

Tuesday 13 October 2020 – Morning

A Level Chemistry B (Salters)

H433/02 Scientific literacy in chemistry

Time allowed: 2 hours 15 minutes

You must have:

- a clean copy of the Advance Notice Article (inside this document)
- the Data Sheet for Chemistry B

You can use:

- · a scientific or graphical calculator
- an HB pencil



Please write clea	arly in black	k ink. Do n	ot writ	te in the barcodes.		
Centre number				Candidate number		
First name(s)						
Last name						

INSTRUCTIONS

- Use black ink. You can use an HB pencil, but only for graphs and diagrams.
- Write your answer to each question in the space provided. If you need extra space use the lined pages at the end of this booklet. The question numbers must be clearly shown.
- · Answer all the questions.
- Where appropriate, your answer should be supported with working. Marks might be given for using a correct method, even if your answer is wrong.

INFORMATION

- The total mark for this paper is **100**.
- The marks for each question are shown in brackets [].
- Quality of extended response will be assessed in questions marked with an asterisk (*).
- This document has 24 pages.

ADVICE

· Read each question carefully before you start your answer.



Answer all the questions.

1 Aspartame is a dipeptide that is used as an artificial sweetener. The structure of aspartame is shown in Fig. 1.1.

aspartame

Fig. 1.1

- (a) Circle a secondary amide group on the structure of aspartame in Fig. 1.1. [1]
- **(b)** Aspartame is hydrolysed by boiling with acid. One of the products is phenylalanine which is shown in **Fig. 1.2**. 'Ph' represents a benzene ring.

$$\begin{array}{c|c} & & O \\ & H & \parallel \\ & & C \\ & C & C \\ & & NH_2 \end{array}$$

phenylalanine

Fig. 1.2

(i) Phenylalanine has a chiral centre. Draw the three dimensional formulae to show the enantiomers of phenylalanine.



[2]

(ii) Phenylalanine in solution exists mainly as a zwitterion.

Complete the diagram in Fig. 1.3 to show the structure of this zwitterion.

Fig. 1.3

[1]

(iii) A student says that acid hydrolysis of aspartame will only produce phenylalanine and one other amino acid.

Comment on the accuracy of this statement, giving the structures of any compounds you include in your answer.

.....[3]

(c) Aspartame tastes sweet. One theory is that the sweetness is caused by hydrogen bonds forming between sweetener molecules and sweetness receptors in the body.

On the structure of aspartame in **Fig. 1.4** circle **all** the atoms that could hydrogen bond with a receptor.

Fig. 1.4

[1]

(d) The enzyme pepsin catalyses the hydrolysis of protein chains. Some students read that pepsin breaks the peptide bonds adjacent to aromatic amino-acids, for example phenylalanine. This is because the active site is specific to benzene rings.

Suggest the type of bonds that form between a benzene ring and the active site.

.....[1]

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(e)	The students mix excess aspartame and pepsin in aqueous solution at room temperature.
	After 30 minutes they place a spot of the mixture on some chromatography paper.

(i)	Describe the	rest	of their	method,	including	how	they	could	tell	whether	hydrolysis	has
	occurred.											

You may include a diagram in your answer.

 [4

(11)	Other students boil the mixture of aspartame and pepsin to attempt to speed up the hydrolysis reaction.
	State and explain whether or not this would work.
	[1]
(iii)	Another student says that using a higher concentration of aspartame at room temperature will speed up the hydrolysis reaction.
	Comment on this statement.
	[2]

2 Some students use a method based on an electrochemical cell to measure the concentration of silver ions in solution.

They set up a cell as shown in Fig. 2.1.

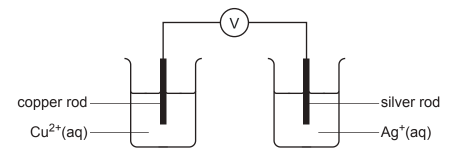


Fig. 2.1

(a) (i) There is something missing from the diagram.

State what it is and describe what it is made of.

 	[2]

(ii) State **two** conditions necessary for the cell to measure electrode potentials under **standard** conditions.

2

(b) Table 2.1 shows the electrode potential data for the cell in Fig. 2.1.

Half-equation	E [⊕] /V
$Cu^{2+}(aq) + 2e^{-} \rightleftharpoons Cu(s)$	+0.34
$Ag^{+}(aq) + e^{-} \rightleftharpoons Ag(s)$	+0.80

Table 2.1

Equation 2.1 shows the reaction that occurs when the standard cell delivers a current.

$$2Ag^{+}(aq) + Cu(s) \rightarrow Cu^{2+}(aq) + 2Ag(s)$$
 Equation 2.1

(i) Identify the oxidising agent in equation 2.1.

.....[1]

(ii)	State and explain which way electrons flow in the wire when the standard cell is delivering a current.
	[1]

(c) Silver bromide, AgBr, is usually described as 'insoluble' but a small amount can dissolve in water.

The students have a saturated solution of AgBr. This contains as much AgBr as will dissolve. They place this saturated solution in the right-hand beaker of the cell in **Fig. 2.1**.

They measure the cell potential against a standard copper electrode. Their result is 0.09 V (with the same electrode polarity as in the standard cell in **(b)**).

The students are told that the relationship between their measured cell potential, $E_{\rm cell}$, and the concentration of silver ions is given by:

$$E_{\text{cell}} = E_{\text{cell}}^{\Theta} + 0.06 \log[\text{Ag}^{+}]$$

Calculate the silver ion concentration in the saturated solution of AgBr.

$$[Ag^{+}] = \dots mol dm^{-3} [4]$$

(d) Some other students make another cell, using two different half cells as shown in Table 2.2.

Half-reaction	E°/V
$Cu^{2+}(aq) + 2e^{-} \rightleftharpoons Cu(s)$	+0.34
$2IO_3^-(aq) + 12 H^+(aq) + 10e^- \rightleftharpoons I_2(aq) + 6H_2O(I)$	+1.19

Table 2.2

(i) Write an equation for the reaction that occurs when the cell delivers a current.

(ii)	Draw a labelled diagram to show how the ${\rm IO_3}^-/{\rm I_2}$ electrode is made up. You do not need to give concentrations.
	[2]
(iii)	The students are told that in a standard IO_3^-/I_2 electrode $[IO_3^-]^2 = [I_2]$.
	They plan to prepare $40\mathrm{cm^3}$ of the solution for the electrode by mixing $20\mathrm{cm^3}$ of 0.08 mol dm ⁻³ $I_2(aq)$ with $20\mathrm{cm^3}$ of $IO_3^-(aq)$.
	Calculate the concentration of $\mathrm{IO_3}^-$ needed so that $[\mathrm{IO_3}^-]^2 = [\mathrm{I_2}]$.
	$[IO_3^-] = \dots mol dm^{-3} [3]$

3

	onopotassium phosphate', KH ₂ PO ₄ , is added to fertilisers. ₂ PO ₄ acts as a buffer and supplies phosphorus.	
(a)	The H ₂ PO ₄ ⁻ ion has two OH groups.	
	Draw a 'dot-and-cross' diagram for this ion.	
		2]
(b)	$\mathrm{KH_2PO_4}$ when used in fertilisers is said to be equivalent to 52% $\mathrm{P_2O_5}$ and 34% $\mathrm{K_2O}$ by mass	s.
	Show that the mole ratio of potassium to phosphorus in 52% P_2O_5 and 34% K_2O is the sar as in KH_2PO_4 .	ıе
		21
(-)		2]
(c)	In a solution of KH ₂ PO ₄ , the equilibrium in equation 3.1 occurs:	
	$H_2PO_4^- \iff H^+ + HPO_4^{2-}$ $K_a = 6.2 \times 10^{-8} \text{mol dm}^{-3}$ Equation 3.1	
	(i) Identify the base in this reaction and explain your choice.	
		1]
	(ii) Calculate the pK_a for the equilibrium in equation 3.1.	

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pK_a =[1]

Ed	uation	3.1	is	repeated	again
_ ~	laatioii	U. I	10	repeated	agaii i.

$H_2PO_4^- \rightleftharpoons H^+ + HPO_4^{2-}$	$K_a = 6.2 \times 10^{-8} \text{mol dm}^{-3}$	Equation 3.1
1121 04 \ 11 \ 111 04	Na - 0.2 ·· 10 morani	Equation 5.1

(iii) Calculate the pH of a $1.0 \times 10^{-3} \, \mathrm{mol \, dm^{-3}}$ solution of $\mathrm{KH_2PO_4}$.

	pH =[2]
(d)	The acidity of KH ₂ PO ₄ reduces the loss of ammonia from ammonium ions (NH ₄ ⁺) in a fertiliser.
	Explain how this happens when the fertiliser is in solution.
	[2]
(e)	A mixture of $\mathrm{HPO_4^{2-}}$ and $\mathrm{H_2PO_4^{-}}$ ions in solution makes a 'phosphate buffer'.
	Calculate the mass of $\rm KH_2PO_4$ that must be added to $1.0\rm dm^3$ of a $0.10\rm moldm^{-3}$ solution of $\rm HPO_4^{\ 2^-}$ to make a solution with a pH of 6.50.

The volume of the solution does not change when the solid is added.

mass of KH_2PO_4 = g [4]

$H_0PO_A \rightleftharpoons H^+ + HPO_A^2 \qquad K_0 = 6.2 \times 10^{\circ} \text{moldm} \circ \text{Equation}$	$H_2PO_4^- \rightleftharpoons H^+ + HPO_4^{2-}$	$K_a = 6.2 \times 10^{-8} \text{mol dm}^{-3}$	Equation 3.1
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_	•	
(f)	(i)	A student says that a solution of $HPO_4^{\ 2^-}$ alone will act as a buffer when acid is added. The student says that this is because the position of equilibrium in equation 3.1 moves to the left to remove H^+ ions.
		Discuss the student's statements.
		[2]
	(ii)	A student is doing calculations involving solutions of hydrochloric acid.
		The student says that the concentration of hydrogen ions can be taken as the concentration of the acid.
		Discuss this statement.
		[1]

(iii) Calculate the **change** in pH when 1 drop $(0.05\,\mathrm{cm^3})$ of $0.01\,\mathrm{mol\,dm^{-3}\,HC}\,l$ is added to $1.0\,\mathrm{dm^3}$ of water at pH 7.0.

pH change =[2]

4 'Fumaric acid' is used as an acidity regulator in food. 'Maleic acid' is a stereoisomer of fumaric acid. The structure of the two acids are shown in **Fig. 4.1.**

Fig. 4.1

(a)	(i)	Describe chemical tests that can be carried out in a laboratory to identify the two functional groups in both acids.
		[2]
	(ii)	A solution has 2.32 g dm ⁻³ of maleic acid.
		Calculate the volume (in cm ³) of 2.0 mol dm ⁻³ NaOH(aq) that would completely neutralise 250 cm ³ of the solution.
		volume = cm ³ [3]
(b)	(i)	Give the empirical formula of both acids.
		[1]
	(ii)	Give the systematic name for maleic acid .
		[2]
(c)	Exp	plain why maleic acid and fumaric acid are different compounds.
		[21

(d)	The mass spectra of maleic and fumaric acid both have a peak at 71. Suggest a reason for this.
	[1]

(e) Fumaric acid is made from maleic acid by an acid-catalysed isomerisation reaction.

A student suggests the mechanism shown in Fig. 4.2 for the reaction.

(i) Draw 'curly arrows' on Fig. 4.2 to complete the mechanism.

Fig. 4.2

[2]

- (ii) Name the **type** of reaction that is occurring in **step 2**.
- (iii) The student then reads that if ${}^2H^+$ ions are used for the isomerisation, then there isn't any 2H found in the fumaric acid formed.

Does this support the student's mechanism in Fig. 4.2?

Explain your answer.

.....

.....[2]

(f) Maleic acid can be made from 'maleic anhydride', $\rm C_4H_2O_3$.

Suggest a skeletal formula for maleic anhydride.

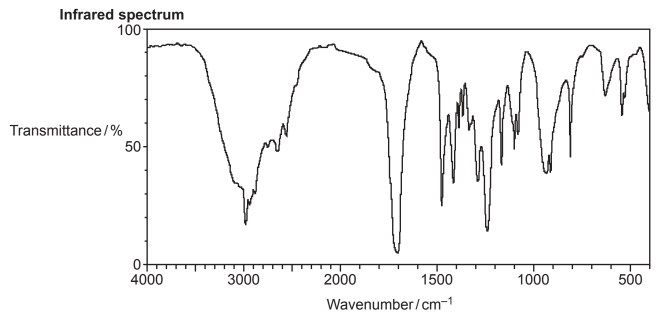
(g)	Mal	eic anhy	dride ca	n be mad	de industr	rially by o	xidising bu	utane or be	nzene.		
	C ₄ H	I ₁₀ + 3.	$50_2 \rightarrow$	C ₄ H ₂ O ₃	+ 4H ₂ O)		Reaction	on 1		
	C ₆ H	l ₆ + 4.	$50_2 \rightarrow$	C ₄ H ₂ O ₃	+ 2H ₂ O	+ 2CO ₂	2	Reaction	on 2		
	(i)	Use ca	lculated	atom eco	onomies t	o sugges	t, with a re	eason, whic	ch reaction	ı is used r	nore.
											[2]
	(ii)	Sugges	st anothe	r reason	why the	reaction s	selected in	ı (g)(i) is pı	eferred.		
											[1]
	(iii)	Calculation 15	ite the model	naximum utane (m	mass of easured a	maleic ar at RTP).	nhydride, (C ₄ H ₂ O ₃ , (ii	n kg) that	could be i	made
		Give yo	our answ	er to an a	appropri	ate numb	er of signi	ficant figur	es.		
				mass	s of malei	ic anhydr	ide =			h	kg [2]

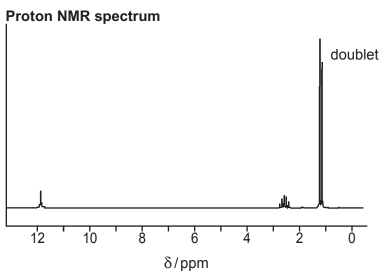
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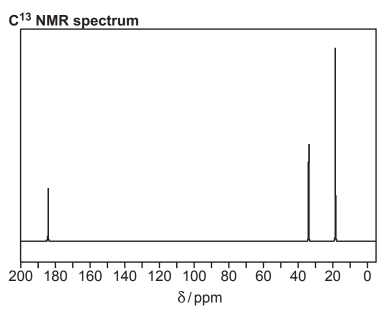
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(h)* Compound A has spectra as shown below.







You may do working on this page but it will not be marked

Compound A has four carbon atoms in its molecule.	
Identify compound A , giving evidence from each spectrum.	[6]
Additional answer space if required	

- 5 This question refers to the Advance Notice Article 'Platinum metal complexes in medicine' that is included inside this document.
 - (a) Cisplatin is a neutral molecule.

Explain why the oxidation state of the platinum in cisplatin is +2.
r4

(ii) Suggest the electronic configuration of the outermost sub-shell in a Pt^{2+} ion.

[1]

(iii) Is platinum a transition metal?

Give a reason.	

.....[1]

(b) Some ligands are described as bidentate.

Give the **full** structural formula of a bidentate ligand from a compound in the article.

[2]

(c) Fig. 5.1 is a diagram from page 3 of the article. It shows a possible way in which a compound derived from cisplatin is thought to attach to a base in DNA. Two bonds have been labelled B and C.

Fig. 5.1

(i)	What is the name of the base in Fig. 5.1? Use the Data Sheet to help you.	
		[1]
(ii)	What is the name of the type of bond at B in Fig. 5.1 ?	
		[1]
(iii)	Name the type of bond at C in Fig. 5.1 and describe how it is formed.	
		[2]
(d) Sat	raplatin was designed to be taken by mouth as it is soluble.	
(i)	Name the shape of a satraplatin molecule around the Pt atom.	
		[1]
(ii)	Give the co-ordination number of Pt in satraplatin.	
		[1]
(iii)	Suggest, in terms of intermolecular bonds, why satraplatin is soluble whereas cispla not.	atin is
		[3]

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END OF QUESTION PAPER

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ADDITIONAL ANSWER SPACE

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