

Monday 15 June 2015 - Afternoon

A2 GCE CHEMISTRY A

F325/01 Equilibria, Energetics and Elements

Candidates answer on the Question Paper.

OCR supplied materials:

Data Sheet for Chemistry A (inserted)

Other materials required:

Scientific calculator

Duration: 2 hours



Candidate forename				Candidate surname			
Centre numb			Candidate nu	ımber			

INSTRUCTIONS TO CANDIDATES

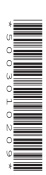
- The Insert will be found inside this document.
- Write your name, centre number and candidate number in the boxes above. Please write clearly and in capital letters.
- Use black ink. HB pencil may be used for graphs and diagrams only.
- Answer all the questions.
- Read each question carefully. Make sure you know what you have to do before starting your answer.
- Write your answer to each question in the space provided. If additional space is required, you should use the lined page at the end of this booklet. The question number(s) must be clearly shown.
- Do not write in the bar codes.

INFORMATION FOR CANDIDATES

- The number of marks is given in brackets [] at the end of each question or part question.
- Where you see this icon you will be awarded marks for the quality of written communication in your answer.

This means, for example, you should:

- ensure that text is legible and that spelling, punctuation and grammar are accurate so that meaning is clear;
- organise information clearly and coherently, using specialist vocabulary when appropriate.
- You may use a scientific calculator.
- A copy of the Data Sheet for Chemistry A is provided as an insert with this question paper.
- You are advised to show all the steps in any calculations.
- The total number of marks for this paper is 100.
- This document consists of **24** pages. Any blank pages are indicated.

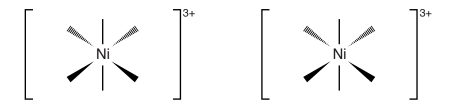


Answer all the questions.

This	s que	estion looks at properties of transition elements, ions and complexes.
(a)	Wh	at is the oxidation number of Cr in the complex ion $[CrOCl_5]^{2-}$?
		[1]
(b)	Wri	te the equation for a reaction catalysed by a named transition element, compound or ion.
	Equ	uation:
	Cat	alyst:[1]
(c)	An	octahedral complex ion ${\bf A}$, ${\bf C_9H_{30}N_6Ni^{3+}}$, exists as two optical isomers.
	In c	complex ion A , Ni ³⁺ is bonded to three molecules of a bidentate ligand B .
	(i)	State what is meant by a bidentate ligand.
		[1]
	(ii)	What is the molecular formula of the bidentate ligand B ?
		[1]
	(iii)	Draw a possible structure for B and explain how B is able to act as a bidentate ligand.
		[2]
	(iv)	What is the coordination number of complex ion A ?
		[1]

1

(v)	Complete the 3-D diagrams of the shapes of the optical isomers of complex ion ${\bf A}$
	You can show the bidentate ligand simply as



[1]

- (d) Describe the reactions of **EITHER** aqueous copper(II) ions **OR** aqueous cobalt(II) ions with:
 - aqueous sodium hydroxide
 - excess aqueous ammonia

In your answer you should link observations with equations.

• hydrochloric acid.

[6]

[Total: 14]

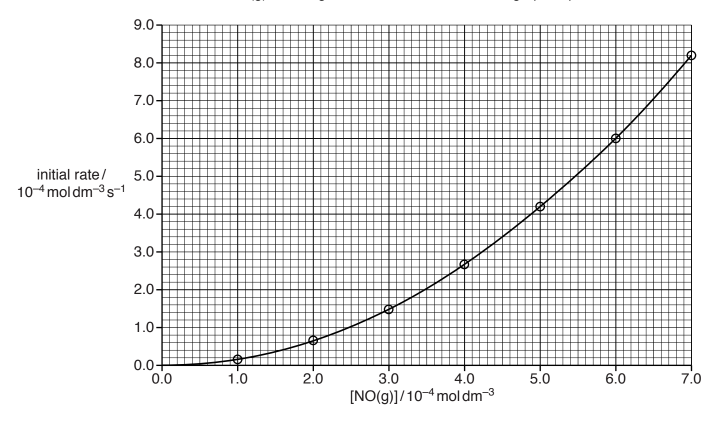
2 Hydrogen, H₂, reacts with nitrogen monoxide, NO, as shown below:

$$2\mathsf{H}_2(\mathsf{g}) + 2\mathsf{NO}(\mathsf{g}) \longrightarrow \mathsf{N}_2(\mathsf{g}) + 2\mathsf{H}_2\mathsf{O}(\mathsf{g})$$

(a) The rate equation for this reaction is:

$$rate = k[H_2(g)][NO(g)]^2$$

The concentration of NO(g) is changed and a rate-concentration graph is plotted.



The chemist uses $H_2(g)$ of concentration $2.0 \times 10^{-2} \, \text{mol dm}^{-3}$.

Using values from the graph, calculate the rate constant, k, for this reaction.

Give your answer to two significant figures and in standard form.

Show your working.

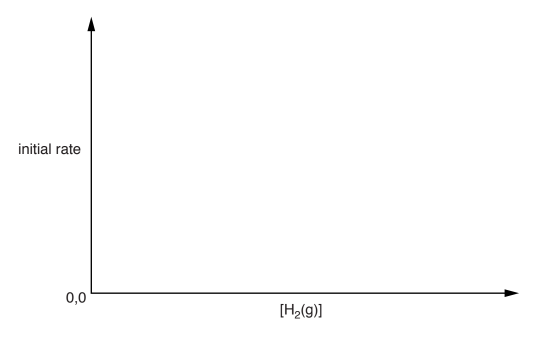
(b) A chemist investigates the effect of changing the concentration of H₂(g) on the initial reaction rate at two different temperatures.

The reaction is first order with respect to $H_2(g)$.

(i) Using the axes below, sketch two graphs of the results.

Label the graphs as follows:

- L for the lower temperature
- **H** for the higher temperature.



[2]

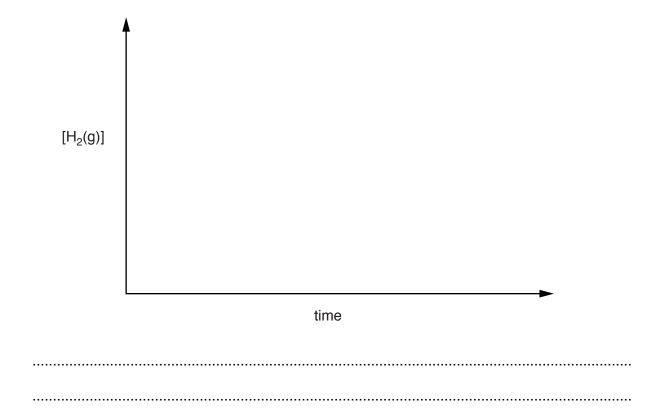
(ii) State the effect of the higher temperature on the rate constant, k.

.....[1]

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(c)	The reaction	can also	be shown	n as being	first order	with	respect to	$H_2(g)$ by	continuous
	monitoring of	[H ₂ (g)] dı	uring the c	ourse of th	e reaction.			_	

- Using the axes below, sketch a graph to show the results.
- State how you would use the graph to show this first order relationship for $H_2(g)$.



(d) The chemist proposes a three-step mechanism for the reaction:

$$2H_2(g) + 2NO(g) \longrightarrow N_2(g) + 2H_2O(g)$$

.....[2]

On the dotted line below, write the equation for step 3.

step 1:	$2NO \rightarrow N_2O_2$	fast	
step 2:	$H_2 + N_2O_2 \rightarrow N_2O + H_2O$	slow	
step 3:		fast	[1]

Explain why this mechanism is consistent with the rate equation $rate = k[H_2(g)][NO(g)]^2$.

[Total: 11]

This question looks at two reactions involving sulfur compounds.

(a)	Hyd	lrogen rea	acts with carbon dis	ulfide as shov	vn below.			
			$4H_2(g) + 0$	$CS_2(g) \rightarrow CF$	I ₄ (g) + 2H ₂ S	6(g)		
	For	this reacti	ion, $\Delta H = -234 \mathrm{kJm}$	$_{ m lol}^{-1}$ and $_{ m }$ $_{ m S}$ =	= -164JK ⁻¹	mol ^{−1} .		
	(i)	Why doe	s the reaction have	a negative e	ntropy chan	ge?		
							[1]
	(ii)	Standard	d entropies are show	wn in the table	e below.			
			substance	CS ₂ (g)	CH ₄ (g)	H ₂ S(g)		
			S ^o /JK ⁻¹ mol ⁻¹	238	186	206		
		Calculate	e the standard entro	opy for H ₂ .				
							JK ⁻¹ mol ⁻¹ [2]
	(iii)	Explain, v	with a calculation, v	whether this re	eaction is fe	asible at 25	°C.	
		Show you	ur working.					
							[3]
	(iv)		with a calculation, t				_	
		rodollorii						
		•••••						···
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(b) A chemist investigated methods to improve the synthesis of sulfur trioxide from sulfur dioxide and oxygen.

$$2SO_2(g) + O_2(g) \rightleftharpoons 2SO_3(g)$$

The chemist:

- mixed together 1.00 mol $\rm SO_2$ and 0.500 mol $\rm O_2$ with a catalyst at room temperature compressed the gas mixture to a volume of 250 cm³
- allowed the mixture to reach equilibrium at constant temperature and without changing the total gas volume.

At equilibrium, 82.0% of the SO₂ had been converted into SO₃.

Determine the concentrations of SO_2 , O_2 and SO_3 present at equilibrium and calculate $K_{\rm c}$ for this reaction.

K_{c}	=	units	[6]
---------	---	-------	----	---

(ii)	Explain what would happen to the pressure as the system was allowed to reach equilibrium.
	[1]
(iii)	The value of $K_{\rm c}$ for this equilibrium decreases with increasing temperature.
	Predict the sign of the enthalpy change for the forward reaction. State the effect on the equilibrium yield of ${\rm SO_3}$ of increasing the temperature at constant pressure.
	Δ <i>H</i> :
	Effect on SO ₃ yield:[1]
(iv)	The chemist repeated the experiment at the same temperature with 1.00 mol ${\rm SO_2}$ and an excess of ${\rm O_2}$. The gas mixture was still compressed to a volume of 250 cm ³ .
	State and explain, in terms of $K_{\rm c}$, how the equilibrium yield of ${\rm SO_3}$ would be different from the yield in the first experiment.
	[3]
	[Total: 19]

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A student is supplied with 0.500 mol dm $^{-3}$ potassium hydroxide, KOH, and 0.480 mol dm $^{-3}$ propanoic acid, $\rm C_2H_5COOH$.
The acid dissociation constant, K_a , for C_2H_5COOH is $1.35 \times 10^{-5}\mathrm{moldm^{-3}}$.
(a) C ₂ H ₅ COOH is a weak Brønsted–Lowry acid.
What is meant by a weak acid and Brønsted-Lowry acid?
[1]
(b) Calculate the pH of 0.500 mol dm ⁻³ potassium hydroxide.
pH =[2]
(c) The student dilutes 25.0 cm ³ 0.480 mol dm ⁻³ C ₂ H ₅ COOH by adding water until the tota volume is 100.0 cm ³ .
(i) Write the expression for K_a for C_2H_5COOH .
[1]
(ii) Calculate the pH of the diluted solution.
(ii) Calculate the pirior the diluted solution.

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pH =[3]

(d)	Aqu	leous propanoic acid, $\mathrm{C_2H_5COOH}$, reacts with carbonates and alkalis.
	(i)	Write the full equation for the reaction of aqueous propanoic acid with sodium carbonate.
		[1]
	(ii)	Write the ionic equation for the reaction of aqueous propanoic acid with aqueous potassium hydroxide.
		[1]
(e)	A st	tudent prepares a buffer solution containing propanoic acid $\rm C_2H_5COOH$ and propanoate s, $\rm C_2H_5COO^-$. The concentrations of $\rm C_2H_5COOH$ and $\rm C_2H_5COO^-$ are both 1.00 mol dm ⁻³ .
	The	following equilibrium is set up.
		$C_2H_5COOH(aq) \rightleftharpoons C_2H_5COO^-(aq) + H^+(aq)$
	The	acid dissociation constant, K_a , for C_2H_5COOH is $1.35 \times 10^{-5} \mathrm{mol dm^{-3}}$.
	(i)	Calculate the pH of this buffer solution.
		Give your answer to two decimal places.
		pH =[1]
	(ii)	A small amount of aqueous ammonia, NH ₃ (aq), is added to the buffer solution.
		Explain, in terms of equilibrium, how the buffer solution would respond to the added $\mathrm{NH_3}(\mathrm{aq}).$
		[2]

(iii)	The student adds 6.075 g Mg to 1.00 dm ³ of this buffer solution.
	Calculate the pH of the new buffer solution.
	Give your answer to two decimal places

pH =[4]

[Total: 16]

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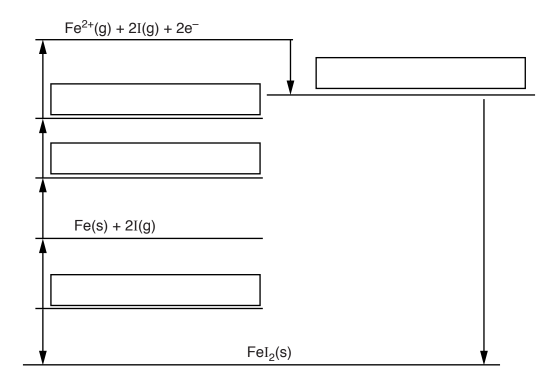
- 5 Iron(II) iodide, FeI_2 , is formed when iron metal reacts with iodine.
 - (a) The table below shows enthalpy changes involving iron, iodine and iron(II) iodide.

	Enthalpy change / kJ mol ⁻¹	
Formation of iron(II) iodide	-113	
1st electron affinity of iodine	-295	
1st ionisation energy of iron	+759	
2nd ionisation energy of iron	+1561	
Atomisation of iodine	+107	
Atomisation of iron	+416	

(i) The incomplete Born–Haber cycle below can be used to determine the lattice enthalpy of iron(II) iodide.

In the boxes, write the species present at each stage in the cycle.

Include state symbols for the species.



Define the term lattice enthalpy.	
	. [2]
Calculate the lattice enthalpy of iron(II) iodide.	

lattice enthalpy = kJ mol⁻¹ [2]

(b) Some electrode potentials for ions are shown below.

Fe ²⁺ (aq) + 2e ⁻	$\stackrel{\textstyle \longleftarrow}{\rightleftharpoons}$	Fe(s)	$E^{\circ} = -0.44 \text{V}$
Fe ³⁺ (aq) + e ⁻		Fe ²⁺ (aq)	$E^{\circ} = +0.77 \text{V}$
$\frac{1}{2}I_{2}(aq) + e^{-}$	=	I [–] (aq)	$E^{\bullet} = +0.54V$
$\frac{1}{2}Br_{2}(aq) + e^{-}$		Br [–] (aq)	$E^{\bullet} = +1.09V$
$\frac{1}{2}Cl_{2}(aq) + e^{-}$		C <i>l</i> [–] (aq)	$E^{\bullet} = +1.36V$

(i)	Complete the electron configurations for Fe ²⁺ and Br ⁻ .	
	Fe ²⁺ : 1s ²	
	Br ⁻ : 1s ² [2) 1
(ii)	Predict the products of reacting Fe(s) separately with $I_2(aq)$, $Br_2(aq)$ and $Cl_2(aq)$.	.,
(,	Explain your predictions using the electrode potential data above.	
	Explain your productions doing the discusses potential data above.	
		•
	[3	;]

(c) Fe $^{2+}$ ions can be used to test for NO $_3^-$ ions. In this test, aqueous iron(II) sulfate is added to a solution containing NO $_3^-$ ions, followed by slow addition of concentrated sulfuric acid. The sulfuric acid forms a layer below the aqueous

In the presence of NO_3^- ions, a brown ring forms between the two layers.

Two reactions take place.

Reaction 1: In the acid conditions ${\rm Fe^{2+}}$ ions reduce ${\rm NO_3^-}$ ions to NO. ${\rm Fe^{2+}}$ ions are oxidised to ${\rm Fe^{3+}}$ ions.

Water also forms.

Reaction 2: A ligand substitution reaction of $[Fe(H_2O)_6]^{2+}$ takes place in which one NO ligand exchanges with one water ligand. A deep brown complex ion forms as the brown

[Total: 16]

Construct equations for these two reactions.

eaction 1:
eaction 2:
[3

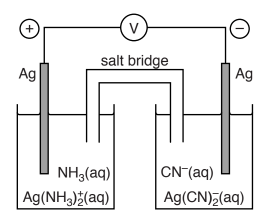
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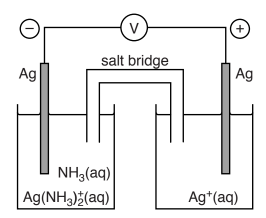
6 Three redox systems, C, D and E are shown in Table 6.1.

С	Ag(NH ₃) ₂ +(aq) + e ⁻	$H_3)_2^+(aq) + e^- \iff Ag(s)$	
D	Ag+(aq) + e-	\rightleftharpoons	Ag(s)
Е	Ag(CN) ₂ ⁻ (aq) + e ⁻	\rightleftharpoons	Ag(s) + 2CN ⁻ (aq)

Table 6.1

The two cells below were set up in an experiment to compare the standard electrode potentials of redox systems **C**, **D** and **E**. The signs on each electrode are shown.





(a) List the three redox systems in order by adding the labels C, D and E to the table below.

E♥	redox system
Most negative	
1	
Least negative	

[1]

	tandard cell is set up between redox system ${\bf D}$ in ${\bf Table~6.1}$ and a standard hydrogen half. The standard cell potential of redox system ${\bf D}$ is +0.34 V.
	cell delivers a current for a length of time. pH of the solution in the standard hydrogen half-cell decreases.
(i)	What is the pH of the solution in a standard hydrogen half-cell?
	pH =[1]
(ii)	Explain, in terms of electrode potentials and equilibrium, why the pH of the solution in the hydrogen half-cell decreases as this cell delivers current.
(iii)	Write the equation for the overall cell reaction that takes place in this cell.
()	[1]
The	CN ⁻ ion is the conjugate base of a very toxic weak acid.
In a	queous solutions of CN ⁻ ions, an acid-base equilibrium is set up.
(i)	Complete the equation for this equilibrium and label the conjugate acid-base pairs.
	CN^- + H_2O \Longrightarrow +
	[1]
(ii)	Explain, in terms of equilibrium, why acidic conditions should not be used with cells containing CN ⁻ (aq) ions.
	cell. The The (i) (iii)

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.....[1]

		20
(d)		ct-ethanol fuel cells (DEFCs) are being developed in which the fuel is ethanol rather than rogen.
	The	half-equation for the reaction at the ethanol electrode of the DEFC is shown below:
		$C_2H_5OH + 3H_2O \longrightarrow 2CO_2 + 12H^+ + 12e^-$
	(i)	State one important difference between a fuel cell and a modern storage cell.
		[1]
	(ii)	Suggest one advantage of using ethanol, rather than hydrogen, in a fuel cell for vehicles.
		[1]

	[[1]
(iv)	Deduce the half-equation for the reaction at the oxygen electrode in a DEFC.	

Write the equation for the overall reaction in a DEFC.

(iii) The overall reaction in a DEFC is the same as for the complete combustion of ethanol.

		. [1]
(v)	Using oxidation numbers, show that oxidation and reduction take place in a DEFC.	
	Oxidation:	
	Reduction:	
		[2]

[Total: 13]

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Turn over for the next question

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Chromite is the main ore of chromium. The chromium-containing compound in chromite is Fe(CrO₂)₂. The percentage of chromium in a sample of chromite can be determined using the method below.

Step 1

A 5.25 g sample of chromite ore is heated with sodium peroxide, Na₂O₂.

$$2Fe(CrO_2)_2 + 7Na_2O_2 \rightarrow 2NaFeO_2 + 4Na_2CrO_4 + 2Na_2O_2$$

Water is added to the resulting mixture.

Na₂CrO₄ dissolves in the water forming a solution containing CrO₄²⁻ ions.

Step 2

The mixture from Step 1 is filtered and the filtrate is made up to 1.00 dm3 in a volumetric flask.

A 25.0 cm³ sample of this alkaline solution is pipetted into a conical flask and an excess of aqueous potassium iodide is added.

- A redox reaction takes place between $\rm I^-$ ions, $\rm CrO_4^{~2-}$ ions and $\rm H_2O.$ In this reaction 1 mol $\rm CrO_4^{~2-}$ forms 1.5 mol $\rm I_2.$

Step 3

The resulting mixture is titrated with 0.100 mol dm⁻³ sodium thiosulfate, Na₂S₂O₃(aq) to estimate the I2 present:

$$I_2(aq) + 2S_2O_3^{2-}(aq) \rightarrow 2I^{-}(aq) + S_4O_6^{2-}(aq)$$

The average titre of $Na_2S_2O_3(aq)$ is 25.5 cm^3 .

(a) In Step 1 Na₂O and NaFeO₂ react with water forming an alkaline solution containing a brown precipitate. This is **not** a redox reaction.

Write equations for:

- the reaction of Na₂O with water
- the reaction of NaFeO₂ with water.

(D)	Determine the percentage, by mass, of chromium in the ore.	
	Give your answer to one decimal place.	
		[6]
(c)	This part refers to Step 2 of the method.	
	In the redox reaction between I ⁻ ions, CrO ₄ ²⁻ ions and H ₂ O: • CrO ₄ ²⁻ ions, are reduced to chromium(III) ions, Cr ³⁺	
	 I⁻ ions are oxidised to iodine, I₂ Construct an overall equation for the redox reaction and write half equations for oxidation and reduction. 	the
	Overall equation:	
	Half equations:	
	[Total:	[3] 111
	[10tal.	

ADDITIONAL ANSWER SPACE

If additional answer space is required, you should use the following lined page. The question number(s) must be clearly shown in the margins.	
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