

# *Entropy and 2nd Law of Thermodynamics*

CHEM 102  
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## *Einstein's view*

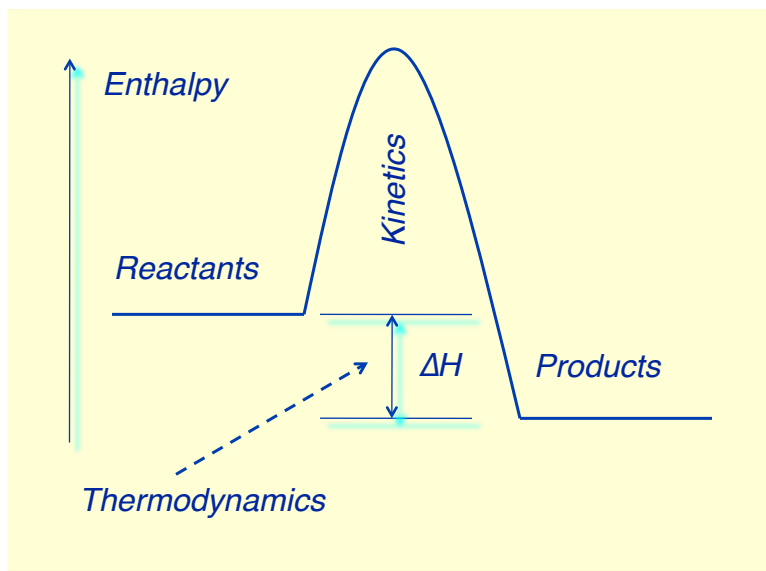
“ [Thermodynamics is] the only physical theory of universal content concerning which I am convinced that, within the framework of the applicability of its basic concepts, it will never be overthrown.”

## Thermodynamics

### Review of some Thermodynamic Concepts

- *First law of thermodynamics*: The law of conservation of energy; energy cannot be created or destroyed.
- *State Function*: Quantity in which its determination is *path independent*.
- $\Delta U = q + w$ : The change in internal energy of a system is a function of heat and work done on or by the system.
- $\Delta H$ : Heat transferred at constant pressure.
- Exothermic Process:  $\Delta H < 0$
- Endothermic Process:  $\Delta H > 0$

## Thermodynamics & Kinetics



## Entropy & the Second Law of Thermodynamics

### *Spontaneity*

- Spontaneous: “Occurring without outside intervention.”
- A reaction or change of state is said to be spontaneous if it is thermodynamically allowed.
- For a chemist, prediction of spontaneity is a major goal of thermodynamics.

## *Factors Affecting Spontaneity*

- Energy or Enthalpy:  $\Delta U$  or  $\Delta H$ 
  - Not a good predictor. Both endothermic and exothermic reactions can occur.
- Temperature
  - Some processes are spontaneous only at certain temperatures.
- Concentrations, pressures, etc.

## *Entropy*

- Entropy (S) is a thermodynamic state function which can be described qualitatively as a measure of the amount of disorder present in a system.
- From a chemical perspective, we usually mean *molecular* disorder.
- Increases in the entropy of a system are usually (not always) accompanied by the flow of heat into the system.

## *Entropy and Disorder*

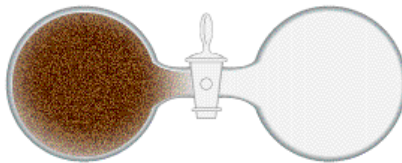
- Entropy is a measure of disorder.  
**more disorder → greater entropy**
- Entropy of a substance depends on physical state.  $S_{\text{gas}} \gg S_{\text{liquid}} > S_{\text{solid}}$
- Entropy depends on temperature.  
Increasing T will increase entropy due to increase in molecular motion.

## *Entropy & Spontaneity*

In many spontaneous processes, entropy of a system increases. ( $\Delta S > 0$ )

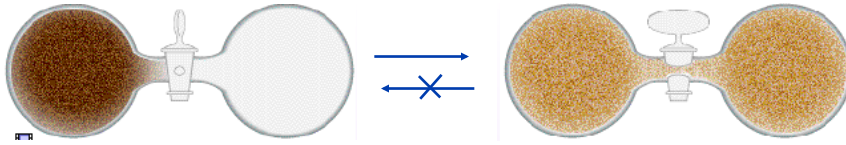
■ Examples:

- expansion of a gas into vacuum



- mixing of gases or generation of a gas from solid or liquid reactants

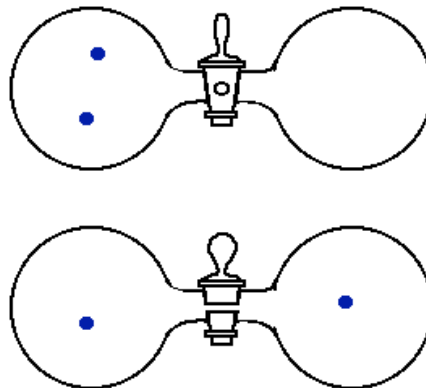
## *Expansion of a Gas*



- This process obviously has a preferred direction, but why?
- Once the valve is open, the probability that all molecules will return to one side is astronomically small.
- $\Delta S > 0$  for this spontaneous change

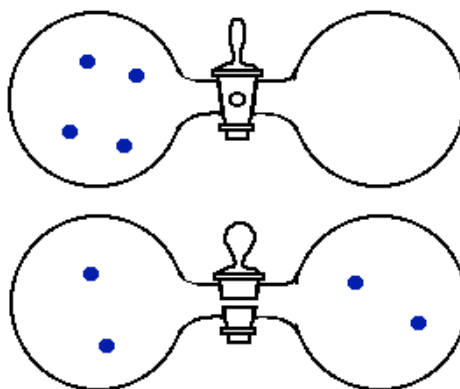
## *Probabilities and Entropy*

Probability that  
both molecules  
will be on left  
hand side:  $(1/2)^2$   
 $= 1/4$ .



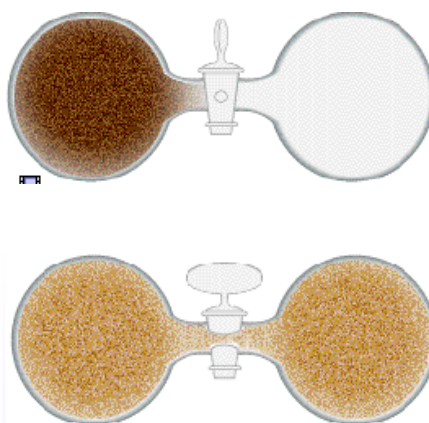
## *Probabilities and Entropy*

Probability that  
both molecules  
will be on left  
hand side:  $(1/2)^4$   
 $= 1/16$ .



## *Probabilities and Entropy*

Probability that  
both molecules  
will be on left  
hand side:  $(1/2)^N$



**= A very small number when N is  
Avogadro's number!**

## *Dispersal of Energy: Entropy*

Boltzmann: As the number of microstates increases, so does the entropy of the system.

$$S = k \cdot \ln W$$

$k$  = Boltzmann's constant ( $1.381 \times 10^{-23}$  J/K)

$W$  = the number of microstates corresponding to the observed macroscopic state of a system

## *Extra Details*

Number of ways of having  $N/2$  molecules on each side:

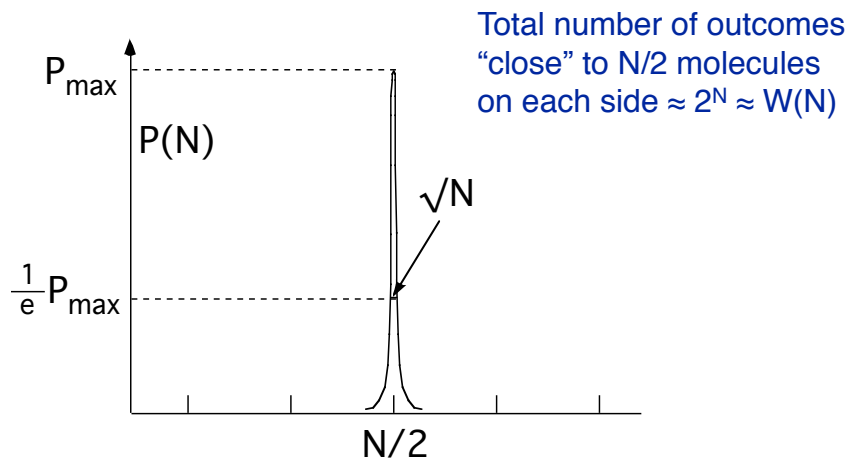
$$W_{N/2} = \binom{N}{N/2} = \frac{N!}{[(N/2)!]^2}$$

Number of ways of having  $N/2 - \sqrt{N}$  molecules on one side:

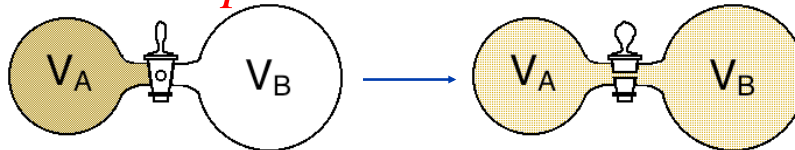
$$\frac{\binom{N}{N/2 - \sqrt{N/2}}}{\binom{N}{N/2}} = \frac{N!}{(N/2 - \sqrt{N/2})!(N/2 + \sqrt{N/2})!}$$
$$\left( \frac{\binom{N}{N/2 - \sqrt{N/2}}}{\binom{N}{N/2}} \right) \rightarrow \frac{1}{e}, \text{ as } N \rightarrow \infty; 0.369 \text{ for } N = 200$$

use Stirling's Approximation:  $N! \approx \sqrt{2\pi N} \left(\frac{N}{e}\right)^N$  and definition of  $e = \lim_{x \rightarrow \infty} \left(1 + \frac{1}{x}\right)^x$

## *Distribution of Probabilities*



## *Gas Expansion & Probabilities*



Entropy change on expansion:

$$\Delta S = k \ln W_f - k \ln W_i = k \ln \left( \frac{W_f}{W_i} \right)$$

$$\Delta S = k \ln \left( \frac{V_f}{V_i} \right)^N = R \ln \left( \frac{V_f}{V_i} \right) \quad (Nk = R)$$

When  $V_A = V_B$ ,  $V_f = 2V_i$ , and  $\Delta S = R \ln 2$

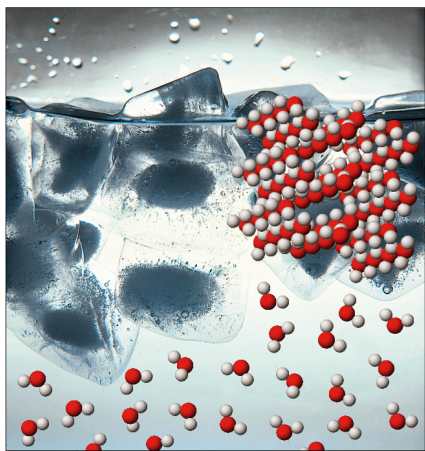
## *Entropy, Probabilities, Disorder*

- For systems with equal energy content, those that are most disordered also turn out to be most statistically likely (most probable).
- **Entropy increases as the statistical likelihood increases.**
- The entropy of the universe tends to increase:  $(\Delta S)_{\text{universe}} > 0$  in all changes

## *Popular Misconceptions*

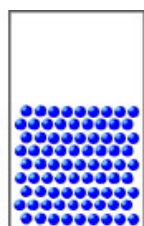
- Sometimes you see it said that “in principle, the universe could show a decrease in entropy, it is just highly unlikely” – True, but means astronomically unlikely EVER!
- The entropy of system *can* decrease, if the entropy of the universe increases. (For example, the “order” we see in life on earth is the direct result of a massive increase in the entropy of the sun.)

## *Dispersal of Energy: Entropy*

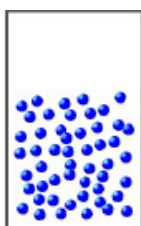


- The entropy of liquid water is greater than the entropy of solid water (ice) at 0° C.
- Energy is more dispersed in liquid water than in solid water due to the lack of an ordered network as in the solid state.

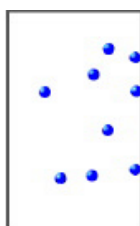
## *Entropy & States of Matter*



**Solid**



**Liquid**



**Gas**

$S^\circ$  (J/K·mol)

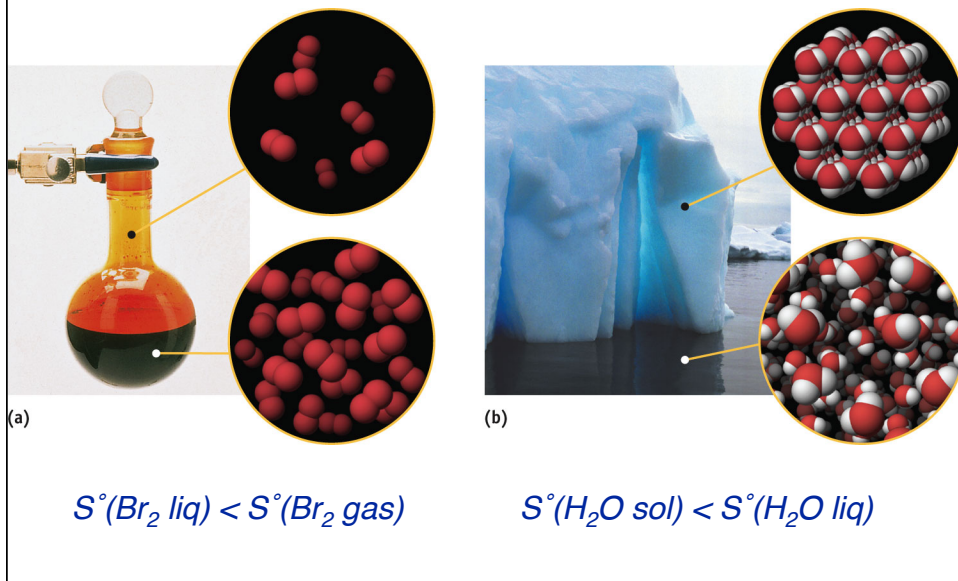
$H_2O(liq)$  69.95

$H_2O(gas)$  188.8

$$S(\text{solids}) < S(\text{liquids}) < S(\text{gases})$$

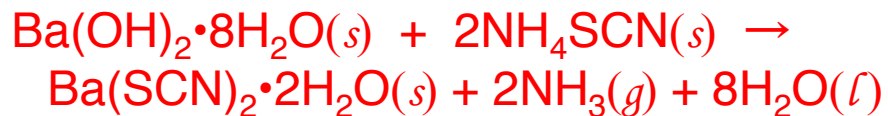
Energy dispersal →

## Entropy & States of Matter



## Generation of a Gas (Demo)

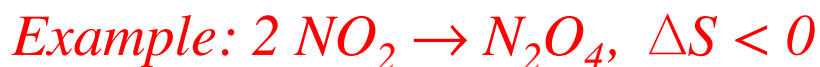
- “Frozen Beaker” demo ( $\Delta H > 0$ ):



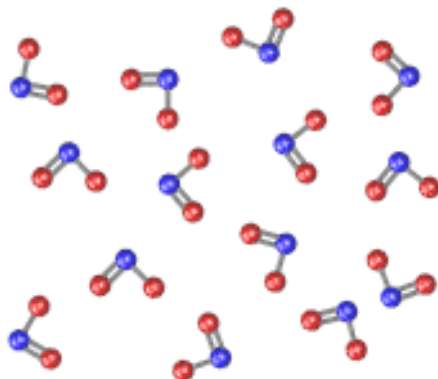
- Solid reactants  $\rightarrow$  solid, liquid, & gas in products
- $\Delta S_{\text{system}} \gg 0$

## *Entropy & Spontaneity*

- In some spontaneous processes, the entropy of a system decreases. ( $\Delta S < 0$ )
- Examples:
  - freezing of a liquid
  - condensation of a gas
  - formation of a solid product from gas or liquid reactants
- These are common events, and they clearly have  $\Delta S < 0$ .



- The conversion of 2 moles of gas into one mole of gas decreases the entropy of the system.



## *Second Law of Thermodynamics*

- Entropy change of the *system* does not correctly predict spontaneity.
- One correct statement of the second law of thermodynamics is:  
“**The entropy of the universe is always increasing.**”
- In equation form, this is:  $\Delta S_{\text{universe}} > 0$ .  
where  $\Delta S_{\text{universe}} = \Delta S_{\text{system}} + \Delta S_{\text{surroundings}}$

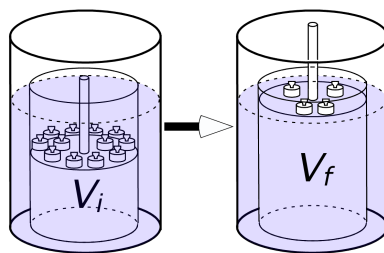
## *Entropy and Heat*

- Simplest case is a process which occurs at constant T. Phase changes are good examples.
- For the case of constant T (isothermal):  
$$\Delta S = q_T/T$$
- $q_T$  is heat at constant T, signs as usual
- From this equation,  $\Delta S$  has units of J/K

*How do we reconcile the “heat transfer”  
and statistical definitions of entropy?*

*Consider a Reversible Isothermal  
Gas Expansion:*

### *Reversible Isothermal Expansion*

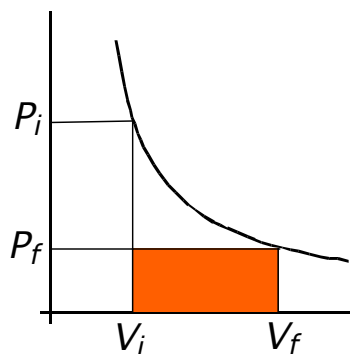


*Gradually  
decrease  
pressure, by  
removing one  
weight at a  
time.*

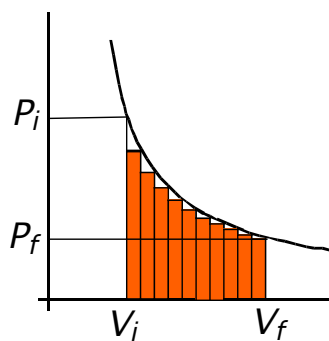
*Note, since  $T = \text{const.}$ ,  
 $P_i V_i = P_f V_f$*

- If we gradually decrease the pressure while keeping the apparatus in a large bath, expansion will be nearly reversible and nearly isothermal.

## Irreversible vs. (nearly) Reversible



Pressure lowered in one step (remove all weights)



Pressure lowered in incremental steps

## Entropy Changes for Phase Changes



For a phase change,

$$\Delta S = \frac{q_{\text{rev}}}{T}$$

where  $q$  = heat transferred in the phase change

For  $\text{H}_2\text{O (liq)} \rightleftharpoons \text{H}_2\text{O(g)}$

$$\Delta H = q = +40,700 \frac{\text{J}}{\text{mol}}$$

$$\Delta S = \frac{q_{\text{rev}}}{T_{\text{boil}}} = \frac{40,700 \text{ J/mol}}{373.15 \text{ K}} = +109 \frac{\text{J}}{\text{mol} \cdot \text{K}}$$

### *Illustrative Problem*

- Dry ice is solid  $\text{CO}_2$ . At  $T = 195 \text{ K}$ , dry ice sublimates (is converted directly from a solid into a gas).  $\Delta H_{\text{sublim.}} = 25.2 \text{ kJ/mol}$
- Suppose 27.5 g of dry ice are allowed to sublime in a room with  $T_{\text{room}} = 26.5^\circ \text{C}$ .
  - Without doing calculations, predict the signs of  $\Delta S_{\text{CO}_2}$ ,  $\Delta S_{\text{room}}$ , and  $\Delta S_{\text{universe}}$ .
  - Calculate  $\Delta S_{\text{CO}_2}$ ,  $\Delta S_{\text{room}}$ , and  $\Delta S_{\text{universe}}$ .

### *Spontaneity*

- Second law says that a process is spontaneous if  $\Delta S_{\text{universe}} > 0$ .
- From problem, realize that the phase change will be spontaneous as long as the temperature of the room is above the sublimation temperature.
- At  $T_{\text{sublim}}$ ,  $\Delta S_{\text{universe}} = 0$ , so both phases can exist in any ratio.  $\rightarrow$  EQUILIBRIUM

## *Absolute Entropy*

- We said earlier that the entropy of any material will increase as  $T$  increases, due to increased molecular motion.
- Molecular motion, and entropy, decrease as  $T$  is lowered.
- $T = 0 \text{ K}$  is the (theoretical) limit to how far we can lower the temperature.
- Thus *minimum* entropy is at  $0 \text{ K}$ .

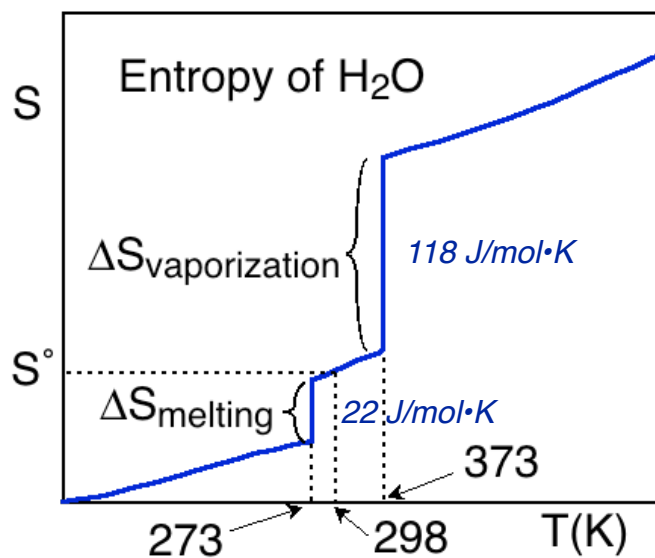
## *The Third Law*

- The third law of thermodynamics says:  
“**The entropy of a pure, perfect crystal at  $0 \text{ K}$  is zero.**”
- Applies to a crystal of any substance.
- Note that this is a fairly hypothetical limit: “pure,” “perfect,” and  $0 \text{ K}$  are all ideals.
- Third law means that we can find absolute entropy. (unlike  $E$ ,  $H$ )

## $S^\circ$ and $\Delta S^\circ$

- Thermo. Tables (Appendix E) give  $S^\circ$  for many substances.  
(Units of  $\text{J mol}^{-1}\text{K}^{-1}$ )
- This is *absolute entropy* at  $25^\circ\text{C}$ , 1 atm.
- Notice that  $S^\circ$  is *never* equal to zero, but approaches zero as the temperature approaches zero.  $S^\circ$  is *never* less than zero (but  $\Delta S_{\text{system}}$  can be less than zero).

## Entropy as a Function of Temp. An increasing function



## *$S^\circ$ and $\Delta S^\circ$*

- Easy to find  $\Delta S^\circ$  from the tabulated  $S^\circ$ 's:

$$\Delta S^\circ = S^\circ_{\text{products}} - S^\circ_{\text{reactants}}.$$

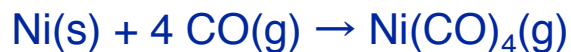
- Useful form is:

$$\Delta S^\circ_{\text{rxn}} = \sum n S^\circ_{\text{products}} - \sum n S^\circ_{\text{reactants}}$$

- Similar in appearance to Hess's law.
- But this uses absolute  $S^\circ$  values, where for  $\Delta H^\circ_{\text{rxn}}$  we used  $\Delta H^\circ_f$ 's.

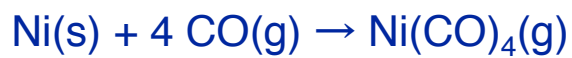
## *Problem*

- The “Mond Process” is used commercially to produce pure nickel.



- First, try to predict the signs of  $\Delta S^\circ$  and  $\Delta H^\circ$ .
- Then use data on next slide to calculate values.

*Data for Problem*



$\Delta S^\circ$ ,  $\Delta H^\circ = ??$

	Ni(s)	CO(g)	Ni(CO) <sub>4</sub> (g)
$\Delta H_f^\circ$	0	-110.52	-602.9
$S_{298}^\circ$	29.87	197.56	410.6

$\Delta H_f^\circ$  in kJ mol<sup>-1</sup>,  $S_{298}^\circ$  in J K<sup>-1</sup> mol<sup>-1</sup>.

Watch units!