Write your name here Surname		Other names	
Pearson Edexcel International Advanced Level	Centre Number		Candidate Number
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Chemistry Advanced Unit 5: General Principl Metals and Orga (including synop	es of Chemis Inic Nitrogen	Ćhemis	
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## Instructions

- Use **black** ink or **black** ball-point pen.
- **Fill in the boxes** at the top of this page with your name, centre number and candidate number.
- Answer **all** questions.
- Answer the questions in the spaces provided
  - there may be more space than you need.

# Information

- The total mark for this paper is 90.
- The marks for **each** question are shown in brackets
  - use this as a guide as to how much time to spend on each question.
- Questions labelled with an asterisk (\*) are ones where the quality of your written communication will be assessed
  - you should take particular care with your spelling, punctuation and grammar, as well as the clarity of expression, on these questions.
- A Periodic Table is printed on the back cover of this paper.

# **Advice**

- Read each question carefully before you start to answer it.
- Check your answers if you have time at the end.
- Show all your working in calculations and give units where appropriate.

Turn over ▶



P51938A
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### **SECTION A**

Answer ALL the questions in this section. You should aim to spend no more than 20 minutes on this section. For each question, select one answer from A to D and put a cross in the box  $\boxtimes$ . If you change your mind, put a line through the box  $\boxtimes$  and then mark your new answer with a cross  $\boxtimes$ .

- In the complex ion [Fe(CN)<sub>6</sub>]<sup>4−</sup>, the oxidation number of the iron is
   A +2
  - **B** +3

(Total for Question 1 = 1 mark)

- 2 Aluminium reacts with aqueous alkali to produce hydrogen gas. By considering the changes in the oxidation numbers of aluminium and hydrogen, it can be deduced that, in this reaction, 1 mol of aluminium produces
  - **A** 1 mol of hydrogen gas.
  - ☑ B 1.5 mol of hydrogen gas.

  - ☑ D 3 mol of hydrogen gas.

(Total for Question 2 = 1 mark)

The standard reduction potentials of two systems are

$$Sn^{4+}(aq) + 2e^{-} \rightleftharpoons Sn^{2+}(aq)$$
  $E^{\oplus} = +0.15V$ 

$$Fe^{3+}(aq) + e^{-} \implies Fe^{2+}(aq) \qquad E^{\oplus} = +0.77V$$

(a) What are the components of the half-cell required to measure the standard reduction potential of the  $Fe^{3+}/Fe^{2+}$  system?

(1)

	lon(s) in the solution	Metal electrode
⊠ A	Fe <sup>2+</sup>	iron
<b>⋈</b> B	Fe³+	iron
<b>⋈</b> C	Fe <sup>2+</sup> and Fe <sup>3+</sup>	iron
☑ D	Fe <sup>2+</sup> and Fe <sup>3+</sup>	platinum

(b) What is the value of  $E_{\text{cell}}^{\ominus}$  for the system below?

$$Sn^{4+}(aq) \ + \ 2Fe^{2+}(aq) \ \to \ Sn^{2+}(aq) \ + \ 2Fe^{3+}(aq)$$

(1)

- B -0.62 V
- ☑ D +1.39 V

# (Total for Question 3 = 2 marks)

- **4** The value of  $E_{\text{cell}}^{\oplus}$  for a reaction is positive. It can be deduced that the
  - $\square$  **A** entropy change for the system,  $\Delta S_{\text{system}}^{\ominus}$ , must be positive.
  - $\ \square$   $\ B$   $\$  entropy change for the surroundings,  $\Delta S^{\ominus}_{surroundings}$  , must be positive.
  - $\hfill \square$  C total entropy change,  $\Delta S^{\ominus}_{total}$  , must be positive.
  - $\square$  **D** entropy change for the system,  $\Delta S_{\text{system}}^{\ominus}$ , and the entropy change for the surroundings,  $\Delta S_{\text{surroundings}}^{\ominus}$ , must **both** be positive.

(Total for Question 4 = 1 mark)



- 5 In a hydrogen-oxygen fuel cell, the oxygen is
  - A oxidised at the negative electrode.
  - **B** oxidised at the positive electrode.
  - C reduced at the negative electrode.
  - **D** reduced at the positive electrode.

(Total for Question 5 = 1 mark)

**6** Iodide ions are oxidised by peroxodisulfate ions in aqueous solution:

$$2I^{-} + S_{2}O_{8}^{2-} \rightarrow I_{2} + 2SO_{4}^{2-}$$

This reaction is catalysed by Fe<sup>3+</sup> ions. This catalysis is effective because

- $\blacksquare$  **A** Fe<sup>3+</sup> is readily reduced to Fe<sup>2+</sup> which is then oxidised to Fe<sup>3+</sup>.
- $\blacksquare$  **B** Fe<sup>3+</sup> has many electrons in its outermost subshells.
- $\square$  **C** Fe<sup>3+</sup> has many active sites on which the reaction can occur.
- **D** Fe<sup>3+</sup> will oxidise peroxodisulfate ions to sulfate ions.

(Total for Question 6 = 1 mark)

- 7 Which has the greatest number of unpaired electrons in each of its atoms?
  - A Vanadium
  - **B** Chromium

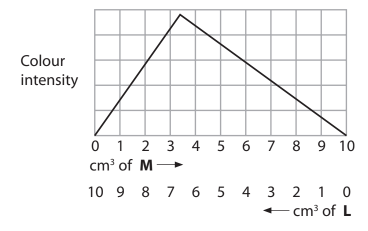
  - D Iron

(Total for Question 7 = 1 mark)

8 A transition metal ion, M, forms a coloured complex with a monodentate ligand L.

Portions of a 0.05 mol dm<sup>-3</sup> solution of **M** are mixed with portions of a solution containing 0.05 mol dm<sup>-3</sup> of **L** so that the total volume of the resulting mixture is always 10 cm<sup>3</sup>. The colour intensity of the complex in each mixture is measured using a colorimeter.

The graph of the results is



What is the formula of the complex?

- $\square$  A ML
- B ML₂
- C ML<sub>3</sub>
- $\square$  D ML<sub>4</sub>

(Total for Question 8 = 1 mark)

**9** What is the total number of different oxidation numbers of vanadium that occurs in the white solid and the three solutions shown in the following sequence?

NH <sub>4</sub> VO <sub>3</sub> (s)	add	pale		dark	add zinc	pale
white solid	water	yellow	add H <sub>2</sub> SO <sub>4</sub>	yellow	powder	purple
write solid		solution		solution		solution

- **■ B** 2
- **C** 3
- □ 4

(Total for Question 9 = 1 mark)

**10** The enthalpy change of hydrogenation of but-1-ene is -126.9 kJ mol<sup>-1</sup> and the enthalpy change of hydrogenation of buta-1,3-diene is -236.4 kJ mol<sup>-1</sup>.

The delocalisation structure of buta-1,3-diene is more stable than the molecule with two isolated C—C bonds by

- **■ B** 109.5 kJ mol<sup>-1</sup>

(Total for Question 10 = 1 mark)

- **11** Aluminium chloride catalyses the reaction of benzene and ethanoyl chloride. This is because aluminium chloride
  - A has significant covalent character.
  - **B** exists as a dimer in non-polar solvents.
  - C is an electron pair acceptor.
  - **D** is rapidly hydrolysed.

(Total for Question 11 = 1 mark)

12		lamine reacts rapidly with dilute sulfuric acid. The reaction is
	<b>⊠</b> A	electrophilic addition.
	■ B	electrophilic substitution.
	<b>⊠</b> C	hydrolysis.
	⊠ D	neutralisation.
_		(Total for Question 12 = 1 mark)
13		e is an amino acid. It is a crystalline solid at room temperature because the est forces between the molecules of alanine are
13	strong	e is an amino acid. It is a crystalline solid at room temperature because the
13	strong	e is an amino acid. It is a crystalline solid at room temperature because the est forces between the molecules of alanine are
13	strong	e is an amino acid. It is a crystalline solid at room temperature because the est forces between the molecules of alanine are dipole-dipole forces.
13	strong  A  B	e is an amino acid. It is a crystalline solid at room temperature because the est forces between the molecules of alanine are dipole-dipole forces.  London forces.

**14** Aspartame is an artificial sweetener. The structure of aspartame is

$$\begin{array}{c|c} H & O \\ \hline H & N \\ \hline HO & H \\ O \\ \end{array}$$

(a) Which functional group is **not** present in aspartame?

(1)

- A Amine
- **B** Ester
- C Ketone
- ☑ D Phenyl
- (b) How many asymmetric carbon atoms are there in a molecule of aspartame?

(1)

- A Two
- B Three
- ☑ D Six

(Total for Question 14 = 2 marks)

**15** The repeat unit of the polymer Kevlar has the structure

(Total for Question 15 = 1 mark)

- **16** Which of these isomers would be expected to have mass spectrum peaks at m/e = 30 and m/e = 73?
  - A CH<sub>3</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>NH<sub>2</sub>
  - ☑ B CH<sub>3</sub>CH(NH<sub>2</sub>)CH<sub>2</sub>CH<sub>3</sub>
  - $\square$  **C** (CH<sub>3</sub>)<sub>3</sub>CNH<sub>2</sub>
  - ☑ D CH<sub>3</sub>CH<sub>2</sub>NHCH<sub>2</sub>CH<sub>3</sub>

(Total for Question 16 = 1 mark)

17			portant advantage of the use of polymer supports in organic synthesis by the aceutical industry is that
	X	A	the equilibrium yield of the reaction is usually higher.
	X	В	polymeric support materials are always inexpensive to prepare.
	X	C	a range of side reactions occur on the polymer support.
	×	D	the products are easily separated from the polymer support.
			(Total for Question 17 = 1 mark)
18	Wł	nich	piece of apparatus is normally used in solvent extraction?
	×	A	Buchner funnel
	X	В	Evaporating basin
	X	C	Liebig condenser
	X	D	Separating funnel
			(Total for Question 18 = 1 mark)

**TOTAL FOR SECTION A = 20 MARKS** 



### **SECTION B**

# Answer ALL the questions. Write your answers in the spaces provided.

**19** Salicylic acid (2-hydroxybenzoic acid) is a starting material in the manufacture of aspirin; it is synthesised from phenol in the Kolbe-Schmitt process outlined.

- (a) Step 2 of the Kolbe-Schmitt process involves an electrophilic substitution.
  - (i) Using your knowledge of electrophilic substitutions, draw the mechanism for the reaction in Step 2.
    - Show the displayed formula of the electrophile CO<sub>2</sub>, and all relevant curly arrows.

(4)

(ii) Identify Reagent Y.

(1)



\*(b) The Kolbe-Schmitt process is carried out at 125 °C and high pressure. State how, if at all, these conditions for electrophilic substitution would have to change if the Kolbe-Schmitt process was carried out with benzene in place of phenol.

Fully justify your answer in terms of the relative reactivity of benzene and phenol.



(c) In the final stage of the synthesis, 2-hydroxybenzoic acid is converted into aspirin in a single step:

2-hydroxybenzoic acid

aspirin

(i) Name the type of reaction occurring.

(1)

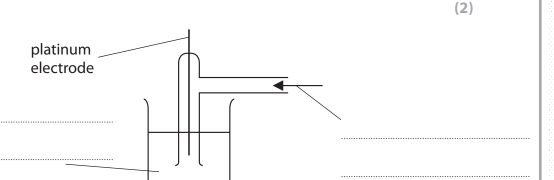
*(ii)	One student suggested that ethanoic acid would be a suitable reagent for this final reaction. A second student suggested using ethanoyl chloride. Compare the suitability of these two reagents, stating any advantages or disadvantages of each.	(3)
	(Total for Question 19 = 13	marks)



(2)

(1)

- **20** Standard electrode potentials are used to predict the thermodynamic feasibility of chemical reactions.
  - (a) Standard electrode potentials can be determined by using the standard hydrogen electrode as a reference standard.
    - (i) Label the diagram of the standard hydrogen electrode, indicating the essential conditions associated with each substance used.



(ii) The platinum electrode of the standard hydrogen electrode is coated in finely divided platinum ('platinum black'). Suggest why platinum is used as the electrode and why it is coated in platinum black.

(iii) State the value assigned to the electrode potential of the standard hydrogen electrode.

(iv) Explain why a reference electrode is needed to measure the electrode potentials of chemical systems. (1)

(v) Sometimes the calculated value of  $E_{\text{cell}}^{\ominus}$  for a reaction shows it to be thermodynamically feasible but, in the laboratory, no reaction occurs. Give **two** possible explanations for this.

(2)

(b) Potassium dichromate(VI) in acid conditions is a laboratory oxidising agent widely used in organic chemistry. Some electrode potentials relating to the use of this reagent are given in the table below.

Electrode reaction	E/V
$CH_3COOH(aq) + 2H^+(aq) + 2e^- \rightleftharpoons CH_3CHO(aq) + H_2O(I)$	-0.94
$CH_3CHO(aq) + 2H^+(aq) + 2e^- \rightleftharpoons C_2H_5OH(aq)$	-0.61
$Cr_2O_7^{2-}(aq) + 14H^+(aq) + 6e^- \implies 2Cr^{3+}(aq) + 7H_2O(I)$	+1.33

(i) Deduce the ionic equation for the oxidation of ethanol to ethanal by potassium dichromate(VI) in acid conditions. State symbols are not required.

(2)

- (ii) Calculate  $E_{cell}$  for the oxidation by acidified potassium dichromate(VI) of
  - ethanol to ethanal
  - ethanal to ethanoic acid.

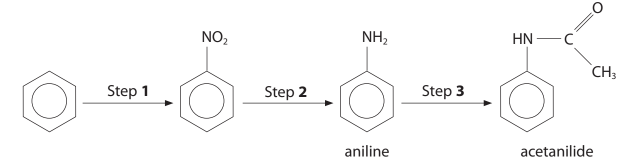
(2)



	oxidation of ethanol.	(1)
ch ab	tassium manganate(VII) is another laboratory oxidising agent used in organic emistry. Use the data on page 17 of your Data Booklet to answer the questions out manganese compounds.  Suggest why potassium manganate(VII) is <b>not</b> used to oxidise alcohols such as ethanol, whereas potassium dichromate(VI) is used. Give the relevant electrode reaction and its standard electrode potential. No calculation is required.	(2)
(ii)	Write the equation for the disproportionation of manganate(VI) ions, $MnO_4^{2^-}$ , and calculate whether or not the reaction is thermodynamically feasible. State symbols are not required.	(3)



- 21 Acetanilide (*N*-phenylethanamide) was the first aniline (phenylamine) derivative discovered to have analgesic properties. It was first used as a painkiller in 1886 but quickly discarded because of a range of toxic effects. Nowadays it is important in the manufacture of 'sulfa' antibacterial drugs.
  - (a) A laboratory synthesis of acetanilide from benzene is outlined below.



Give the reagents for each step and the conditions for Step 1 and Step 2.

(5)

Step <b>1</b>	 	
Step <b>2</b>		
Stan 3		 



	_				_		_	
(h)	The macc	spectrum o	of acetanilide	contains no	acks at m/a	-13 m/a	o – 77 and	$m/\rho - 135$
\UI	1116 111033	3DECHUIII U	n acciainnac i	LUITTAILIS DE	ans at III / c	— <del>1</del> 3,111/c	. — // anu	III/ E — 133.

(i) Give the **molecular formulae** of the species that produce the peaks at m/e = 135 and m/e = 77.

(2)

(ii) Give the **structure** of the species responsible for the peak at m/e = 43.

(1)

(c) Use your Data Booklet to suggest the wavenumber range of **two** peaks that would be present in the infrared spectrum of acetanilide but not in the infrared spectrum of aniline. Identify the bond stretches responsible for each of your chosen peaks.

(2)

(Total for Question 21 = 10 marks)

**22** A volatile organic compound **X** has the percentage by mass composition:

$$C = 61.0\%$$
;  $H = 15.3\%$ ;  $N = 23.7\%$ .

0.105 g of **X** was vaporised and the volume of dry air displaced measured. After adjustment to room temperature and pressure, the volume of air displaced was 42.7 cm<sup>3</sup>.

(a) Calculate the empirical formula of **X**. You **must** show your working.

(3)

(b) Calculate the molar mass of **X** and hence deduce its molecular formula. You **must** show your working.

Molar volume of gas at room temperature and pressure =  $24.0\,\mathrm{dm^3\,mol^{-1}}$ 

(2)

(c) Draw two of the possible structures for **X**.

(2)

(d) The low resolution proton nmr spectrum of **X** has three peaks of relative area 6:2:1. Identify the structure of **X** and explain how your structure is consistent with the nmr spectrum.

(2)

(Total for Question 22 = 9 marks)

**TOTAL FOR SECTION B = 50 MARKS** 



### **SECTION C**

# Answer ALL the questions. Write your answers in the spaces provided.

23 The importance of the element copper (Z = 29) is due to its limited chemical reactivity and high thermal and electrical conductivity. Copper compounds have significant biological activity. Like silver and gold, copper can occur in nature as the pure element, although it is mostly obtained from one of the minerals that contain copper compounds.

The physical properties of copper have resulted in its widespread use in electrical wiring and in plumbing. Due to its cost and relative softness, the use of pure copper in coinage has largely been superseded by alloys, some of which contain no copper at all.

Copper is placed in the d-block of the Periodic Table and forms compounds with two stable oxidation states: copper(I), which resembles silver(I), and copper(II). The characteristics of copper(II) lead to the classification of copper as a transition element. Both copper(I) and copper(II) form a range of complex ions; copper(I) complexes are colourless but copper(II) complexes are coloured, often blue.

(a) (i) Complete the electronic structures of Cu, Cu<sup>+</sup> and Cu<sup>2+</sup>, using the 'electrons-in-boxes' notation.

 Cu
 (Ar)

 Cu<sup>+</sup>
 (Ar)

(ii)	Explain	why	copper is	classified	as a	d-block	element.
------	---------	-----	-----------	------------	------	---------	----------

(2)

(1)

(iii) Explain why the electronic structure of Cu <sup>2+</sup> leads to copper being classified as a transition element.	
	(1)
(b) Anhydrous copper(II) sulfate is a white crystalline solid which dissolves in water to form a solution that has a pale blue colour.	
(i) Explain fully why it becomes coloured when it is dissolved in water.	(5)
	(3)
(ii) Explain why anhydrous copper(II) sulfate has no colour.	
	(1)



(c) The percentage of copper in a piece of copper tubing was determined by a titration method. The tubing was weighed, dissolved in concentrated nitric acid and the solution with all the washings transferred to a 250.0 cm³ volumetric flask. The pH of the solution was adjusted before it was made up to the mark with deionised water and mixed thoroughly. 25.0 cm³ portions of this solution were titrated with a solution of edta until concordant results were obtained. The equation for the reaction in the titration is

$$[Cu(H_2O)_6]^{2+} + edta^{4-} \rightarrow [Cu(edta)]^{2-} + 6H_2O$$

The results of the experiment are summarised in the table.

Measurement	Value
Mass of tubing / g	3.63
Concentration of edta solution / mol dm <sup>-3</sup>	0.205
Mean titre / cm³	27.50

(i) Suggest what might be added to the solution to adjust the pH.

(1)

(ii) Calculate the percentage by mass of copper in the tubing.

(3)



(1)
e
and omplexes
(2)

(ii) Draw a diagram showing how **one** molecule of cysteamine complexes with a  $Cu^{2+}$  ion.

(1)



*(iii) Use your knowledge of the shape of copper(I) complexes to sugger cysteamine does <b>not</b> act as a bidentate ligand with Cu <sup>+</sup> .	est why (2)
(Total for Question	n 23 = 20 marks)
TOTAL FOR SECTION	N C = 20 MARKS

**TOTAL FOR PAPER = 90 MARKS** 



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# The Periodic Table of Elements

0 (8)	(18) 4.0 <b>He</b> helium 2	20.2	neon 10	39.9	Ar argon 18	83.8	궃	krypton 36	131.3	×	xenon 54	[222]	R	radon 86		ted	
7	(17)	19.0	fluorine 9	35.5	CI chlorine 17	6.62	Ŗ	bromine 35	126.9	Ι	iodine 53	[210]	Αt	astatine 85		seen repor	
9	(16)	16.0	oxygen 8	32.1	Sulfur 16	79.0	Se	selenium 34	127.6	Ъ	tellurium 52	[509]	8	polonium 84		116 have t	iticated
2	(15)	14.0	nitrogen 7	31.0	P phosphorus 15	74.9	As	arsenic 33	121.8	Sb	antimony 51	209.0	Bi	bismuth 83		nbers 112-	but not fully authenticated
4	(14)	12.0	carbon 6	28.1	Silicon 14	72.6	g	germanium 32	118.7	S	20 th	207.2	ď	lead 82		atomic nun	but not fr
ъ	(13)	10.8	boron 5	27.0	AI aluminium 13	69.7	g	gallium 31	114.8	Ę	mnipui 49	204.4	F	thallium 81		Elements with atomic numbers 112-116 have been reported	
					(12)	65.4	Zu	zinc 30	112.4	8	cadmium 48	200.6	Ĕ	mercury 80		Elem	
					(11)	63.5	3	copper 29	107.9	Ag	silver 47	197.0	Ρη	gold 79	[272]	æ	oentgenium 111
					(10)	58.7	ź	nickel 28	106.4	В	palladium 46	195.1	풉	platinum 78	[271]	õ	darmstactrium r 110
					(6)	58.9	ပိ	cobalt 27	102.9	윤	rhodium 45	192.2	<u> </u>	iridium 77	[368]		meitnerium 109
	1.0 hydrogen				(8)	55.8	Fe	iron 26	101.1	æ	ruthenium 44	190.2	õ	osmium 76	[277]		hassium 108
					(2)	54.9	W	manganese 25	[86]	բ	technetium 43	186.2	Re	rhenium 75	1 —	监	bohrium 107
		mass	umber		(9)	52.0	ხ	vanadium chromium manganese 23 24 25	95.9	Wo	molybdenum technetium 42 43	183.8	>	tungsten 74	[592]	Sg	seaborgium 106
	Key	relative atomic mass	atomic (proton) number		(5)	50.9	>	vanadium 23	92.9	g	niobium 41	180.9	Тa	tantalum 73	۱_		dubnium 105
		relati	atomic		(4)	47.9	ï	titanium 22	91.2	Zr	zirconium 40	178.5	Ξ	hafnium 72	[261]	₹	nutherfordium 104
					(3)	45.0	S	scandium 21	88.9	>	yttrium 39	138.9	La*	lanthanum 57	[227]		actinium 89
2	ē	9,0	beryllium 4	24.3	Mg magnesium 12	40.1	S	calcium 20	97.6	'n	strontium 38	137.3		barium 56	[526]	Ra	radium 88
-	$\varepsilon$	6.9	Lithkish 3		Ma sodkun 11	39.1	¥	potassium 19	85.5	<b>&amp;</b>	rubidium 37	132.9	ర	caesium 55	[223]	ድ	francium 87

Lanthanide series

<sup>\*</sup> Actinide series

	141	144	[147]	150	152	157	159	163	165	167	169	173	175
	P.	Ž	Pa	S	교	В	4	δ	운	Ъ	ᆵ	Ą	2
8	eodymium)	neodymium	promethium	samarinm	europium	gadolinium	terbium	dysprosium	holmium	erbinm	thulium	ytterbium	lutetium
	26	9	61	62	63	64	65	99	- 67	68	69	70	71
匚	231]	238	[237]	[242]	[243]	[247]	[245]	[251]	[254]	[253]	[356]	[254]	[257]
	Pa	_	δ	P	Αm	క్ర	쓢	ង	E	F	ΡW	£	۲
Ď.	tactinium	uranium	neptunium	plutonium	americium	arrium	berkelium	catifornium	einsteinium	fermium	mendelevium	nobelium	lawrencium
	7	92	93	94	95	96	46	86	66	9	101	102	103