

MARK SCHEME for the May/June 2012 question paper
for the guidance of teachers

9701 CHEMISTRY

9701/41

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

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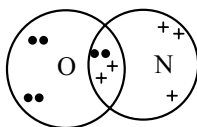


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- 1 (a) (i) the enthalpy change/released when **1 mole is formed** [1]
of ionic lattice **from the gas phase ions** [1]
(ii) $\text{Mg}^{2+} + \text{O}^{2-} \longrightarrow \text{MgO}$ [1]
[3]
- (b) measurements needed:
volume/mass/weight of water (in calorimeter) [1]
initial + final temperature/temperature change/temperature rise (of the water) [1]
mass of Mg (used)/mass MgO [1]
Not volume/moles/mass of oxygen used [3]
- (c) $\Delta H = 148 + 736 + 1450 + 496/2 - 141 + 798 - 3791$
= **-552 kJ mol⁻¹** [3]
[3]
- (d) $\text{Na}_2\text{O(s)} + \text{H}_2\text{O(aq/l)} \longrightarrow 2\text{NaOH(aq)}$ [1]
 $\text{MgO(s)} + \text{H}_2\text{O(aq/l)} \longrightarrow \text{Mg(OH)}_2\text{(s) or Mg(OH)}_2\text{(aq)}$ [1]
pH 12.5-14 [NaOH] **AND** 8-10.5 [Mg(OH)₂] respectively [1]
[3]

[Total: 12]

2. (a) (i)



- [1]
- (ii) -180 kJ mol^{-1} [1]
- (iii) (formation of NO is endothermic) so high T **and** equilibrium pushed over to NO side.
or high T **and** needed to break N-N bond in N₂ [1]
- (iv) $-180 = 2 E(\text{NO}) - 994 - 496$ [1]
 $E(\text{NO}) = \mathbf{+655 \text{ kJ mol}^{-1}}$ [1]
[5]
- (b) (i) (from 1 and 2:) as p(NO) halves, rate decreases to 1/4, **so order = 2** [1]
(from 1 and 3:) as p(H₂) halves, so does rate, **so order = 1** [1]
- (ii) $\text{rate} = k p_{\text{NO}}^2 \cdot p_{\text{H}_2}$ [1]
units (of k) are $\text{atm}^{-2} \text{ s}^{-1}$ [1]

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- (iii) add all three equations:
 $\text{NO} + \text{NO} + \text{H}_2 + \text{O} + \text{H}_2 + \text{N}_2\text{O} \rightarrow \text{N}_2\text{O} + \text{O} + \text{H}_2\text{O} + \text{N}_2 + \text{H}_2\text{O}$ [1]
 cross out all species common to both sides:
 $\text{NO} + \text{NO} + \text{H}_2 + \text{O} + \text{H}_2 + \text{N}_2\text{O} \rightarrow \text{N}_2\text{O} + \text{O} + \text{H}_2\text{O} + \text{N}_2 + \text{H}_2\text{O}$ [1]
 $(\Rightarrow 2\text{NO} + 2\text{H}_2 \rightarrow \text{N}_2 + 2\text{H}_2\text{O})$

- (iv) *either: step 2* since it involves H_2 [1]
 O formed from NO [1]
or: step 3 since it involves H_2 [1]
 N_2O formed from NO [1]
[8]

- (c) (i) NO [1]

- (ii) $3\text{Fe}^{2+} + 4\text{H}^+ + \text{NO}_3^- \longrightarrow 3\text{Fe}^{3+} + \text{NO} + 2\text{H}_2\text{O}$ [1]
 (allow $\text{Fe}^{2+} + \text{H}^+ + \text{HNO}_2 \longrightarrow \text{Fe}^{3+} + \text{NO} + \text{H}_2\text{O}$)

- (iii) dative/coordinate bonding [1]

- (iv) $[\text{Fe}(\text{H}_2\text{O})_{6-n}(\text{NO})_n]^{2+}$ ($n = 1-6$) [1]
[4]

[Total:17]

3. (a) (i) $\text{C}_{16}\text{H}_{10}\text{N}_2\text{O}_2$ [1]
 (ii) ketone, alkene, amine, aryl (benzene/arene/phenyl) (any 3) [2]
[3]

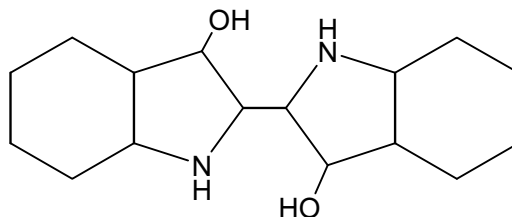
- (b) (i) reduction or redox [1]

- (ii) NaBH_4 or LiAlH_4 (**NOT** H_2 + Ni) [1]
[2]

- (c) 1. 2,4-DNPH [1] red/yellow-orange/orange ppt. [1] no reaction
 2. Na metal [1] no reaction gas given off/fizzing [1]
or $\text{PCl}_5/\text{SOCl}_2$ [1] no reaction steamy fumes/fizzing [1]
or PCl_3 + warm misty/white fumes
 2 x "no reaction" must be linked to "correct reagent" [1]
[5]

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(d) (i)



[1]

(ii) $M_r = 262$, so $2.5 \text{ g} = 2.5/262 = 9.54 \times 10^{-3} \text{ mol}$

[1]

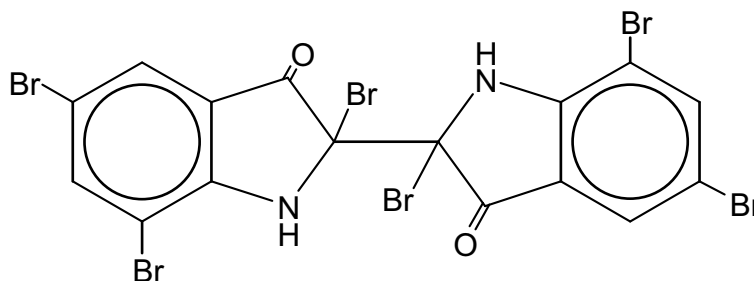
(1 mol indigo absorbs 9 mol of H_2)

so volume of $\text{H}_2 = 9 \times 24 - 9.54 \times 10^{-3} = \mathbf{2.06 \text{ dm}^3}$ (2060 cm^3)

[1]

[3]

(e)



2 x Br on **C=C** [1]

a Br on each ring [1]

TWO non-adjacent Br on **each ring** [1]

[3]

[Total: 16]

4 (a) (i) volatilities decrease down the group

[1]

due to greater van der Waals (VDW) forces (*intermolecular is not sufficient*)

[1]

due to larger no of electrons

[1]

(ii) CCl_4 does not react with water

[1]

CCl_4 unreactive due to no **d**-orbitals

[1]

GeCl_4 **and** PbCl_4 hydrolyse/react

[1]

$\text{MCl}_4 + 2\text{H}_2\text{O} \longrightarrow \text{MO}_2 + 4\text{HCl}$ (M = Ge or Pb)

[1]

[7]

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(b) (i) B is PbSO_4 and C is PbCl_2 [1]

(ii) $\text{SnO}_2 + 2\text{H}_2\text{SO}_4 \longrightarrow \text{Sn}(\text{SO}_4)_2 + 2\text{H}_2\text{O}$ [1]

$\text{PbO}_2 + \text{H}_2\text{SO}_4 \longrightarrow \text{PbSO}_4 + \text{H}_2\text{O} + \frac{1}{2} \text{O}_2$ [1]

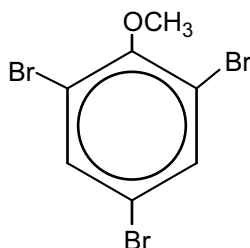
$\text{PbO}_2 + 6\text{HCl} \longrightarrow \text{H}_2\text{PbCl}_6 + 2\text{H}_2\text{O}$ [1]

$\text{H}_2\text{PbCl}_6 \longrightarrow \text{PbCl}_2 + 2\text{HCl} + \text{Cl}_2$ [1]

[5 max 4]

[Total: 11]

5 (a) (i)

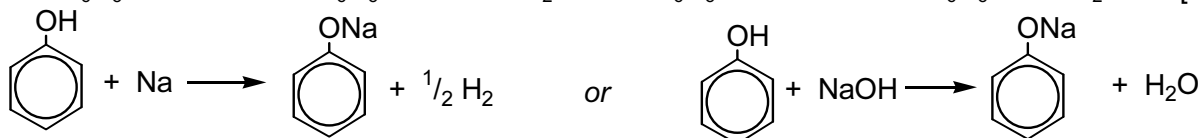


[1]

(ii) Na metal or NaOH [1]

Fizzes/gas given off with phenol or phenol dissolves (anisole doesn't) [1]

$\text{C}_6\text{H}_5\text{OH} + \text{Na} \rightarrow \text{C}_6\text{H}_5\text{ONa} + \frac{1}{2} \text{H}_2$ or $\text{C}_6\text{H}_5\text{OH} + \text{OH}^- \rightarrow \text{C}_6\text{H}_5\text{O}^- + \text{H}_2\text{O}$ [1]



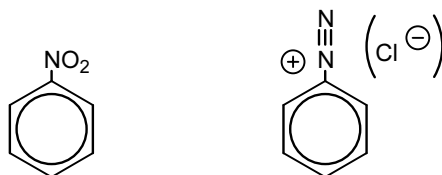
(neutral) iron(III) chloride [1]

Solution goes purple/violet [1]

$3\text{C}_6\text{H}_5\text{OH} + \text{FeCl}_3 \rightarrow \text{Fe}(\text{OC}_6\text{H}_5)_3 + 3\text{HCl}$ [1]

[4]

(b) (i)



D

E

[1] + [1]

(ii) step 2: $\text{Sn} + \text{HCl}$ NOT LiAlH_4 , NaBH_4 [1]

conc. + reflux (warm is insufficient) [1]

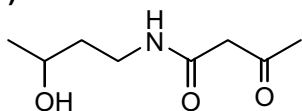
step 4 is conditional of structure E

step 4: warm + in H_2O [1]

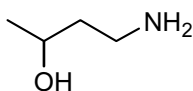
[5 max 4]

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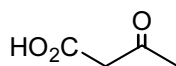
(c) (i)



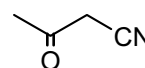
F



G



H



J

F must be an **amide**

[4]

- (ii) reaction 1: $\text{H}_2 + \text{Ni}$ or LiAlH_4
 reaction 2: heat + aqueous HCl

[1]

[1]

[6]

[Total: 14]

6 (a) (i) Condensation

[1]

(ii) ala-ala, gly-gly, ala-gly

[2]

[3]

(b) (i) Correct sugar-phosphate backbones
 (with **two sugars and one phosphate attached**)

[1]

C – G pair correct **or** A – T pair correct

[1]

deoxyribose label **and** all bases coming from sugars

[1]

(ii) Replication would be slower/difficult
 because the DNA/strands could not be separated

[1]

[4]

(c) (i) Some amino acids have more than one (triplet) code

[1]

(ii) loss/disruption of ionic bonding/hydrogen bonding

[1]

(iii) There would be a potential loss of all tertiary structure
 or
frameshift – deletion of a base changes protein structure

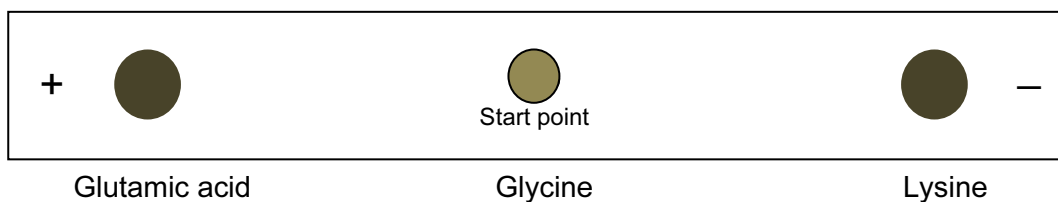
[1]

[3]

[Total: 10]

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7 (a)



- Glutamic acid between + and start point [1]
 Lysine between – and start point [1]
 Glycine at, or very close to, start point [1]
[3]

(b) (i) Ratio of the concentration of a solute in each of two solvents
 or equilibrium constant representing the distribution of a solute between two solvents. [1]

(ii) illustration of some method of getting into our body via the food chain [1]

They dissolve preferentially in fats/oils [1]
[3]

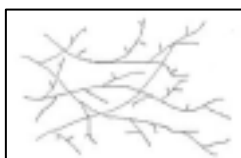
(c) (i) $156 = \text{C}_3\text{H}_6^{35}\text{Cl}^{79}\text{Br}^+$ [1]
 $158 = \text{C}_3\text{H}_6^{37}\text{Cl}^{79}\text{Br}^+$ [1]
 $158 = \text{C}_3\text{H}_6^{35}\text{Cl}^{81}\text{Br}^+$ [1]
 $160 = \text{C}_3\text{H}_6^{37}\text{Cl}^{81}\text{Br}^+$ [1]

(ii) $m/e = 15$ Species = CH_3^+ [1]
[5 max 4]

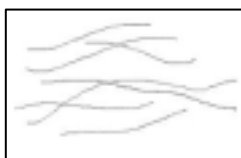
[Total: 10]

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8 (a)



LDPE



HDPE

minimum of 2 chains suitable sketches [1]

(The close packing of unbranched side chains means)

LDPE **more space** between the chains/polymers or HDPE less empty space between the chains [1]

[2]

(b) van der Waals' (VDW) forces
are weaker

[1]

[1]

[2]

(c)

Addition OR	condensation
requires C=C/double bond	does not need C=C/double bond
uses the same functional group	needs two different functional groups
same general (empirical) formula as monomer	different formula
no loss of small molecule/H ₂ O/HCl	small molecule /H ₂ O/HCl is formed

Any two differences

[1]

[2]

(d) (i) (through its long chain of) delocalised electrons/mobile electrons
free electrons is not sufficient

[1]

(ii) planar

[1]

the π bonds/p-orbitals overlap (with each other)

[1]

(iii) C₈H₆
C₄H₃

[2]

[5 max 4]

[Total: 10]