MARK SCHEME for the October/November 2010 question paper

for the guidance of teachers

9701 CHEMISTRY

9701/41

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

This mark scheme is published as an aid to teachers and candidates, to indicate the requirements of the examination. It shows the basis on which Examiners were instructed to award marks. It does not indicate the details of the discussions that took place at an Examiners' meeting before marking began, which would have considered the acceptability of alternative answers.

Mark schemes must be read in conjunction with the question papers and the report on the examination.

• CIE will not enter into discussions or correspondence in connection with these mark schemes.

CIE is publishing the mark schemes for the October/November 2010 question papers for most IGCSE, GCE Advanced Level and Advanced Subsidiary Level syllabuses and some Ordinary Level syllabuses.



| | Page 2 | | k Scheme: Teachers' version | Syllabus | Paper |
|---|-------------------|---|---|----------|-------------|
| | | GCE A | LEVEL – October/November 2010 | 9701 | 41 |
| 1 | (a) PC <i>l</i> 5 | + $4H_2O \rightarrow H_3PO$ | D ₄ + 5HC <i>l</i> (1) | | |
| | SiCl | $_{1} + 2H_{2}O \rightarrow SiO_{2}$ | + 4HC l (or giving H ₂ SiO ₃ , Si(OH) ₄ etc.) (7 | 1) | [2] |
| | (b) bonc | C <i>l</i> -C | = 264 kJ mol ⁻¹ $l = 244 kJ mol^{-1}$ = 250 kJ mol ⁻¹ | | |
| | ΔH : | = 8 × 264 + 8 × 2 | 44 – 16 × 250 = +64 kJ mol ⁻¹ (2) | | [2] |
| | (c) (i) | +2 (1) | | | |
| | | (half) the sulfur go (the other half) go | bes up by +2, (1) bes down by –2 (1) | | |
| | (iii) | HC <i>l</i> (can be read | into (iv)) (1) | | |
| | (iv) | $2SCl_2 + 2H_2O \rightarrow$ | $S + SO_2 + 4HCl(1)$ | | |
| | (v) | (+ AgNO ₃) w (+ K ₂ Cr ₂ O ₇) so | hite ppt. (1) olution turns green (1) | | [7] |
| | | | | | [Total: 11] |
| | | | | | |

2 (a) (i) A ligand is a species that contains a <u>lone pair of electrons</u>, *or* that can form a <u>dative bond</u> (to a transition element) (1)

(ii)

| species | can be a ligand | cannot be a ligand |
|---------------------------------|-----------------|--------------------|
| OH⁻ | \checkmark | |
| NH_4^+ | | \checkmark |
| CH₃OH | \checkmark | |
| CH ₃ NH ₂ | \checkmark | |

 $(4 \times \frac{1}{2})$ [3]

- (b) (i) C is $[Cu(NH_3)_6]^{2+} SO_4^{2-}$ (allow $[Cu(NH_3)_4]^{2+} SO_4^{2-}$ (1) D is CuO (1) E is Na₂SO₄ (1) F is BaSO₄ (1)
 - (ii) acid-base or neutralisation (1)
- (c) (i) any two from: brown fumes or vapour evolved / gas relights glowing splint / black solid formed (2)
 - (ii) $2Cu(NO_3)_2 \rightarrow 2CuO + 4NO_2 + O_2(1)$ [3]

[Total: 11 max 10]

[5]

| | Page 3 | 3 | | | | | | me: T | | | | | | | S | Syllal | | | Paper | |
|---|---------------------|----------------------------|-----------------------|---------------------|-----------------|------------------|--------|------------------|-------|--------------------|-------------------------------|--------------------|--------------------|-----------------|---------------|------------------|---------|---------|----------|--------------------|
| | | | | G | ice A | LE | /EL - | – Oct | tobeı | r/Nov | vemb | ber 2 | 010 | | | 970 | 1 | | 41 | |
| 3 | (a) (i) | Cu(s | s) – 2 | 2e | $ \rightarrow$ | Cu ²⁺ | (aq) | allov | w ele | ctror | ns on | RHS | 6 (1) | | | | | | | |
| | (ii) | | - | - | - | | | hich i I (owt | | • | ositiv | ve tha | an +0. | .34V | for | Cu ²⁺ | /Cu, (′ | 1) | | |
| | (iii) | E ^e fo Ni is | | | | | `` | goes | into | solu | tion a | as Ni ² | ²⁺ (aq) | (1) | [] | Mark | (ii) an | d (iii) | to ma | ıx 3] |
| | (iv) | Cu ²⁺ | ⁺(aq) | 1) + | 2e- | → C | u(s) (| (1) | | | | | | | | | | | | |
| | (v) | E ^e fo | or Zr | '.n ²⁺ / | ′Zn is | nega | ative | / = _ | -0.76 | V, so | o Zn ^{2[·]} | + is n | ot ea | sily r | edu | iced. | (1) | | | |
| | (vi) | | | | coloui ease: | | es be | ecau | se C | u ²⁺ (a | ıq) is | s beir | ng re | place | ed I | oy Zi | n²⁺(aq |) or 1 | Ni²⁺(aq |) or [7] |
| | (b) am am | ount o | | | | | | | • | , | • • • | | 087) | mol (| 1) | | | | | |
| | | of co of mo | | | | | | | | | | | .46 m | ol (1 |) | | | | | |
| | per | centa | ige " | "wa | isted' | ' = 1 | 100 × | (7.40 | 61 – | 7.08 | 7)/7.4 | 461 : | = 5.0 |)1 (5 . | . 0)% | % (ac | cept 4 | .98–5 | 5.10) (′ | 1) [4] |
| | (c) E ^e | | | | = . e = . | | |) | | | | | | | | | | | | |
| | Be | cause | e the | e Fe | e pote | ential | is m | ore n | egati | ive th | nan th | he Ni | pote | ntial, | the | iron | will di | issolv | e (1) | [2] |
| | | | | | | | | | | | | | | | | | | [| Total: | 13] |
| 4 | (a) (i) | SnC |) ₂ | (| Can b | e rea | ad int | to equ | uatio | n (1) | | | | | | | | | | |

- 4 (a) (i) SnO₂ Can be read into equation (1) 2NaOH + SnO₂ \rightarrow Na₂SnO₃ + H₂O (1)
 - (ii) PbO Can be read into equation (1) PbO + 2HC $l \rightarrow$ PbC l_2 + H₂O (1)
 - (b) moles of oxygen = 9.3/16 = 0.581 molmoles of lead = 90.7/207 = 0.438 mol (both 3 s.f.) (1)so formula is Pb₃O₄ (1)
 - (c) (i) $K_{sp} = [Pb^{2^+}][Cl^-]^2(1)$ units = mol³ dm⁻⁹(1)
 - (ii) if $[Pb^{2^+}] = x$, $K_{sp} = 4x^3$, so $x = {}^3\sqrt{\{K_{sp}/4\}}$ $[Pb^{2^+}] = {}^3\sqrt{\{2 \times 10^{-5}/4\}} = 1.71 \times 10^{-2} \text{ mol dm}^{-3} (1)$
 - (iii) $[Pb^{2+}] = 2 \times 10^{-5} / (0.5)^2 = 8.0 \times 10^{-5} \text{ mol dm}^{-3} (1)$
 - (iv) common ion effect, or increased $[Cl^{-}]$ forces solubility equilibrium over to the left (1)

[Max 4]

[4]

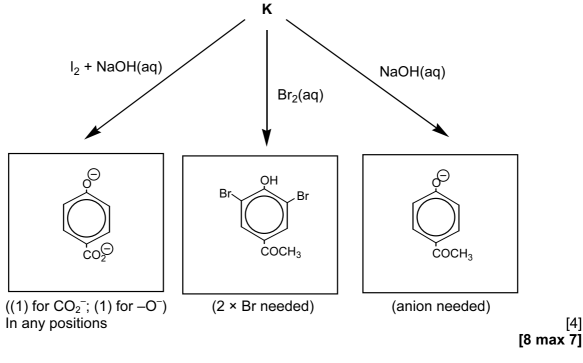
[2]

[Total: 10]

| Page 4 | Mark Scheme: Teachers' version | Syllabus | Paper | |
|--------|-------------------------------------|----------|-------|--|
| | GCE A LEVEL – October/November 2010 | 9701 | 41 | |

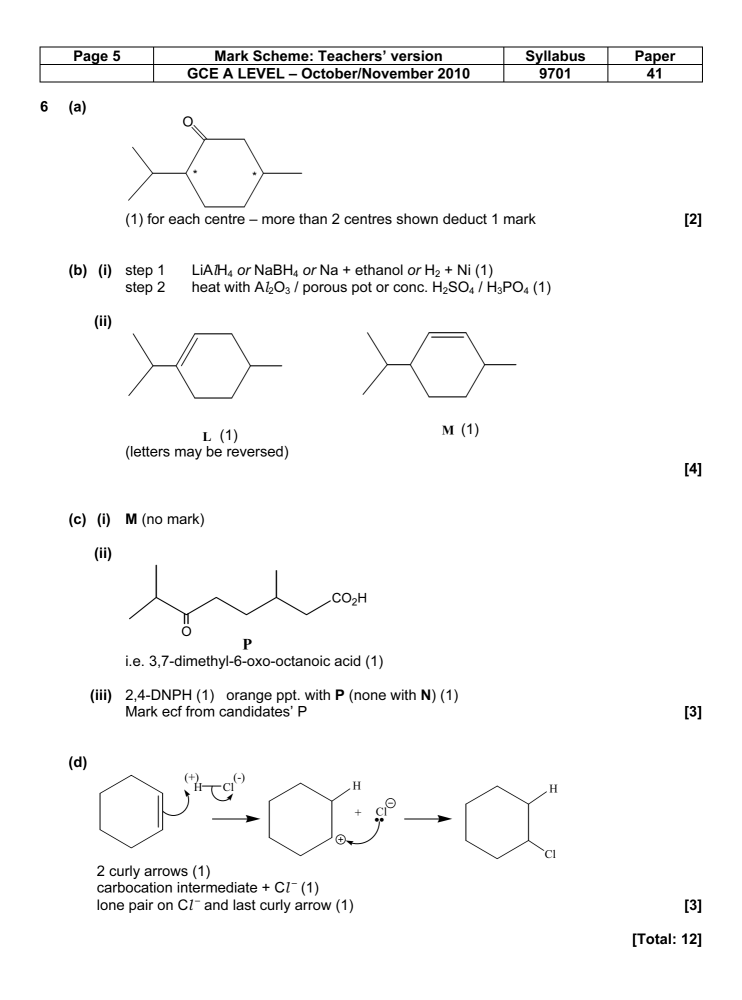
- **5 (a) (i)** ester (1)
 - (ii) H is nitrobenzene structure needed here (1)J is phenyldiazonium chloride structure needed here (1)
 - (iii) step 2 Sn/Zn + HC l/H_2 + named cat / NaBH₄ / LiA lH_4 / Na + ethanol (1) step 3 HNO₂/NaNO₂ + HCl at T = 10°C or less (1) step 4 heat/warm to T > 10°C (1) step 5 CH₃COCl/ CH₃COCOCOCH₃ (1)
 - (b) (i) compounds that have the same molecular formula, but different structures (1)
 - (ii) phenol (NOT hydroxy) (1) (methyl) ketone *or* carbonyl (1)
 - (iii) K is 4-ethanoylphenol, $HO-C_6H_4$ -COCH₃ (must be 1,4- disubstituted isomer) (1)







[7]



| | Page 6 | 6 | Mark Scheme: Teachers' version GCE A LEVEL – October/November 2010 | Syllabus 9701 | Paper 41 | |
|---|-------------|--------------------------|---|---|---------------------|--|
| 7 | (a) (i) | Disu | Ilfide bond / group / bridge (1) | | | |
| • | (ii) | | tertiary structure (1) | | | |
| | (iii) | The | substrate will no longer bond to / fit into the active site hape of active site is changed | (1) | [3] | |
| | (b) (i) | Acid | l-base / proton donor / neutralisation / salt formation (1 |) | | |
| | (ii) | The | ability of the $-CO_2H$ group to form hydrogen bonds (1) | and ionic intera | ctions (1) | |
| | | The | $-CO_2H/-CO_2^-$ group is no longer able to interact with - | -NH ₂ /-NH ₃ ⁺ (1) | | |
| | | The | Ag ⁺ forms a strong bond with $-COO^-(1)$ | | [5] max [4] | |
| | (c) (i) | 8 bu | t allow $4O_2$ if specified as molecules (1) | | | |
| | (ii) | Dati | ve / co-ordinate (1) | | | |
| | (iii) | Octa | ahedral / 6 co-ordinate (1) | | [3] | |
| | | | | | [Total: 10] | |
| 8 | Ele in 〉 | NMR, ectron: X-ray | energy is absorbed due to the two spin states (1) s (1) crystallography, X-rays are diffracted (by regions of hig | h electron densi | ity) (1) [4] | |
| | (b) (i) | The Alco | no mark spectrum of alcohol / Y contains different peaks hol / Y contains different chemical environments ctrum 2 contains only one peak (1) | | | |
| | (ii) | Spe | ctrum 2 only shows 1 peak so Z must be a ketone (1) | | | |
| | | Hen | ce Y must be a 2° alcohol (1) | | | |
| | | Num | hber of carbon atoms present $=\frac{0.6 \times 100}{17.6 \times 1.1} = 3$ (1) | | | |
| | | Thu | s Z must be CH_3COCH_3 (1) | | | |
| | | Hen | ce Y must be propan-2-ol, $CH_3CH(OH)CH_3$ (1) | | | |
| | (iii) | Y is | $ \begin{array}{c} H\\ H\\ CH_3 - C - CH_3\\ H\\ OH \end{array} $ (1) | | | |
| | (iv) | | f the protons in Z are in the same chemical environme | nt (1) | [8] max [7] | |
| | | | | | [Total: 11] | |
| | | | | | | |

| Page 7 | Mark Scheme: Teachers' version | Syllabus | Paper | |
|--------|-------------------------------------|----------|-------|--|
| | GCE A LEVEL – October/November 2010 | 9701 | 41 | |
| | | | | |

- **9** (a) (i) A few nanometres (accept 0.5–10 nm) (1)
 - (ii) Graphite/graphene (1)
 - (iii) van der Waals' (1)
 Carbon atoms in the nanotubes are joined by covalent bonds (1)
 (as are the hydrogen atoms in a hydrogen molecule)
 or no dipoles on C or H₂ or the substances are non-polar
 [4]
 - (b) More hydrogen can be packed into the same space/volume (1) [1]
 - (c) If a system at equilibrium is disturbed, the equilibrium moves in the direction which tends to reduce the disturbance (owtte) (1)

When H_2 is removed the pressure drops and more H_2 is released from that adsorbed (1)

The equilibrium $H_{2adsorbed} \rightleftharpoons H_{2gaseous}(1)$

Equilibrium shifts to the right as pressure drops (1)

[4]

[Total: 9]