


Chemistry 163B, Winter 2020
Lecture 1- Introductory Lecture

Chemistry 163B
Thermodynamics
Winter 2020

1

1

Menu: for TODAY



- Introduction and Class Resources, etc.
- Careful definition of thermodynamic variables (P, V, T , etc)
- Ideal gas ENERGY DEPENDS ONLY ON TEMPERATURE (*derivation*)
- Van der Waals equation (*example of equation of state for 'real' gas*)

2

2

Thermodynamics is a
really beautiful scientific
story !!

3

3

"rules" of science

- observations

↓

- guiding principles

↓

- predictions and applications based on principles

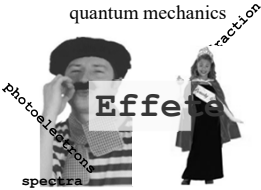
4

4

Observations (QM vs Thermo)

Thermodynamics is very 'working class' in its origins:


quantum mechanics



photoelectric effect
spectra

$H \psi = E \psi$

thermodynamics



Blue Collar
disorder

$\Delta S_{\text{universe}} > 0$

5

5

observations: thermo = heat

- Count Rumford, 1799
- observed water turning into steam when canon barrel was bored
- $\text{work} \leftrightarrow \text{heat}$

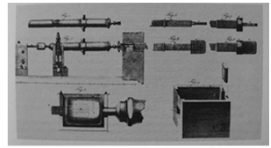




FIGURE 11. An illustration from Rumford's paper, "An Inquiry Concerning the Source of the Heat Which is Excited by Friction," showing the apparatus used by him in the cannon boring experiment. Figure 1, upper left, shows the cannon as received from the foundry, and Figure 2, below, shows it mounted in the machine used for boring. (Reproduced with the permission of Harvard University Press.)

6



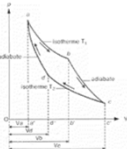
6

Chemistry 163B, Winter 2020

Lecture 1- Introductory Lecture

observations: mechanical efficiency of steam engine

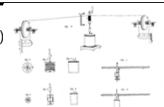


- Sadi Carnot, 1824
- efficiency of engines

7

guiding principles

- Conservation of heat and work (Joule, 1845)
1st LAW OF THERMODYNAMICS
- Clausius, 1860 Entropy
2nd LAW OF THERMODYNAMICS
- Boltzmann, late 19th century,
molecular picture of entropy

the thermodynamic functions U, H, and S
(1st and 2nd laws)

8


7

8

"Applications" U,H,S,G

How does knowledge about efficiencies of steam engines, mechanical systems, etc, relate to processes in chemical, biological, and geological systems?

ANSWERED BY:

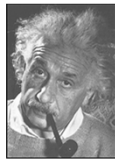


J. W. Gibbs- arguably the first great American scientist who combined the concepts of heat and entropy and proposed "[Gibbs] Free Energy", G, a thermodynamic state function that leads to a whole spectrum of applications

9

9

from: Einstein's "Autobiographical Notes"



A theory is the more impressive the greater the simplicity of its premises, the more different kinds of things it relates, and the more extended its area of applicability. Hence the deep impression that classical thermodynamics made upon me. It 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 (for the special attention of those who are skeptics on principle).

10


10

types of problems that thermodynamics addresses (reactions)

might

1. Under what conditions ~~will~~ a reaction occur ?

EXAMPLE:
 $C(\text{graphite}) \rightarrow C(\text{diamond})$ 1 atm 398K *nada*
 30,000 -100,000 atm *jHurra!*
 1000 K - 3000 K
 1st artificially produced diamonds, 1954 at General Electric Labs




11


11

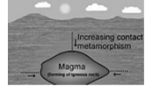
types of problems that thermodynamics addresses (equilibria)

2. How far will a reaction proceed (given enough time)?
 How do the thermal and entropic properties determine EQUILIBRIUM ?
 How do the EQUILIBRIUM conditions depend on T,P ?



$N_2(g) + 3H_2(g) \rightleftharpoons 2NH_3(g)$
 what are best T,P for NH₃ products?
The Haber Process





$SiO_2 + CaCO_3 \rightleftharpoons CaO \cdot SiO_2 + CO_2(g)$
 quartz calcite wollastonite
geologic thermometer:
 The fraction of wollastonite in a rock sample can be used to estimate the temperature at which the rock metamorphosis occurred [the T at which the equilibrium was rapidly frozen]
Geologic Thermometer

12

12

Chemistry 163B, Winter 2020

Lecture 1- Introductory Lecture

types of problems that thermodynamics addresses (biology)

3. Chemical and physical changes in biological systems

[K⁺]
inside

[K⁺]
outside

cell

what difference in [K⁺]_{out} vs [K⁺]_{in} can be tolerated before cell wall bursts or collapses?

low hypotonic (rupture) ← extracellular salt normal isotonic extracellular salt → high hypertonic (collapse)

<http://www.vivo.colostate.edu/books/cmb/cells/pmemb/osismosis.html>

types of problems that thermodynamics addresses (biology)

4. Why does an egg hard boil?
(protein conformation)

5. Membrane potentials and ion concentrations in neurons.
(electrochemistry and thermodynamics)

13

14

types of problems that thermodynamics addresses (ecology)

6. Thermodynamic feasibility of SO₂ removal

$\text{SO}_2(\text{g}) \rightleftharpoons \text{S}(\text{s}) + \text{O}_2(\text{g})$

<http://healthandenergy.com/images/magnifica%20smoke%20stacks.jpg>

15

what thermodynamics can't answer

1. How fast a reaction proceeds (kinetics, catalysts, enzymes; in chem 103, BMB 100, chem 163C)
2. Macroscopic thermodynamics does not prove or require hypotheses about molecular structure; however we will use our knowledge of molecular structure to get an atomic "picture" of thermodynamic processes. The quantitative connection is made by statistical thermodynamics.

$\text{chem 163A} \rightleftharpoons \text{chem 163B}$
 chem 163C
3. Although in chemistry 163B we will study how thermodynamics put limits on processes at equilibrium, there exists a whole other field of *non-equilibrium thermodynamics*.

16

aims of class

1. Clear conceptual picture of thermodynamics
2. Ability to relate and apply thermodynamics to chemical and biological systems
3. **PROBLEM SOLVING: Chemistry + Logic + Mathematics**
4. How to do independent and advanced reading/research in areas that utilize thermodynamics.
5. THE GRAND PICTURE of how thermodynamics and quantum mechanics fit into our picture of 'nature'.
6. Advanced mathematical techniques

17

class administration

www.chemistry.ucsc.edu ⇒ course pages ⇒ Chemistry 163B
<http://switkes.chemistry.ucsc.edu/teaching/CHEM163B/>

- lectures: **A MUST**
- homework: **A MUST**
- sections: required (**A MUST**)
- tutorial EVENT & Office Hours & LSS (for YOU!)
- midterms: 31st January
28th February
- final: 19th March, 4:00-7:00 PM
(last class 13th March)


18

Chemistry 163B, Winter 2020

Lecture 1- Introductory Lecture

Learn Thermodynamics

- **Lectures**
- **Sections** (start Monday, Jan 13; HW#1 Probs 1-8)
- **Tutorial Event** (starts TOMORROW Tues, Jan 7, HW#1 Probs1-2)
- **Office hours** (start TODAY Mon, Jan 6)
- **LSS Tutor**



19

19

first Tutorial Tuesday 7th January, NS Annex 102 , 5:25PM-6:30PM

Chemistry 163B Winter 2020 Chemistry 163B

Comments on Mathematical Techniques for Chemistry 163B

Our best (50% 40% 30% 20% 10% 0%) contains detailed information on the topics of differential and integral calculus relevant to our study of thermodynamics. It is only a reference and integral calculus relevant to our study of thermodynamics, other than just as abstract mathematical exercises. However it is useful for categories from below. In your first discussion section the TA will also comment on the techniques.

1. Partial derivatives.
 - a. Know physical meaning of partial derivatives.
 - b. Know how to evaluate from mechanically.
 - c. Know chain rule.
2. You should have "at hand" the definitions and integrals of functions like e^x , x^n , $\ln(x)$, $\sin(x)$, $\cos(x)$. (I don't expect your assistance here!)
3. Differential expressions (5th ed 2.8, pp 39-41, 43) $y = x + \dots$.
 - a. $\int v dx$ is inexact differential if $\int \frac{\partial v}{\partial x} dx$ depends on the path from initial \rightarrow final. This implies there is no "underlying" well behaved, function v that changes like $v dx$.
 - b. $\int v dx$ is exact differential if $\int \frac{\partial v}{\partial x} dx$ does not depend on the path from initial \rightarrow final.
4. Implications that arise from $v dx$ being an exact differential.
 - a. There is a well behaved function v whose change is described by $v dx$ or in more general notation $\int v dx = \int_{x_1}^{x_2} v dx = \phi(x_2) - \phi(x_1)$.
 - b. $\oint v dx = 0$ (integral for any cyclic path, i.e. the initial and final states are identical).
 - c. $\frac{\partial}{\partial x} \left(\frac{\partial v}{\partial y} \right) = \frac{\partial}{\partial y} \left(\frac{\partial v}{\partial x} \right)$.
 - d. $\frac{\partial}{\partial x} \left(\frac{\partial v}{\partial y} \right) = \frac{\partial}{\partial y} \left(\frac{\partial v}{\partial x} \right)$.

20

20

Winter 2020 Chemistry 163B

Tutorial Tuesday 7th January

Homework Problems (#1-#8)

1. For an ideal gas $P\bar{V} = RT$ ($\bar{V} \equiv V_m \equiv \frac{V}{n}$ molar volume) evaluate:

- a. $\left(\frac{\partial P}{\partial \bar{V}} \right)_T$
- b. $\left(\frac{\partial \bar{V}}{\partial T} \right)_P$
- c. $\left(\frac{\partial T}{\partial P} \right)_P$
- d. $\left(\frac{\partial P}{\partial \bar{V}} \right)_T \left(\frac{\partial \bar{V}}{\partial T} \right)_P \left(\frac{\partial T}{\partial P} \right)_P$

2. For a Van der Waals gas: $\left(P + \frac{a}{\bar{V}^2} \right) (\bar{V} - b) = RT$ evaluate:

- a. $\left(\frac{\partial P}{\partial \bar{V}} \right)_T$
- b. $\left(\frac{\partial^2 P}{\partial \bar{V}^2} \right)_T$
- c. $\left(\frac{\partial P}{\partial T} \right)_P$
- d. $\left[\frac{\partial}{\partial T} \left(\frac{\partial P}{\partial \bar{V}} \right)_T \right]_P$
- e. $\left[\frac{\partial}{\partial \bar{V}} \left(\frac{\partial P}{\partial T} \right)_P \right]_T$

21

21

Winter 2020 Chemistry 163B

3. Engel & Reid problems P1-38 (4th edition)

to HW#1 listing on CANVAS

Van der Waals gas

4. (optional) The Van der Waals and virial expressions are two commonly used equations of state as approximations to real gas behavior:

$$P = \frac{RT}{(\bar{V}-b)} - \frac{a}{\bar{V}^2} \quad \text{Van der Waals}$$

$$P = RT \left[C_0 \left(\frac{T}{T^*} \right) + C_1 \left(\frac{T}{T^*} \right)^{-1} + C_2 \left(\frac{T}{T^*} \right)^{-2} + \dots \right] \quad \text{virial}$$

a. Show that the first three virial coefficients are related to the Van der Waals parameters a and b in the following way:

$C_0(T) = 1$
 $C_1(T) = b - a/RT$
 $C_2(T) = b^2$

HINT: note that the virial expansion is just a Taylor (Maclaurin) series in $\rho = \frac{1}{\bar{V}}$

$$\frac{P}{RT} = \sum_{i=0}^{\infty} C_i(T) \left(\frac{1}{\bar{V}} \right)^i = \sum_{i=0}^{\infty} C_i(T) \rho^i = f(\rho, T)$$

with coefficient $C_i(T) = \frac{1}{i!} \left(\frac{\partial^i P}{\partial \rho^i} \right)_{\rho=0, T}$

b. Why does the Van der Waals "a" only appear in $C_1(T)$?




22

22

enough hellos

and NOW TO BEGIN !!

our thermodynamics trek

23

23


system and surroundings (careful definitions) E&R_{4th} sec 1.3

A thermodynamic **system** consists of all the materials involved in the process under study. This material could be the contents of an open beaker containing reagents, the electrolyte solution within an electrochemical cell, or the contents of a cylinder and movable piston assembly in an engine.

In thermodynamics, the rest of the universe is referred to as the **surroundings**.

system + surroundings = the UNIVERSE

If a system can exchange matter with the surroundings, it is called an **open system**; if not, it is a **closed system**.




24




24

Chemistry 163B, Winter 2020

Lecture 1- Introductory Lecture


Variables of state: V, T, P (careful definitions)  E&R_{4th} sec 1.2

STATE VARIABLES:
properties whose values are required to specify the **STATE** of a system

- **VOLUME:** MEASURED WITH A RULER 
- **PRESSURE:** FORCE/AREA (physics 5-6A)
- **TEMPERATURE:** SEE IDEAL GAS THERMOMETER HANDOUT 
- **EQUATION of STATE:** RELATES P, V, T (more later) 

25

25

Pressure, Kinetic Energy, and Temperature SEE HANDOUT 

$PV = nRT$ $n = \text{moles, } R = \text{gas constant}$
 $PV = (nN) \frac{R}{N} T$ $N = \text{Avogadro's number } \left[\frac{\text{molecules}}{\text{mol}} \right]$
 $PV = n^* k T$ $nN = n^* = \text{number of molecules}$
 $k = \frac{R}{N}$ Boltzmann's constant


Fundamental and Defined Constants
 Engel & Reid [front cover, Table 1.1 (p13_{en}, [8]_{en}), Table 1.2 (p13_m, [9]_m)]

$R = 8.3145 \text{ J mol}^{-1} \text{ K}^{-1}$
 $= .083145 \text{ L bar mol}^{-1} \text{ K}^{-1}$
 $= .082058 \text{ L atm mol}^{-1} \text{ K}^{-1}$

$0.986923 \text{ atm} = 1 \text{ bar} = 10^5 \text{ kg m}^{-1} \text{ s}^{-2} = 10^5 \text{ Pa (pascal)}$
 $1 \text{ L} = 1 \text{ dm}^3 = 10^{-3} \text{ m}^3$


26

26


Ideal Gas Energy: Game Plan Handout #6  [E&R pp 7-8]_{en}

for monatomic ideal gas

from $PV = n^* k T$ and $P = F/A$



use physics to relate pressure \leftrightarrow energy of gas

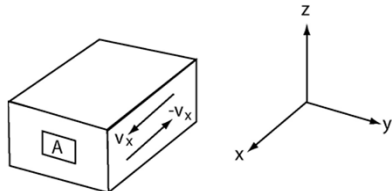


show $E = \frac{3}{2} n^* k T$ (n^* atoms gas) or $E = \frac{3}{2} n R T$ (n moles gas)

and thus for monatomic ideal gas E depends only on T !!!
[in thermodynamic notation $E \equiv U$ (internal energy)] 27

27

heuristic derivation of $E = E(T)$ for ideal gas [$U = U(T)$] E&R pp.7-8_{en}




RESULT: for ('monatomic') ideal gas
 (how) **energy depends only on temperature**

28

28

heuristic derivation

1. molecules all with same $|v_x|$ (all same v_x is 'heuristic')
2. elastic collision with wall velocity of mass goes $v_x \rightarrow -v_x$
3. from physics $p = \frac{F}{A}$ $p = \text{pressure}$
4. from physics $F = \frac{dp}{dt}$ $p = mv$, momentum; $t = \text{time}$
5. $dp \approx \Delta p = 2m v_x$ per collision (m is mass of particle)



6. total Δp in given time Δt , $\left(\frac{\Delta p}{\Delta t} \right) \approx \left(\frac{dp}{dt} \right)$

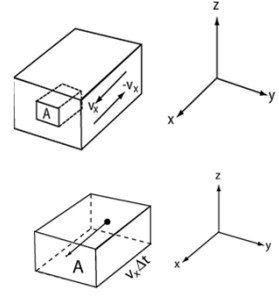
would depend on number of collisions in that interval

29

29

heuristic derivation

7.



30

30

Chemistry 163B, Winter 2020

Lecture 1- Introductory Lecture

heuristic derivation

8. $\frac{n^*}{V}$ is density of molecules. $Av_x \Delta t = V$ is volume of rectangular box,

9. $\left(\frac{1}{2} \frac{n^*}{V}\right) Av_x \Delta t$ is total number of molecules colliding with area A in Δt

10. total $\Delta p = (2mv_x) \left(\frac{1}{2}\right) \left(\frac{n^*}{V}\right) (Av_x \Delta t) = (mv_x^2) \left(\frac{n^*}{V}\right) (A \Delta t)$

31

31

heuristic derivation

11. after some algebra and equating

$$P = \frac{F}{A} = \frac{\Delta p}{A \Delta t} = (mv_x^2) \left(\frac{n^*}{V}\right) = \frac{n^* kT}{V}$$

$$P = \left(\frac{mv_x^2}{2}\right) \left(\frac{n^*}{V}\right) = \left(\frac{2}{V}\right) \frac{kT}{2}$$

$\left(\frac{mv_x^2}{2}\right) = \frac{kT}{2}$ for each molecule of 1-D ideal gas

12. in 3D with $|v_x| = |v_y| = |v_z|$

$$\text{total KE} = \left(\frac{1}{2}mv_x^2 + \frac{1}{2}mv_y^2 + \frac{1}{2}mv_z^2\right) n^* = \frac{3}{2}n^* kT = \frac{3}{2}nRT$$

$\frac{1}{2}kT$ for each degree (x,y,z) of 'classical freedom'

32

32

TAKE HOME MESSAGES

- Good warm up of physics and equation derivation
- For a molecule with only kinetic energy (e.g. monatomic species), and ideal gas (no intermolecular forces)

$$E = \frac{3}{2}kT \text{ per molecule or } E = \frac{3}{2}RT \text{ per mole}$$

- For monatomic ideal gas, E is function of **only T**;

T constant \leftrightarrow E constant

33

33

beyond PV=nRT: 'real' gases and equations of state (E&R sec 1.5)

Ideal
Prof S ??

vs

Real
Prof S ✓✓

34

34

various equations of state (Raff Table 1.2) $\frac{V}{n} \equiv \bar{V} \equiv V_{\text{molar volume}}$

Equation of State	Functional Form	Number of Parameters
Ideal gas	$PV_m = RT$	0
van der Waals	$(V_m - b)(P + a/V_m^2) = RT$	2
Dieterici	$P(V_m - b) \exp(a/RTV_m) = RT$	2
Berthelot	$(V_m - b)(P + a/TV_m^2) = RT$	2
Virial	$P = RT \left[V_m^{-1} + \sum_{i=2}^{\infty} C_i(T) V_m^{-i} \right]$	∞
Beattie-Bridgman	$PV_m^2 = (1 - \gamma)RT(V_m + \beta) - \alpha$ with $\gamma = c_m/T^2 V_m$, $\beta = b_m(1 - b/V_m)$, and $\alpha = a_m(1 + a/V_m)$	5
Redlich-Kwong	$P = \frac{RT}{(V_m - b)} - \frac{a}{T^{1/2} V_m(V_m + b)}$	2
Reichsantalt	$PV = RT + AP + BP^2 + CP^3 + \dots$	∞

35

35

Some comments on the
Van der Waals equation of state

36

36

Slide 32

ES2

Eugene Switkes, 1/6/2020

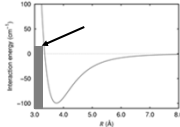
Chemistry 163B, Winter 2020

Lecture 1- Introductory Lecture

van der Waals equation the parameter b

ideal gas law $P_{ideal} \bar{V}_{ideal} = RT$

van der Waals $\left(P + \frac{a}{\bar{V}^2}\right)(\bar{V} - b) = RT$



interpretation of parameters:

- b is correction for actual volume of atoms/molecules
 - volume available to molecules $(\bar{V} - b)$ smaller than \bar{V}
 - b is associated with repulsive forces

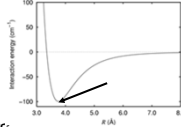
37

37

van der Waals equation the parameter a

$P_{ideal} \bar{V}_{ideal} = RT$

$\left(P + \frac{a}{\bar{V}^2}\right)(\bar{V} - b) = RT$



interpretation of parameters.

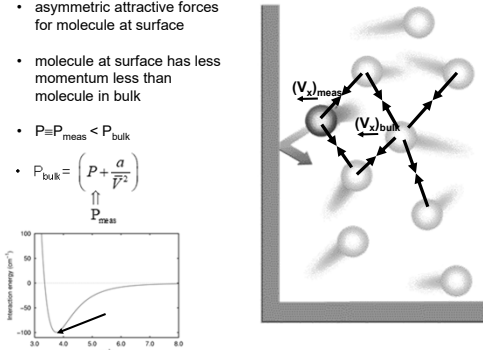
- a is correction for attractive forces of atoms/molecules
 - a is associated with attractive forces
 - $P_{bulk} \equiv P_{ideal}$ is greater than P_{meas} measured at surface
 - $P_{bulk} = \left(P_{meas} + \frac{a}{\bar{V}^2}\right)$ $P_{meas} \rightarrow P_{bulk}$ $\bar{V} \rightarrow \infty$

38

38

heuristic justification for attractive constant a

- asymmetric attractive forces for molecule at surface
- molecule at surface has less momentum less than molecule in bulk
- $P_{meas} < P_{bulk}$
- $P_{bulk} = \left(P + \frac{a}{\bar{V}^2}\right)$



39

39

van der Waals equation

$$\left(P + \frac{a}{\bar{V}^2}\right)(\bar{V} - b) = RT$$

Table 1.1 van der Waals parameters		
Gas	b (L mol ⁻¹)	a (L ² bar mol ⁻²)
He	0.0238	0.0346
Ne	0.01672	0.208
Ar	0.03201	1.355
Kr	0.0396	2.325
Acetylene	0.0522	4.516
N ₂	0.0387	1.37
H ₂ O	0.03049	5.537
CO ₂	0.04286	3.658

Source: Handbook of Chemistry and Physics, 78th edition, CRC Press, Boca Raton, FL, 1997-98

40

40

van der Waals equation

$$\left(P + \frac{a}{\bar{V}^2}\right)(\bar{V} - b) = RT$$

Gas	b (L mol ⁻¹)	a (L ² bar mol ⁻²)
He	0.0238	0.0346
Ne	0.01672	0.208
Ar	0.03201	1.355
Kr	0.0396	2.325
Acetylene	0.0522	4.516
N ₂	0.0387	1.37
H ₂ O	0.03049	5.537
CO ₂	0.04286	3.658

Source: Handbook of Chemistry and Physics, 78th edition, CRC Press, Boca Raton, FL, 1997-98

41

41

some trash talk on VDW eqn: inaccurate but instructive !!! (from other texts)

Validity

However, the Van der Waals model is not appropriate for rigorous quantitative calculations, remaining useful only for **teaching** and qualitative purposes.^[1]

Nowadays, Eq. 2.9 belongs to “**pedagogical physics**.” it is the simplest equation that illustrates several important concepts, but its accuracy is not satisfactory.

42

42

Chemistry 163B, Winter 2020

Lecture 1- Introductory Lecture

Menu: for TODAY



- ✓ Introduction and Class Resources, etc.
- ✓ Careful definition of thermodynamic variables (P , V , T , etc)
- ✓ Ideal gas ENERGY DEPENDS ONLY ON TEMPERATURE (T)
- ✓ Van der Waals equation (example of equation of state for 'real' gas)

43

43

End of Lecture 1

44

44

various equations of state (Raff Table 1.2)

Table 1.2 Some commonly used equations of state

Equation of State	Functional Form	Number of Parameters
Ideal gas	$PV_m = RT$	0
van der Waals	$(V_m - b)(P + a/V_m^2) = RT$	2
Dieterici	$P(V_m - b)\exp[a/RTV_m] = RT$	2
Berthelot	$(V_m - b)(P + a/TV_m^2) = RT$	2
Virial	$P = RT \left[V_m^{-1} + \sum_{n=2}^{\infty} C_n(T) V_m^{-n} \right]$	∞
Beattie-Bridgman	$PV_m^2 = (1 - \gamma)RT(V_m + \beta) - \alpha$ with $\gamma = c_1/T^2 V_m$, $\beta = b_1(1 - b_2/V_m)$, and $\alpha = a_1(1 + a_2/V_m)$	5
Redlich-Kwong	$P = \frac{RT}{(V_m - b)} - \frac{a}{T^{1/2}V_m(V_m + b)}$	2
Reichsanstalt	$PV = RT + AP + BP^2 + CP^3 + \dots$	∞

45

45

The Haber Process (thermodynamics and kinetics)

Previously the problem had been that N_2 is a very stable molecule, and so most attempts to convert it to less stable molecules, such as NH_3 , failed because of thermodynamic or entropy problems. The secret to the Haber-Bosch process proved to be a catalyst of iron with a small amount of aluminium added (aluminium was at the time an exotic and expensive metal that probably attracted Haber's attention as a novelty). The Haber-Bosch process operates at high pressure so as to shift the equilibrium to the right, and high temperature to increase the rates of the reaction. Of course, operating at high temperature actually shifted the reaction to the left, but the trade-off for faster rates was accepted. By removing the ammonia as liquid ammonia, the equilibrium is continuously shifted to the right.

46

46

heuristic

heu-ris-tic [h'voo-ris-tik or, often, yoo-] adjective

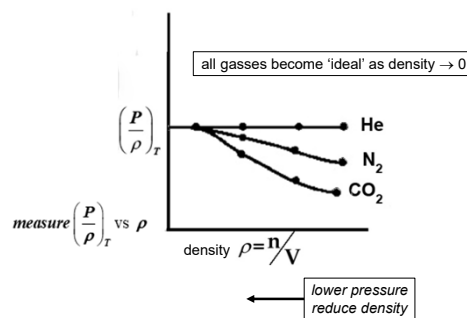
1. serving to indicate or point out; stimulating interest as a means of furthering investigation.



47

47

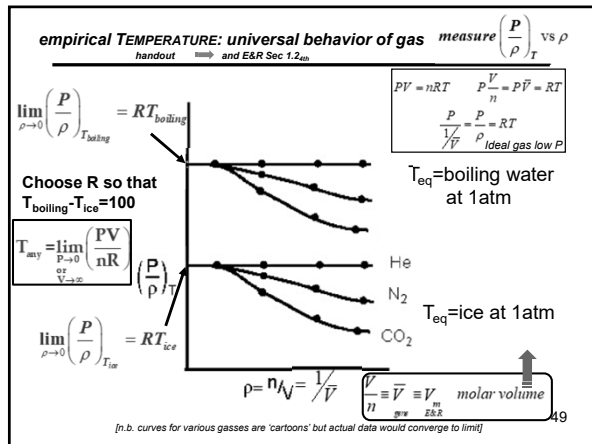
universal behavior of gas



48

48

Chemistry 163B, Winter 2020 Lecture 1- Introductory Lecture



49