) Order w.r.t Order w.r.t	[CH ₃ CHO] = 1 . [CH ₃ OH] = 1					
	Order w.r.t	[H ⁺] = 1					[3]
(ii) rate = k[CH	H ₃ CHO][CH ₃ O	H][H+]				[1]
(i	ii) units = mo	l ⁻² dm ⁶ s ⁻¹					[1]
(i	v) rate will be	$2 \times 4 = 8$ time	es as fast as re	eaction 1 (relat	ive rate = 8)		[1]
						[T	otal: 16]



4

[1]

[Total: 13]

4	(a)	$K_2Cr_2O_7 + H^+ + heat under reflux$	[1]
	(b)	nucleophilic substitution	[1]
	(c)	heat under reflux + aqueous HCl	[1]

- (d) alkene [1]
- (e) amide or ester [1]
- (f)

(f)



ecf 5 × [1]

[5]

[Total: 10]

(a)	(i)	C = C double bonds / alkenes
(a)	(I)	C = C double bolius / alkelles

- (ii) -OH groups / accept alcohols or acids [1]
- (iii) CH₃CO- or CH₃CH(OH)- groups [1]
- (iv) carbonyl, >C=0, groups / accept aldehydes and ketones [1]



[2]

[1]

(c) isomers of G

5



correct structure (excluding stereochemistry) (1) cis and trans drawn correctly (1) type of isomerism is cis-trans or geometrical isomerism (1) [3]

[Total: 9]

- 6 (a) (i) A is Cl_2 /chlorine (1) **B** is NaC*l* or HC*l* or C*l*⁻ [or words] (1) **C** is salt bridge or $KCl/KNO_3(1)$ **D** is platinum/Pt (1) **E** is $Fe^{2+} + Fe^{3+}$ or mixture of Fe(II) + Fe(III) salts (1) [5]
 - $E^{\bullet} = E^{\bullet}_{R} E^{\bullet}_{L} = 0.77 1.36 = (-)0.59$ (V) (ignore sign) (1) (ii) (since R.H. electrode is negative electrons flow (from right) to left or to the chlorine electrode *or* anticlockwise *or* from (beaker) **E** to (beaker) **B** (1) [2]

(b) (i)
$$\Delta H^{6} = 3 \times (-167.2) + (-48.5) - (-399.5) (1)$$

 $= -150.6 \text{ or } 151 \text{ (kJ mol}^{-1}) (1)$
correct answer only (2) [2]
(ii) $2\text{Fe}^{3+} + \text{Cu} \rightarrow 2\text{Fe}^{2+} + \text{Cu}^{2+} (1)$
(or molecular: $2\text{Fe}\text{C}l_{3} + \text{Cu} \rightarrow 2\text{Fe}\text{C}l_{2} + \text{Cu}\text{C}l_{2})$
 $\text{E}^{6} = 0.77 - 0.34 = (+) 0.43 \text{ (V)} (1)$ [2]

(or molecular:
$$2FeCl_3 + Cu \rightarrow 2FeCl_2 + CuCl_2$$
)
 $E^{\circ} = 0.77 - 0.34 = (+) 0.43$ (V) (1)
(no mark for -0.43V)

[Total: 11]

6

7 (a)

process	sign of ΔS
$NaBr(s) + (aq) \rightarrow NaBr(aq)$	+
$H_2O(I) \rightarrow H_2O(g)$	+
$2H_2(g) + O_2(g) \rightarrow 2H_2O(g)$	_
$CoCl_2(s) + 6H_2O(I) \rightarrow CoCl_2.6H_2O(s)$	_

2 correct, (1) mark 4 correct, (2) marks

[2]

- (b) $\Delta S^{o} = (214 \times 2) + (70 \times 3) (161 \times 1) (205 \times 3)$ = -138 J K⁻¹ mol⁻¹ [2]
- (c) As temperature increases $T \Delta S$ is more negative or $-T \Delta S$ increases (1) At high temperature $T \Delta S$ is more negative than ΔH (so ΔG is positive) (1) [2]
- (d) the reaction is feasible, ΔG is negative so T > $\Delta H/T \Delta S$ or use of T = $\Delta H/T \Delta S$ (1)
 - T = 178000/159 (1) T = 1119.5 K units required or T>1120 K (1)

[3]

[Total: 9]

(a) X is

allow –N₂— and –ONa [1]

(b) reaction I: $\underline{C}l_2 + \text{light}(1) (\text{not } aq)$ reaction II: $Br_2 + AlBr_3 \text{ or } Fe \text{ or } FeBr_3 (1) (\text{not } aq)$ reaction III: NaOH, heat in ethanol (1) (allow aqueous EtOH) reaction IV: $HNO_3 + H_2SO_4 (1) \text{ conc } and 60 \,^\circ\text{C} (1)$ reaction V: $KMnO_4 + H^+/OH^- + \text{heat } (1)$ reaction VI: Sn + HCl (1)reaction VII: $HNO_2 + HCl < 10 \,^\circ\text{C} (1)$

OH

[Total: 9]

[8]

8

9 (a) time for a component between injection and travelling to the detector [1]

(b) (i) No. of carbon atoms present in J is
$$\frac{100 \times 1.3}{1.1 \times 23.5} = 5$$
 carbons (must show working) [1]

(ii) 4 different carbon environments (1)

$$\delta$$
 210 is C = O carbon (1)
 δ 15-45 are alkyl carbons/C–C (1) [3]
(iii) Y is

[Total: 8]

10 (a) (many) monomers add together to form a polymer and small molecule (such as H_2O , HCl) [1]

(b)

bonding type	secondary structure	tertiary structure
hydrogen bonding	\checkmark	\checkmark
ionic bonding		\checkmark
van der Waals'		\checkmark

2 correct [1]; all correct [2]

C

ĊН₃ **С**

CH₃

(1) Isomer A would show 5 absorptions/peaks (1) Isomer **B** would only show 3 absorptions/peaks (1)

H₃C 、

- (c) (i) pH of the buffer solution
 - (ii)

amino acid	Identity of amino acid (any one of)
А	Asp, Glu
В	Gly, Val, Phe, Ala
С	Lys

2 correct [1]; 3 correct [2]

[2]

[2]

[1]

[3]