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MARK SCHEME

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Question	Answer	Marks
1(a)	N +2 to +3 (and oxidised)	1
	Br ₂ /Br 0 to –1 (and reduced)	1
1(b)	$\begin{pmatrix} x & x & \\ x & 0 & \begin{pmatrix} x & \bullet & \\ x & \bullet & \end{pmatrix} & \begin{pmatrix} x & x & \\ x & \bullet & \\ x & \bullet & \end{pmatrix} & \begin{pmatrix} x & x & \\ x & x & \\ x & x & \end{pmatrix}$	
	3 bonding pairs around N (in a structure involving NOBr)	1
	rest of molecule correct	1
1(c)(i)	the power to which a concentration of a reactant is raised in the rate equation	1
1(c)(ii)	using expt. 2 and 3 a = 2 or [NO] 2nd order and conc × 3 rate × 9 or $6.1 \times 10^{-2}/6.8 \times 10^{-3} = (0.09/0.03)^a$	1
	using expt. 1 and 2 b = 1 or [Br ₂] 1 st order and conc × 2 rate × 2 or $6.8 \times 10^{-3}/3.4 \times 10^{-3} = (0.04/0.02)^b$	1
(c)(iii)	initial rate = 0.16(32)	1
1(c)(iv)	$(0.0034 = k(0.03)^2(0.02))$ k = 188.9	1
	$mol^{-2}dm^{6}s^{-1}$	1
1(c)(v)	k decreases (as rate decreases)	1

Question	Answer	Marks
1(d)	m = 2 and n = 0	1

Question	Answer	Marks
2(a)	it/solubility decreases down the group and K_{sp} decreases	1
2(b)(i)	$MgCO_3(s) \rightleftharpoons Mg^{2+}(aq) + CO_3^{2-}(aq)$	1
2(b)(ii)	(white) solid appears / precipitation (of MgCO ₃)	1
	as [CO ₃ ²⁻] increases shifting equilibrium to the LHS (precipitating out MgCO ₃)	1
2(c)	solubility = $\sqrt{1.0 \times 10^{-5}}$ = 3.16 × 10 ⁻³ mol dm ⁻³	1
	solubility= $3.2 \times 10^{-3} \times 84.3 = 0.27 \text{g dm}^{-3}$	1
2(d)(i)	Mg ²⁺ ion is smaller than Ba ²⁺ ion or ionic radii increase down group ora	1
	(Mg ²⁺) distorts/polarises/the anion/nitrate group/nitrate ion /NO ₃ ⁽¹⁾⁻ /NO ₃ ion more easily (than Ba ²⁺) ora	1
2(d)(ii)	$Ba(NO_3)_2 \rightarrow BaO + 2NO_2 + \frac{1}{2}O_2$	1
2(d)(iii)	$BaO + H_2O \rightarrow Ba(OH)_2$	1
	$Ba(OH)_2 + H_2SO_4 \rightarrow BaSO_4 + 2H_2O$	1

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Question	Answer	Marks
3(a)	the potential difference between two half-cells/two electrodes (in a cell)	1
	under standard conditions of 1 atm., 298 K, (all) solutions being 1 mol dm ⁻³	1
3(b)(i)	8 marking points, any 2 points for each mark H ₂ / hydrogen correct delivery system for H ₂ Pb ²⁺ (aq) Pb electrode Pt electrode H*(aq) solution salt bridge voltmeter/V labelled	4
3(b)(ii)	more negative	1
	shifts Pb^{2+} (+ $2e^{-}$) \Rightarrow Pb equilibrium/reaction to the left	1

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Question	Answer	Marks
3(c)(i)	3(c)(i) Q = $0.4 \times 80 \times 60 = 1920$ C and use of $96500/193000$ Moles of Pb = $1920/193000 = 9.95 \times 10^{-3}$ Mass of Pb = $207.2 \times 9.95 \times 10^{-3} = 2.1$ g	
	OR $Q = 0.4 \times 80 \times 60 = 1920 \text{C}$ and use of $1.6 \times 10^{-19} / 1.2 \times 10^{22}$ atoms Pb = 6×10^{21} ; moles of Pb = $6 \times 10^{21} / 6 \times 10^{23} = 0.01$ Mass of Pb = $207.2 \times 0.01 = 2.1 \text{g}$	
3(c)(ii)	$PbO_2(s) + SO_4^{2-}(aq) + 4H^+ + 2e^- \rightarrow PbSO_4(s) + 2H_2O$	1
3(d)	reagents/PbO ₂ /H ₂ SO ₄ and used up/concentration decreases	1
	as fuel/hydrogen is being continuously supplied/fuel has not run out	1

Question	Answer	Marks			
4(a)	density is higher and melting point is higher				
	(density) due to A_r being larger and smaller atomic radii or (Co) atoms/ions heavier and smaller				
	(melting point) due to stronger attraction to cations as more delocalised electrons	1			
4(b)	(a molecule or ion) formed by a central metal atom/ion surrounded by (one or more) ligands	1			
4(c)(i)	same number and type of atoms and different structural formula	1			

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Question			Answer		Marks
4(c)(ii)	octahedral AND 3D structure of e.g. NH ₃ H ₃ N Co Br H ₃ H ₃ H ₃	NH_3 N_{IIII_1} NH_3 N_3 N_4 N_4 N_4 N_4 N_4 N_5 N	NH ₃ Co Br NH ₃		1
4(c)(iii)	co-ordinate / dative covalent				1
4(c)(iv)	+3 for both				1
4(d)	(HNO ₃) Ag ⁺ /AgNO ₃ cream(–ye	ellow) ppt. (of AgBr) and n	o reaction/white pp	t. for other isomer	1
	Ba(OH) ₂ /Ba ²⁺ (aq)/BaCl ₂ /Ba	(NO ₃) ₂ white ppt. (of BaSC	D ₄) and no reaction f	or other isomer	1
4(e)	(d-d) energy gap / ∆E is differe	nt			1
	absorb different wavelength/	frequency (of light)			1
4(f)			heterogeneous	homogeneous	2
	F	e in the Haber process	✓		
		Fe ²⁺ in the I ⁻ /S ₂ O ₈ ²⁻ reaction		√	
		NO ₂ in the oxidation of SO ₂		√	
		V ₂ O ₅ in the Contact process	√		

Question	Answer	Marks
5(a)	nitrile; alkene; chloro; benzene / arene	
5(b)		1
	addition (polymerisation)	1

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Question		Answer		Marks
5(c)		I		- Nanc
	reagent	structure of product	type of organic reaction	
	excess Br ₂ (aq)	CI H CN CN Br Br [1]	(electrophilic) addition	
	excess hot, conc. MnO ₄ ⁻ (aq)	CI HO O CN COOH O COOH [1] + [1]	oxidation	
	excess hot, aqueous HC <i>l</i>	С <i>I</i> Н С СООН СООН [1]	hydrolysis	
	excess H₂/Pt catalyst	both CH ₂ NH ₂ formed [1] both arene and alkene reduced [1]	reduction/ hydrogenation	
		structures [6]	2 correct for 1 mark total [2]	

Question	Answer	Marks
6(a)(i)	CH ₃ NO ₂	1
6(a)(ii)	$HNO_3 + 2H_2SO_4 \rightarrow H_3O^+ + NO_2^+ + 2HSO_4^-$	1
6(a)(iii)	any three from:	3
	 Point 1: bonds/electrons are partially delocalised in T or delocalised /π system/π bonding extends over only five carbons Point 2: four π-electrons in the (delocalised system of T) or methylbenzene has (two) more π-electrons/(two) more delocalised electrons Point 3: contains a carbon that is sp³ hybridised in T or (all the) carbons are sp² hybridised in methylbenzene Point 4: one carbon has a bond angle of 109.5°/tetrahedral (in T) or (C-C) bond strengths/lengths are not all the same or not all the bond angles are 120° (in T) 	
6(b)(i)	4-aminobenzoic acid	1
6(b)(ii)	step 1 Sn + HC l [1] concentrated/reflux/heat [1] step 2 CH ₃ COC l [1] step 3 KMnO ₄ /manganate(\underline{VII})/MnO ₄ ⁻ (acidified/alkaline) and heat [1] step 4 aqueous HC l and heat [1] step 5 ethanol, H ₂ SO ₄ , concentrated/reflux/heat [1]	6

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Question		Ansv	ver		Marks
6(c)	(benzocaine) is less (basi	c than ethylamine) AND vailable to accept a proton/H ⁺			
	since (lone pair on N) is d or phenyl ring is electron				
	OR ethylamine is more basic lone pair (on N) is more a	(than benzocaine) AND available to accept a proton/H ⁺			
	since ethyl/alkyl group is	electron-donating group			
6(d)(i)	7 peaks				
6(d)(ii)	CDC l ₃ will produce no sig or CHC l ₃ would produce a	nal in the spectrum a signal/would be detected			
6(d)(iii)	δ/ppm	group responsible for the peak	number of H atoms responsible for the peak	splitting pattern	
	1.2	CH ₍₃₎	3	triplet	
	3.5	CH ₍₂₎ O	2	quartet	
	5.5	NH ₂	2	singlet (broad)	
	7.1–7.4	H attached to aromatic/benzene ring	4	multiplet	
6(d)(iv)	neighbouring/adjacent carbon atom has two protons/H (attached to it) or there is an adjacent CH ₂ (O) group				
6(d)(v)	peak at 5.5/NH ₂ peak will and NH ₂ /protons exchange/s	• •			

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Question	Answer	Marks
6(e)(i)	NaNO ₂ + HC <i>l</i> or HNO ₂	1
6(e)(ii)	CO ₂ C ₂ H ₅ CO ₂ C ₂ H ₅ N R S OH	
	structure of diazonium salt R	1
	structure of azo dye S	1

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Question	Answer	Marks
7(a)	Fe atom= (1s ² 2s ² 2p ⁶)3s ² 3p ⁶ 3d ⁶ 4s ²	1
	Fe^{3+} ion= $(1s^22s^22p^6)3s^23p^63d^5$	
7(b)	$([H^+]^2 = 8.9 \times 10^{-4} \times 0.25 \text{ or } 2.225 \times 10^{-4})$ $[H^+] = 0.0149$	1
	pH = -log(0.0149) = 1.83	1
7(c)(i)	(K _{stab} is) the equilibrium constant for the formation of a complex (ion) (in a solvent from its constituent ions/molecules)	1
7(c)(ii)	$[Fe(H_2O)_5F]^{2+}$ and $[Hg(H_2O)_5Cl]^+$	1
7(d)	$K_{\text{stab}} = \frac{[\text{Fe}(\text{ed})_2 \text{C}l_2^{3-}]}{[\text{Fe}(\text{H}_2\text{O})_4 \text{C}l_2^+][\text{ed}]^2}$	1
	$\mathrm{mol}^{-2}\mathrm{dm}^{6}$	1
7(e)(i)	CIMINATION OCCUPANTION OCCUPAN	3

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Question	Answer	Marks
7(e)(ii)	any cis isomer and the trans isomer identified	1
7(e)(iii)	both correct cis isomers identified	1
7(e)(iv)	trans isomer identified	1

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