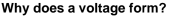
1.11 Redox Equilibria

Electrochemical cells

A cell has two half-cells.The two half cells have to be connected with a salt bridge.

•Simple half cells will consist of a metal (acts an electrode) and a solution of a compound containing that metal (eg Cu and $CuSO_4$).

•These two half cells will produce a small voltage if connected into a circuit. (i.e. become a Battery or cell).



In the cell pictured above

When connected together the zinc half-cell has more of a tendency to oxidise to the Zn^{2+} ion and release electrons than the copper half-cell. ($Zn \rightarrow Zn^{2+} + 2e^{-}$)

More electrons will therefore build up on the zinc electrode than the copper electrode.

A potential difference is created between the two electrodes. The zinc strip is the negative terminal and the copper strip is the positive terminal.

This potential difference is measured with a high resistance voltmeter, and is given the symbol **E**. The E for the above cell is E = +1.1V.

Why use a high resistance voltmeter?

Salt bridge

copper

1M

copper

sulfate

solution

electrode

Electron flow

Zinc

1M zinc

sulfate

solution

electrode

The voltmeter needs to be of very high resistance to stop the current from flowing in the circuit. In this state it is possible to measure the maximum possible potential difference (E).

The reactions will not be occurring because the very high resistance voltmeter stops the current from flowing.

Salt bridge

The salt bridge is used to connect up the circuit. The **free moving ions** conduct the charge.

A salt bridge is usually made from a piece of filter paper (or material) soaked in a salt solution, usually **potassium nitrate**.

The salt should be **unreactive with the electrodes and electrode** solutions. E.g. potassium chloride would not be suitable for copper systems because chloride ions can form complexes with copper ions.

A wire is not used because the metal wire would set up its own electrode system with the solutions.

What happens if current is allowed to flow?

If the voltmeter is removed and replaced with a bulb or if the circuit is short circuited, a current flows. The reactions will then occur separately at each electrode. The voltage will fall to zero as the reactants are used up.

The most positive electrode will always undergo reduction. $Cu^{2+}(aq) + 2e^{-} \rightarrow Cu(s)$ (positive as electrons are used up)

The most negative electrode will always undergo oxidation. $Zn(s) \rightarrow Zn^{2+}(aq) + 2e^{-}$ (negative as electrons are given off)

• The solid vertical line represents the boundary Electrochemical cells can be represented by a cell diagram: between phases e.g. solid (electrode) and solution (electrolyte) $Zn(s) | Zn^{2+}(aq) | | Cu^{2+}(aq) | Cu (s) E = +1.1V$ •The double line represents the salt bridge R between the two half cells Most oxidised form is put next to the double line the voltage produced is indicated •the more positive half cell is written on the right if possible (but this is not essential)

Systems that do not include metals.

If a system does not include a metal that can act as an electrode, then a platinum electrode must be used and included in the cell diagram. It provides a conducting surface for electron transfer. A platinum electrode is used because it is **unreactive** and **can conduct electricity.**

If the system contains several species e.g. for Fe²⁺ (aq) \rightarrow Fe³⁺ (aq) + e⁻ there is no e.g. $MnO_4^- + 8H^+ + 5e^- \rightarrow Mn^{2+} + 4H_2O$ solid conducting surface, a Pt electrode must be then in the cell diagram the balancing numbers, H⁺ used. ions and H₂O can be left out. The cell diagram is drawn as: | | Fe³⁺ (aq), Fe²⁺ (aq) |Pt | | MnO₄- , Mn²⁺ |Pt or if on left hand side Still with more oxidised form near the double line Pt | Mn²⁺ , MnO₄⁻ || A comma separates the oxidised from the reduced species.

If a half equation has several physical states then the solid vertical line should be used between each different state boundary.

4e⁻ + 2H₂O (I) +O₂ (g) \rightarrow 4OH⁻ (aq) \rightarrow || O₂ | H₂O, OH⁻ | Pt

 $Cl_2(g) + 2e^- \rightarrow 2Cl^-(aq)$ || $Cl_2 | Cl^- | Pt$

As the phase line also separates the oxidised and reduced terms a comma is not necessary here.

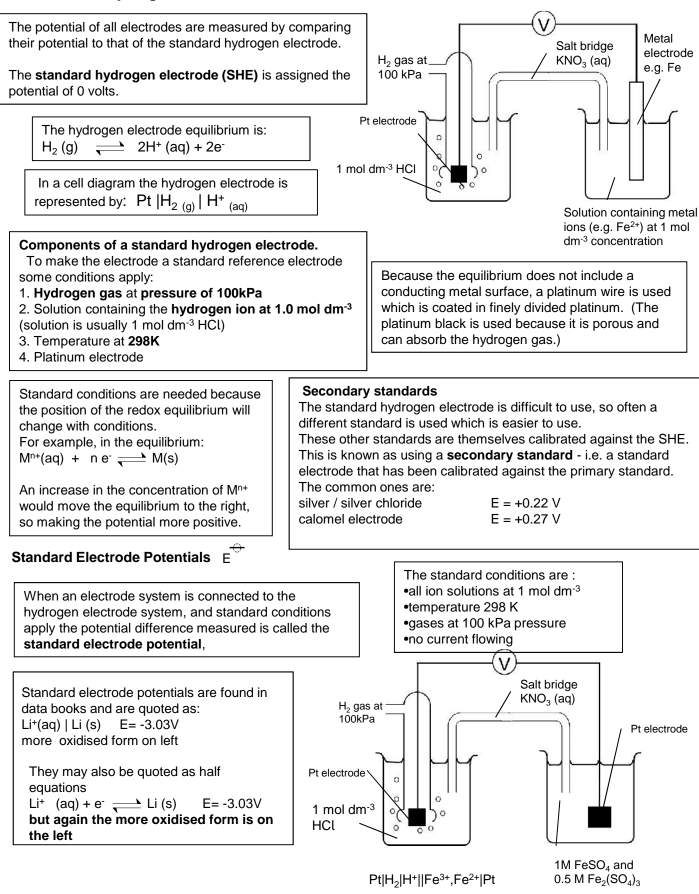
Measuring the electrode potential of a cell

 It is not possible to measure the absolute potential of a half electrode on its own. It is only possible to measure the potential difference between two electrodes.

• To measure it, it has to be connected to another half-cell of known potential, and the potential difference between the two half-cells measured.

 by convention we can assign a relative potential to each electrode by linking it to a reference electrode (hydrogen electrode), which is given a potential of zero Volts

The Standard Hydrogen Electrode



Note: in the electrode system containing two solutions it is necessary to use a platinum electrode and **both** ion solutions must be of a 1 mol dm⁻³ concentration, so $[Fe^{2+}] = 1$ mol dm⁻³ and $[Fe^{3+}] = 1$ mol dm⁻³.

Calculating the EMF of a cell

 $Mg(s) | Mg^{2+}(aq) | | Cu^{2+}(aq) | Cu (s) = +1.1V$

use the equation

Ecell= Erhs - Elhs

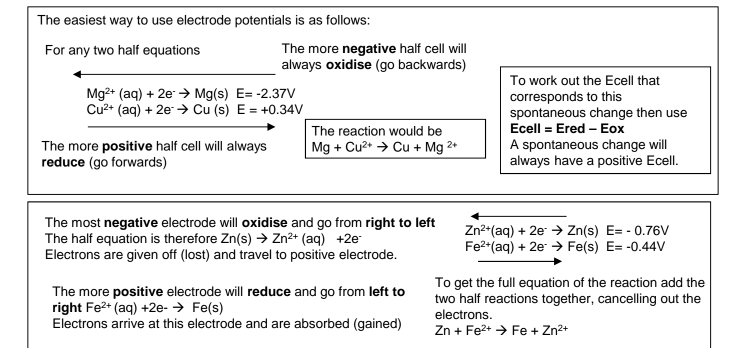
For the cell diagram above Ecell = 0.34 - -2.37= + 2.71 V

Using Electrode Potentials

The most useful application of electrode potentials is to show the direction of spontaneous change for redox reactions.

In order to calculate the Ecell, we must use '**standard** electrode potentials' for the half cells.

Each half cell has a **standard electrode potential** value Mg^{2+} (aq)| Mg(s) E= -2.37V Cu^{2+} (aq) | Cu (s) E = +0.34V



Using series of standard electrode potentials

	oxidation	uio	Most strong reducing agents found here
As more +ve increasing tendency for species on left to reduce, and act as oxidising agents	Li⁺ + e⁻ → Li Mn²+ + 2e⁻ → Mn	-3.03V -1.19V	As more -ve increasing tendency for species on right
	2H ⁺ + 2e ⁻ → H ₂	0V	to oxidise , and act as reducing
	Ag ⁺ + e ⁻ → Ag $F_2 + 2e^- → 2F^-$	+0.8V +2.87	agents
Most strong oxidising agents found here	reduction		To work out the Ecell from two standard electrode potentials then use Ecell = Ered – Eox

The most **powerful reducing agents** will be found at the most **negative** end of the series on the right (i.e. the one with the lower oxidation number). The most **powerful oxidising agents** will be found at the most **positive** end of the series on the left (i.e. the one with the higher oxidation number).

Example 1

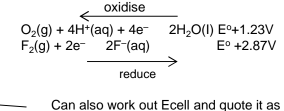
Explanation to write

 $2F_2(g) + 2H_2O(I)$

Use electrode data to explain why fluorine reacts with water. Write an equation for the reaction that occurs.

First apply idea that more positive E° will reduce (go forward) and more negative E° will oxidise (go backwards)

 $4F^{-}(aq) + O_{2}(g) + 4H^{+}(aq)$



Can also work out Ecell and quote it as part of your answer Ecell = Ered - Eox = 2.87-1.23 = 1.64V

Remember to cancel out electrons in full equation

2Cl⁻(aq)

Conventional Cell diagram for above example

As $E^{\circ} F_2/F^- > E^{\circ} O_2/H_2O$, F_2 will oxidise H_2O to O_2

Pt|H₂O|O₂ || F₂|F⁻|Pt

Example 2

Equation

Use data from the table to explain why chlorine should undergo a redox reaction with water. Write an equation for this reaction.

First select relevant half equations by considering the E° values and applying the idea that more positive E° will reduce (go forward) and more negative E° will oxidise (go backwards).

red	uce	
Cl ₂ (aq) + 2e-	2Cl ⁻ (aq)	E°+1.36V
O ₂ (g) + 4H ⁺ (a	lq) + 4e⁻ 2H	₂ O(I) E ^o +1.23V

 $2H_2O(I)$

 $2H_2O(I)$

oxidise

Explanation to write As E° $Cl_2/Cl^{-} > E^{\circ}$ O_2/H_2O , Cl_2 will oxidise H_2O to O_2

Equation $2Cl_2(g) + 2H_2O(I)$ $4Cl^-(aq) + O_2(g) + 4H^+(aq)$

Example 3

Suggest what reactions occur, if any, when hydrogen gas is bubbled into a solution containing a mixture of iron(II) and iron(III) ions. Explain your answer.

First select relevant half equations by considering the E^o values and applying the idea that more positive E^o will reduce (go forward) and more negative E^o will oxidise (go backwards)

Explanation to write

Fe³⁺ will be reduced to Fe²⁺ by H₂ oxidising to H⁺ because E^o Fe³⁺/Fe²⁺ > E^o H⁺/H₂

Example 4

Use the half-equations to explain in terms of oxidation states what happens to hydrogen peroxide when it is reduced.

Explanation to write

As $E^{\circ} H_2O_2/H_2O > E^{\circ} O_2/H_2O_2$, H_2O_2 disproportionates from -1 oxidation state to 0 in O_2 and -2 in H_2O

Fe³⁺ (aq) + e⁻ Fe²⁺ (aq) E° +0.77V 2H+(ag) + 2e-E° 0.00V $H_2(g)$ Fe²⁺ (aq) + 2e⁻ Fe(s) E°-0.44V reduce Fe³⁺ (aq) + e⁻ Fe²⁺ (aq) E°+0.77V 2H+(aq) + 2e- $H_2(g)$ E° 0.00V oxidise

Equation

 $Cl_{2}(aq) + 2e^{-}$

2HOCl(aq) + 2H⁺(aq) + 2e⁻

 $H_2O_2(aq) + 2H^+(aq) + 2e^-$

 $O_2(g) + 4H^+(aq) + 4e^-$

$$2Fe^{3+}(aq) + H_2(g) = 2Fe^{2+}(aq) + 2H^+(aq)$$

 $\begin{array}{c} \xrightarrow{\text{reduce}} \\ H_2O_2(aq) + 2H^+(aq) + 2e^- & 2H_2O(I) \\ O_2(g) + 2H^+(aq) + 2e^- & H_2O_2(aq) \\ \leftarrow & \text{oxidise} \end{array} \xrightarrow{E^\circ + 0.68V}$

Equation

$$2H_2O_2(aq) = 2H_2O(I) + O_2(g)$$

E°+1.36V

E° +1.77V

E° +1.23V

 $Cl_2(aq) + 2H_2O(I) E^{\circ} + 1.64V$

The effects of changing conditions on E cell can be made by applying le Chatelier's principle.

If current is allowed to flow, the cell reaction will occur and the Ecell will fall to zero as the reaction proceeds and the reactant concentrations drop.

Effect of concentration on Ecell

Looking at cell reactions is a straight forward application of le Chatelier. So increasing concentration of 'reactants' would increase Ecell and decreasing them would cause Ecell to decrease.

Effect of temperature on Ecell

Ecell is a measure of how far from equilibrium the cell reaction lies. The more positive the Ecell the more likely the reaction is to occur.

> $Zn^{2+}(aq) + 2e^{-} \rightarrow Zn(s) = -0.76V$ $Fe^{2+}(aq) + 2e^{-} \rightarrow Fe(s) = -0.44V$

 $Zn + Fe^{2+} \rightarrow Fe + Zn^{2+} E= +0.32$

Increasing the concentration of Fe^{2+} and decreasing the concentration of Zn^{2+} would cause Ecell to increase.

Most cells are exothermic in the spontaneous direction so applying Le Chatelier to a temperature rise to these would result in a decrease in Ecell because the equilibrium reactions would shift backwards.

If the Ecell positive it indicates a reaction might occur. There is still a possibility, however, that the reaction will not occur or will occur so slowly that effectively it does not happen. If the reaction has a high activation energy the reaction will not occur.

Cells

Electrochemical cells can be used as a commercial source of electrical energy Cells can be non-rechargeable (irreversible), rechargeable and fuel cells.

You should be able to work out Ecell for given half reactions.

You do not need to learn the details of most of these cells. Relevant cell information will be given. You should be able to convert between standard electrode potential half cells, full cell reactions and cell diagrams and be able to calculate potentials from given data.

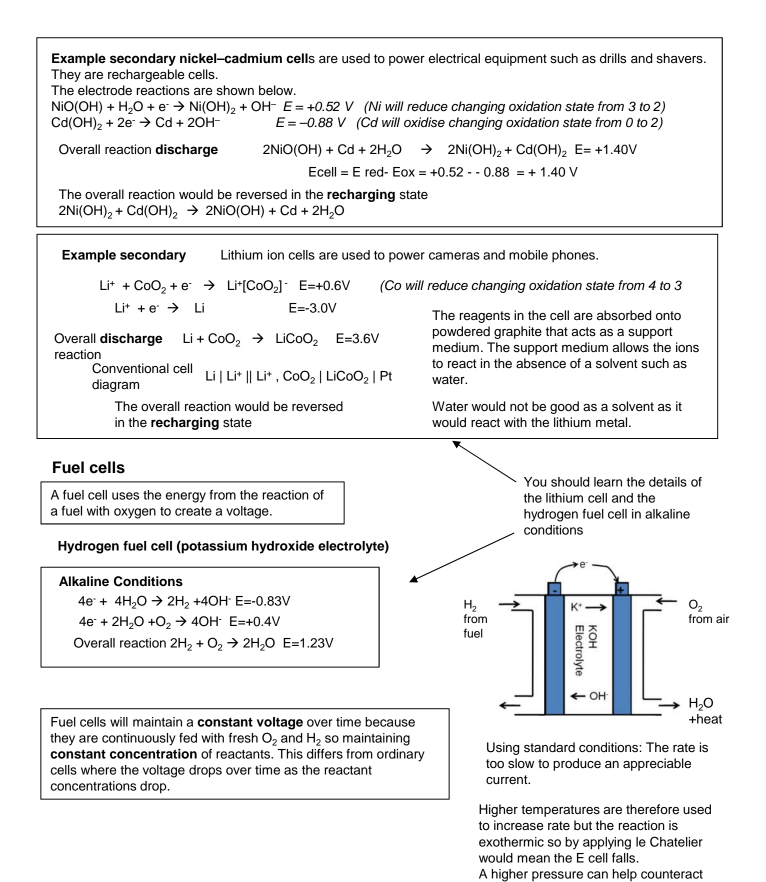
Example primary non rechargeable cells		Cells are non-rechargeable when the reactions that occur with in them are non-reversible.		
Dry Cell	Zn ²⁺ (<i>aq</i>) + 2e ⁻ → Zn(<i>s</i>) E = - 0 2MnO ₂ (<i>s</i>) + 2NH ₄ ⁺ (<i>aq</i>) + 2 e ⁻	0.76 V More negative half equation will oxidise $Mn_2O_3(s) + 2NH_3(aq) + H_2O(l) = 0.75 V$		
Overall reaction	$2MnO_2 + 2NH_4^+ + Zn = Mn_2O_3$	+ 2NH ₃ + H ₂ O + Zn ²⁺ Ecell =+1.51V		

Example primary Lithium -manganese dioxide cell- non rechargeable

Li_{aq}^{+} +e ⁻ \rightarrow Li (s) E = - 3.04 V		More negative half equation will oxidise		
$\text{Li}_{aq}^{+} + \text{MnO}_{2(s)}^{-} + e^{-} \rightarrow \text{LiMnO}_{2}^{-}$	_(s) E = - 0.13 V	(Mn will reduce changing oxidation state from $+4$ to $+3$)		
Overall reaction Li + MnO ₂	LiMnO ₂			
		Ecell = E red- Eox = -0.13 3.04 = 2.91 V		

Conventional cell diagram $\text{Li}_{(s)}$ | Li_{aq} | | Li_{aq} | $\text{MnO}_{2(s)}$, $\text{LiMnO}_{2(s)}$ | Pt(s) Ecell =+2.91V

				The forward reaction occurs on discharge giving out charge.	
Lead acid cell	$PbSO_4 + 2e^- \rightarrow Pb + SO_4^{2-}$	E= -0.356V	Charging causes the reaction to reverse		
Overall reaction	$PbO_{2} + SO_{4}^{2-} + 4H^{+} + 2e^{-} \rightarrow$ $PbO_{2} + Pb + 2SO_{4}^{2-} + 4H^{+} \rightarrow$	• =		Reversible cells only work if the product stays attached to the electrode and does not disperse	



this.

Advantages of fuel cells over conventional petrol or diesel-powered vehicles (i) less pollution and less CO_2 . (Pure hydrogen emits only water whilst hydrogen-rich fuels produce only small amounts of air pollutants and CO_2).

(ii) greater efficiency

Hydrogen is readily available by the electrolysis of water, but this is expensive. To be a green fuel the electricity needed would need to be produced from renewable resources

Ethanol fuel cells

Ethanol fuel cells have also been developed. Compared to hydrogen fuel cells they have certain advantages including. Ethanol can be made from renewable sources in a carbon neutral way.

Raw materials to produce ethanol by fermentation are abundant.

Ethanol is less explosive and easier to store than hydrogen. New petrol stations would not be required as ethanol is a liquid fuel.

Limitations of hydrogen fuel cells (i) expensive

(ii) storing and transporting hydrogen, in terms of safety, feasibility of a pressurised liquid and a limited life cycle of a solid 'adsorber' or 'absorber'
(iii) limited lifetime (requiring regular replacement and disposal) and high production costs,

(iv) use of toxic chemicals in their production

Hydrogen can be stored in fuel cells (i) as a liquid under pressure, (ii) adsorbed on the surface of a solid material, (iii) absorbed within a solid material:

Equation that occurs at oxygen electrode $4e^{-} + 4H^{+} + O_2 \rightarrow 2H_2O$ E=1.23V

Equation that occurs at ethanol electrode $C_2H_5OH + 3H_2O$ $2CO_2 + 12H^+ + 12e^-$

Overall equation $C_2H_5OH + 3O_2 = 2CO_2 + 3H_2O$