CAMBRIDGE INTERNATIONAL EXAMINATIONS

Cambridge International Advanced Level

MARK SCHEME for the May/June 2015 series

9701 CHEMISTRY

9701/42

Paper 4 (Structured Questions), maximum raw mark 100

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1 (a) fluorine: $1s^22s^22p^5$ [1]

sulfur: 1s²2s²2p⁶3s²3p⁴

(b) (i)
$$2HCl \longrightarrow H_2 + Cl_2$$
 [1]

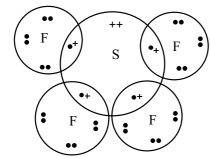
(ii) bond energies: HF (562) is **stronger** than HC
$$l$$
 (431) or F₂ (158) is **weaker** than C l_2 (244)

(c) electronegativity: [2]

The attraction by an atom/nucleus/element of the electrons in a bond *or* a shared pair *or* a molecule *bond polarity:*

..is due to atoms/elements of different electronegativities at each end of a bond

(d) (i)



(ii) Yes, it will have a dipole moment, either because it has an uneven distribution of electrons or because it contains a lone pair

or the S–F dipoles don't cancel or molecule is not symmetrical or diagram of see-saw shape.

(allow an ecf for "no dipole" if their structure in (d)(i) has **no** lone pair)

- (e) Sulfur can use its d-orbitals or has low-lying/accessible/available d-orbitals or can expand its octet.
 (allow reverse argument for oxygen; do NOT allow just "sulfur has d-orbitals")
- **(f) (i)** Burning of **fossil** fuels *or* coal/oil/petrol/natural gas (NOT methane *or* hydrocarbons) *or* volcanoes *or* roasting/burning sulfide ores

(ii) Acid rain [2]

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2 (a)
$$A_r = 204 \times 0.019 + 206 \times 0.248 + 207$$
 [2] = 207.21 (correct ans = [2])

The **last** answer written by the candidate needs to be written with 2 d.p. to get the last mark.

(b) (i)
$$Tin(II)$$
 oxide is more basic than $tin(IV)$ oxide or $tin(II)$ oxide is less acidic than $tin(IV)$ oxide

(ii) e.g. SnO + 2HC
$$l$$
 \longrightarrow SnC l_2 + H2O(or ionic or with H₂SO₄) [2]
SnO₂ + 2NaOH \longrightarrow Na₂SnO₃ + H₂O (or ionic or with KOH etc.)

PbO₂ changes colour (from brown/black to yellow/orange/red)

$$PbO_2 \longrightarrow PbO + \frac{1}{2} O_2$$
 or $3PbO_2 \longrightarrow Pb_3O_4 + O_2$

[Total: 8]

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(b) Solubility decreases (from Mg to Ba *or* down the group) [4]

Both lattice energy/ ΔH_{latt} and enthalpy change of hydration/ ΔH_{hyd} are involved enthalpy change of hydration **decreases more** than lattice energy

So enthalpy change of solution $/\Delta H_{sol}$ becomes more endothermic *or* more positive *or* less exothermic *or* less negative (NOT ΔH_{sol} decreases, or increases)

(c) precipitate/solid CaSO₄ would form due to the **common ion effect** or K_{sp} is exceeded or the following equilibrium shifted over to the right $Ca^{2+(aq)} + SO4^{2-}(aq) \rightleftharpoons CaSO_4(s)$ [2]

(d) charge passed =
$$1.8 \times 40 \times 60$$
 (= 4320 C) [4]
 $n(e^{-})$ = $4320/96500$ (= $4.477 \times 10^{-2} \text{ mol}$) ecf
 $n(Cr)$ = $0.776/52$ (= $1.492 \times 10^{-2} \text{ mol}$) ecf
 n = $4.477 \times 10^{-2}/1.492 \times 10^{-2} = 3.00$ (=3)

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- **4 (a) (i)** a solution that resists/minimises a change in its pH *or* **helps** maintain its pH..... [2] (NOT any of: "maintains pH"; "keeps pH constant"; "no change in pH")when small amounts of acid/H⁺ or base/OH⁻ are added (**both** acid and base are needed)
 - (ii) HCO_3^- reacts with H^+ ions as follows: $HCO_3^- + H^+ \longrightarrow H_2CO_3 (or H_2O + CO_2)$ and with OH^- ions thus: $HCO_3^- + OH^- \longrightarrow CO_3^{2-} + H_2O$

(the equation arrows can be equilibrium arrows, as long as HCO₃⁻ is on the left)

(iii)
$$(pK_a = -log(K_a) = 7.21)$$
 [2]
 $pH = pK_a + log([base]/[acid] = 7.21 + log(0.5/0.3)$
 $= 7.43 (7.4)$

- (b) (i) $K_{sp} = [Ag^{+}]^{3}[PO_{4}^{3-}]$ and units: $mol^{4}dm^{-12}$ [1]
 - (ii) call $[PO_4^{3-}] = x$, then $[Ag^+] = 3x$, and $K_{sp} = 27x^4$ [3] $x = (K_{sp}/27)^{1/4} = (1.25 \times 10^{-20}/27)^{1/4} = 4.64 \times 10^{-6} \text{ mol dm}^{-3}$ $[Ag^+] = 3x = 1.39 \times 10^{-5} \text{ (mol dm}^{-3}\text{)} \qquad \text{(allow } 1.4 \times 10^{-5}\text{)}$

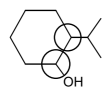
(c)
$$H_3PO_3 + 2Fe^{3+} + H_2O \longrightarrow H_3PO_4 + 2Fe^{2+} + 2H^+$$
 [2]
 $E\Theta_{cell} = 0.77 - (-0.28) = (+)1.05 \text{ V}$
or $3H_3PO_3 + 3H_2O + 2Fe^{3+} \longrightarrow 3H_3PO_4 + 6H^+ + 2Fe$
 $E\Theta_{cell} = -0.04 - (-0.28) = (+)0.24 \text{ V}$

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5 (a) (i)
$$H_2 + Pt \ or \ H_2 + Ni/Pd + heat/warm \ or \ 50^\circ < T < 500^\circ C$$
 [1]

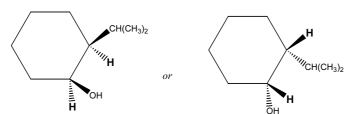
(ii)



[1]

(iii)
$$2^2 = 4$$
 [1]

(iv)



2 Hs have to be on the **same side** of the ring. Allow $-C_3H_7$ or -R for $-CH(CH_3)_2$ [1]

(b) (i) $C \qquad \qquad Or \qquad \bigvee_{N_2^+} Or \qquad \bigvee_{N^+}$

(ii) step 1: conc HNO₃ + H₂SO₄ (@ 25 °C < T < 60 °C – see below) ("aq" negates) step 2: Sn/Fe + HC*l* step 3: HNO₂ or NaNO₂ + HC*l* (@ T< 10 °C – see below) both temperatures correct for steps 1 + 3 (temperature not required for step 2) (inclusion of the word "heat" or "reflux" in step 3 negates the temperature mark)

(c) [5]

| HBr | no reaction | Br |
|----------|-------------|-------------|
| Na | ONa | ONa |
| NaOH(aq) | ONa | no reaction |

[Total: 14]

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6 (a) There are three acceptable alternatives – follow each column down vertically:

| (i) D is | RCOC1 | RCOOCH₂CH₃ | RCO ₂ ⁻ NH ₄ ⁺ |
|-----------------|--|--|--|
| | | | |
| (ii) step 1 | $SOCl_2$ (or PCl_3 or PCl_5) | ethanol (e.g.) + conc H ₂ SO ₄ | NH ₃ |
| (ii) step 2 | NH ₃ (NaOH negates th | nis mark) | heat |
| (ii) step 3 | LiAlH ₄ (aq) negates(NOT NaBH ₄ ; Sn + HCl etc.) | | |

(b) (i) amine (other groups negate)

[1]

(ii) phenol and carboxylic acid (both needed)

[1]

(iii)

[4]

| compound | first functional group | second functional group |
|----------|------------------------|-------------------------|
| Е | amide | alcohol |
| F | amine | carboxylic acid |
| G | amine | ester |
| Н | amide | phenol |

- (iv) Mark this in the following way. For each structure of E, F, G and H: [4]
 - check whether the structure fits the molecular formula C₈H₉NO₂, i.e. that it has: **one** nitrogen, **two** oxygens and **eight** carbons.
 - check that it contains the two groups that the candidate's answers to part (ii) says it contains.

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| (a) | | it is the only compound that is an amino acid <i>or</i> can form (NOT <i>con</i> H–CO– / amide / peptide linkages / bonds | tain) | [1] |
| | it c | contains an N atom/NH ₂ group/CO ₂ H group | | |
| | | | | |
| (b) | M1 M2 M3 M4 M5 | codons (base triplets) codes for one amino acid mRNA binds to/associates with the ribosome tRNAs are specific to their amino acids tRNA contains an anticodon <i>or</i> bonds to the codon/mRNA throutranslates the RNA code into the amino acid sequence | | [4] airing <i>or</i> |
| (c) | M1 M2 M3 M4 M5 M6 | fewer hydrogen bonds <i>or</i> more van der Waals' (id–id) forces fewer ionic bonds form the tertiary structure/folding/(3D) shape (of the protein) would c | hange | [3] |

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| 8 | (a) (i) | The nucleus/proton of a hydrogen atom has spin | | [1] |
| | (ii) | Hydrogen doesn't have enough electrons/electron density | | [1] |
| | (iii) | S/sulfur – it has the greatest number of electrons or highest electrons | on density | [1] |
| | (b) (i) | 12 protons (=9+2+1) | | [1] |
| | (ii) | The group responsible for this peak is –OH (allow NH) The D in D ₂ O exchanges with the H in –OH or H is replaced by D or "–OH \rightarrow –OD", | | [2] |
| | (iii) | The adjacent carbon atom has no hydrogen atoms bonded to it | | [1] |
| | (iv) | Methyl/CH ₃ group | | [1] |
| | (v) | P is (CH ₃) ₃ C–CH ₂ OH | | [1] |
| | (c) (i) | $n = \frac{100 \times (M+1)}{1.1 \times M} = \frac{100 \times 0.5}{1.1 \times 9.3} = 50/10.23$ $= 4.89 \text{ hence } 5 \text{ carbons}$ | | [1] |
| | (ii) | (Ratio of ⁷⁹ Br: ⁸¹ Br is 1 : 1), hence ratio of M : M+2 : M+4 is 1 : 2 : 1 | | [1] |
| | (iii) | Molecular formula of ${f R}$ is $C_5H_{10}Br_2$ | | [1] |

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9 (a)_______ [3]

| monomer | addition | condensation | both |
|--------------------|----------|--------------|------|
| H H OH | | ✓ | |
| HC=CH | √ | | |
| H ₃ C H | ✓ | | |

(b) polythene is non-polar or its bonds are non-polar so not (easily) hydrolysed

(Allow displayed, skeletal, part-skeletal, structural etc.)

(ii) The ester (or –COO–) linkage/bond is hydrolysed *or* reacts with water [1]

(d) Polythene has (weak) van der Waals' (or id-id) forcesPVC has stronger van der Waals' forces or additional dipole forcesNylon has (strong) hydrogen bonding

[Total: 10]