PMT

# MARK SCHEME for the May/June 2013 series

# 9701 CHEMISTRY

9701/42

Paper 4 (A2 Structured Questions), maximum raw mark 100

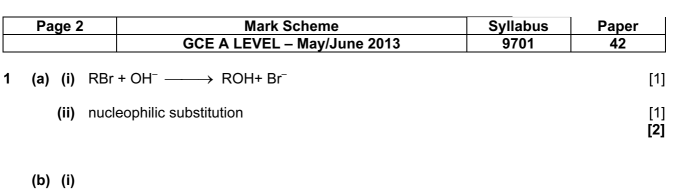
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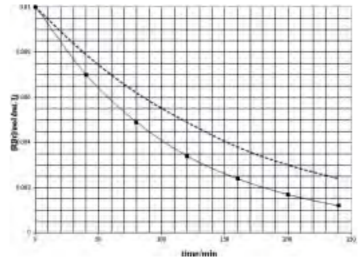
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plotting of all points (plotted to within  $\frac{1}{2}$  small square) [1]

good line of best fit [1]

(ii) t<sub>1/2</sub> = 118 min or 79 min (± 5 min) or construction lines for two half-lives and mention that half-life is constant or calculate the ratio of two rates at two different concentrations [1]
(iii) either ratio of initial rates (slopes) or ratio of t<sub>1/2</sub> or ratio of times for [RBr] to fall to the same level: all should be = 1.5 [1]

therefore reaction is first order w.r.t. [OH⁻]

(iv) rate = k[RBr][OH<sup>-</sup>]

initial rate =  $0.01 / 185 = 5.4 \times 10^{-5} \text{ (mol dm}^{-3} \text{ min}^{-1}\text{)}$  [1]

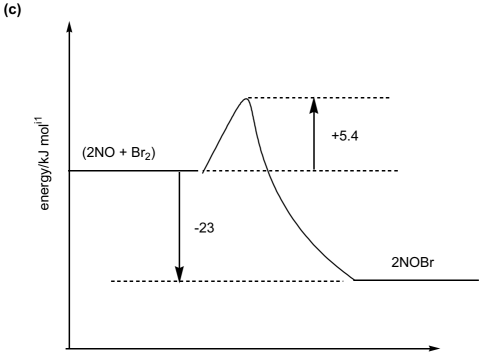
$$k = 5.4 \times 10^{-5} / (0.01 \times 0.1) = 0.054 \text{ (mol}^{-1} \text{ dm}^3 \text{ min}^{-1})$$
[1]

[8 max 7]

[1]

[1]

Page 3	Mark Scheme	Syllabus	Paper
	GCE A LEVEL – May/June 2013	9701	42



extent of reaction

 $\begin{array}{l} \mbox{four marking points: one activation "hump"} \\ \underline{2} \mbox{NOBr (not just NOBr)} \\ \Delta H \mbox{ labelled correctly (arrow down, or double headed, or just a line)} \\ E_a \mbox{ labelled correctly (arrow up, or double headed, or just a line)} \\ \mbox{ all four points [2]} \\ \mbox{ three or two points [1]} \\ \mbox{ [2]} \end{array}$ 

[Total: 11]

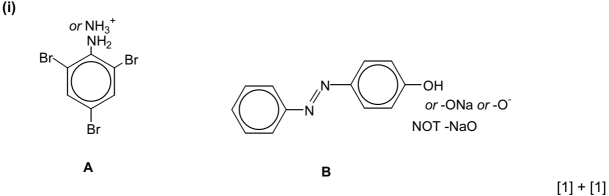
4	Mark Scheme	Syllabus	Paper
	GCE A LEVEL – May/June 2013	9701	42
•	at 1 atm Pt $H^+/HCl at 1 mol dm^{-3} and 298K$ $H_2(g) going$	ition, with H <sub>2</sub> bubl	
	solution at 1 mol dm <sup>-3</sup> (or 0.5 M if H		
<i>E</i> ° =	1.33 – (-0.41) = 1.74 V		[1]
Cr <sub>2</sub> O	$P_7^{2^-}$ + 14H <sup>+</sup> + 6Cr <sup>2+</sup> $\longrightarrow$ 8Cr <sup>3+</sup> + 7H <sub>2</sub> O		[1]
Colo	ur would change from orange		[1]
to gro	een		[1] <b>[8]</b>
ere are a = –lo	two ways of calculating the ratio: $g_{10}(K_a) = -\log_{10}(1.79 \times 10^{-5}) = 4.747 (4.75) \text{ or } [\text{H}^+] =$	10 <sup>-5.5</sup> = 3.16 x 10 <sup>-</sup>	<sup>6</sup> [1]
J₁₀([B] /	/ [A]) = pH – pK <sub>a</sub> = 0.753 (0.75) <i>or</i> [salt] / [acid] = K <sub>a</sub> /	[H <sup>+</sup> ]	[1]
= 1.79 <sup>-</sup> [A] / [E	$x 10^{-5} / 3.16 \times 10^{-6} = 5.66$ B] = <u>0.177</u> )		[1] io = [3] marks)
	hydi gas : $E^{\circ} =$ $Cr_2C$ Colo to gr $e^{re}$ are a = -lo $h_{10}([B], [A])$ [B] / [A]	GCE A LEVEL – May/June 2013hydrogen gas at 1 atmmathematical formula (figure (fig	GCE A LEVEL – May/June 20139701hydrogen gas at 1 atmImage: state of the state o

	Pa	ige 5	5	Mark Scheme	Syllabus	Paper
				GCE A LEVEL – May/June 2013	9701	42
	(c)	.,		$CO_2Na + HCl \longrightarrow CH_3CO_2H + NaCl$ $CO_2H + NaOH \longrightarrow CH_3CO_2Na + H_2O$		[1] [1]
	(d)	e.g or	hydr hydr nitra dehy (or ti	rolysis of esters $\text{RCO}_2\text{R'}$ (+ $\text{H}_2\text{O}$ ) $\longrightarrow$ $\text{RCO}_2\text{H}$ + $\text{R'OH}$ rolysis of amides: $\text{RCONH}_2$ (+ $\text{H}_3\text{O}^+$ ) $\longrightarrow$ $\text{RCO}_2\text{H}$ + $\text{H}_2$ rolysis of nitriles: $\text{RCN}$ (+ $\text{H}_3\text{O}^+$ + $\text{H}_2\text{O}$ ) $\longrightarrow$ $\text{RCO}_2\text{H}$ tion of benzene (or any arene): $\text{C}_6\text{H}_6$ + $\text{HNO}_3$ $\longrightarrow$ $\text{C}_3$ (dration of alcohols, e.g. : $\text{CH}_3\text{CH}(\text{OH})\text{CH}_3$ $\longrightarrow$ $\text{CH}_3$ he reverse) genation of ketones, e.g. : $\text{CH}_3\text{COCH}_3$ + $\text{X}_2$ $\longrightarrow$ $\text{CH}_3$	$NH_4^+$ + $NH_4^+$ ${}_6H_5NO_2 (+ H_2O)$ CH=CH <sub>2</sub> + H <sub>2</sub> O	[2]
3	(a)	(i)		D <sub>3</sub> + H₂SO₄ c (both acids) <b>and</b> 30°C < T < 60°C <i>or</i> warm		<b>[Total: 17]</b> [1] [1]
		(ii)		e HNO₃ <i>or</i> HNO₃(aq) room temp. (allow T ≤ 30ºC)		[1] <b>[3]</b>
	(b)			termediate from methylbenzene)		[1] <b>[1]</b>

(c) Sn/tin (or SnC $l_2$ , Fe) + HCl (NOT H<sub>2</sub>SO<sub>4</sub> or H<sup>+</sup>, Zn, or LiAlH<sub>4</sub>.) [1] [1]

Page 6	Mark Scheme	Syllabus	Paper
	GCE A LEVEL – May/June 2013	9701	42

(d) (i)



- (ii)  $NaNO_2 + HCl or H_2SO_4 or H^+ or HNO_2$  [1]  $T \le 10^{\circ}C$  [1] [4 max 3]
- (e) (i) amide
  - (ii)  $M_r = 108 + 11 + 14 + 16 = 149$

%N = (14 x 100)/149 = 9.4%

(iii) NHCOC<sub>2</sub>H<sub>5</sub>

[1] **[3]** 

[1]

[1]

- [Total: 11]
- (a) (i) Many electrons of similar energy in a valence-shell orbital or successive ionisation energies rise steadily (no big jumps) or ability to form bonds with ligands can stabilise very low or very high oxidation states or 4s + 3d orbitals/shells/energy levels have similar / same energies

(ii) 
$$VO_2^+: +5$$
  
 $CrF_6^{2-}: +4$   
 $MnO_4^{2-}: +6$ 
[3 × 1]  
[4]

PMT

Page 7	Mark Scheme	Syllabus	Paper
	GCE A LEVEL – May/June 2013	9701	42

#### (b)

• (colour due to) absorption of light/photons/frequencies/wavelengths or

colour seen is complement of colour absorbed.

- d-orbitals/d-subshell split (by ligand field)
- (when photon is absorbed), electron is promoted *or* moves (from lower) to higher (d–)orbital
- energy difference/gap or  $\Delta E$  or splitting corresponds to photon/frequency/wavelength in visible region
- in s-block elements the energy gap is too large (to be able to absorb visible light)

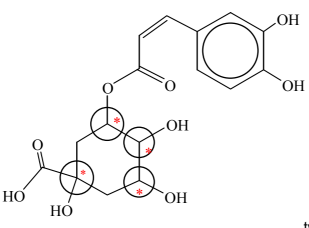
[any four 4 × 1] [4]

(c) (i)	$2MnO_4^{-} + 2H_2O + 5SO_2 \longrightarrow 2Mn^{2+} + 5SO_4^{2-} + 4H^+$	[1]
(ii)	solution will go from purple	[1]
	to colourless	[1] <b>[3]</b>
<b>(d)</b> (pa	le) blue solution	[1]
giv	es a (pale) blue ppt.	[1]
wh	ich re-dissolves, <i>or</i> forms a solution, which is dark/deep blue <i>or</i> purple	[1] <b>[3]</b>

## [Total: 14]

Page 8	Mark Scheme	Syllabus	Paper
	GCE A LEVEL – May/June 2013	9701	42

5 (a) (i)



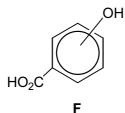
two or three centres correctly identified [1]

- four centres correctly identified [2]
- (ii) C<sub>16</sub>H<sub>18</sub>O<sub>9</sub>
   [1]

   (iii) 3 moles of H<sub>2</sub>
   [1]

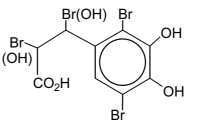
   (iv) in cold: 3 moles of NaOH
   [1]
  - ,
  - on heating: 4 moles of NaOH [1]
- (b) (i) hydrolysis
  - (ii) alkene or C=C [1]
  - (iii) with Na2CO3(aq): carboxylic acid[1]with Br2(aq): phenol[1]

(iv)



(OH can be at the 3, 4, or 5 positions, but not the 2 or 6 positions) [1]

(v) geometrical or cis-trans or E-Z

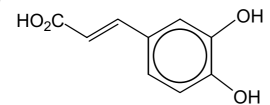


- G(ring subst. allow 2 or 3 Br in ring)[1](addition to C=C: allow one of the aliphatic Br<br/>to be OH, but not both)[1]
  - [1]

[1]

Page 9	Mark Scheme	Syllabus	Paper
	GCE A LEVEL – May/June 2013	9701	42

(vi)



skeletal or structural [1] [9 max 8]

(c)  $M_r(\mathbf{E}) = 180$ , so 0.1 g = 1/1800 (5.56 x 10<sup>-4</sup>) mol

3 mol NaOH react with 1 mol of **E**, so  $n(NaOH) = 3/1800 = 1/600 \text{ mol} = 1.67 \times 10^{-3} \text{ mol}$  [1]

volume of 0.1M NaOH = 1000/(600 x 0.1) = 16.7 cm<sup>3</sup>

[1] **[3]** 

[1]

[Total: 17]

Page 10	Mark Scheme	Syllabus	Paper
	GCE A LEVEL – May/June 2013	9701	42

6 (a)

Γ		
substance	protein synthesis	formation of DNA
cysteine	$\checkmark$	
cytosine		$\checkmark$
glutamine	$\checkmark$	
guanine		$\checkmark$

		[3] <b>[3]</b>
(b) (i)	Hydrogen bonding	[1]
	Between bases or between A,T, C and G (all four needed)	[1]
(ii)	Bonds are (relatively) weak or easily broken	[1]
	This enables strands to separate or DNA to unzip/unwind/unravel.	[1] <b>[4]</b>

### (c) changes / mutations in DNA

• by the addition / insertion /deletion / substitution / replacement of a base

• adds / deletes / replaces an amino acid or changes the amino acid sequence

• this causes a loss of function or changes the shape / tertiary structure of the protein

any three points [3]

[3]

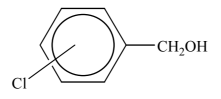
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Page 11	Mark Scheme	Syllabus	Paper
	GCE A LEVEL – May/June 2013	9701	42

7. (a) (i)  $\frac{43.3}{3.35} = \frac{100}{1.1 \text{ x n}}$ 

n = 
$$\frac{100 \times 3.35}{43.3 \times 1.1}$$
 = 7.03 = 7 (calculation must be shown) [1]

- (ii) The M and M+2 peaks are in the ratio 3 : 1 hence the halogen is chlorine/Cl [1]
- (iii) L contains 7 hydrogen atoms *or* there are 3 types/environments of proton/H [1]
- (iv) The multiplet with 4 hydrogens *or* peaks at δ 7.3 suggests a benzene ring The singlet with 2 hydrogens *or* peak at δ 4.7 suggests a –CH<sub>2</sub>– group The singlet with 1 hydrogen *or* peak at δ 2.3 suggests an –OH group *or* reaction with Na suggests an OH group OH must be an alcohol, not a phenol (due to its δ value) Since L also contains 7 carbon atoms and chlorine, this accounts for 126 of the 142 mass, the remaining atom must be oxygen Thus L is



(allow the 2-, 3- or 4- isomer)

[6] **[9 max 7]** 

- (b) (i) we expect propene to have a CH<sub>3</sub> peak *or* a peak at m/e 15 *or* cyclopropane would have fewer peaks
  - (ii) cyclopropane would have 1 peak (ignore splitting) propene would have 2 (*or* 3, *or* 4) peaks (ignore splitting) *or* propene would have peaks in the  $\delta$  4.5-6.0 (alkene) region no splitting of cyclopropane peak (any two points)

[2] **[3]** 

[1]

[Total: 10]

Page 12	2	Mark Scheme	Syllabus	Paper
		GCE A LEVEL – May/June 2013	9701	42
8 (a) (i)	(a) (i) $CH_2 = CH - CO_2H$ or $CH_2 = CH - CO_2R$ or $CH_2 = CH - COCl$			
(ii)	addi	tion (polymerisation)		[1]
(iii)	C(CI	H <sub>2</sub> OH) <sub>4</sub>		[1]
(iv)	wate	r		[1] <b>[5]</b>
(b) (water is bonded to the polymer by) hydrogen bonding hydrogen bonds are weak or easily broken				[1] [1] <b>[2]</b>
(c) (i)		s-linking causes no reduction in the number of –OH gr oss-linking molecules also have –OH groups	oups	[1]
(ii)	<ul> <li>(ii) property e.g. becomes harder / more rigid / less flexible / stronger / higher melting point.</li> <li>because the chains are more strongly / tightly held</li> </ul>			melting [1] [1] <b>[3]</b>
	[Total:			