

F325: Equilibria, Energetics and Elements

5.2.3 Electrode Potentials and Fuel Cells

59 marks

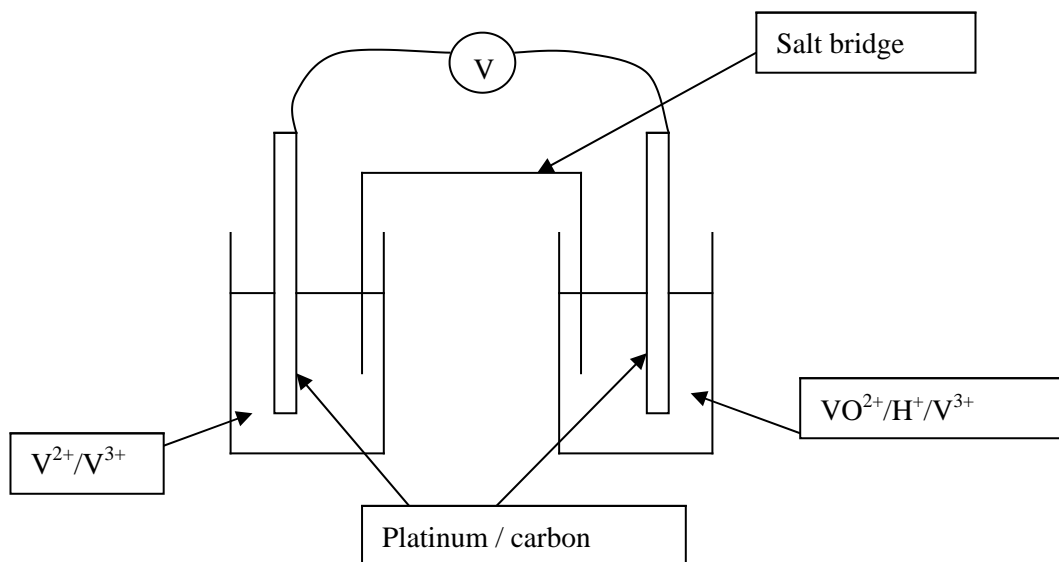
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|----|------|---|-------------|-----|
| 1. | (i) | oxidation: $\text{Fe} \rightarrow \text{Fe}^{2+} + 2\text{e}^-$ (1)
reduction: $\text{V}^{3+} + \text{e}^- \rightarrow \text{V}^{2+}$ (1) | 2 | |
| | (ii) | $E_{\text{cell}} = 0.18 \text{ V}$ (1) | 1 | [3] |
| | | | | |
| 2. | (i) | system III $\times 2$ and reversed + system IV (1)
$2\text{H}_2 + \text{O}_2 \rightarrow 2\text{H}_2\text{O}$ /
$\text{H}_2 + \frac{1}{2}\text{O}_2 \rightarrow \text{H}_2\text{O}$ (1) | 2 | |
| | (ii) | advantages:
only H_2O formed/ non-polluting
greater efficiency (1)

disadvantages:
H_2 difficult to store (1)
H_2 difficult to manufactured initially /
limited life cycle of H_2 adsorber/absorber (1) | 4 | [6] |
| | | | | |
| 3. | (a) | Emf/voltage/potential difference (of electrochemical cell)
comprising a (Cu/Cu^{2+}) half cell combined with a standard
hydrogen electrode | 1 | |
| | | 1 atm, 1 mol.dm ⁻³ , 298K (all 3 needed but can transfer mark if stated in (b)) | 1 | |
| | (b) | Salt bridge and voltmeter
Platinum electrode dipping into 1 mol dm ⁻³ H ⁺
Hydrogen gas feed
(Accept a suitable alternative standard electrode) | 1
1
1 | [6] |
| | | | | |
| 4. | (a) | (i) Stainless steel + corrosion resistance or alloys for tools
+ hardness or other named alloy/use/property
Allow chrome plating with attractive or barrier to corrosion | 1 | |
| | (ii) | Chromium $1\text{s}^2 2\text{s}^2 2\text{p}^6 3\text{s}^2 3\text{p}^6 3\text{d}^5 4\text{s}^1$ (allow...4s ¹ 3d ⁵) | 1 | |

- (b) (i) $\text{Cr}_2\text{O}_7^{2-} + 14\text{H}^+ + 6\text{Fe}^{2+} \rightarrow 2\text{Cr}^{3+} + 6\text{Fe}^{3+} + 7\text{H}_2\text{O}$ 1
 $\text{Cr}_2\text{O}_7^{2-} / \text{Cr}^{3+}$ has more positive electrode potential 1
Therefore $\text{Cr}_2\text{O}_7^{2-}$ is the stronger oxidising agent which
oxidises Fe^{2+} to Fe^{3+} (ora) 1
(ii) Emf = (+) 0.56 V 1

[6]

5. (a) VO_2^+ 1
(b) (i) B and D 1
(ii)



Allow ecf from (b) (i)
Solutions can be reversed. 4

- (iii) 298 K / 25 °C temperature
all solutions 1 mol dm⁻³
Both needed for 1 mark. Ignore any reference to pressure 1

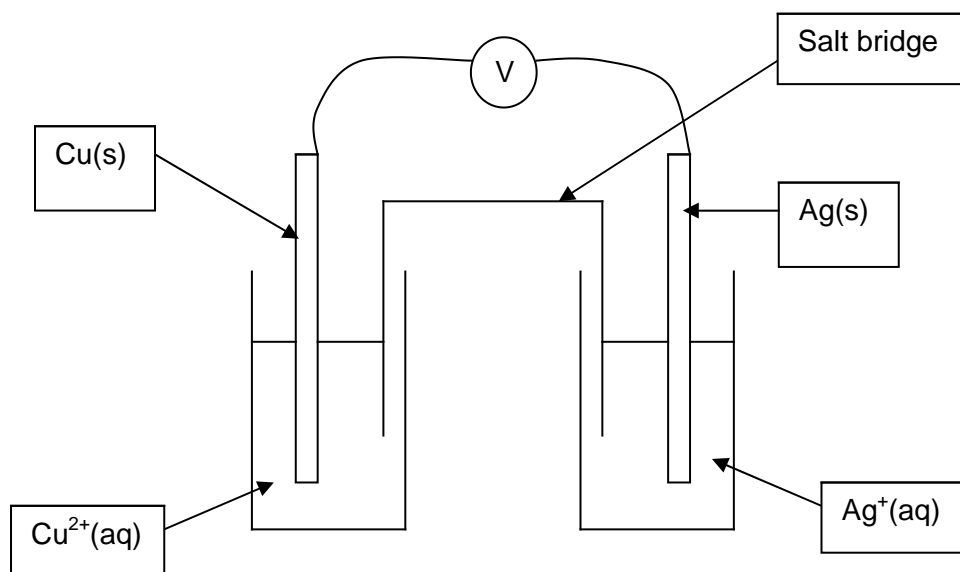
[7]

6. (a) A = Platinum(electrode)
B = $\text{H}^+(\text{aq}) / \text{HCl}(\text{aq}) /$ other suitable acid
C = Voltmeter / galvanometer
D = $\text{Cl}_2(\text{g})$
State symbols needed for B and D
All correct = 2, 3 correct = 1 2

- (b) (i) Arrow marked on or close to wire via voltmeter pointing from hydrogen half cell to chlorine half cell 1
Electrons flow to half cell with more +ve standard electrode potential 1
- (ii) Pressure = 1 Atm / 100 kPa
Temp = 298 K / 25°C
Concentration = 1 mol dm⁻³
All 3 correct = 2 marks 2 correct = 1 mark 2
- (c) The standard electrode potential for ClO₃⁻ / ½Cl₂ is more positive than that of ½Cl₂ / Cl⁻ 1
ClO₃⁻ has a greater tendency to gain electrons than Cl₂ / ClO₃⁻ is a better oxidising agent than Cl₂ 1
Alternative:
Because E° is positive, the reaction will go from left to right therefore ClO₃⁻ is reduced so it must be a better oxidising agent than chlorine.

[8]

7. (a) Emf / voltage / potential difference 1
Half cell combined with standard hydrogen electrode 1
Standard conditions 298K, 1 mol dm⁻³, 1 atm 1
(all 3 required for 1 mark)
- (b) (i) Diagram shows:
Voltmeter + salt bridge + complete circuit 1
Solution labelled Cu²⁺ and electrode labelled Ag 1



- (ii) Direction from Cu(s) to Ag(s) (must be in / close to wire) 1

(iii) $0.80 - 0.34 = 0.46 \text{ V}$ 1

(iv) $\text{Cu} + 2\text{Ag}^+ \rightarrow \text{Cu}^{2+} + 2\text{Ag}$ 1

(c) Standard Electrode Potential for chlorine is more positive than Fe^{3+} therefore it is a better oxidising agent than Fe^{3+} (do not accept E^\ominus is larger or smaller) 1

Standard Electrode Potential for iodine is less positive than Fe^{3+} therefore it is a poorer oxidising agent than Fe^{3+} 1
(Accept release of electrons/equilibrium arguments)

[10]

8. $4\text{NO}_2 + \text{O}_2 + 2\text{H}_2\text{O} \rightarrow 4\text{HNO}_3$ (1)

N from +4 to +5

O from 0 to -2 (1) Could be below equation 2

[2]

9. (a) Emf of a cell / voltage / potential difference / cell potential 1

Comprising half cell combined with standard hydrogen electrode 1

Conc = 1 mol.dm^{-3} ; Pressure (of H_2) = 1 atm; Temp = 298K 1

(all of above = 1 mark)

(b) +0.16 V (unit required) 1

[4]

10. (a) (i) $2\text{MnO}_4^- + 10\text{Cl}^- + 16\text{H}^+ \rightarrow 2\text{Mn}^{2+} + 5\text{Cl}_2 + 8\text{H}_2\text{O}$ 1

correct species on both sides of equation equation balanced 1
(ignore electrons for first mark, penalise for balance)

(ii) Chlorine -1 \rightarrow 0 1

Manganese +7 \rightarrow +2 1

Link to (i) and allow ecf

(iii) Chloride ion oxidised (not chlorine) 1

Manganate(VII) ion reduced (not manganese) 1

(b) 0.16 V too small/rate too slow/insufficient activation energy/not standard conditions 1

[7]