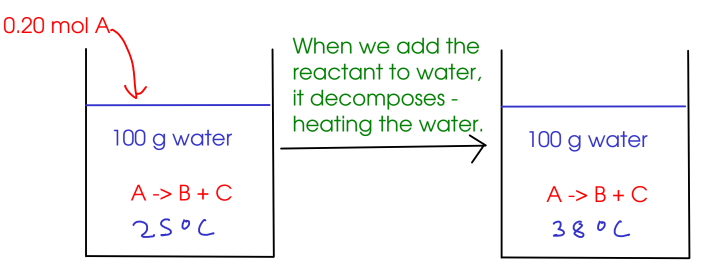
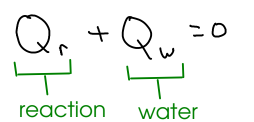
CALORIMETRY

- the measurement of heat. But how do we measure heat?



... what is Q for this reaction?

Assuming that no heat is lost from the water to the surrounding air,



Conservation of energy. The terms add to zero because they have opposite signs.

... if we knew something about the WATER, we could use that to find the heat of the REACTION!

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- a measured quantity. The amount of energy required to change the temperature of one gram of a particular substance by one degree Celsius.

- Specific heat information for common substances is readily available. For water,

$$4.184 \frac{5}{5^{\circ}C} \stackrel{o}{=} 1.000 \frac{Cal}{5^{\circ}C}$$

$$Q = M \times 5 \times \Delta T$$

$$m = mass$$

$$s = specific heat$$

$$\Delta T = Tfinal - Tinitial$$

$$M = mass$$

$$M = m$$

- For objects, like reaction vessels, you might know the HEAT CAPACITY, which is the amount of energy required to change the temperature of an object by one degree Celsius

Units: 
$$J/o_c$$
 or  $cal/o_c$   
 $Q = C \times \Delta T$   
 $c = heat capacity$ 

<sup>155</sup> 0.20 mol A  
When we add the  
reactant to water,  
it decomposes -  
heating the water.  
A -> B + C  

$$2.5 \circ C$$
  
When we add the  
reactant to water,  
it decomposes -  
heating the water.  
A -> B + C  
 $3 \& \circ C$   
Specific heat of water:  
H - 1 &  $\frac{3}{2} \circ C$ 

$$Q_r + Q_w = 0$$
  
= S439.2J  
 $Q_r + S439.2J = 0; Q_r = -5439.2J$ 

To report the energy change in this reaction to others, we should express it in terms of heat transfer per mole of something. A different amount of reactant would have a different Q

Qrxn = 
$$\frac{Qr}{moles A} = \frac{-5439.25}{0.20 \text{ mol} A} = -27000 \frac{\text{k}}{\text{mol} A} = \frac{-27 \frac{\text{k}}{\text{mol} A}}{\sqrt{1000 \text{ mol} A}}$$
  
This number is usually called the "heat of reaction"...

One problem ...

PATH. The amount of energy required for a process depends on how the process is carried out.

Example: Driving from Florence to Columbia. How much energy is required? (gas)

2000 Jeep Cherokee vs 2008 Toyota Prius. The Jeep will use much more fuel than the Prius even though they start and end from exactly the same place. So the fuel usage is what we call a <u>PATH FUNCTION</u>, while the location is a STATE FUNCTION.

- so the heat of reaction depends on how the reaction is done.

- we need (for reporting) some kind of standard condition. At constant pressure, we can define a state function called ENTHALPY (H)

H = U + PV $\triangle H = Q constant pressure$ 

 $\Lambda H_{r}$ 

... we record the "enthalpy change of reaction" in our data books.

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<sup>57</sup> SINCE the enthalpy change does NOT depend on path, this means that we can use standard values for enthalpy to predict the heat change in reactions that we have not tested in a calorimeter.

## THERMOCHEMICAL EQUATIONS

- is like a regular chemical equation, except that phase labels are REQUIRED and the enthalpy for the reaction is given along with the equation.

$$CH_3(O(H_3(l) + 4O_2(g) \rightarrow 3(O_2(g) + 3H_2O(l); AH = -1800 kJ$$

- Why are phase labels required? Because phase changes either absorb or release energy.

 $\Delta H = -1600 \text{ kJ} \dots \text{ what does this mean}?$ 

 $\frac{1}{4} \mod (H_{8}COCH_{3} = -1800 \text{ kJ}) + 4 \mod 02 = -1800 \text{ kJ}}{3 \mod 02} = -1800 \text{ kJ}}$   $\frac{3}{2} \mod 120 = -1800 \text{ kJ}}{3 \mod 120} = -1800 \text{ kJ}}$ 

We treat the enthalpy change as if it's another product of the reaction!

$$CH_{3}(O(H_{3}|l) + 4O_{2}(g) \rightarrow 3(O_{2}(g) + 3H_{2}O(l); A = -1800 kJ$$
What would be the enthapy change when 25 g of water are produced by the reaction?  
1 - Convert 25 g water to moles. Use FORMULA WEIGHT.  
2 - Convert moles water to enthalpy change. Use THERMOCHEMICAL EQUATION.  
()  $H_{2}O - H: 2 \times 1 - 008$   
 $O: \frac{1 \times 16.05}{18 \cdot 016g H_{2}O = mol H_{2}O}$   
2 3 mol  $H_{2}O = -1800 \text{ KJ}$   
1) This is an EXOTHERMIC  
process! (not too  
surprising! This is a combustion  
reaction!)

2) Provided we do this reaction at constant pressure, Q = -830 kJ, too!

A few more terms related to enthalpy:

- Enthalpy of vaporization / heat of vaporization: The enthalpy change on vaporizing one mole of a substance. (from liquid to vapor)

- Enthalpy of fusion / heat of fusion: The enthalpy change when a mole of liquid changes to the solid state.

Phase changes require energy, too!

<sup>159</sup> Example problems:

$$2H_{2}(g) + O_{2}(g) \longrightarrow 2H_{2}O(g); \Delta H = -484 kJ$$

Calculate the enthalpy change for the combustion of 1000. g of hydrogen gas.

FORMULA WEIGHTS in g/mol

1- Convert 1000. g hydrogen gas to moles. Use FORMULA WEIGHT.2- Convert moles hydrogen gas to enthalpy. Use THERMOCHEMICAL EQUATION.

160 propane

$$(_{3}H_{8}(g) + 50_{2}(g) \rightarrow 3(0_{2}(g) + 4H_{2}0(g); \Delta H = -2043 kJ$$

Calculate the volume of propane gas at 25.0 C and 1.08 atm required to provide 565 kJ of heat using the reaction above.

1 - Convert heat requirement (565 kJ) to moles PROPANE. Use THERMOCHEMICAL EQUATION. 2 - Convert moles propane to volume. Use IDEAL GAS EQUATION, PV=nRT.

$$\begin{array}{l} (1 - 2043 \ \text{KJ} = \text{mol} \ (_{3}\text{Hg} \\ -565 \ \text{KJ} \times \frac{\text{mol} \ (_{3}\text{Hg} \\ -2043 \ \text{KJ} = 0.2765540871 \ \text{mol} \ (_{3}\text{Hg} \text{energy!} \\ \end{array} \right) \\ \hline \\ (2 \ \text{PV} = nRT \\ V = \frac{nRT}{P} \\ R = 0.2765540871 \ \text{mol} \ (_{3}\text{Hg} \\ T = 25.0^{6}(1 - 298.2\text{K}) \\ T = 25.0^{6}(1 - 298.2\text{K}) \\ \hline \\ (1.08 \ \text{atm}) \\ = 6.27L \ (_{3} \ \text{Hg} \ \text{at} 25.0^{6}(1 - 1.08 \ \text{atm}) \\ \end{array}$$

## END OF CHAPTER 6