

Cambridge International AS & A Level

CANDIDATE NAME						
CENTRE NUMBER				CANDIDATE NUMBER		

CHEMISTRY 9701/42

Paper 4 A Level Structured Questions

May/June 2023

2 hours

You must answer on the question paper.

No additional materials are needed.

INSTRUCTIONS

- Answer all questions.
- Use a black or dark blue pen. You may use an HB pencil for any diagrams or graphs.
- Write your name, centre number and candidate number in the boxes at the top of the page.
- Write your answer to each guestion in the space provided.
- Do not use an erasable pen or correction fluid.
- Do not write on any bar codes.
- You may use a calculator.
- You should show all your working and use appropriate units.

INFORMATION

- The total mark for this paper is 100.
- The number of marks for each question or part question is shown in brackets [].
- The Periodic Table is printed in the question paper.
- Important values, constants and standards are printed in the question paper.

This document has 24 pages.

- 1 (a) Group 2 carbonates decompose when heated to form the metal oxide and carbon dioxide.
 - (i) Suggest a mechanism for the decomposition of the carbonate ion by adding **two** curly arrows in Fig. 1.1.



Fig. 1.1

[1]

	(ii)	Describe the variation in the thermal stability of Group 2 carbonates. Explain your answer.
		[3]
(b)	(i)	Define lattice energy.
		[2]
	(ii)	The lattice energy of the Group 2 carbonates, $\Delta H_{\rm latt}^{\rm e}({\rm MCO_3})$, becomes less exothermic down the group.
		The lattice energy of the Group 2 oxides, $\Delta H_{\rm latt}^{\Theta}({\rm MO})$, also becomes less exothermic down the group.
		$\Delta H_{\rm latt}^{\Theta}({\rm MCO_3})$ and $\Delta H_{\rm latt}^{\Theta}({\rm MO})$ change by different amounts going down the group.
		Suggest how the standard enthalpy change of the decomposition reaction for Group 2 carbonates changes down the group.
		Explain your reasoning in terms of the relative sizes of the anions and the relative changes in lattice energy down the group.
		AHEAD

(c) Potassium sulfite, K₂SO₃, is used as a food additive.

The concentration of sulfite ions, SO_3^{2-} , can be determined by titration using aqueous acidified manganate(VII) ions, MnO₄-.

A 250 cm³ solution contains 3.40 g of impure $\rm K_2SO_3$. 25.0 cm³ of this solution requires 22.40 cm³ of 0.0250 mol dm⁻³ acidified $\rm MnO_4^-$ to reach the end-point. All the $\rm SO_3^{2-}$ ions are oxidised. None of the other species in the impure $\rm K_2SO_3$ are oxidised.

The reaction occurs as shown by the two half-equations.

$$H_2O + SO_3^{2-} \rightarrow SO_4^{2-} + 2H^+ + 2e^ MnO_4^- + 8H^+ + 5e^- \rightarrow Mn^{2+} + 4H_2O$$

(i)	(i) Give the ionic equation for the reaction between SO_3^{2-} and acidified MnO_4^{-} .					
		[1				

(ii) Calculate the percentage purity of the sample of K₂SO₃. Show your working.

(d) Potassium disulfite, $K_2S_2O_5$, is another food additive. The disulfite ion, $S_2O_5^{2-}$, has the displayed formula shown in Fig. 1.2.

$$S \longrightarrow S \longrightarrow S \longrightarrow O$$

Fig. 1.2

Deduce the geometry (shape) around the $S(\alpha)$ atom in $S_2O_5^{2-}$.

geometry around $S(\alpha)$



2	(a)	State two typical chemical properties of a transition element.	
		1	
		2	[1]
	(b)	Aqueous solutions of cobalt(II) salts contain the complex ion $[Co(H_2O)_6]^{2+}$.	
		(i) Define complex ion.	

(ii) Samples of $[Co(H_2O)_6]^{2+}$ are reacted separately with an excess of aqueous ammonia, with an excess of concentrated HCl and with an excess of aqueous sodium hydroxide, as shown in Fig. 2.1.

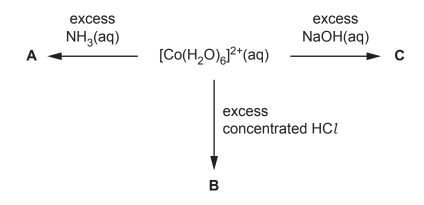


Fig. 2.1

Complete Table 2.1 about the reactions shown by $[Co(H_2O)_6]^{2+.}$

Table 2.1

reagent added to [Co(H ₂ O) ₆] ²⁺ (aq)	formula of cobalt species formed	colour and state of cobalt species formed	type of reaction
an excess of NH ₃ (aq)	A =		
an excess of concentrated HC1	B =		
an excess of NaOH(aq)	C =		



(c)	The			
	(i)	Explain what is meant by a bidentate ligand.		
			[2]	
	(ii)	The complex $[Co(H_2O)_2(C_2O_4)BrCl]^-$ exists as stereoisomers.		
		Complete the three-dimensional diagrams in Fig. 2.2 to show four s $[Co(H_2O)_2(C_2O_4)BrCl]^-$.	tereoisomers of	
		The $\mathrm{C_2O_4^{2-}}$ ligand is represented using O O .		
		Milling Commo		
		Milling Country Countr		
Fig. 2.2				
	(iii)	State the oxidation state of cobalt in this complex and a type of stereoisc	merism shown.	
		oxidation state of cobalt		
		type of stereoisomerism		
			[1]	

[Total: 12]



3 (a) Complete Table 3.1 by placing **one** tick (✓) in each row to indicate the sign of each type of energy change under standard conditions.

Table 3.1

energy change	always positive	always negative	can be either negative or positive
bond energy			
enthalpy change of atomisation			
enthalpy change of formation			
		ı	[1]

(b)	Define standard enthalpy change of atomisation.	
		[2]

(c) Table 3.2 shows some energy changes.

Table 3.2

energy change	value/kJ mol ⁻¹
standard enthalpy change of atomisation of silver	+285
first ionisation energy of silver	+731
second ionisation energy of silver	+2074
bond energy of O=O	+496
bond energy of O-O	+150
first electron affinity of oxygen	-141
second electron affinity of oxygen	+798
first ionisation energy of oxygen	+1314
standard enthalpy change of formation of silver oxide, Ag ₂ O(s)	-31



Calculate the lattice energy, $\Delta H_{\text{latt}}^{\Theta}$, of Ag ₂ O(s) using relevant data from Table 3.2.
It may be helpful to draw a labelled energy cycle.
Show your working.

	$\Delta H_{\text{latt}}^{\Theta} \text{ of Ag}_2 O(s) = \dots kJ \text{ mol}^{-1}$	[3]
(d)	Suggest the trend in the magnitude of the lattice energies of the silver compounds ${\rm Ag_2O}$ and ${\rm Ag_2Se}$.	յ _շ Տ,
	Explain your answer.	
	least exothermic most exothermic	
		[2]



(e)	Silver sulfite, Ag ₂ SO ₃ (s), is sparingly soluble in water.			
	(i)	Give an expression for the solubili	ty product, $K_{\rm sp}$, of ${\rm Ag_2SO_3}$.	
		K _{sp} =		
			[1]	
	(ii)	Calculate the equilibrium concentr	ation of Ag ⁺ in a saturated solution of Ag ₂ SO ₃ at 298 K.	
		$[K_{\rm sp}: {\rm Ag}_2{\rm SO}_3, 1.50 \times 10^{-14}{\rm mol}^3{\rm dn}]$	ı ^{–9} at 298K]	
			$[Ag^{+}] = \dots mol dm^{-3} [1]$	
(f)	The	e standard enthalpy change of solut	ion, $\Delta H_{\text{sol}}^{\Theta}$, of AgNO ₃ (s) in water is +22.6 kJ mol ⁻¹ .	
	Sug	ggest how the feasibility of dissolvin	g AgNO ₃ (s) in water changes with temperature.	
	Exp	olain your answer.		



[Total: 12]

4 (a) In aqueous solution, iron(III) ions react with iodide ions, as shown.

$$2Fe^{3+} + 2I^{-} \rightarrow 2Fe^{2+} + I_{2}$$

A series of experiments is carried out using different concentrations of ${\rm Fe^{3+}}$ and ${\rm I^-}$, as shown in Table 4.1.

Table 4.1

experiment	[Fe ³⁺]/moldm ⁻³	[I ⁻]/moldm ⁻³	initial rate/moldm ⁻³ s ⁻¹
1	0.0400	0.0200	2.64 × 10 ⁻⁴
2	0.1200	0.0200	7.92 × 10 ⁻⁴
3	0.0800	0.0400	2.11 × 10 ⁻³

(i)	Explain what is meant by overall order of reaction.
	[1]
(ii)	Use the data in Table 4.1 to deduce the order of reaction with respect to ${\rm Fe^{3+}}$ and with respect to ${\rm I^-}$.
	Explain your reasoning.
	[2]
(iii)	Use your answer to (a)(ii) to construct the rate equation for this reaction.
	rate = [1]
(iv)	Use your answer to (a)(iii) and the data from experiment 1 to calculate the rate constant, k , for this reaction. Include the units of k .
	k = units [2]
(v)	Describe qualitatively the effect of an increase in temperature on the rate constant and on the rate of this reaction.
	ZEA F
	AHEAD [1]

(b) In aqueous solution, iodide ions react with acidified hydrogen peroxide, as shown.

$$2I^- + H_2O_2 + 2H^+ \rightarrow I_2 + 2H_2O$$

The initial rate of reaction is found to be first order with respect to I^- , first order with respect to H_2O_2 and zero order with respect to H^+ .

Fig. 4.1 shows a possible four-step mechanism for this reaction.

$$\begin{array}{lll} \text{step 1} & \text{H}_2\text{O}_2 + \text{I}^- \rightarrow \text{IO}^- + \text{H}_2\text{O} \\ \\ \text{step 2} & \text{H}^+ + \text{IO}^- \rightarrow \text{HIO} \\ \\ \text{step 3} & \text{HIO} + \text{I}^- \rightarrow \text{I}_2 + \text{OH}^- \\ \\ \text{step 4} & \text{OH}^- + \text{H}^+ \rightarrow \text{H}_2\text{O} \\ \end{array}$$

Fig. 4.1

(i)	Suggest which of the steps, 1, 2, 3 or 4, in this mechanism is the rate-determining step.
	Explain your answer.
	[1]
(ii)	Identify a step in Fig. 4.1 that involves a redox reaction.
	Explain your answer in terms of oxidation numbers.
	[1]
(iii)	Suggest the role of HIO in this mechanism.
	Explain your reasoning.
	[1]



[Total: 10]

5 (a) Methylbenzene can undergo different reactions, as shown in Fig. 5.1.

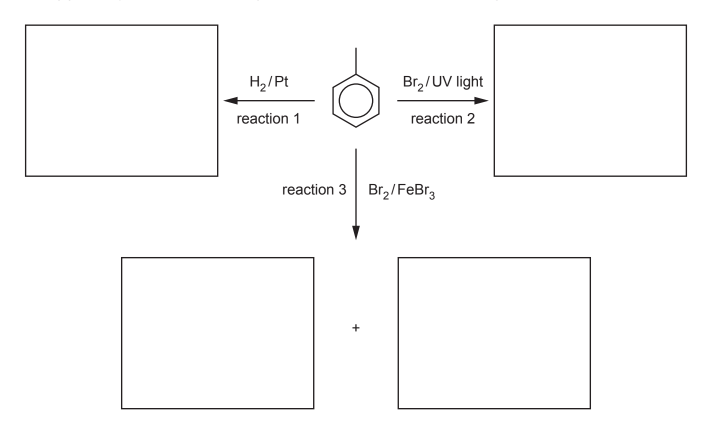


Fig. 5.1

- (i) Draw structures in Fig. 5.1 for the possible organic products of the three reactions shown. [3]
- (ii) Complete Table 5.1.

Table 5.1

	type of reaction	mechanism
reaction 1		
reaction 2		

[2]

(b) When methylbenzene reacts with an electrophile, a substitution reaction occurs. No addition reaction takes place under these conditions.

Explain why no addition reaction takes place.



(c) The reaction of methylbenzene with thionyl bromide, SOBr₂, in the presence of an iron(III) bromide catalyst, FeBr₃, is shown in Fig. 5.2.

Fig. 5.2

The mechanism of this reaction is similar to that of the bromination of benzene.

The first step of the mechanism generates the SOBr⁺ electrophile, as shown.

$$\mathsf{SOBr}_2^+ \mathsf{FeBr}_3 \! \to \! \mathsf{SOBr}^+ + \mathsf{FeBr}_4^-$$

(i) The reaction of methylbenzene with SOBr⁺ ions is shown in Fig. 5.3. Complete the mechanism in Fig. 5.3.

Include all relevant curly arrows and charges.

Draw the structure of the organic intermediate.

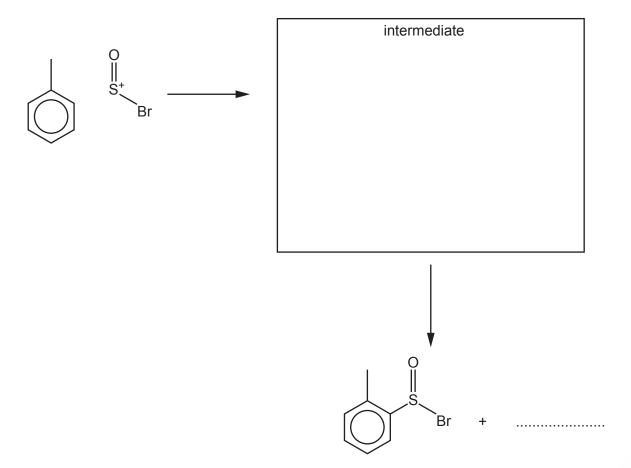


Fig. 5.3



	(ii)	The reaction shown in Fig. 5.2 produces a small amount of a by-product, ${\bf P}$, with the molecular formula ${\rm C_{14}H_{14}OS}.$
		Suggest a structure for by-product P .
		[1]
(d)	Acy	I bromides, RCOBr, can be synthesised by the reaction of a carboxylic acid and SOBr ₂ .
	This	s is a similar reaction to the synthesis of acyl chlorides using $\mathrm{SOC}\mathit{l}_{2}.$
	(i)	Give an equation for the reaction between ethanoic acid and SOBr ₂ .
		[1]
	(ii)	Suggest the relative ease of hydrolysis of acyl bromides, RCOBr, acyl chlorides, RCOC \it{l} , and alkyl chlorides, RC \it{l} .
		Explain your answer.
		easiest to hydrolyse > > hardest to hydrolyse
		[3]
		[Total: 14]

[Total: TT]



6 (a) Perindopril is a drug used to treat heart disease.

perindopril

Fig. 6.1

	(i)	State the number of chiral carbon atoms present in one molecule of perindopril.	
			[1]
	(ii)	Suggest one benefit and one disadvantage of producing a drug such as perindopril single pure optical isomer.	as a
		benefit	
		disadvantage	
			[2]
(b)	(i)	Name all the functional groups in perindopril.	
			[2]



(ii)

A sample of perindopril is hydrolysed with hot	aqueous acid.
Draw the structures of the three organic properindopril.	oducts of the complete acid hydrolysis of

[3]

[Total: 8]



7	(a)	Explain why phenol is brominated much more easily than benzene is brominated.	
			[3]
	(b)	lodine monobromide, I–Br, reacts with benzene in the presence of an $\mathrm{A}\mathit{l}\mathrm{Br}_3$ catalyst.	
		Predict whether the organic product will be bromobenzene or iodobenzene. Explain your answer.	
	(c)	Fig. 7.1 shows some reactions of phenol.	[1]
		$Q \qquad \qquad \begin{array}{c} \text{Na(s)} \\ \text{OH} \end{array} \qquad \begin{array}{c} \text{excess Br}_2(\text{aq}) \\ \text{R} \end{array}$	
		reaction 1 reaction 2	
		O_2N OH OH	
		Fig. 7.1	
		(i) Give an equation for the reaction of phenol with Na(s).	
			[1]



(11)	of Br ₂ (aq).	an excess
(iii)	State the reagents and conditions for reaction 1 and reaction 2 in Fig. 7.1.	[1]
	reaction 1	
	reaction 2	[2]
		[Total: 8]



8 (a)	Describe the difference in reactive	vity between chloroethane a	nd chlorobenzene with OH ⁻ (aq).
	Explain your answer.		
			[2]
(b)	Compound T , $C_5H_9O_2Cl$, is a us		
	Fig. 8.1 shows some reactions of	of T .	
	Т	W	Х
01/	excess NaOH(aq)		+
Cl	heat		
	step 1		
	\	V	
NC	excess HCl(aq)	Y	+ OH
140	heat step 2		
	V		
	Z , C ₆ H ₁₃ NO ₂		

Fig. 8.1



(1)	Give the systematic name for 1.	
		[1]
(ii)	Draw the structures of W , X , Y and Z in Fig. 8.1.	[4]
(iii)	State the reagents and conditions for steps 1 and 2 in Fig. 8.1.	
	step 1	
	step 2	
		121



(c) The proton (¹H) NMR spectrum of compound T, $C_5H_9O_2Cl$, in CDC l_3 is shown in Fig. 8.2.

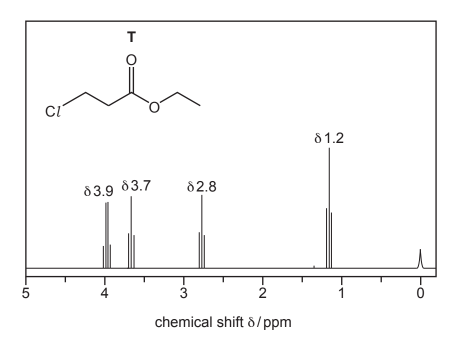


Fig. 8.2

Table 8.1

environment of proton	example	chemical shift range δ/ppm
alkane	-CH ₃ , -CH ₂ -, >CH-	0.9–1.7
alkyl next to C=O	CH ₃ -C=O, -CH ₂ -C=O, >CH-C=O	2.2–3.0
alkyl next to aromatic ring	CH ₃ -Ar, -CH ₂ -Ar, >CH-Ar	2.3–3.0
alkyl next to electronegative atom	CH ₃ -O, -CH ₂ -O, -CH ₂ -C <i>l</i>	3.2-4.0
attached to alkene	=CHR	4.5–6.0
attached to aromatic ring	H –Ar	6.0–9.0
aldehyde	HCOR	9.3–10.5
alcohol	ROH	0.5–6.0
phenol	Ar–O H	4.5–7.0
carboxylic acid	RCOOH	9.0–13.0
alkyl amine	R-N H -	1.0–5.0
aryl amine	Ar-NH ₂	3.0-6.0
amide	RCONHR	5.0–12.0



(i) Suggest why ${\rm CDC} l_3$ is used as a solvent instead of ${\rm CHC} l_3$ for the proton (^1H) NMR spectrum.

	Tabl	e 8.2	
nemical ft δ/ppm	environment of proton	splitting pattern	number of ¹ H atoms responsible for the peak
1.2			
2.8			
3.7			
3.9			



9	(a)	Define standard cell potential, E_{cell}^{Θ} .
		[1
	(b)	An electrochemical cell is set up to measure $E_{\rm cell}^{\Theta}$ of a cell consisting of an Fe ³⁺ /Fe ²⁺ half-cell and a C l_2 /C l^- half-cell.

Draw a labelled diagram of this electrochemical cell.

Include all necessary substances. It is **not** necessary to state conditions used.

[3]

(c) The cell reaction for the electrochemical cell in (b) is shown.

$$Cl_2 + 2Fe^{2+} \rightarrow 2Fe^{3+} + 2Cl^ E_{cell}^{\Theta} = +0.59V$$

Calculate ΔG^{Θ} , in kJ mol⁻¹, for this cell reaction.

$$\Delta G^{\Theta} = \dots kJ \text{ mol}^{-1} [2]$$



(d) Another experiment is set up using the same electrochemical cell.

In this experiment the Fe²⁺ concentration is 0.15 mol dm⁻³. All other concentrations remain at their standard values.

The Nernst equation is shown.

$$E = E^{\oplus} + (0.059/z) \log \frac{\text{[oxidised species]}}{\text{[reduced species]}}$$

(i) Use the Nernst equation to calculate the electrode potential, E, for the Fe³⁺/Fe²⁺ half-cell in this experiment.

$$[E^{\Theta}: Fe^{3+}/Fe^{2+} = +0.77V]$$

(ii) Use your answer to (d)(i) to calculate $E_{\rm cell}$ for this electrochemical cell.

[Total: 8]

Important values, constants and standards

molar gas constant	$R = 8.31 \mathrm{J}\mathrm{K}^{-1}\mathrm{mol}^{-1}$
Faraday constant	$F = 9.65 \times 10^4 \mathrm{C}\mathrm{mol}^{-1}$
Avogadro constant	$L = 6.022 \times 10^{23} \mathrm{mol}^{-1}$
electronic charge	$e = -1.60 \times 10^{-19}$ C
molar volume of gas	$V_{\rm m} = 22.4 {\rm dm^3 mol^{-1}}$ at s.t.p. (101 kPa and 273 K) $V_{\rm m} = 24.0 {\rm dm^3 mol^{-1}}$ at room conditions
ionic product of water	$K_{\rm w} = 1.00 \times 10^{-14} \rm mol^2 dm^{-6} (at 298 K (25 ^{\circ}C))$
specific heat capacity of water	$c = 4.18 \mathrm{kJ kg^{-1} K^{-1}} (4.18 \mathrm{J g^{-1} K^{-1}})$



The Periodic Table of Elements

	18	2	<u>е</u>	lium 1.0	10	e	90n 0.2	18	 ;-	argon 39.9	36	ン	rpton 3.8	74	é	non 11.3	36	٦	uop I	18	g	nesson	
			_	P P																			+
	17				6	ш	fluorine 19.0	17	Cl	chlorine 35.5	35	Ŗ	bromine 79.9	53	Н	iodine 126.9	85	Ą	astatine -	117	<u>⊼</u>	tennessine	-
	16				80	0	oxygen 16.0	16	S	sulfur 32.1	34	Se	selenium 79.0	52	<u>e</u>	tellurium 127.6	84	Ъо	polonium –	116	_	livermorium	
	15				7	z	nitrogen 14.0	15	۵	phosphorus 31.0	33	As	arsenic 74.9	51	Sb	antimony 121.8	83	<u>B</u>	bismuth 209.0	115	Mc	moscovium	1
	14				9	O	carbon 12.0	14	S	silicon 28.1	32	Ge	germanium 72.6	20	Sn	tin 118.7	82	Pb	lead 207.2	114	Εl	flerovium	ı
	13				2	В	boron 10.8	13	Al	aluminium 27.0	31	Ga	gallium 69.7	49	I	indium 114.8	84	11	thallium 204.4	113	R	nihonium	1
										12	30	Zu	zinc 65.4	48	ပ္ပ	cadmium 112.4	80	Я	mercury 200.6	112	ပ်	copernicium	1
										7	29	D C	copper 63.5	47	Ag	silver 107.9	62	Αu	gold 197.0	111	Rg	roentgenium	1
dn										10	28	z	nickel 58.7	46	Pd	palladium 106.4	78	₫	platinum 195.1	110	Ds	darmstadtium	
Group										6	27	ပိ	cobalt 58.9	45	돈	rhodium 102.9	77	'n	iridium 192.2	109	₩	meitnerium	1
		-	I	hydrogen 1.0						80	26	Ьe	iron 55.8	4	Ru	ruthenium 101.1	9/	SO	osmium 190.2	108	R	hassium	ı
										7	25	Mn	manganese 54.9	43	ပ	technetium -	75	Re	rhenium 186.2	107	В	bohrium	ı
						loc	SS			9	24	ပ်	chromium 52.0	42	Mo	molybdenum 95.9	74	>	tungsten 183.8	106	Sg	seaborgium	ı
				Key	atomic number	atomic symbo	name relative atomic mass			2	23	>	vanadium 50.9	41	g	niobium 92.9	73	<u>⊾</u>	tantalum 180.9	105	op O	dubnium	1
						ato	rela			4	22	ı	titanium 47.9	40	Zr	zirconium 91.2	72	Ξ	hafnium 178.5	104	Ŗ	rutherfordium	
										က	21	Sc	scandium 45.0	39	>	yttrium 88.9	57-71	lanthanoids		89-103	actinoids		
	2				4	Be	beryllium 9.0	12	Mg	magnesium 24.3	20	Ca	calcium 40.1	38	ပွဲ	strontium 87.6	26	Ba	barium 137.3	88	Ra	radium	
	_				က	=	lithium 6.9	=	Na	sodium 23.0	19	\prec	potassium 39.1	37	Rb	rubidium 85.5	55	S	caesium 132.9	87	ъ́	francium	1

r ₇	lutetium 175.0	103	۲	lawrencium	1
° X	ytterbium 173.1	102	%	nobelium	1
69 Tm	thulium 168.9	101	Md	mendelevium	ı
88 F	erbium 167.3	100	Fm	ferminm	ı
67 Ho	holmium 164.9	66	Es	einsteinium	1
99	dysprosium 162.5	86	ರ	californium	ı
65 Tb	terbium 158.9	26	Ř	berkelium	ı
[⊉] D	gadolinium 157.3	96	Cm	curium	ı
63 Eu	europium 152.0	92	Am	americium	ı
Sm	samarium 150.4	26	Pu	plutonium	I
Pm	promethium	93	ď	neptunium	ı
9 PZ	neodymium 144.4	92	\supset	uranium	238.0
_{ගී} උ	praseodymium 140.9	91	Ра	protactinium	231.0
Se O	cerium 140.1	06	Т	thorium	232.0
57 La	lanthanum 138.9	89	Ac	actinium	ı

lanthanoids

actinoids

To avoid the issue of disclosure of answer-related information to candidates, all copyright acknowledgements are reproduced online in the Cambridge Assessment International Education Copyright Acknowledgements Booklet. This is produced for each series of examinations and is freely available to download at www.cambridgeinternational.org after the live examination series.