

# **GCE MARKING SCHEME**

## CHEMISTRY AS/Advanced

**SUMMER 2012** 

#### CH5

### SECTION A

1. (a) $1 \text{ dm}^3$ at 20°C contains 52.9 g and at 0°C it contains 17.5 g (1) $\therefore$ amount crystallised = 52.9 - 17.5 = 35.4 g (1)	[2]
(b) (i) 2 mol of $K_2S_2O_8$ give 1 mol of $O_2$ 2 mol of $K_2S_2O_8$ give 29.0 dm <sup>3</sup> of $O_2$ (1) $\therefore$ 0.1 mol of $K_2S_2O_8$ gives 29.0/20 = 1.45 dm <sup>3</sup> of oxygen (1)	[2]
<ul> <li>(ii) Measure the volume of oxygen produced at specified time intervals / Measure the pH of the solution at specified time intervals</li> </ul>	[1]
(c) (i) An (inert) electrode that is used to carry the charge / current / electron flow	[1]
<ul> <li>(ii) A comment on the relative values (e.g. the persulfate system is the more posit of the two systems) (1)</li> <li>The more positive 'reagent' / persulfate ions acts as the oxidising agent, accep electrons via the external circuit (1)</li> <li>must have the first mark to get second</li> </ul>	
(d) (i) The experiments show that both the concentrations of iodide and persulfat have doubled (1) therefore the initial rate should increase four times $4 \times 8.64 \times 10^{-6} = 3.46 \times 10^{-5}$ (1)	e [2]
(ii) Rate = $k [S_2 O_8^{2-}] [I^-]$ (1)	
$\therefore k = \frac{8.64 \times 10^{-6}}{0.0400 \times 0.0100}$	
$= 0.0216 (1) dm^3 mol^{-1} s^{-1} (1)$	[3]
(iii) In the rate equation one $S_2O_8^{2-}$ ion reacts with one I <sup>-</sup> ion. The rate-determining step therefore has to have 1 mole of each reacting, as (only) seen in step 1	[1]

Total [14]

2.	(a)	- 705 (kJ mol <sup>-1</sup> ) (1) for correct sign (1) for correct number	[2]
	(b) (i)	hydration	[1]
	(ii)	e.g. add a small 'amount' of an alkali / sodium hydroxide / NaOH / OH <sup>-</sup> ions (1 this would remove / react with hydrogen ions giving water, shifting the position of equilibrium to the left (removing iodine) (1) add $P6^{2+}$ / $Ag^+$ ect.	
	(c) (i)	Any TWO from white / misty fumes (of HI) yellow solid / solution (of sulfur) brown / black solid / purple vapour (of iodine) bubbles / effervescence / fizzing	
		One mark for each correct response	[2]
	(ii)	The values show that chlorine is the best oxidising agent, as it has the most positive $E^{\theta}$ value and therefore iodide is the better reducing agent (1)	
		and is 'strong' enough to reduce the sulfuric acid. / OWTTE (1)	[2]
	(d) (i)	$2 \text{ NaOH} + \text{Cl}_2 \rightarrow \text{NaOCl} + \text{NaCl} + \text{H}_2\text{O}$	[1]
	(ii)	e.g. bleach, kills bacteria	[1]
		Tot	tal [11]

3. (a) (i)

Number of moles of EDTA = 
$$\frac{19.20 \times 0.010}{1000}$$
 =  $1.92 \times 10^{-4} / 0.000192$  [1]

- error carried forward throughout (a)

(ii) 
$$1.92 \times 10^{-4} / 0.000192$$
 [1]

(iii) Concentration = 
$$\frac{1.92 \times 10^{-4} \times 1000}{50}$$
 =  $3.84 \times 10^{-3} / 0.00384 \text{ mol dm}^{-3}$  (1)

Concentration = 
$$3.84 \times 10^{-3} \times 63.5 = 0.244 \text{ g dm}^{-3}$$
 (1) [2]

(iv) % Cu = 
$$0.244 \times 100 = 2.11$$
 [1]

- (b) Transition elements have either a partly filled 3d sub-shell or form ions that have a partly filled 3d sub-shell (1) However copper forms  $Cu^{2+}$  ions that are '3d<sup>9</sup>' / partly filled 3d sub-shell (1) whereas  $Zn^{2+}$  ions are '3d<sup>10</sup>' / full 3d sub-shell (1) - any 2 from 3 [2]
  - *QWC Organisation of information clearly and coherently; use of specialist vocabulary where appropriate.* [1]

(c)

Complex ion	Shape	Colour
$\left[\mathrm{CuCl}_4\right]^{2-}$	tetrahedral	yellow / lime green
$[Cu(NH_3)_4(H_2O)_2]^{2+}$	octahedral	deep blue

Any two correct (1) all correct (2)

[2]

- (d) The more negative the  $\Delta H_f$  value the more stable the oxide (1) PbO is relatively the more stable / CuO is relatively the less stable (1) - must have the first mark to get second [2]
- (e) (i) Any TWO from variable oxidation states partially filled 3d energy levels ability to adsorb 'molecules' ability to form complexes with reacting molecules / temporary / co-ordinate bonds
   One mark for each correct response [2]
   (ii) e.g. to allow lower pressures / temperatures use recyclable catalysts - needs qualifying [1] longer lasting / less toxic catalysts

Total [15]

#### **SECTION B**

4. (a)	$CO \rightarrow C +2 \qquad CO_2 \rightarrow +4 \qquad (1)$	
	Increase of (positive) oxidation number = oxidation / reducing agents themselves are always oxidised are always oxidised (1)	
	OR $I_2O_5 \rightarrow I +5$ $I_2 \rightarrow I_2 0$ (1)	
	Oxidation number of iodine reduced, reduction occurring, CO reducing agent (1)	[2]
(b)	+2 state becomes mores stable down the group and +4 becomes less stable.	[1]
<ul> <li>(c) (i) Add (a little) sodium hydroxide solution (1) to each solution.</li> <li>A white precipitate (1) of aluminium / lead(II) hydroxide (1) is seen.</li> </ul>		
	When more sodium hydroxide solution is added these precipitates (dissolve giving a colourless solution). (1)	[4]
	<i>QWC</i> Legibility of text: accuracy of spelling, punctuation and grammar; clarity of meaning.	[1]
	(ii) Yellow precipitate (1) $Pb^{2+} + 2I^- \rightarrow PbI_2$	[2]

(d) (i) The bonding of aluminium in the monomer has not completed the octet / suitable diagram / 6 electrons in its outer shell (1)
 When the dimer is formed this octet of bonded electrons is formed (1)

$$\begin{array}{c} CI \\ | \\ CI \\ | \\ | \\ CI \\ \\ CI \\ \\ CI \end{array}$$
(1) [3]

(ii) (As a catalyst) in the chlorination of benzene / making ionic liquids [1]

(iii) I The number of (gaseous) species is increasing, leading to less order [1]

II For the reaction to be just spontaneous  $\Delta G = 0$  (1)

substituting  $0 = 60\ 000 - 88\ T$ 

$$T = 60\ 000\ /\ 88 = 682\ K\ /\ 409^{\circ}C \qquad (1) \qquad [2]$$

$$K_{c} = [[\underline{Al(H_{2}O)_{5}(OH)}]^{2+}(aq)][[\underline{H^{+}}](aq)]} \\ [[Al(H_{2}O)_{6}]^{3+}(aq)]$$

$$\therefore 1.26 \times 10^{-5} = [H^{+}]^{2} / 0.10$$

$$\therefore [H^{+}]^{2} = 1.26 \times 10^{-6} [1]$$

$$\therefore [H^{+}] = \sqrt{1.26 \times 10^{-6}} = 1.12 \times 10^{-3} / 0.00112 (1)$$

$$- \text{ error carried forward}$$

$$pH = -\log_{10}[H^{+}] = -\log_{10} 1.12 \times 10^{-3} = 2.95 (1) [3]$$

Total [20]

(e)

5. (a)	(i) $K_p = \frac{pSO_3(g) \times pNO(g)}{pSO_2(g) \times pNO_2(g)}$ (1) there are no units (1)	[2]		
(i	i) The line for SO <sub>3</sub> / NO at equilibrium should be above the SO <sub>2</sub> / NO <sub>2</sub> line (1) as $K_p$ has a value of 2.5, the partial pressures of SO <sub>3</sub> and NO at equilibrium will be greater than the partial pressures of SO <sub>2</sub> and NO <sub>2</sub> . (1)			
	- accept answer in terms alternative calculated K			
	The line for equilibrium should start at 9 hours. (1) as at equilibrium the concentrations is unchanged as time progresses. (1)	F 4 J		
	There may be other acceptable forms of explanation to be discussed at the conference	[4] e		
(iii)	) If the temperature rises then the position of equilibrium will move to the left, (reducing the quantities of SO <sub>3</sub> and NO). (1) This will make the value of K <sub>p</sub> smaller. (1)	[2]		
(b) (	i) Nitric acid is a strong acid and its pH is low $/ <2 / 1.0$ (1)	[-]		
	As aqueous ammonia is added the pH slowly rises (1) until a pH of $\sim$ 3 is reached, when it rises rapidly (1)			
	At a pH of 8-9, it tails off slowly as ammonia is a weak base. (1) Accept any 3 from 4	[3]		
	Selection of a form and style of writing which is appropriate to purpose and to complexity of subject matter	[1]		
(ii)	) The equivalence point is reached when $20.0 \text{ cm}^3$ of ammonia solution has been added as this is at the mid point of the more vertical section. (1)	1		
	Since both reagents have the same concentration and the volumes used are both 20 cm the same, the number of moles of each are the same (1)	n <sup>3</sup> /		
	OR the number of moles of both nitric acid and aqueous ammonia are calculated (0.0 and shown to be the same (1)	0020)		
	$\therefore$ Mole ratio must be 1 : 1 (1)	[2]		
(iii)	<ul> <li>I Ammonium nitrate is the salt of a strong acid and weak base / there is a buffering in operation.</li> <li>II ~ 5.5</li> </ul>	effect [1] [1]		
(iv)	Blue, as bromophenol blue is blue at a pH of 4.7 and above	[1]		
(c)	Number of moles of ammonium nitrate $= \frac{40}{80} = 0.50$ (1)			
	- error carried forward Concentration of ammonium nitrate solution = $\frac{0.5 \times 1000}{200}$ = 2.5 mol dm <sup>-3</sup>	(1)		
	$\therefore \text{ Temperature drop} = 2.5 \times 6.2 = 15.5^{\circ}\text{C}  (1)$	[3]		
	Total [20]			
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