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Candidate Number

Centre Number

ADVANCED General Certificate of Education 2018

Chemistry

Assessment Unit A2 2

assessing

Analytical, Transition Metals, Electrochemistry and Further Organic Chemistry



[ACH22]

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TUESDAY 12 JUNE, AFTERNOON

TIME

2 hours.

INSTRUCTIONS TO CANDIDATES

Write your Centre Number and Candidate Number in the spaces provided at the top of this page.

Answer all eighteen questions.

Answer **all ten** questions in **Section A**. Record your answers by marking the appropriate letter on the answer sheet provided. Use only the spaces numbered 1 to 10. Keep in sequence when answering.

Answer all eight questions in Section B.

You must answer the questions in the spaces provided.

Do not write outside the boxed area on each page or on blank pages.

Complete in black ink only. Do not write with a gel pen.

INFORMATION FOR CANDIDATES

The total mark for this paper is 110.

Quality of written communication will be assessed in Questions 12(b) and 13(a)(vi).

In Section A all questions carry equal marks, i.e. one mark for each question.

In Section B the figures in brackets printed down the right-hand side of pages indicate the marks awarded to each question or part question.

A Periodic Table of Elements, containing some data, is included with this question paper.



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Section A

For each of the following questions only one of the lettered responses (A–D) is correct.

Select the correct response in each case and mark its code letter by connecting the dots as illustrated on the answer sheet.

1 When hydrogen peroxide is added to acidified potassium dichromate(VI) the following reaction occurs:

$$\mathrm{Cr_2O_7^{2-} + 3H_2O_2 + 8H^+ \rightarrow 2Cr^{3+} + 3O_2 + 7H_2O}$$

Which statement is **not** correct?

- A Hydrogen peroxide is acting as an oxidising agent
- B The solution changes colour from orange to green
- C The oxidation state of oxygen changes in the reaction
- D The oxidation state of chromium changes in the reaction
- 2 Which compound has only singlet peaks in its proton nmr spectrum?
 - A Ethyl ethanoate
 - B Methyl propanoate
 - C Methyl ethanoate
 - D Propyl ethanoate



- **3** A painkiller, ibuprofen, produced effervescence when added to a solution of sodium carbonate. Which functional group is present in ibuprofen?
 - A -CHO
 - B -COOH
 - C -COOR
 - D -CONH₂
- 4 Which compound is the weakest base?
 - A Ammonia

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- B Methanamide
- C Methylamine
- D Phenylamine
- 5 What is the oxidation number of cobalt in $[Co(H_2O)_4en]Cl_2$?
 - A -2
 - B 0
 - C +2
 - D +3

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6	Wh	nich atom has an unpaired electron in an s-orbital in the ground state?
	Α	Chromium
	В	Cobalt
	С	Iron
	D	Manganese
7	Wh	nich compound will react most rapidly with 1,6-diaminohexane to form nylon?
	Α	Hexane-1,6-diol
	В	Hexanedioic acid
	С	Hexanedioyl dichloride
	D	Sodium hexanedioate
8		sample of chlorine gas was placed in a mass spectrometer. How many molecular peaks would be observed?
	Α	2
	В	3
	С	4
	D	5



- **9** What mass of butanamide is required to synthesise 6.90 g of butanenitrile if the yield is 80%?
 - A 6.97g
 - B 8.70g
 - C 10.9g
 - D 12.2g

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- **10** Some glycine is dissolved in a buffer solution of pH 11. What is the structure formed at this pH?
 - A HOOCCH₂NH₃⁺
 - B NH₂CH₂COOH
 - C NH₂CH₂COO⁻
 - D NH-CH₂COO-

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Section B

Answer all eight questions in the spaces provided

- 11 Transition metal complexes are often coloured.
 - (a) State the colour of the following aqueous complexes:

aqueous complex	colour
[Mn(H ₂ O) ₆] ²⁺	
[Ni(H ₂ O) ₆] ²⁺	
[Co(H ₂ O) ₆] ²⁺	
[V(H ₂ O) ₆] ³⁺	
[Ni(NH ₃) ₆] ²⁺	

[5]

(b) Aqueous hexaaquacopper(II) ions can undergo ligand replacement with concentrated hydrochloric acid to form tetrachlorocuprate(II) ions.

(i)	Write the electronic configuration of the copper ion in hexaaquacopper(II)
	ions and use this to explain why copper can be described as a transition
	metal.

[2]

(ii)	Define the term ligand .

[1]

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ions react with concentrated hydrochloric acid.	
Write the equation for the ligand substitution reaction which occurs when hexaaquacopper(II) ions form tetrachlorocuprate(II) ions.	[′
	[2
Write the colour change observed when this reaction occurs.	[′
Explain why this ligand replacement is thermodynamically feasible.	
	[2
oper can form complexes with ammonia or ethylamine. State and explain ch of these would be the stronger ligand.	
	[2
	Write the equation for the ligand substitution reaction which occurs when hexaaquacopper(II) ions form tetrachlorocuprate(II) ions. Write the colour change observed when this reaction occurs. Explain why this ligand replacement is thermodynamically feasible.

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12 Two standard electrode potentials are given below:

(a)	half-cell	E [⊕] /V
	$Zn^{2+}(aq) + 2e^{-} \rightleftharpoons Zn$	-0.76
	$Cu^{2+}(aq) + 2e^{-} \rightleftharpoons Cu$	+0.34

(i)	Define standard electrode potential.

		[2]



[Turn over

(b)	Describe how you would set up a standard hydrogen electrode and use it to measure the electrode potential for a half-cell.
	In this question you will be assessed on using your written communication skills including the use of specialist scientific terms.
	[6]

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- 13 A method of synthesising aspirin is given below using the following steps:
 - Add 7.5 cm³ (an excess) of ethanoic anhydride to 3.0 g of salicylic acid in a conical flask
 - 2. Add eight drops of concentrated phosphoric acid
 - 3. Heat, with stirring, for 20 minutes in a water bath
 - 4. Add 3 cm³ of deionised water to the flask
 - 5. Add 30 cm³ of deionised water and cool to room temperature, allowing the aspirin to crystallise
 - 6. Filter the crystals by Buchner filtration; continue to suck air through the Buchner funnel for five minutes after completion of the filtration

(a) (i)	preference to ethanoic acid.
	[2]

(ii) Write the equation for the reaction in step 1.

[2]



(iii)	Why is concentrated phosphoric acid added?	[1]
(iv)	Suggest why water is added in step 4.	
		[1]
(v)	Explain why air is sucked through the apparatus for five minutes.	
		[1]

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((vi)	Describe, giving full experimental detail, the TLC method which can be used to determine whether the reaction is complete after step 3. The solvent is ethyl ethanoate.	
		In this question you will be assessed on using your written communication skills including the use of specialist scientific term	s.
			[6
(b) (Cald	culate the percentage yield if 2.3g of aspirin is obtained (answer to one	
		imal place).	
-			
-			[3]
-			_ [3]



- (c) Salicylic acid is a bifunctional molecule with a carboxylic acid group and a hydroxyl group attached to the benzene ring. The hydroxyl group displays acidic behaviour.
 - (i) Suggest why the hydroxyl group attached to a benzene ring is more acidic than the hydroxyl group in aliphatic alcohols.

_____[2]

(ii) Write an equation for the reaction of salicylic acid with excess aqueous sodium hydroxide.

______[2]

- (iii) State why it is preferable to use the sodium salt of aspirin.

 [1]
- (d) Bromine will give an electrophilic substitution reaction with salicylic acid as shown below:

$$\begin{array}{c|c} \text{OH} & \text{OH} & \text{OH} \\ + \text{ Br}_2 \rightarrow & \text{COOH} & + \text{ HBr} \end{array}$$

Salicylic acid will react with bromine without a catalyst being present. This differs from benzene, which requires a metal halide catalyst.

Name a catalyst which is used in the bromination of benzene.

_____[1]

[Turn over



14 Mesalazine, an anti-inflammatory drug which is used to treat bowel disease, can be synthesised from 2-hydroxybenzoic acid using the flow scheme below.

2-hydroxybenzoic acid

mesalazine

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(a) (i) State the reagents that could be used for Step 1 and Step 2.

(ii) Mesalazine can be converted into azo dyes through reaction with naphth-1-ol or resorcinol:

Draw the structure of the ion mesalazine must be converted into before the azo dye can be formed.

[2]



	(iii)	Draw the structure of the azo dye produced by the reaction with resort	inol.
			[2]
	(iv)	Explain why azo dyes are coloured and suggest why the azo dyes pro by resorcinol and naphth-1-ol have slightly different colours.	duced
			[4]
(b)		/droxybenzoic acid can be converted into an ester by reacting with an imolar amount of ethane-1,2-diol. Write an equation for this reaction.	
			[2]
(c)	Poly	vethylene terephthalate can be produced from ethane-1,2-diol.	
	(i)	State a use for polyethylene terephthalate.	
			[1]
	(ii)	Explain why polyethylene terephthalate is biodegradable.	
			[1]
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15 An ester, with the molecular formula $C_7H_{14}O_2$, produced three signals when analysed by proton nmr. The data is provided in the table below:

signal	а	b	С
chemical shift	4.1	1.2	1.1
integration ratio	2	3	9
splitting pattern	quartet	triplet	singlet

(a)	(i)	Explain why solvents which contain hydrogen atoms should not be used in nmr spectroscopy. Suggest a suitable solvent which could be used.	n
			[2]
	(ii)	Give the name and formula of the molecule used in nmr spectroscopy as a standard.	[2]
	(iii)	State two reasons why the molecule identified in part (ii) is used.	
			[2]



b)	(i)	Explain which alkyl group in the ester produces signals a and b mal reference to the spin-spin splitting pattern and the integration ratios	
		reference to the spin-spin spilling pattern and the integration ratios	•
			[3]
	/ii\	Draw the alkyl group that would give rise to signal c	
	(11)	Draw the alkyl group that would give rise to signal c .	
			[4 ⁻
			[1]
c)	Dra	w the possible structure of the ester based upon the nmr data given.	
			[2]

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(a)	(i)	Write the ionic equation for the reaction of bromine with sodium hydrox
	(ii)	State the colour change observed for the above reaction.
(b)	Acid	dified bromate(V) ions will oxidise iodide ions to iodine.
	(i)	Write a half-equation for the reduction of bromate(V) ions to bromide.
	(ii)	Write a half-equation for the oxidation of iodide ions to iodine.
	(iii)	Write the overall equation for this reaction.

(c)	libe con bro resi 0.10	e iodine produced can then be reduced by thiosulfate ions. Titrations of the trated iodine with sodium thiosulfate solution can be used to determine the incentration of bromate(V) ions. A 20.0 cm ³ solution containing acidified mate(V) ions was added to a solution containing excess iodide ions and the ulting mixture made up to 1.0 dm ³ . A 25.0 cm ³ aliquot was titrated against 0 M sodium thiosulfate, adding starch indicator just before the end point. The e was found to be 23.8 cm ³ .									
	(i)	Explain why it is necessary to add the starch indicator just before the end point.	t								
			[1]								
	(ii)	Calculate, to two significant figures, the concentration of the original bromate(V) solution.									
			 _ [4]								

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17 Aspartame is used as a sweetener in many food products. It is a methyl ester of the dipeptide produced in the condensation reaction between aspartic acid and phenylalanine.

Aspartame

(a) Use the structure of aspartame to suggest structures for aspartic acid and phenylalanine.

aspartic acid

phenylalanine

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(b) Another amino acid derivative that is used as a food additive is monosodium glutamate which can be synthesised from glutamic acid.

monosodium glutamate

- (i) Circle on the structure above any chiral centre present in monosodium glutamate. [1]
- (ii) Draw the structure of the zwitterion formed by glutamic acid.

(iii) Write an equation for the reaction of glutamic acid with sodium carbonate to form monosodium glutamate.

[2]

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[1]

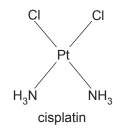


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18 Cisplatin was first described by Peyrone in 1845 and was approved for use in the treatment of testicular and ovarian cancers in the USA in 1978.



(a) Explain why cisplatin is effective in acting as an anticancer drug.

______[1]

(b) Cisplatin has a number of undesired side-effects which are believed to be caused by the drug activating before it reaches the targeted tumour. Attempts to reduce these side-effects have included modifying the structure to give derivatives such as carboplatin.

Give the molecular formula of carboplatin.

____ [1]



(c) Another method of developing more targetable platinum-based anticancer drugs has been through the development of photoactivable drugs which are activated through photoreduction by light. Early examples of these were diiodo complexes.



(i) Explain, using oxidation states, why converting ${\bf A}$ into ${\bf B}$ is regarded as a reduction.

_____[2]

(ii) Describe the change in both shape and co-ordination number in converting A into B.

______ [4

THIS IS THE END OF THE QUESTION PAPER



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Periodic Table of the Elements

For the use of candidates taking
Advanced Subsidiary and Advanced Level
Chemistry Examinations

Copies must be free from notes or additions of any kind. No other type of data booklet or information sheet is authorised for use in the examinations.

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I	II		TH	E PEI	RIODI	C TAE Gro		III	IV	V	VI	VII	0				
Hydrogen Hydrogen Li Lithium Na Sodium	9 Be Beryllium 4 24 Mg Magnesium	occu Plan Gas	upies a v ick Cons Consta	volume stant = 0 int = 8.3	is at 20° of 24 dn 6.63 × 1 31 J mol ⁻ = 6.02 :	1 ³ . 0 ⁻³⁴ Js - ¹ K ⁻¹		ire of 1	atmosph	iere (10) ⁵ Pa)	11 B Boron 5 27 Al Aluminium	12 Carbon 6 28 Si Silicon	14 N Nitrogen 7 31 P	16 Oxygen 8 32 Sulfur	19 Fluorine 9 35.5 Cl Chlorine	4 Helium 2 20 Neon 10 40 Ar Argon
11	12 40	45	48	51	52	55	56	59	59	64	65	13 70	14 73	15 75	16 79	17 80	18 84
Potassium	Calcium 20	Sc Scandium 21	Titanium 22	Vanadium 23	Cr Chromium 24	Mn Manganese 25	Fe	Co Cobalt 27	Nickel 28	Cu Copper 29	Zn Zinc 30	Gallium 31	Germanium	As	Se Selenium	Bromine 35	Kr Krypton 36
85	88	89	91	93	96	99	101	103	106	108	112	115	119	122	128	127	131
Rb Rubidium	Sr Strontium	Yttrium 39	Zr Zirconium 40	Nb Niobium 41	Molybdenum	Technetium 43	Ru Ruthenium	Rhodium	Palladium 46	Ag Silver	Cadmium 48	Indium 49	Sn 50	Sb Antimony 51	Tellurium 52	lodine 53	Xe Xenon 54
133	137 R 2	139 I 3 *	178	181 T 3	184 W	186 D O	190	192	195 Pt	197	201	204 T	207 Pb	209 Bi	210 D O	210 ^+	222 D n
Caesium 55	Ba Barium 56	Lanthanum 57	Hafnium 72	Tantalum	Tungsten	Re Rhenium 75	Osmium 76	Iridium 77	Platinum 78	Gold 79	Hg Mercury 80	Thallium 81	Lead 82	Bismuth 83	Po Polonium 84	At Astatine 85	Radon 86
Francium	Radium 88	Actinium 89		•	•		•	•	•	•		•	•	•	•	•	

* 58–71 Lanthanum series † 90–103 Actinium series

a b x

a = relative atomic mass (approx.) **x** = atomic symbol **b** = atomic number

140	141	144	147	150	152	157	159	162	165	167	169	173	175
Ce	Pr	Nd	Pm	Sm	Eu	Gd	Tb	Dy	Но	Er	Tm	Yb	Lu
Cerium	1 '		Promethium		Europium	Gadolinium		Dysprosium		Erbium	Thulium	Ytterbium	Lutetium
58	59	60	61	62	63	64	65	66	67	68	69	70	71
232	231	238	237	242	243	247	245	251	254	253	256	254	257
Th	Pa	U	Nρ	Pu	Am	Cm	Bk	Cf	Es	Fm	Md	No	Lr
	Protactinium	Uranium	Neptunium		Americium	Curium	Berkelium	Californium	Einsteinium		Mendelevium	Nobelium	Lawrencium
90	91	92	93	94	95	96	97	98	99	100	101	102	103