Mark Scheme - 1.7 Equilibria and Acid-Base Reactions

1. Benefits: (a) Stops fossil fuels from running out Reduces CO₂ emissions / greenhouse emissions / global warming / effect of global warming Reduces SO₂ emissions / acid rain There will be an investment in new technology Difficulties: Dependence on fossil fuel/Unlikely to meet current demand Renewable energy currently more expensive Reliability of supply from renewables Major development in energy efficiency technologies required Opposition by vested interests (Maximum3 marks from list, but need examples of both) (3) Consideration and discussion of benefits/difficulties (1) [4] QWC Legibility of text; accuracy of spelling, punctuation and grammar, clarity of meaning (b) I As temperature increases yield decreases As pressure increases yield decreases [1] II As temperature is increased, equilibrium moves to the left (1) Therefore forward reaction is exothermic (1) As pressure is increased, equilibrium moves to the left (1) Therefore more gas moles in products (1) [4] QWC The information is organised clearly and coherently, using specialist vocabulary where appropriate [1] (ii) If temperature is too low, then reaction is too slow (1) If temperature is too high, yield is too low (1) Compromise temperature – acceptable rate and yield (1) (Accept any two points) [2] Heterogenous catalyst (iii) [1] Lower temperatures could be used (1) (iv) Less energy consumption/increased yield (1) Equilibrium could be reached more quickly (1) (Accept any two points) [2] (v) curve (1) Energy [2] $\Delta H(1)$ Extent of reaction [1] $\Delta H = E_t - E_b$ (vi)

Total [19]

(a)		ction in which) the rate of the forward reaction is equal to the rate backward reaction	[1]			
(b)	goes darker / more brown (1)					
	because the (forward) reaction has a +ve ΔH / is endothermic (1)					
	goes paler / less brown (1)					
	becau	because there are more moles / molecules on RHS (1)				
	no ch	ange (because catalysts do not affect the position of an equilibrium)	(1) [5]			
(c)	(i)	moles $N_2H_4 = 14000/32.04 = 437.0$ (1)				
		this produces $437.0 \times 3 = 1311$ moles of gas (1)				
		volume = $1311 \times 24 = 3.15 \times 10^4 \text{ dm}^3$ (1) [minimum 2 sf]	[3]			
	(ii)	(large volume of) gas produced	[1]			
(d)	(i)	an acid is a proton / H+ donor	[1]			
	(ii)	$\rightarrow NO_2^- + H_3O^+$	[1]			
	(iii)	sulfuric acid is behaving as the acid / nitric acid is behaving as a base (1)				
		as it donates a proton / as it accepts a proton (1)	[2]			

Total [14]

(a)	(i)	1	burette / (graduated) pipette	[1]	
		11	volumetric / graduated / standard flask	[1]	
	(ii)	0.0	0064	[1]	
	(iii)	1.2	20 g / 100 cm ³ solution	[1]	
	(iv)	12	.0 g / 100 cm ³ solution	[1]	
(b)	(i)		ne rate of the forward reaction is equal to t		
	(ii)		action.	[1]	
	(ii)	C	H ₄ O	[1]	
4.					
Ac	id: Pr	oto	n donor (1)		
1990			quilibrium: Reversible reaction v equal (1)	where the rate of forward and reverse	[2]
5.					
(a)	(sod	um) chlorate(I) as bleach/ (sodium)	mportant chlorine containing compound e.g chlorate(V) as weedkiller/ aluminium chlori	
	С	atai	yst in halogenation - do not accept CFCs		[1]
(b)	(i	i)	$K_c = \frac{[HI]^2}{[H_2][I_2]}$ must be	square brackets	[1]
	(i	ii)	$K_c = \frac{0.11^2}{3.11^2} = 1.25 \times 10^{-3}$	follow through error (ft)	[1]
	(i	iii)	K _c has no units	ft	[1]
	(i	iv)	when temperature increases $K_{\rm c}$	ncreases (1)	
			this means equilibrium has move / increasing temperature favours		
			therefore ΔH for forward reaction (mark only awarded if marking p		[3]

(a) (i) Cold
$$Cl_2 + 2NaOH$$
 NaCl + NaClO + H_2O (1) Warm $3Cl_2 + 6NaOH$ 5NaCl + NaClO₃ + $3H_2O$ (1)

[2]

(ii) Disproportionation [1]

(b) (i)
$$K_c = [CH_3COOCH_3][H_2O]$$
 (1) $[CH_3COOH][CH_3OH]$

No units

(1) [2]

(ii) moles =
$$\frac{1.25 \times 32.0}{1000}$$
 = 0.04(0) [1]

(iii) [CH₃COOH] = 0.04, therefore 0.06 used in reaction and

$$[CH_3COOCH_3] = 0.06, [H_2O] = 0.06$$
and

$$[CH3OH] = 0.083 - 0.06 = 0.023$$
 (1)

$$K_c = \frac{0.06 \times 0.06}{0.04 \times 0.023} = 3.91$$
 (1) [2]

 (iv) Value of K_c decreases since the equilibrium shifts to the left / the forward reaction is exothermic [1]

7.

(a)

(4)						
	1	2	3	4		
Volume used / cm ³	20.75	20.20	20.10	20.30		

[1]

(b) 20.20 cm³ [1]

(a)	(i)	$K_p = \frac{pSO_3(g) \times pNO(g)}{pSO_2(g) \times pNO_2(g)}$ (1) there are no units (1) [2]]
(ii)	(ii)	The line for SO ₃ / NO at equilibrium should be above the SO ₂ / NO ₂ line (1) as K _p has a value of 2.5, the partial pressures of SO ₃ and NO at equilibrium will be greater than the partial pressures of SO ₂ and NO ₂ . (1) - accept answer in terms of alternative calculated K ₂ with	
		The line for equilibrium should start at 9 hours. (1) as at equilibrium the concentrations is unchanged as time progresses. (1)	
		[4] There may be other acceptable forms of explanation to be discussed at the conference	
(i	ii)	If the temperature rises then the position of equilibrium will move to the left, (reducing the quantities of SO ₃ and NO). (1)	
		This will make the value of K_p smaller. (1) [2]	
(b)	(i)	Nitric acid is a strong acid and its pH is low / <2 / 1.0 (1) As aqueous ammonia is added the pH slowly rises (1) until a pH of ~3 is reached, when it rises rapidly (1) At a pH of 8-9, it tails off slowly as ammonia is a weak base. (1) Accept any 3 from 4 [3]	
		Selection of a form and style of writing which is appropriate to purpose and to complexity of subject matter [1]	
(ii)	The equivalence point is reached when 20.0 cm ³ of ammonia solution has been added as this is at the mid point of the more vertical section. (1)	
		Since both reagents have the same concentration and the volumes used are both 20 cm ³ / the same, the number of moles of each are the same (1)	
		OR the number of moles of both nitric acid and aqueous ammonia are calculated (0.0020 and shown to be the same (1))
		: Mole ratio must be 1:1 (1) [2]	
(ii		I Ammonium nitrate is the salt of a strong acid and weak base / there is a buffering effection operation. [1] II ~ 6.4 [1]	
(it	V)	Blue, as bromophenol blue is blue at a pH of 4.7 and above [1]	
(c)		Number of moles of ammonium nitrate = $\frac{40}{80}$ = 0.50 (1)	
		- error carried forward Concentration of ammonium nitrate solution = $\frac{0.5 \times 1000}{200}$ = 2.5 mol dm ⁻³ (1)	
		:. Temperature drop = $2.5 \times 6.2 = 15.5$ °C (1) [3]	
		Total [20]	

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x = 10
                                                                                                [1]
(a)
               number of moles = 250 x 0.200 ÷ 1000 = 0.05 mol (1) - ft
(b)
       (i)
               mass of sodium carbonate = 0.05 \times M_r(Na_2CO_3) = 0.05 \times 286.2
                                                                                                [2]
                                      = 14.31g(1)
       (ii)
               any two points from:
               weigh by difference (1)
               add less water initially (1)
               wash out beaker / glass rod / funnel and put water into volumetric flask (1)
               add water up to mark in volumetric flask (1)
                                     - 2 max
                                                                                                [2]
(c)
       add few drops of indicator (1)
                                             do not accept 'universal indicator'
       take initial and final reading on burette (1)
       swirl the conical flask (1)
       add acid until the indicator changes colour (1)
                                                                                                [4]
       QWC: organisation of information clearly and coherently; use of specialist vocabulary
       where appropriate.
                                                                                          QWC [1]
                                                                                         Total [10]
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(a)	$K_w = [H^+][OH^-]$ Units = mol ² dm ⁻⁶		(1) (1)	[2]
(b)	(i)	In pure water [H ⁺] = [OH ⁻] or [H ⁺] = $\sqrt{1.0 \times 10^{-14}}$	(1)	
		$pH = -log \ 10^{-7} = 7$	(1)	[2]
	(ii)	Final volume of solution is 1000 cm ³ so acid has l factor of 100 so final concentration of acid is 0.00		by a
		or moles acid = $0.1 \times 10 = 0.001$	(1)	
		$pH = -\log 0.001 = 3$	(1)	[2]
(c)	1.78	$\times 10^{-5} = [\underline{H^+}] \times 0.02$ 0.01	(1)	
	[H ⁺]	$= 8.90 \times 10^{-6}$	(1)	
	pH	= 5.05 allow 5 or 5.1	(1)	[3]
(d)	Thes	solution is a buffer	(1)	
	Solu	tion contains a large amount of CH3COOH and CH3	COO-ions	
	(Acc	eept correct equations)	(1)	
	Whe	n an acid is added, the CH3COO-ions react with the	H ⁺ ions, rem	noving
	them	from solution and keeping the pH constant	(1)	[3]
			1000	

Total [12]

(a) an acid is a proton / H⁺ donor [1] pH = -log[H⁺] / negative log of hydrogen ion concentration (b) [1] a low pH corresponds to a high concentration of H⁺ (1) (c) a strong acid is totally dissociated whilst a weak acid is partially dissociated (1) need to consider concentration (of acid solution) as well as strength of the acid (1) a concentrated solution of a weak acid could have a lower pH than a dilute solution of a strong acid (1) [4] QWC Accuracy of spelling, punctuation and grammar QWC [1] $K_a = \underline{[HCOO^-][H^+]}$ [HCOOH] (d) [1] $1.75 \times 10^{-4} = \frac{x^2}{0.1} (1)$ (ii) $x = 4.183 \times 10^{-3}$ (1) pH = 2.38(1)[3] buffer (e) (i) [1] RCOOH

RCOOT + H⁺ and RCOONa → RCOOT + Na⁺ (1) (ii) added H⁺ removed by salt anion/ A⁻+ H⁺ → HA (1) added OH removed by acid/ OH + HA → A + H₂O (1) [3] Total [15] 12. (dissociates to) release H⁺ ions [1] (a) (b) 2.5-6.0 [1]