

General Physics I

Lect.25 Applications of Themodynamics



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Final Exam

- Time: Dec. 26, 2023 9:00-11:00* (extension possible based on difficulty)
- Room: E10-201
- Type: Close-book
- Scope: Lecture 13 onwards, similar to midterm, 4-5 problems



Applications of Kinetic Theory

Previously, we saw that the probability of finding a particle in different places, per unit volume, varies as e^{-potential energy/kT} based on kinetic theory.

We can use such basic observation to explain phenomena such as:

- a liquid evaporating, or
- electrons in a metal coming out of the surface, or
- a chemical reaction with a large number of atoms involved.

In such cases it is no longer possible to make from the kinetic theory, to describe above processes with 100% accuracy, because the situation is too complicated. But we can understand *more or less* how things ought to behave.

In the cases we will discuss, potential energy needs to be larger than kT for it to be the dominant effect.

Evaporation

Suppose we have a box with a large volume, partially filled with liquid in equilibrium and with the vapor at a certain temperature. The ratio of the molecules in the vapor phase, compared to the ones in the liquid at temperature T can be written as:

$$nV_a = e^{-W/kT}$$

where n is the number of molecules per unit volume in the vapor, and $1/V_a$ is the number of atoms per unit volume in the liquid, given the volume occupied to be V_a . The energy difference in the two phases is W=E_{vapor}-E_{liquid.} To understand this, simply recall the Maxwell-Boltzmann distribution we discussed in L17.

$$n = n_0 e^{-mgh/kT}$$
 $n = ({
m constant}) e^{-{
m P.E.}/kT}$

(generalize)

(specific)





Evaporation



<u>Alternatively</u>, consider that the vapor molecules continually bombard the surface of the liquid, at a given moment a certain number of atoms condense onto the surface of the liquid, given by $N_c=nv$, where v is the mean velocity related to kT. Molecules in a liquid can also be kicked out when they acquire excess energy due to

strong attraction forces, e.g., <u>Van der Waals</u>. The probability of an atom picking up more than W work is $e^{-W/kT}$.

$$nv = (v/V_a)e^{-W/kT}$$

Consider a surface of the liquid occupies a certain cross-sectional area A, and the molecule with average speed v have to move one molecular diameter D (the thickness of the first layer). Then the number evaporating in a chunk of D*A liquid is

$$(1/A)(v/D)e^{-W/kT}$$

To evaluate how fast it will evaporate per volume, one can assume an unknown reflection coefficient R, as we add a correction term (1-R) in the above relation.

$$N_e=nv(1-R)=(v(1-R)/V_a)e^{-W/kT}$$

Thermionic emission



A heated <u>tungsten filament</u> with a positively <u>charged plate</u> to attract the electrons, any electron that escapes from the surface of the tungsten is immediately swept away to the plate. How many electrons per second can we get out of a piece of tungsten? The only difference is the $W=q_e\phi$, where ϕ is the so-called work function. The current per unit area is equal to the charge on each times the number that arrive per second per unit area, which is the number per unit volume times the velocity:

$$I=q_env=(q_ev/V_a)e^{-q_e\phi/kT}$$

Now one electron volt corresponds to kT at a temperature of 11,600 K, so at the operating at a temperature of 1000 K, this exponential factor is about e^{-10} . The constant factors in front is quite wrong, as a limit for the classical theory. And if W is a linear function of T, like $W=W_0+\alpha kT$, then it will appear as a correction of the constant factor in front.

$$e^{-W/kT} = e^{-(W_0+lpha kT)/kT} = e^{-lpha} e^{-W_0/kT}$$

Thermal ionization



Now let's consider ionization. Suppose that in a gas total of N atoms are in the neutral state, but the gas is hot and the atoms can become ionized. The number of atoms and positive ions follows the <u>number conservation</u> $n_a+n_i=N$, and <u>charge conservation</u> $n_e=n_i$ for electrons. Now we call the energy to lift an electron out of the atom, the ionization energy, W. Now the "Boltzmann factor" (i.e., $e^{-W/kT}$) represents the ratio of the number of free electrons per unit volume in the "vapor" over the number of bound electrons per unit volume in the atoms.

The total number of places for electrons is n_i (two electron cannot occupy the same atom state) and we will suppose that they are bound within a certain volume V_a . So the total amount of volume which is available to electrons which would be bound is n_iV_a . Equating this to the number of free electrons per unit volume, n_e , we have

$$rac{n_e n_i}{n_a} = rac{1}{V_a} \, e^{-W/kT}$$
 (Saha ionization equation)

or
$$\frac{f^2}{1-f} \frac{N}{V} = \frac{e^{-W/kT}}{V_a}$$
 with $n_e = fN/V = n_i$ and $n_a = (1-f)N/V$

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If V goes up, fraction f of electrons and ions also goes up.



Chemical kinetics

Without loss of generality, let us now consider chemical reaction where A and B combine into a compound AB, with a work W released. And the same relation holds in terms of the respective density $n_{A/B/AB}$, up to an unknown constant c. W governs the proportions of A/B/AB.

$$rac{n_A n_B}{n_{AB}} = c e^{-W/kT}$$



In reality, A and B need to hit each other with sufficient energy above activation energy, A*. So the reaction of A+B=AB is $e^{-A^*/kT}$ suppressed, while $R_f=R_r$ at equilibrium.

$$R_f = n_A n_B v \sigma_{AB} e^{-A^{m{*}/kT}} \qquad R_r = c' n_{AB} e^{-(W+A^{m{*}})/kT}$$
 .

A* also controls the rate of the reaction. Using intermediate state, or a special surface of the reaction can effectively decrease the energy barrier to increase the rates. These are called *catalysts*.

Einstein's laws of radiation



Consider the mth level and the nth level of an atom, Einstein assumed that there are three kinds of processes: an <u>absorption</u> proportional to the intensity of light, an emission proportional to the intensity of light, called <u>induced emission</u> or sometimes <u>stimulated</u> <u>emission</u>, and a <u>spontaneous emission</u> independent of light. And $E_m - E_n = \hbar \omega$. Now suppose that we have, in equilibrium at temperature T, a certain number of atoms N_n in the state n and another number N_m in the state m, then each second

$$R_{n
ightarrow m} = N_n B_{nm} I(\omega) \quad R_{m
ightarrow n} = N_m [A_{mn} + B_{mn} I(\omega)].$$

At equilibrium, equating the two rates gives

$$B_{nm}I(\omega)e^{\hbar\omega/kT}=A_{mn}+B_{mn}I(\omega)$$
 .

Substitute using $N_m = N_n e^{-\hbar\omega/kT}$, the intensity $I(\omega) = rac{A_{mn}}{B_{nm}e^{\hbar\omega/kT}-B_{mn}}$

Compare with Planck's result, $I(\omega) d\omega = \frac{\hbar \omega^3 d\omega}{\pi^2 c^2 (e^{\hbar \omega/kT} - 1)}$, we have $A_{mn}/B_{mn} = \hbar \omega^3/\pi^2 c^2$

More about $I(\omega)$ see L18

LASER

Laser is an acronym for "light amplification by the stimulated emission of radiation."

Imagine we can prepare the m metastable state far above equilibrium than e^{-(Em-En)/kT}. Then they are all dumped down together by induced emissions. To increase the efficiency of the induced effect, nearly perfect mirrors can be placed on each side of a "box", allowing light to be emitted multiple times. Although the mirrors are almost 100% reflecting, there is a slight transmission, resulting in some light being emitted.

In the end, from the conservation of energy, all the light goes out in a nice uniform straight direction which makes the strong light beams that we call laser.



By exciting, say by blue light, a higher state h, which may emit a photon leaving atoms in state m, the number in this state m becomes sufficiently large to start laser action.

Time constant and mean free path

Now let's discuss what happens when things are near, but not exactly in, equilibrium, e.g., the <u>diffusion of ions in a gas</u>. In a gas with a small ion concentration, an electric field causing the ion to accelerate only until it collides with another molecule, losing its momentum. We can define a <u>time constant</u>, τ , which is the average time between collisions, and a total number N₀. The number/probability of each collision follows

$$N(t + dt) = N(t) - N(t) \frac{dt}{\tau} \longrightarrow \frac{dN(t)}{dt} = -\frac{N(t)}{\tau} \longrightarrow N(t) = N_0 \frac{e^{-t/\tau}}{probability \text{ of no collision}}$$

Rather than the time between collisions, we now about how far the particle moves between collisions, the mean free path unit area

 $l = \tau v$

Consider a moving particle which travels a distance dx through a gas which has n_0 scatterers per unit volume, then we shall have $\sigma_c n_0 l = 1$, which is saying there should be one collision, on the average, when the particle travel a distance l, where $\sigma_c = \pi (r_1 + r_2)^2$ is the classical cross section.



Total area covered is $\sigma_c n_0 dx$



Drift speed

Consider special molecule is acted on by some force F (e.g., electric currents in batteries, sedimentation, centrifugal separation), plus the collisions with the background molecules. First let's assume each collision gets a completely "fresh" start. **Drift speed** is the net progress in the direction of the acceleration F/m, proportional to τ , mind that we do not have ½ factor here:

$$v_{
m drift} = rac{F au}{m} = rac{ extsf{\phi}}{\mu}F$$



In the case of electrical field, F=Eq. The drift velocity of atomic ions in krypton and xenon at normal conditions, where we find v_{drift} ~E.

Suppose our special molecule is a heavy object in a background of lighter molecules, i.e., it will not lose its "forward" momentum in each collision. It would take several collisions before its motion was "randomized" again. We can define a new time constant τ , which corresponds to the average time it loses a certain fraction of its momentum. Such a longer τ means the system has a longer average "forgetting time".



Diffusion: spreading by molecular motions



$$J = rac{n_- v \, \Delta T - n_+ v \, \Delta T}{\Delta T} \longrightarrow J = (n_- - n_+) v$$

We say $n_{-}(n_{+})$ is the number per unit volume to the left (right), evaluated one mean free path *l* away from the imaginary surface. So we relate their difference by a gradient

$$(n_{+} - n_{-}) = \frac{dn_{a}}{dx} \Delta x = \frac{dn_{a}}{dx} \cdot 2l \cdot \frac{\text{substitute}}{\text{average v/3}} \quad J_{x} = -\frac{lv}{3} \frac{dn_{a}}{dx}$$
Now substitute l=vt and τ =µm, so $J_{x} = -\frac{1}{3}mv^{2}\mu \frac{dn_{a}}{dx}$

$$Use mv^{2}/2=3kT/2 \quad J_{x} = -\mu kT \frac{dn_{a}}{dx}$$

$$J_{x} = -\mu kT \frac{dn_{a}}{dx}$$

$$D, \text{ the diffusion coefficient, is just kT times µ, the mobility coefficient}$$



Derivation from stat mech



Imagine we have a diffusion current proportional to the density gradient. We now apply a force field F in the x-direction. According to the definition of the mobility μ there will be a drift velocity given by $v_{\text{drift}} = \mu F$, and thus the drift current is $J_{\text{drift}} = n_a \mu F$. Making J_x+J_{drift}=0, so the drift current from F balance the diffusion, we have

$$D \ rac{dn_a}{dx} = n_a \mu F_{\cdot} \longrightarrow rac{dn_a}{dx} = rac{n_a \mu F}{D}$$

Since we are describing an equilibrium condition, the number density n_a follows the Boltzmann distribution w.r.t. potential energy U: $n_a = n_0 e^{-U/kT}$

$$rac{dn_a}{dx} = -n_0 e^{-U/kT} \cdot rac{1}{kT} rac{dU}{dx} \stackrel{\mathrm{dU/dx=-F}}{\longrightarrow} rac{dn_a}{dx} = rac{n_a F}{kT}$$

Comparing the above two equations, we have again reached $D=\mu kT$. Thus this relation is in general true.

Similarly, we can derive the relation of <u>thermal conductivity</u>, as an exercise for homework.

Dark Matter Detector: a drift (time projection) chamber



- **Drift time** is the time for electrons to travel up the detector under the electrical field, which tells about the z position of the event.
- Ring of electron signal undergoes diffusion as it drift: bigger radius with longer drift time
- Phase transition is a core piece of xenon distillation, purification and slow control
- Some unknown effect trapping electrons in the DM detectors still bothers us nowadays, the answer may be thermodynamical...