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.

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

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

$$1 \text{ mol CH}_3 \text{ COCH}_3 = -1800 \text{ kJ}$$

$$4 \text{ mol } 02 = -1800 \text{ kJ}$$

$$3 \text{ mol } 02 = -1800 \text{ kJ}$$

$$3 \text{ mol } 420 = -1800 \text{ kJ}$$

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

CH3 (O CH3 (l) + 402(g) -> 3 (O2(g) + 3H20(l); AH = -1800 KJ

What would be the enthapy change when 25 g of water are produced by the reaction?

- 1 Convert 25.0 g water to moles using formula weight.
- 2 Convert moles water to enthalpy change using themochemical equation.

18.016g
$$H_{20} = mol H_{20}$$
 $3 mol H_{20} = -1800 kJ$
25.0g $H_{20} \times \frac{mol H_{20}}{18.016g H_{20}} \times \frac{-1800 kJ}{3 mol H_{20}} = -830 kJ$

- This reaction is EXOTHERMIC
- Energy is released from the reaction mixture, heating the surroundings. (This is true for all combustions...)
- Provided that the reaction takes place at constant pressure, this enthalpy change should be equal to Q (measurable).

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.



- A reaction that forms exactly one mole of the specified substance from its elements at their STANDARD STATE at 25C and 1 atm pressure.

(O2(g):
$$(s,graphite) + O_2(g) \rightarrow (O_2(g);\Delta H = -393.5 kJ)$$

heat of formation of carbon dioxide $\Delta H_{\epsilon}^{\circ}$ or ΔH_{ϵ}
"enthalpy of formation"
 $(o(g): (s,graphite) + \frac{1}{2}O_2(g) \rightarrow (o(g);\Delta H = -110.5 kJ)$

you may see fractional coefficients in these formation reactions, because you MUST form exactly one mole of the product!

- The heat of formation for an element in its standard state at 25C and 1 atm is ZERO.

- What are formation reactions good for?

... finding enthalpies for more interesting reactions!

Let's say we would like to find the enthalpy of reaction for this equation: $2(2H_2(g) + 5O_2(g) \longrightarrow 4(O_1(g) + 2H_2O(g); \Delta H = 1)$ Hess' Law: If you add two reactions to get a new reaction, their enthalpies also add. DH = 226.7 (242: 2((s) + H2(g) -> (2H2(g)) (O2 (g) DH = -393,5 CO2 1 $((s) + O_2(y) \longrightarrow$ DH= -241,8 H20(g) $H_2(g) + \frac{1}{2}O_2(g)$ H20 1 $\rightarrow 2(s) + H_2(s)$ (2H2lg) DH = -226.7 \rightarrow 2((s) + H_2 (g) (2H2lg) DH = -226.7 $((s) + O_2(g))$ $\rightarrow (O_2(g))$ DH = -393.5 ((5) + O2 (g) $\longrightarrow (O_2(q)$ DH = -393,5 $((s) + O_2(g))$ $\longrightarrow (O_2(q)$ DH = -393,5 $\longrightarrow (O_2(q)$ DH = -343.5 $H_2(q) + \frac{1}{2}O_2(q)$ H20 (9) DH= -241,8 $2(q) + \frac{1}{2}O_2(q)$ H20 (9) DH= -241,8

$$2(2H_2(g) + So_2(g) \longrightarrow 4co_2(g) + 2H_2o(e)$$

Hess' Law using enthalpy of formation:

$$\Delta H = \sum \Delta H_{F,pNJ-uts} - \sum \Delta H_{F,mnchnts}$$

 $\Delta H = 4(-393.5) + 2(-241.8) - (2(226.7) + S(0))$
= 2511 KJ

See Appendix C in the textbook for enthalpy of formation data: p A-8 to A-11

* Remember:

- Multiply each enthalpy by its stoichiometric coefficient from the reaction
- Enthalpy of formation of an element at its standard state is zero
- Watch phase labels. You will usually find SEVERAL enthalpies of formation for a given substance in different phases!
- For ionic substances in solution, remember that they exist as free ions, so look up the aqueous IONS!

$$2.016$$
 32.00 16.02 in purple $2H_2(g) + O_2(g) \longrightarrow 2H_2O(g)$; $\Delta H = -464 kJ$

Calculate the enthalpy change for the combustion of 1.00 kg of hydrogen gas.

- 1 Convert 1.00 kg of hydrogen gas to moles using formula weight.
- 2 Convert moles hydrogen gas to enthalpy change using thermochemical equation.

1.00 kg H2 x
$$\frac{10^3g}{kg}$$
 x $\frac{mol H2}{2.016g H2}$ x $\frac{-484 kJ}{2 mol H2} = -120000 kJ$ per kg H2