Law of Thermodynamics

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1 The fundamental postulation and Liouville's theorem

1.1 The Fundamental postulate

An isolated system in equilibrium is equally likely to be found in any of the microstates accessible to it.

- **System** an part of universe that is only weakly coupled to the rest of the universe so that its dynamic is dominated by internal interactions.
- Equilibrium the measurement of quantities are time independent.
- Microstate a complete microscope specification of coordinates of every particles (position and velocity).
- Ensemble a collection of the system that are macroscopically the same but microscopically different.

1.2 Motivation for the fundamental postulation

From the above fundamental postulation and use the ergodic hypothesis, we can relate the macroscopic properties of a system that we can measure to the statistic (probabilistic) description of microstate.

Ergodic hypothesis assumes that 1) the system's internal dynamics are such that the microstates of the system are constantly changing and 2) the system will visit all possible microstate and spend an equal time in each of them. As a result, as we carry out measurement, the system will likely to be found in a configuration (macroscopic properties) that is represented by the most microstates¹.

The following example illustrate this notation: consider a box of 100 identical coins, shaked hard and we measure the number of the coins facing up as we open the box (macroscopic result). We do not really care the actual configuration of the outcome (coin 1 face up, \cdots coin N face down) which is a microscopic property since we know each of these configurations is equally possible. It's easy to see that the most probable result is 50 up and 50 down, but let's see the possibility of outcome that deviate from this average value:

50 up and 50 down
$$=$$
 $\frac{100!}{(50!)^2} \approx 4 \times 10^{27}$
53 up and 47 down $=$ $\frac{100!}{53!47!} \approx 3 \times 10^{27}$
90 up and 10 down $=$ $\frac{100!}{90!10!} \approx 10^{13}$
100 up and 0 down $=$ 1 (1)

where each result is determined by counting their configurations. We see that the probability that result deviate far from the average decay exponentially. For actual physic system, the number of particles are $\propto 10^{23}$, which essentially mean that the possibility of deviation is ignorable and when we measure the macroscopic properties of physical system, we almostly certainly obtain the value given from probabilistic calculation.

The macroscopic propertie of the system can thus be calculated as following: For N particles we have in total 6N coordinates $(q_1, q_2, \dots, q_{3N}, p_1, p_2, \dots, p_{3N})$ which completely define a microscope state. We define the "phase density" as:

$$\rho(q_1, q_2, \cdots, q_{3N}, p_1, p_2, \cdots, p_{3N}, t)$$
 (2)

$$\rightarrow$$
 Probability of finding a system near $(q_1, q_2, \dots, q_{3N}, p_1, p_2, \dots, p_{3N})$ at time t (3)

¹Blundell, p35-37

If property of this system is given by a function f(q, p), then the ensemble average of f at time t will be given as:

$$\langle f(t) \rangle = \frac{\int \int \cdots \int f(q,p)\rho(q,p,t)dq^{3N}dp^{3N}}{\int \int \cdots \int \rho(q,p,t)dq^{3N}dp^{3N}}$$
(4)

With the above definition, $\rho(q, p, t)dq^{3N}dp^{3N}$ gives the number of states (points) that are included in the phase space volume $dq^{3N}dp^{3N}$ near (q, p).

1.3 Liouville's theorem

Liouville's theorem states that the evolution of ρ is given by:

$$\frac{d\rho}{dt} = 0\tag{5}$$

which is to say that if we follow the trajectory of a state (q, p) as it evolve over time, its phase space density will not change (total derivative): $\rho(q(0), p(0), t = 0) = \rho(q(t'), p(t'), t = t')$.

Proof 1 In this proof, we consider the phase space points inclosed by a volume at t = 0 at (q_1, p_1) , at a later time δt , we locate those phase space points agian and we show that the volume of phase space that enclose these points are the same. This will thus mean the phase (point) density do not change following the trajectory.

let's consider an area in a two dimensional phase space that is a rectangle specified by its 2 diagonal points $(q_1, p_1), (q_2, p_2)$ at some initial time t, then at time $t + \delta t$, the points changed to $(q_1 + \dot{q}_1 \delta t, p_1 + \dot{p}_1 \delta t)$ and $(q_2 + \dot{q}_2 \delta t, p_2 + \dot{p}_2 \delta t)$. The volume difference, to first order in δt is:

$$\Delta V = (q_2 + \dot{q}_2 \delta t - q_1 - \dot{q}_1 \delta t)(p_2 + \dot{p}_2 \delta t - p_1 - \dot{p}_1 \delta t) - (q_2 - q_1)(p_2 - p_1)
= (\dot{q}_2 - \dot{q}_1)(p_2 - p_1) + (\dot{p}_2 - \dot{p}_1)(q_2 - q_1)
= \frac{1}{V} \left(\frac{\dot{q}_2 - \dot{q}_1}{q_2 - q_1} + \frac{\dot{p}_2 - \dot{p}_1}{p_2 - p_1} \right)
= \frac{1}{V} \left(\frac{\partial \dot{q}}{\partial q} + \frac{\partial \dot{p}}{\partial p} \right) \delta t$$
(6)

If a system envolve under Hamiltonian dynamics:

$$\dot{q}_i = \frac{\partial H}{\partial p_i}; \quad \dot{p}_i = -\frac{\partial H}{\partial q_i}$$
 (7)

then

$$\frac{\partial \dot{q}}{\partial q} + \frac{\partial \dot{p}}{\partial p} = \frac{\partial^2 H}{\partial p \partial q} - \frac{\partial^2 H}{\partial q \partial p} = 0 \tag{8}$$

which shows that the enclosing volume of those phase space points do not change as the system evolve, and therefore, the phase space density in this volume do not change over time, giving the result:

$$\frac{d\rho}{dt} = \frac{\partial\rho}{\partial t} + \sum_{i} \left(\frac{\partial\rho}{\partial q_i} \frac{\partial q_i}{\partial t} + \frac{\partial\rho}{\partial p_i} \frac{\partial p_i}{\partial t} \right) = 0 \tag{9}$$

Where the first equality is given merely by the definition of total derivative.

Proof 2 In this proof, we compute the partial derivatives first and show that they result in the result of Liouville's theorem 2 .

We first compute $\partial \rho/\partial t$. Consider the flow the phase space points in and out of a cubic volume element in the phase space arount (q, p), The net flow of phase space points is given by:

$$\frac{\partial N}{\partial t} = -\sum_{i} \left(\frac{\partial (\rho \dot{q}_{i})}{\partial q_{i}} + \frac{\partial (\rho \dot{p}_{i})}{\partial p_{i}} \right) dq \cdots dp$$

$$\frac{\partial \rho}{\partial t} = -\sum_{i} \left(\frac{\partial (\rho \dot{q}_{i})}{\partial q_{i}} + \frac{\partial (\rho \dot{p}_{i})}{\partial p_{i}} \right) \tag{10}$$

²Taken from https://hepweb.ucsd.edu/ph110b/110b_notes/node93.html

the total derivative is then:

$$\begin{split} \frac{d\rho}{dt} &= \frac{\partial \rho}{\partial t} + \sum_{i} \left(\frac{\partial \rho}{\partial q_{i}} \frac{\partial q_{i}}{\partial t} + \frac{\partial \rho}{\partial p_{i}} \frac{\partial p_{i}}{\partial t} \right) \\ &= -\sum_{i} \left(\frac{\partial (\rho \dot{q}_{i})}{\partial q_{i}} + \frac{\partial (\rho \dot{p}_{i})}{\partial p_{i}} \right) + \sum_{i} \left(\frac{\partial \rho}{\partial q_{i}} \frac{\partial q_{i}}{\partial t} + \frac{\partial \rho}{\partial p_{i}} \frac{\partial p_{i}}{\partial t} \right) \\ &= -\sum_{i} \left(\rho \frac{\partial \dot{q}_{i}}{\partial q_{i}} + \rho \frac{\partial \dot{p}_{i}}{\partial p_{i}} \right) = 0 \end{split} \tag{11}$$

thus proving the Liouville's theorem.

2 Ensembles

We can define an ensemble as a collection of possible configurations that satisfy a given macroscopic property³. We have already seen an ensemble containing possible configurations of **an isolated** system where energy is known and all configurations are of equal probability. This ensemble is known as the **Microcanonical ensemble**. For a microcanonical ensemble, particle number, volume and energy are specified at the same time, thus it is also called **NVE ensemble**.

Now consider a system that can exchange energy through a contact with a temperature bath and eventually come to an equilibrium. This ensemble is called **Canonical ensemble**. For canonical ensemble, it's dynamic is still governed by its internal interaction but now its energy may vary. Since now the energy of this system can change, it is more appropriate to describe it in terms of temperature T, from which we can find its energy.

Now we want to find the probability distribution of its microstate (probability of finding the system to be in a specific microstate), The system and reservior together is described by a microcanonical ensemble, in which each state is described by $(q_1, \dots, q_n, q_{n+1}, \dots q_N, p_1, \dots, p_n, p_{n+1}, \dots p_N)$ where the first n coordinates describe the system, and the rest coordinates describe the microstate of the reservior. Every microstates of the combined system are equal likely, therefore, the probability to find the microstate of the canonical ensemble $(q'_1, \dots, q'_n, p'_1, \dots, p'_n)$ depend on the number of possible configurations of the coordinates in the reservior, i.e. the number of the microstate of the combined system in which $(q_1, \dots, q_n, p_1, \dots, p_n) = (q'_1, \dots, q'_n, p'_1, \dots, p'_n)$.

We have, with E_i^r denote the energy of the reservior for the microstate *i* of system, and $\Delta E = E_j - E_i$, the relative probability of two distinct microstates of the system(which actually depend on the number of corresponding states in the reservior):

$$\frac{P_j}{P_i} = \frac{\Omega_r(E_i^r - \Delta E)}{\Omega_r(E_i^r)} = \exp\left(\ln \Omega_r(E_i^r - \ln \Delta E) - \Omega_r(E_i^r)\right) \approx \exp\left(-\frac{E_j - E_i}{k_B T}\right)$$
(12)

Where we introduced the statistic definition of temperature

$$\frac{\mathrm{d}\ln\Omega}{\mathrm{d}E} = \frac{1}{k_B T} \tag{13}$$

This give the result, with $\beta = 1/k_BT$

$$P_i \propto e^{-\beta E_i} \tag{14}$$

Define the canonical partition function Q_N

$$Q_N(T, V, N) = \sum_j e^{-\beta E_j} \tag{15}$$

The free energy is defined by:

$$A(T, V, N) = -k_B T \ln Q_N \tag{16}$$

³Blundell, p38

The probability of finding a given microstate of the system is then

$$P_i = \frac{1}{Q_N} e^{-\beta E_i} = e^{\beta (A - E_i)} \tag{17}$$

To specific a canonical potential, we need to specify V, N, T, therefore, canonical potential is also known as the **NVT ensemble**

3 First Law

Suppose we have the function f(x), where x are parameters of the state. If the integral:

$$\Delta f = \int_{x_i}^{x_f} df = f(x_f) - f(x_i) \tag{18}$$

i.e., if Δf is independent of the path chosen, then we call f a function of state, which only depend on the system parameter at a certain instance.

Equation of states are equation that contain only function of states: i.e., for ideal gas, we have pV = RT which depend on p, V, T, all of them are function of states.

First law: energy is conserved in the form of heat and work.

Writting U as the internal energy, The equation of states can be written as

$$\Delta U = \Delta Q + \Delta W \tag{19}$$

$$dU = \tilde{\mathbf{d}}Q + \tilde{\mathbf{d}}W = \tilde{\mathbf{d}}Q - pdV \tag{20}$$

where Δ gives the total change and d represent differential change. $\tilde{\mathbf{d}}$ represent changes that are path dependent. It is clear that the heat absorbed and the work done to the system depend on the specific process, and thus they cannot be exact differentiated with respect to the system parameters, while volume V is a function of states.

3.1 Heat capacity of gas at constant volume or pressure

For ideal gas, we have:

$$\tilde{\mathbf{d}}Q = \mathrm{d}U + p\mathrm{d}V \tag{21}$$

$$\tilde{\mathbf{d}}Q = \left(\frac{\partial U}{\partial T}\right)_V dT + \left(\frac{\partial U}{\partial V}\right)_T dV + pdV \tag{22}$$

$$\frac{\tilde{\mathbf{d}}Q}{\mathrm{d}T} = \left(\frac{\partial U}{\partial T}\right)_V + \left[\left(\frac{\partial U}{\partial V}\right)_T + p\right] \frac{\mathrm{d}V}{\mathrm{d}T} \tag{23}$$

Therefore, at fixed volume, dV = 0 and we have:

$$C_V = \left(\frac{\tilde{\mathbf{d}}Q}{\mathrm{d}T}\right)_V = \left(\frac{\partial U}{\partial T}\right)_V \tag{24}$$

at fixed pressure:

$$C_p = \left(\frac{\tilde{\mathbf{d}}Q}{\mathrm{d}T}\right)_p \tag{25}$$

$$= C_V + \left[\left(\frac{\partial U}{\partial V} \right)_T + p \right] \left(\frac{\mathrm{d}V}{\mathrm{d}T} \right)_p \tag{26}$$

For ideal gas, we have $\left(\frac{\partial U}{\partial V}\right)_T = 0$. and $p\left(\frac{\mathrm{d}V}{\mathrm{d}T}\right)_p = R$, so that we have: $C_p = C_V + R$. For an equilibrium gas at constant pressure, the volume of the gas also expand when the temperature of the gas is increased (gas

do work to the environment). So we in generally need to input more heat to the increase the temperature of the gas, compared to the case of constant volume.

In general gas, we define the ratio γ called **adiabatic index**:

$$\gamma = \frac{C_p}{C_V} \tag{27}$$

4 Second Law

4.1 Reversible process

we define a process to be reversible if at every moment during the process, the system is in equilibrium. As an example, for an ideal gas, if the equation of state pV = RT is hold true at every moment during the process, then it implies that the gas is always in an equilibrium state and the process is reversible: we could not notice if this process happens in reversed. We consider two types of reversible process in terms of gas: First is called **Isothermal process** and the second is called **Adiabatic process**.

In the isothermal process, the temperature of the system is fixed, in contact with a heat reservior:

$$\Delta T = 0 \tag{28}$$

For ideal gas, this implies that $\Delta U = 0$, for reversible process of ideal gas, The equation of states pV = RT is always true. This gives:

$$\Delta U = \tilde{\mathbf{d}}Q - \frac{RT}{V}dV = 0 \tag{29}$$

$$\Delta Q = \int_{V_1}^{V_2} \tilde{\mathbf{d}} Q = \int_{V_1}^{V_2} \frac{RT}{V} dV = RT \ln \frac{V_2}{V_1}$$

In the adiabatic process, the system is not allowed to exchange heat with the environment, thus we have $\tilde{\mathbf{d}}Q = 0$. We find:

$$dU = \tilde{\mathbf{d}}W \tag{30}$$

$$C_V dT = -p dV = -\frac{RT}{V} dV$$
(31)

$$\ln \frac{T_2}{T_1} = -\frac{R}{C_V} \ln \frac{V_2}{V_1} \tag{32}$$

For ideal gas, $C_p = C_V + R$, $\gamma = 1 + R/C_V$ and we find:

$$\ln \frac{T_2}{T_1} = (1 - \gamma) \ln \frac{V_2}{V_1} \tag{33}$$

$$\frac{T_2}{T_1} = \left(\frac{V_2}{V_1}\right)^{1-\gamma} = \left(\frac{V_1}{V_2}\right)^{\gamma-1} \tag{34}$$

$$TV^{\gamma-1} = \text{const}$$
 (35)

$$pV^{\gamma} = \text{const}$$

where we used relationship pV = RT to obtain the last relationship. Equation pV^{γ} enable us to find the pressure of the system if we know the volume of the system during a adiabatic expansion or compression. Temperature can be derived using the ideal gas law then, if we know the pressure and volume.

4.2 Second law

The second law can be stated in the following two equivalent form:

Clausius statement No process whose sole effect is to transfer heat from cold to hot body

Kelvin statement No process is possible whose sole effect is to convert heat into work

It is important to note the word "sole effect". For example, in an isothermal process of ideal gas, $\tilde{\mathbf{d}}Q + \tilde{\mathbf{d}}W = 0$ so all the heat is converted into work done by the system. However, the accompanying effect is that the volume of the gas expanded. Therefore, To study the ability to convert heat into work, we should consider process which has no other effect other than the conversion, such as Carnot engine which work in cycles

4.3 Carnot engine

We define a carnot engine which consist of two reversible adiabatic and two reversible isothermal process, working between temperature T_h and T_l . The carnot engine do work in the following process:

$$\begin{array}{ll} A \to B & \text{isothermal expansion} & Q_h = RT_h \ln \frac{V_B}{V_A} \\ B \to C & \text{adiabatic} & \frac{T_h}{T_l} = \left(\frac{V_C}{V_B}\right)^{\gamma-1} \\ C \to D & \text{isothermal compression} & Q_l = -RT_h \ln \frac{V_D}{V_C} \\ D \to A & \text{adiabatic} & \frac{T_h}{T_l} = \left(\frac{V_D}{V_A}\right)^{\gamma-1} \end{array}$$

The adiabatic process lead to $\frac{V_C}{V_B} = \frac{V_D}{V_A}$, with $V_B > V_A$. Finally, we find the important result for a carnot energy:

$$\left| \frac{Q_h}{T_h} = \frac{Q_l}{T_l} \right|$$

$$Q_h - Q_l = W$$
(36)

where W is the work done in one carnot cycle. We can find the efficiency of the engine ⁴:

$$\eta = \frac{W}{Q_h} = 1 - \frac{T_l}{T_h} \tag{37}$$

We have the following statement related to the (reversible) carnot engines:

- Carnot engine is the most efficient engine
- all reversible engine operating at the same temperature environment have the same efficiency
- Clausius statement of the second law is the same as Kelvin's statement ⁵

4.4 Refrigerator

We consider the above carnot energy run backwards: it absorb heat Q_l from the low temperature side with an isothermal expansion. Input work W is provided during the adiabatic process, and dump heat Q_h into the hot side with an isothermal compression process. The result for such a refrigerator is:

$$Q_l + W = Q_h \tag{38}$$

$$\frac{Q_h}{T_h} = \frac{Q_l}{T_l} \tag{39}$$

$$\eta = \frac{Q_l}{W} = \frac{T_l}{T_h - T_l} \tag{40}$$

⁴This carnot engine seems to convert all the heat absorbed into work, $Q_h - Q_l \to W$. But it has the additional effect to transport heat from one source to another. Therefore, this does not conflict with Kelvin's statement

⁵See book

4.5 Clausius theorem

Suppose we have a seriers of heat exchange in a working cycle of an engine. At each step, the engine exchange heat $\tilde{\mathbf{d}}Q_i$ from a **contact** with temperature T_i . The maximum work this engine can produce is then:

$$\Delta W = \sum_{i} \tilde{\mathbf{d}} Q_i \tag{41}$$

Furthermore, suppose each of these contact are connect to a single environment with temperature T through a carnot cycle. Each of these carnot cycle can be considered as heat pumps or refrigerators, which operates to provide (absorb) heat $\tilde{\mathbf{d}}Q_i$ at each contact. We have the relationship for these carnot cycles:

$$\frac{\tilde{\mathbf{d}}Q}{T} = \frac{\tilde{\mathbf{d}}Q_i}{T_i} \tag{42}$$

$$\tilde{\mathbf{d}}Q = \tilde{\mathbf{d}}Q_i + \tilde{\mathbf{d}}W_i \tag{43}$$

Since this system work in cycles, it cannot absorb heat from a single heat resevior (environment at T) and output work (second law). So the total work output by this system is necessary zero (do nothing) or negative (external work need to be provided to get this system going). So we have the following requirement:

$$\sum_{i} \tilde{\mathbf{d}} W_i + \Delta W \le 0 \tag{44}$$

We have:

$$\tilde{\mathbf{d}}W_i = \tilde{\mathbf{d}}Q - \tilde{\mathbf{d}}Q_i = \tilde{\mathbf{d}}Q_i \left(\frac{T}{T_i} - 1\right) \tag{45}$$

$$\Delta W = \sum_{i} \tilde{\mathbf{d}} Q_i \tag{46}$$

So we can find:

$$\sum_{i} \tilde{\mathbf{d}} Q_i \frac{T}{T_i} \le 0 \tag{47}$$

$$\sum_{i} \frac{\tilde{\mathbf{d}}Q_{i}}{T_{i}} \le 0 \quad \text{or} \quad \oint \frac{\tilde{\mathbf{d}}Q}{T} \le 0$$

The second equation is the result of the Clausius theorem. The equal sign is only achieved for a reversible energy, which we write:

$$\oint \frac{\tilde{\mathbf{d}}Q_{rev}}{T} = 0$$

4.6 Entropy

For an reversible process, we have the relationship:

$$\oint \frac{\tilde{\mathbf{d}}Q_{rev}}{T} = 0$$
(48)

suggesting that the integral:

$$\int_{A}^{B} \frac{\tilde{\mathbf{d}}Q_{rev}}{T} \text{ is path independent}$$
 (49)

We therefore define a value $dS = \tilde{\mathbf{d}}Q_{rev}/T$ and the value of S is an function of state. S is called the (Clausius) entropy. For adiabatic process, $\tilde{\mathbf{d}}Q = 0$ and dS = 0.

In general, we consider a cycle consists of both reversible and irreversible process. According to Clausius theorem, we have:

$$\oint \frac{\tilde{\mathbf{d}}Q}{T} = \int_{A}^{B} \frac{\tilde{\mathbf{d}}Q}{T} + \int_{B}^{A} \frac{\tilde{\mathbf{d}}Q_{rev}}{T} \le 0$$
(50)

so that we have:

$$\int_{A}^{B} \frac{\tilde{\mathbf{d}}Q}{T} \le \int_{A}^{B} \frac{\tilde{\mathbf{d}}Q_{rev}}{T} = \int_{A}^{B} dS$$
 (51)

taking the process from A to B infinitsemial, we thus find:

$$\frac{\tilde{\mathbf{d}}Q}{T} \le \mathrm{d}S \tag{52}$$

and the equality only happens for reversible process.

In thermally isolated system, dQ = 0, and we reach an important conclusion that for such isolated system, the entropy of the system can only increase: $dS \ge 0$.

We take note that in this example, we entropy increase from process A to process B, as indicated by the integral in Eq.51. If we choose the direction to be $B \to A$, then we will have $dS \le 0$. However, for a irreversible process, the direction is fixed and only the direction that lead to an increase of entropy will be naturally occurring $(A \to B)$.

For a reversible process, $\tilde{\mathbf{d}}Q = TdS$ and the first law gives:

$$dU = TdS + \tilde{\mathbf{d}}W \tag{53}$$

$$= T dS - p dV \tag{54}$$

Since value U, T, S, p, V are all function of state. This result is independent of the process and also holds for irreversible process: for an irreversible process, $dU = \tilde{\mathbf{d}}Q + \tilde{\mathbf{d}}W$ but:

$$\tilde{\mathbf{d}}Q \neq T\mathbf{d}S\tag{55}$$

$$\tilde{\mathbf{d}}W \neq -p\mathbf{d}V\tag{56}$$

and Eq.54 still holds

$$\tilde{\mathbf{d}}Q + \tilde{\mathbf{d}}W = \tilde{\mathbf{d}}Q_{rev} - pdV = TdS - pdV$$
(57)

Equation.54 also give the result:

$$p = -\left(\frac{\mathrm{d}U}{\mathrm{d}V}\right)_S \tag{58}$$

$$T = \left(\frac{\mathrm{d}U}{\mathrm{d}S}\right)_V \tag{59}$$

4.7 Joule Expansion

We consider a process of Joule expansion as an example of irreversible process: suppose a container is separate into half, each with volume V_0 . The gas is initially confined one side of the container with pressure p_i . The other side of the container is vacuum. Now we remove the separation and let the gas take up the volume of the whole container.

We have the equation of states:

$$p_i V_0 = RT_i \tag{60}$$

$$p_f 2V_0 = RT_f \tag{61}$$

since the internal energy of the gas does not change, $\Delta U = 0$ and therefore for ideal gas, its temperature remain the same: $T_i = T_f$. We thus have: $p_f = \frac{1}{2}p_i$. and

$$dU = TdS - pdV = 0 (62)$$

$$dS = \frac{p}{T}dV \tag{63}$$

$$\Delta S = \int_{i}^{f} dS = \int_{i}^{f} \frac{p}{T} dV = \int_{i}^{f} \frac{R}{V} dV = R \ln 2$$
 (64)

where we consider the process as a reversible isothermal expansion, but is since S is function of state, the result is true for any process 6 .

4.8 Entropy of macrostates and total entropy

Since we have the statistical definition of temperature

$$\frac{1}{k_B T} = \frac{\mathrm{d} \ln \Omega}{\mathrm{d} E} \tag{65}$$

where Ω is the number of microstates with energy E. Assume a microcanonical ensemble (each microstates have the same energy and are equally probably), then the internal energy coinside with energy E. Using the relation:

$$T = \left(\frac{\partial U}{\partial S}\right)_V \tag{66}$$

We find the result:

$$S = k_B \ln \Omega \tag{67}$$

which we call the **Boltzmann's definition of entropy**. ⁷. The entropy increase of the joule expansion can be obtained from the Boltzmann's entropy formula by considering the number of phase space before and after the expansion: The phase space of velocity remain the same while the phase space of possible particle position is doubled. Giving the result: $\Delta S = R \ln 2$.

The above case correspond to an isolated system with fixed energy (microcanonical ensemble). We now consider the case where the system of interest is connected to a reservior. The number of states microstate of the whole system is N, We have:

$$S_{tot} = S_{sus} + S_{res} = k_B \ln N \tag{68}$$

suppose that the i^{th} microstate of our system correspond to n_i microstate of the reservior, then the probability of our system to be in the i^{th} microstate is then $P_i = n_i/N$ with corresponding entropy of the reservior $S_{res,i} = k_B \ln n_i$. The average entropy of the reservior can then be calculated:

$$S_{res} = \langle S_{res} \rangle = \sum_{i} P_i S_{res,i} = \sum_{i} P_i k_B \ln n_i$$
 (69)

We have than the entropy of our system:

$$S = k_B \left(\sum_i p_i \right) \ln N - \sum_i P_i k_B \ln n_i \tag{70}$$

$$=-k_B\sum_{i}p_i\ln p_i\tag{71}$$

where we used the result $\sum_i p_i = 1$. This is known as the **Gibb expression of entropy**. The reasoning applied here is the same as finding the probability of an canonical system. The probability here can be given, for example, by canonical distribution function $p_i = e^{-\beta E_i}/Z$

Gibbs entropy formula is useful because most of the system we consider are connected to some heat bath at some tmperature T. Similar to why canonical ensemble is more useful than microcanonical ensemble. Also, Gibb's formula only contain probability, which can be measured, compared to number of microstates in Boltzmann formula, which cannot be measured. ⁸

 $^{^6}$ Joule expansion is associated with Maxwell's Demon, which is an interesting read at page 149 of the book

⁷https://physics.stackexchange.com/questions/141321/what-is-the-conceptual-difference-between-gibbs-and-boltzmann-entropies

⁸This is original expression given by the book, which seems to express a slightly different concept related to the macroscopic properties. So I kept it here in the footnote. But I think the reasoning is the same.

Suppose the system have in total N possible microstates of equal probability, which can be divided into i macrostates that we can distinguish by experimental measurement. Each of those macrostates contain n_i microstates. We define the Gibb's

4.9 Entropy of information

In information theory, we define the information content Q of a statement (or a prediction) by:

$$Q = -k \log P \tag{76}$$

where P is the probability that the statement turn out to be true, and k is a positive scaling constants. If we take k = 1, and take \log_2 for logarithm, then the information content is measured in bits.

As an example, consider the following statement: a single coin toss give a result of "face up". The information content is:

$$Q = -\log_2 P = 1 \tag{77}$$

Therefore, this information is 1 bit long: to store the above statement, we only require 1 bit to store whether the result is face up with 0, 1.

Now, to describe(transmit) a probability distribution, we can use(transmit) a series of statements. For example, to describe the probability distribution of weather (sunny, cloudy and rainy) in a day, we have different events with different probabilities and information contents:

- the probability of a rainy day is 20%
- the probability of a sunny day is 40%
- the probability of a cloudy day is 40%

We associate each statement with its information content and define the average information content as:

$$S = \langle Q \rangle = -k \sum_{i} P_i \log P_i \tag{78}$$

information content S is called **Shannon entropy**. Shannon entropy therefore give the average information content of succesive measurements.

As an example, consider a Bernoulli trial with outcome probability p. Using k = 1, the Shannon entropy is calculated by:

$$S = -\sum_{i} P_{i} \log_{2} P_{i} = -p \log_{2} p - (1 - p) \log_{2} (1 - p)$$
(79)

S is maximum when p=0.5, showing that the information content of a Bernoulli distribution is greatest when the uncertainty of output is greatest. When p deviates from 0.5, the outcome become biased and it is very likely that most measurement give the same result. The information content with this probability distribution decrease. In the extreme case, if the event is deterministic with single outcome, than we will gain no information after measurement of the result.

entropy:

$$S_{tot} = S + S_{micro} (72)$$

where $S_{tot} = k_B \ln N$. The microscopic entropy is given by:

$$S_{micro} = \langle S \rangle = \sum_{i} P_{i} S_{i} = \sum_{i} P_{i} k_{B} \ln n_{i}$$
 (73)

where the probability $p_i = n_i/N$ and $\sum_i p_i = 1$. We have:

$$S = k_B \left(\sum_i p_i \right) \ln N - \sum_i P_i k_B \ln n_i \tag{74}$$

$$= -k_B \sum_{i} p_i \ln p_i \tag{75}$$

4.10 Entropy and encoding

The Shannon entropy measures the average information content, which can be interpreted as the **average** bits needed to transit the measurement, when measured in bits. Consider the following two case:

Case 1. Suppose an event have 8 possible outcomes with equal probability, the Shannon entropy is then:

$$S = -8 \times \frac{1}{8} \log_2 \frac{1}{8} = 3 \ bits \tag{80}$$

the distribution is random, so we need to use 3 bits to store each outcome and there is no way to compress the informations.

Case 2. Suppose now that we have different probability of the outcome given by the following distribution:

$$1/2, 1/4, 1/8, 1/16, 1/64, 1/64, 1/64, 1/64$$

Now, the Shannon entropy is:

$$S = -\sum_{i} P_i \log_2 P_i = 2 \ bits \tag{81}$$

It is now possible to compress the result by the following code: 0, 10, 110, 1110, 111100, 111101, 111111 and the average length of the code we need to transmit is:

$$L = 1/2 \times 1 + 1/4 \times 2 + 1/8 \times 3 + 1/16 \times 4 + 4 \times 1/64 \times 6 = 3 \text{ bits}$$
(82)

which agree with the value of the Shannon entropy where the length required to store one result is given by $Q = -\log_2 P$.

5 Third Law

We can measure the change of entropy of a system by measuring its heat capacitiy:

$$C_p = T \left(\frac{\partial S}{\partial T} \right)_n \tag{83}$$

$$S = \int \frac{C_p}{T} dT \tag{84}$$

so that we have:

$$S(T) = S(T_0) + \int_{T_0}^{T} \frac{C_p}{T} dT$$
 (85)

The third law states that the entropy of a system at absolute zero will be zero:

Planck's statement of the third law The entropy of all systems in internal equilibrium is the same at absolute zero and may be taken to be zero.

It is noted that the system should be in a relaxed internal equilibrium for the third law to hold, it is possible, for example, to freeze a metastable glass phase instead of a crystalline phase and go down o $T \to 0$ with non zero entropy.

As a consequence of the third law, Heat capacity, given by Eq.83 will necessarily go to zero as $T \to 0$.

As another implication, Using the statistical definition of the entropy $S = k_B \ln \Omega$ implies that the ground states of a system at absolute zero, is necessary non-degenerate with $\Omega = 1$ so that the entropy will be zero. Such non-degeneracy would be enforced by the third law for real systems, as well as the requirement for $C_p to0$. As an example, the curie's law gives a susceptibility $\chi \to \infty$ as $T \to 0$. But with the third law, we require $\partial \chi/\partial T \to 0$. This problem is caused because for Curie law, we assumed a mean field interaction where the interaction of magnetic moment is ignored⁹.

⁹See discussion in page 204-205

6 Thermodynamic potentials

We call **Thermodynamic potentials** functions that are constructed from the functions of state. We define the following Thermodynamic potentials, The first being the internal energy, the following are the Enthalpy, Helmholtz function and the Gibbs function.

$$\begin{array}{lll} U(S,V) & U & \mathrm{d} U = T \mathrm{d} S - p \mathrm{d} V & T = \left(\frac{\partial U}{\partial S}\right)_V, \, p = -\left(\frac{\partial U}{\partial V}\right)_S \\ H(S,p) & H = U + PV & \mathrm{d} H = T \mathrm{d} S + p \mathrm{d} V & T = \left(\frac{\partial H}{\partial S}\right)_V, \, V = \left(\frac{\partial H}{\partial p}\right)_S \\ F(T,V) & F = U - TS & \mathrm{d} F = -S \mathrm{d} T - p \mathrm{d} V & S = -\left(\frac{\partial F}{\partial T}\right)_V, \, p = -\left(\frac{\partial F}{\partial V}\right)_T \\ G(T,p) & G = H - TS & \mathrm{d} G = -S \mathrm{d} T + V \mathrm{d} p & S = -\left(\frac{\partial G}{\partial T}\right)_p, \, V = \left(\frac{\partial G}{\partial p}\right)_T \end{array}$$

Regarding the thermodynamic potentials, we have the following comments:

- Different thermodynamic potentials are expressed in different **natural variables** because their differential form. For example, because dF = -SdT pdV, the natural variables for F are U, V. The differential form implies that the change in F is related to the change in T or V. if dV = 0, then the change in F is independent of pressure P.
- Each form of the thermodynamic potential gives different formula of thermodynamic quantities, which can be easier to access. For example, $S = -(\partial F/\partial T)_V$ gives the value of entropy with easily accessible quantities F, T.
- Each thermodynamic potentials describe the system subject to different constrains: For a system with fixed pressure and fixed temperature in a process, its Gibbs function will remain the same through the process. While for a system with fixed volume and temperature, it's free energy F will be fixed over the process.

6.1 Maxwell's relationship

Maxwell's relationship can be derived from the above definition of the thermodynamic potential function, and they are stated as below:

$$\left(\frac{\partial T}{\partial V}\right)_S = -\left(\frac{\partial p}{\partial S}\right)_V \tag{86}$$

$$\left(\frac{\partial T}{\partial p}\right)_S = \left(\frac{\partial V}{\partial S}\right)_p \tag{87}$$

$$\left(\frac{\partial S}{\partial V}\right)_T = -\left(\frac{\partial p}{\partial T}\right)_V \tag{88}$$

$$\left(\frac{\partial S}{\partial p}\right)_T = -\left(\frac{\partial V}{\partial T}\right)_V \tag{89}$$

6.2 Legendre transformation

In general, Legendre transformation is a transformation on the real valued convex functions of one of the variable and is usually used to convert functions of one quantity (position, pressure) into function of the conjugate quantity (momentum, volume) Consider function F(x) with dF/dx = s(x), we can write a function G(s) with property dG/ds = x(s). They are related by:

$$d(F+G) = sdx + xds = d(xs) \tag{90}$$

so that F(x) + G(s) = xs. One property of Legendre transformation that the result of the transformation is also convex function. For details of the geometry meaning and convex requirement, see Morin's chapter.

For thermodynamic potentials, we use a non-standard definition dG/ds = -x(s), so that the transformation agree with the thermal dynamic variables. leading to

$$d(F - G) = sdx + xds = d(xs)$$

$$(91)$$

and therefore $F - G = xs^{-10}$.

We have seem that intensity and extensity properties would come in pairs, such as (T, S), (μ, N) and (P, V) due to their definition. We have also see that by defining different ensemble, we can write down different potential function for each ensemble:

$$\begin{array}{l} {\rm NVT~ensemble} \Rightarrow dA(N,V,T) = -SdT - pdV + \mu dN \\ {\rm }\mu {\rm VT~ensemble} \Rightarrow d\Omega(\mu,V,T) = -SdT - pdV - Nd\mu \\ {\rm }(Gibbs)~{\rm NPT~ensemble} \Rightarrow dG(N,p,T) = -SdT + Vdp + \mu dN \end{array}$$

The relationship between different potential function can be expressed with Legendre transformation.

Writting x for an intensive property and S as the conjugating extensive property. For a potential F(x) with the thermodynamic relationship dF = Sdx, if another potential G is related to F(x) by F(x) - G(S) = xS, then, we can find dG(S) = -xdS.

As an example, to find the Legendre transformation of the free energy in canonical ensemble A(N, V, T) in terms of variable V, we have:

$$dA = -pdV$$

The transformed potential is thus:

$$dG(N, p, T) = Vdp$$

corresponding to the Gibbs potential.

6.3 Isothermal magnetization and adiabatic demagnetization

To generalize the above treatment in general system, we can write:

$$\tilde{\mathbf{d}}W = Xdx \tag{92}$$

where X is some intensive generalized force and x is some extensive generalized displacement.

Consider a system of magnetic moments arranged in a lattice. We assume paramagnetic system with no interaction between the magnetic moments. The first law of Thermodynamic for such system is

$$dU = TdS - mdB \tag{93}$$

We define the magnetic susceptibility as:

$$\chi = \lim_{H \to 0} \frac{M}{H} \approx \frac{\mu_0 M}{B} \tag{94}$$

since $B = \mu_0(H + M)$ and $M \ll H$. Curie's law states $\chi \propto 1/T$ and therefore

$$\left(\frac{\partial \chi}{\partial T}\right)_B < 0 \tag{95}$$

Consider the Helmholtz function:

$$dF = -SdT - mdB \tag{96}$$

which gives the Maxwell relation:

$$\left(\frac{\partial S}{\partial B}\right)_T = \left(\frac{\partial m}{\partial T}\right)_B \approx \frac{VB}{\mu_0} \left(\frac{\partial \chi}{\partial T}\right)_B \tag{97}$$

 $^{^{10}}$ Both is obviously correct. An example of the standard usage is Legendre transformation for Lagrangian and Hamiltonian is given by F+G=xs. see "Understanding the transformation in terms of derivatives" section in https://en.wikipedia.org/wiki/Legendre_transformation.

The heat absorbed in an isothermal change of B is:

$$\Delta Q = T \left(\frac{\partial S}{\partial B} \right)_T \Delta B = \frac{TVB}{\mu_0} \left(\frac{\partial \chi}{\partial T} \right)_B \Delta B < 0 \tag{98}$$

So that an isothermal increase of B will release heat to the environment.

The change in temperature in an adiabatic change of B is

$$\left(\frac{\partial T}{\partial B}\right)_S = -\left(\frac{\partial T}{\partial S}\right)_B \left(\frac{\partial S}{\partial B}\right)_T \tag{99}$$

using

$$C_B = T \left(\frac{\partial S}{\partial T} \right)_B \tag{100}$$

we have:

$$\left(\frac{\partial T}{\partial B}\right)_{S} = -\frac{TVB}{\mu_{0}C_{B}} \left(\frac{\partial \chi}{\partial T}\right)_{B} > 0 \tag{101}$$

So that in an adiabatic process, the temperature will decrease with the decrease of magnetization, which is called **adiabatic demagnetization**

7 Equipartition of energy

To derive the equipartition theorem, we first consider a system whose energy is given by a quadratic form:

$$E = \alpha x^2 \tag{102}$$

where α is some positive constant and x is a variable that descript the microscopic configuration of the system. The probability of the system taking configuration x is then given by the canonical distribution:

$$P(x) = \frac{e^{-\beta \alpha x^2}}{\int_{-\infty}^{\infty} e^{-\beta \alpha x^2 dx}}$$
 (103)

The mean energy of the system is then:

$$\langle E \rangle = \int_{-\infty}^{\infty} EP(x)dx$$
 (104)

$$= \frac{\int_{-\infty}^{\infty} \alpha x^2 e^{-\beta \alpha x^2} dx}{\int_{-\infty}^{\infty} e^{-\beta \alpha x^2} dx}$$
 (105)

$$=\frac{1}{2}k_BT\tag{106}$$

For a system with multiple variables, we similarly have:

$$E = \sum_{i=1}^{n} \alpha_i x_i^2 \tag{107}$$

The total energy is then:

$$\langle E \rangle = \frac{\int_{-\infty}^{\infty} \cdots \int_{-\infty}^{\infty} \sum_{i=1}^{n} \alpha_{i} x_{i}^{2} e^{-\beta \sum_{j=1}^{n} \alpha_{j} x_{j}^{2}} dx_{1} \cdots dx_{n}}{\int_{-\infty}^{\infty} \cdots \int_{-\infty}^{\infty} e^{-\beta \sum_{i=1}^{n} \alpha_{i} x_{i}^{2}} dx_{1} \cdots dx_{n}}$$
(108)

$$= \sum_{i=1}^{n} \frac{\int_{-\infty}^{\infty} \cdots \int_{-\infty}^{\infty} \alpha_{i} x_{i}^{2} e^{-\beta \sum_{j=1}^{n} \alpha_{j} x_{j}^{2}} dx_{1} \cdots dx_{n}}{\int_{-\infty}^{\infty} \cdots \int_{-\infty}^{\infty} e^{-\beta \sum_{i=1}^{n} \alpha_{i} x_{i}^{2}} dx_{1} \cdots dx_{n}}$$

$$(109)$$

$$=\sum_{i=1}^{n} \frac{\int_{-\infty}^{\infty} \alpha_i x_i^2 e^{-\beta \alpha_i x_i^2} dx_i}{\int_{-\infty}^{\infty} e^{-\beta \alpha_i x_i^2} dx_i}$$

$$\tag{110}$$

$$=\sum_{i=1}^{n} \frac{1}{2} k_B T = \frac{n}{2} k_B T \tag{111}$$

where n the number of variable of the system (microscope variable that determine the energy of the system), which we can call degree of freedom.

The formal equipartition theorem states: If the energy of a classic system is the sum of the n quadratic variables and the system is in contact with a thermal reservior at temperature T, then its average energy is $n \times \frac{1}{2}k_BT$

7.1 Application of equipartition theorem to heat capacity of crystals

We consider each atom in the crystal are kept at their respective equilibrium positions by a spring. Each atom is therefore described by 6 variables: their kinetic energy is given by their velocity \mathbf{v} and their potential energy by their position \mathbf{r} . Therefore, there are 6N degree of freedom in a solid with N numbers of atoms. The total energy of the crystal at temperature T is then:

$$\langle E \rangle = 3Nk_BT \tag{112}$$

and the Molar heat capacity of a solid is then $3N_Ak_B = 3R$, known as the Dulong-Petit rule.

7.2 Assumption and limitation of equipartition theorem

The critical assumption we made in deriving the equipartition theorem is as follows:

- 1. We have assumed that the system variable x is continuous through the intergration $\int_{-\infty}^{\infty} E(x)P(x)dx$. However, for quantum system, the system are found to take distinct states with quantized energy levels. For example, at low temperature, thermal energy may not be enough to excite the system to the next energy level and the system will be in the ground state with P=1. A classical system in this case assumes that there are always levels for system to distribute to even at low temperature and the above integral is valid.
- 2. We assume that that the energy of the system can be written in a quadratic form. In most case this is valid, since the system will minimize its energy in equilibrium and the energy can therefore be written as an expansion to second order: $E = E_0 + \alpha(x x_0)^2$ around $x = x_0$. However, when temperature is high and x start to deviate largely from x_0 , higher order terms become important and quadratic form is no longer valid.

As a conclusion, we learned that the equipartition theorem is valid in a temperature range high enough so that states are almost continuous ($\Delta E \ll k_B T$ with ΔE the energy difference between states) but low enough that the quadratic form of the energy is valid.

8 The partition function

We define the partition function as:

$$Z = \sum_{\alpha} e^{-\beta E_{\alpha}} \tag{113}$$

which allow us to derive all the important thermodynamic quantities.

The internal energy is given by:

$$U = \langle E \rangle = \frac{\sum_{i} E_{i} e^{-\beta E_{i}}}{\sum_{i} e^{-\beta E_{i}}} = -\frac{\mathrm{d} \ln Z}{\mathrm{d} \beta} = k_{B} T^{2} \frac{\mathrm{d} \ln Z}{\mathrm{d} T}$$
(114)

The entropy is found by:

$$S = -k_B \sum_{i} P_i \ln P_i = k_B \sum_{i} P_i (\beta E_i + \ln Z) = k_B (\beta U + \ln Z)$$
 (115)

or we can write:

$$S = U/T + k_B \ln Z \tag{116}$$

Helmholtz function can be written by F = U - TS, using the above result, we have:

$$F = -k_B T \ln Z \tag{117}$$

$$Z = e^{-\beta F} \tag{118}$$

Other derived quantities can also be found:

$$C_V = \left(\frac{\partial U}{\partial T}\right)_V = k_B T \left[2 \left(\frac{\partial \ln Z}{\partial T}\right)_V + T \left(\frac{\partial^2 \ln Z}{\partial T^2}\right)_V \right]$$
(119)

$$p = -\left(\frac{\partial F}{\partial V}\right)_T = k_B T \left(\frac{\partial \ln Z}{\partial V}\right)_T \tag{120}$$

$$H = U + pV = k_B T \left[\left(T \frac{\partial \ln Z}{\partial T} \right)_V + V \left(\frac{\partial \ln Z}{\partial V} \right)_T \right]$$
 (121)

$$G = F + pV = k_B T \left[-\ln Z + V \left(\frac{\partial \ln Z}{\partial V} \right)_T \right]$$
 (122)

It should be noted that the value of the partition function itself is not uniquely defined, because the zero of the energy is arbitrary, so that partitiona function can be defined up to an arbitrary multiplicative constant. However, In terms of the thermodynamic quantities calculated in the above equations, The result not on Z but on $\ln Z$ and its derivative, so the those quantities are up to a additive constants or show no dependence on the arbitrarness of the partition function.

9 Chemical potential and Grand potential

We consider adding particles to a system, then the internal energy of the system will increase by an amount, which we denote **chemical potential**. For a large system with μ not changing significantly by adding or removing a particle, we have

$$dU = TdS - pdV + \mu dN \tag{123}$$

with N the particle number and μ can be written:

$$\mu = \left(\frac{\partial U}{\partial N}\right)_{S,V} \tag{124}$$

In terms of other constrains, we have:

$$dF = -pdV - SdT + \mu dN \tag{125}$$

$$dG = Vdp - SdT + \mu dN \tag{126}$$

and we have the expression for chemical potential as:

$$\mu = \left(\frac{\partial F}{\partial N}\right)_{T,V} = \left(\frac{\partial G}{\partial N}\right)_{T,p} \tag{127}$$

For an isolated system, the entrope will increase as system go to equilibrium. We write:

$$dS = \left(\frac{\partial S}{\partial U}\right)_{N,V} dU + \left(\frac{\partial S}{\partial V}\right)_{N,U} dV + \left(\frac{\partial S}{\partial N}\right)_{U,V} dN$$
(128)

$$=\frac{\mathrm{d}U}{T} + \frac{p\mathrm{d}V}{T} - \frac{\mu\mathrm{d}N}{T} \tag{129}$$

where the second equality follow from Eq.123. Therefore, we identify:

$$\left(\frac{\partial S}{\partial U}\right)_{N,V} = \frac{1}{T}; \quad \left(\frac{\partial S}{\partial V}\right)_{N,U} = \frac{p}{T}; \quad \left(\frac{\partial S}{\partial N}\right)_{U,V} = -\frac{\mu}{T} \tag{130}$$

We consider two system connected to each other and isolated from from the environment, If heat is allowed to follow, we have:

$$dS = \left(\frac{\partial S_1}{\partial U_1}\right)_{N,V} dU_1 + \left(\frac{\partial S_2}{\partial U_2}\right)_{N,V} dU_2$$
(131)

$$= \left(\frac{1}{T_1} - \frac{1}{T_2}\right) dU_1 \ge 0 \tag{132}$$

Therefore, the equilibrium can be found at $T_1 = T_2$.

Similarly, if the system are allowed to exchange particle, we have:

$$dS = \left(\frac{\mu_1}{T_1} - \frac{\mu_2}{T_2}\right) dN_1 \ge 0 \tag{133}$$

the equilibrium can be found at $\mu_1 = \mu_2$, if the two subsystem have already the same temperature.

9.1 Grand partition function

For a grand canonical ensemble, The probability that a system will be in a state with energy E_i and particle number N_i are given by

$$P_i = \frac{e^{\beta(\mu N_i - E_i)}}{\mathcal{Z}} \tag{134}$$

and the grand partition function \mathcal{Z} is given by:

$$\mathcal{Z} = \sum_{i} e^{\beta(\mu N_i - E_i)} \tag{135}$$

We can find the thermodynamic quantities using \mathcal{Z} in a similar way as with Z, with the result:

$$\langle N \rangle = \sum_{i} N_{i} P_{i} = k_{B} T \left(\frac{\partial \ln \mathcal{Z}}{\partial \mu} \right)_{T}$$
 (136)

$$U = -\left(\frac{\partial \ln \mathcal{Z}}{\partial \beta}\right)_{\mu} + \mu N \tag{137}$$

$$S = -k_B \sum_{i} P_i \ln P_i = \frac{U - \mu N + k_B T \ln \mathcal{Z}}{T}$$
(138)

We further define the grand potential:

$$\Phi_G = -k_B T \ln \mathcal{Z} \tag{139}$$

which is a function of state. We have, using Eq.138:

$$\Phi_G = -k_B T \ln \mathcal{Z} = U - TS - \mu N = F - \mu N \tag{140}$$

$$d\Phi_G = dF - \mu dN - Nd\mu \tag{141}$$

9.2 Different types of particles

If there are different types of particle, we can write:

$$dU = TdS - pdV + \sum_{i} \mu_{i} dN_{i}$$
(142)

$$dF = -pdV - SdT + \sum_{i} \mu_{i} dN_{i}$$
(143)

$$dG = Vdp - SdT + \sum_{i} \mu_{i} dN_{i}$$
(144)

10 Quantum Statistical Mechanics

For a quantum **ensemble** $|\Psi\rangle$ expressed in a complete set of basis $|\phi_i\rangle$ (state of system) as

$$|\Psi\rangle = \sum_{i} c_i |\phi_i\rangle \tag{145}$$

we define the density matrix

$$\rho = |\Psi\rangle\langle\Psi| = \sum_{ij} |\phi_i\rangle\langle\phi_i|\Psi\rangle\langle\Psi|\phi_j\rangle\langle\phi_j| = \sum_{ij} c_i c_j^* |\phi_i\rangle\langle\phi_j|$$
(146)

and therefore

$$\langle \phi_i | \rho | \phi_j \rangle = c_i c_j^* \tag{147}$$

So that the expectation value of observables A is then:

$$\langle A \rangle = \langle \Psi | A | \Psi \rangle = \sum_{ij} \langle \Psi | \phi_i \rangle \langle \phi_i | A | \phi_j \rangle \langle \phi_j | \Psi \rangle = \sum_{ij} c_i^* c_j \langle \phi_i | A | \phi_j \rangle$$
 (148)

while we also have:

$$\operatorname{Tr}[\rho A] = \sum_{i} \langle \phi_{i} | \rho A | \phi_{i} \rangle = \sum_{ij} \langle \phi_{i} | \rho | \phi_{j} \rangle \langle \phi_{j} | A | \phi_{i} \rangle = \sum_{ij} c_{i} c_{j}^{*} \langle \phi_{j} | A | \phi_{i} \rangle = \langle A \rangle \tag{149}$$

So that we find the relationship

$$\langle A \rangle = \text{Tr}[\rho A] \tag{150}$$

Now, we consider the time dependence of the trace operator, using the time evolution operator:

$$-ih\frac{\partial}{\partial t}\rho = -ih\frac{\partial}{\partial t}\sum_{ij}c_{i}c_{j}^{*}|\phi_{i}\rangle\langle\phi_{j}|$$

$$= \sum_{ij}c_{i}c_{j}^{*}\left(-ih\frac{\partial}{\partial t}|\phi_{i}\rangle\langle\phi_{j}| - |\phi_{i}\rangle ih\frac{\partial}{\partial t}\langle\phi_{j}|\right)$$

$$= \sum_{ij}c_{i}c_{j}^{*}\left(H|\phi_{i}\rangle\langle\phi_{j}| - |\phi_{i}\rangle\langle\phi_{j}|H\right)$$

$$= [H\rho - \rho H] = [H, \rho]$$
(151)

In an equilibrium, the density operator will be time dependent: $\partial \rho/\partial t = 0$, suggesting that H and ρ commute: $[H, \rho] = 0$. The eigenstates of the Hamiltonian thus are also the eigenstates of the density operator. Therefore, the density operator in Eq.146 can be written in this form:

$$\rho = \sum_{n} w_n |n\rangle\langle n| \tag{152}$$

where $|n\rangle$ are the eigenstates of the Hamiltonian, and $w_n = c_n^* c_n = |c_n|^2$ is the probability for the system $|\Psi\rangle$ to be in the energy eigenstate $|n\rangle$ For Microcanonical ensemble with Ω accessible states, we simply have:

$$w_n = \frac{1}{\Omega} \tag{153}$$

For Canonical ensemble, we have:

$$w_n = \frac{e^{-\beta E_n}}{\sum_{n'} e^{-\beta E_{n'}}} \tag{154}$$

and the partition function $Q = \text{Tr}e^{-\beta H}$ (an observables). For Grand Canonical ensemble, taking account of the particle number in an system state $|n\rangle$, we have:

$$w_n = \frac{e^{-\beta(E_n - \mu N_n)}}{\sum_{n'} e^{-\beta(E_{n'} - \mu N_{n'})}}$$
(155)

and partition function $Q = \text{Tr}e^{-\beta(H-\mu N)}$, where N is the operator for particle density.

10.1 Distribution function

Suppose the Hamiltonian can be written as a function of particle number $H = \sum_{p} \varepsilon_{p} n_{p}$ with ε_{p} , n_{p} denoting the energy and occupation number of a single particle state p. We start by calculating the partition function of a quantum ensemble, since we wish to study the probability of system containing different particle number, we use the grand canonical potential:

$$Q(\mu, V, T) = \sum_{N=0}^{\infty} \sum_{\{n_p\}_N} e^{-\beta \sum_p (\varepsilon_p - \mu) n_p} = \sum_{\{n_p\}} e^{-\beta \sum_p (\varepsilon_p - \mu) n_p}$$

$$(156)$$

where in the first expression, $\{n_p\}_N$ denotes a many body state with occupation $\{n_{p_1}, n_{p_2}, \dots, n_{p_i}\}$ in each of the single particle with the constraint that the total number of particles sum up to N. In the second expression, the two summation in the first expression are combined. We now take the summation p in the exponential out:

$$Q(\mu, V, T) = \sum_{\{n_p\}} e^{-\beta \sum_p (\varepsilon_p - \mu) n_p} = \prod_p \sum_{n_p} e^{-\beta (\varepsilon_p - \mu) n_p} = \prod_p Q_p$$

$$\tag{157}$$

We can now study the sum of $e^{-\beta(\varepsilon_p-\mu)n_p}$ over all possible occupation number: **Fermion** we can only take $n_p=0$ or 1, so that

$$Q_p = 1 + e^{-\beta(\varepsilon_p - \mu)} \tag{158}$$

$$\Omega_p = -k_B T \ln Q_p \tag{159}$$

and

$$N_p = -\left(\frac{\partial\Omega}{\partial\mu}\right)_{TV} = \frac{1}{e^{\beta(\varepsilon_p - \mu)} + 1} \tag{160}$$

Boson the sum of n_p is from 0 to ∞ , thus:

$$\sum_{n_p=0}^{\infty} e^{-\beta(\varepsilon_p - \mu)n_p} = \frac{1}{1 - e^{-\beta(\varepsilon_p - \mu)}}$$
(161)

giving

$$N_p = \frac{1}{e^{\beta(\varepsilon_p - \mu)} - 1} \tag{162}$$

The partition function and total particle number can be summarized by given by:

$$\Omega = -ak_B T \sum_{p} \ln(1 + ae^{-\beta(\varepsilon_p - \mu)})$$
(163)

$$N = \sum_{p} \frac{1}{e^{\beta(\varepsilon_p - \mu)} + a} \tag{164}$$

with a = 1 for fermion and -1 for boson.

10.2 Boson gas

For an ideal gas of spinless bosons ¹¹, the single particle state will be the momentum eigenstate given by k and kinetic energy $\varepsilon_k = \frac{\hbar^2 k^2}{2m}$. The chemical potential μ will be negative otherwise the state with energy

¹¹for spin S, each momentum eigenstate k will have (2S+1) fold degeneracy

smaller than μ will have negative occupation. We calculate the grand potential:

$$\Omega = k_B T \sum_{k} \ln(1 - e^{-\beta(\varepsilon_p - \mu)})$$

$$= k_B T \int_0^\infty \ln(1 - e^{-\beta(\varepsilon_p - \mu)}) g(\varepsilon) d\varepsilon$$

$$= k_B T \int_0^\infty \ln(1 - e^{-\beta(\varepsilon_p - \mu)}) \frac{V \varepsilon^{1/2}}{(2\pi)^2} \left(\frac{2m}{\hbar^2}\right)^{3/2} d\varepsilon$$

$$= -\frac{2}{3} \frac{V}{(2\pi)^2} \left(\frac{2m}{\hbar^2}\right)^{3/2} \int_0^\infty \frac{\varepsilon^{3/2} d\varepsilon}{e^{\beta(\varepsilon_p - \mu)} - 1} \tag{165}$$

define $z = e^{\beta \mu}$, the particle number and internal energy can be written as:

$$N = \frac{V}{(2\pi)^2} \left(\frac{2m}{\hbar^2}\right)^{3/2} \int_0^\infty \frac{\varepsilon^{1/2} d\varepsilon}{e^{\beta \varepsilon_p}/z - 1}$$
 (166)

$$U = \frac{V}{(2\pi)^2} \left(\frac{2m}{\hbar^2}\right)^{3/2} \int_0^\infty \frac{\varepsilon^{3/2} d\varepsilon}{e^{\beta \varepsilon_p}/z - 1}$$
 (167)

The integral we can write in terms of polylogarithm function:

$$\int_0^\infty \frac{\varepsilon^{n-1} d\varepsilon}{e^{\beta \varepsilon_p}/z - 1} = (k_B T)^n (n-1)! L_n(z)$$
(168)

with $n/2! = n/2 \times (n-2)/2 \times \cdots \times 1/2$ with n an odd number. So that

$$N = \frac{V}{(2\pi)^2} \left(\frac{2m}{\hbar^2}\right)^{3/2} (k_B T)^{3/2} (1/2)! L_{3/2}(z) = \frac{V}{\lambda^3} L_{3/2}(z)$$
 (169)

$$U = \frac{V}{(2\pi)^2} \left(\frac{2m}{\hbar^2}\right)^{3/2} (k_B T)^{5/2} (3/2)! L_{5/2}(z) = \frac{3}{2} \frac{V k_B T}{\lambda^3} L_{5/2}(z)$$
(170)

 λ is defined to absorbed all the numerical factor in Eq.169 and $\lambda \propto T^{-1/2}$. Let's now consider Eq.169

$$n\lambda^3 = L_{3/2}(z) \tag{171}$$

since μ need to be negative, the value of z is bound to (0,1), the function $L_{3/2}$ is monoclinically increaing with z and bound between $(0, L_{3/2}(1)) \approx (0, 2.612)$, however, as we decrease temperature and $z \to 1$, λ will increase without bound, which conflict with Eq.171.

This inconsistency come from the fact that when we convert the summation of k into integral of energy ε , we essentially omitted a single state $(k = 0, \varepsilon = 0)$ since $g(\varepsilon = 0) = 0$. Its density of state $g(\varepsilon)$ is ignorable compared to other states but at low temperature, it's occupation maybe very large. Including the occupation of this ground state explicitly in Eq.169:

$$N = N_0 + N_1 = \frac{1}{1/z - 1} + \frac{V}{\lambda^3} L_{3/2}(z)$$
(172)

if we define

$$n\lambda(T_c)^3 = L_{3/2}(z=1) \tag{173}$$

then

$$\frac{N_0}{N} = \frac{N - N_1}{N} = 1 - \left(\frac{T}{T_c}\right)^{3/2} \tag{174}$$

if the temperature is low enough so that $N_0/N \approx 1$, then almost all particles are condensed in the ground state. T_c for an ideal boson gas can be calculated:

$$k_B T_c \approx 0.061 \frac{\hbar^2}{m} n^{2/3}$$
 (175)

For example, liquid Helium⁴ with a density of $1.5e22cm^{-3}$ gives a $T_c \approx 3K$. Expreiment observed a phase transition to a new phase with superfluid properties around this temperature, associated with the condensation.

11 Thermodynamic Properties of magnetic system

We write the internal energy of an magnetic system as:

$$dU = \tilde{\mathbf{d}}Q + \mu_0 H dM \tag{176}$$

where H is the external field. For reversible process, we have:

$$dU = TdS + \mu_0 HdM \tag{177}$$

and we can express the temperature T and field H by:

$$T = \left(\frac{\partial U}{\partial S}\right)_{M} \quad \text{and} \quad H = \frac{1}{\mu_0} \left(\frac{\partial U}{\partial M}\right)_{S}$$
 (178)

and we obtain the Maxwell relationship with internal energy:

$$\left(\frac{\partial T}{\partial M}\right)_S = \mu_0 \left(\frac{\partial H}{\partial S}\right)_M$$
(179)

Heat absorbed can be written as:

$$\tilde{\mathbf{d}}Q = \left(\frac{\partial U}{\partial T}\right)_M dT + \left[\left(\frac{\partial U}{\partial M}\right)_T - \mu_0 H\right] dM \tag{180}$$

$$= C_M dT + ldM \tag{181}$$

where we identify C_M to be the specific heat at constant magnetization: when dM = 0, we have $\tilde{\mathbf{d}}Q = C_M dT$, and

$$C_M = \left(\frac{\partial U}{\partial T}\right)_M = \left(\frac{\tilde{\mathbf{d}}Q}{\mathrm{d}T}\right)_M = T\left(\frac{\partial S}{\partial T}\right)_M \tag{182}$$

where we have used $\tilde{\mathbf{d}}Q = T dS$. For l, we can find, using Maxwell relation:

$$l = \left(\frac{\tilde{\mathbf{d}}Q}{\mathrm{d}M}\right)_T = T\left(\frac{\partial S}{\partial M}\right)_T = -\mu_0 T\left(\frac{\partial H}{\partial T}\right)_M \tag{183}$$

We introduce thermodynamic quantities called magnetic enthalpy:

$$\mathcal{H} = U - \mu_0 H M \tag{184}$$

$$d\mathcal{H} = TdS - \mu_0 MdH \tag{185}$$

which gives the Maxwell relationship:

$$\left(\frac{\partial T}{\partial H}\right)_S = -\mu_0 \left(\frac{\partial M}{\partial S}\right)_H$$
(186)

Similar to above, we have the heat absorbed:

$$\tilde{\mathbf{d}}Q = \left(\frac{\partial \mathcal{H}}{\partial T}\right)_H dT + \left[\left(\frac{\partial \mathcal{H}}{\partial H}\right)_T + \mu_0 H\right] dM \tag{187}$$

$$= C_H dT + h dM \tag{188}$$

where

$$C_H = \left(\frac{\partial \mathcal{H}}{\partial T}\right)_H = T \left(\frac{\partial S}{\partial T}\right)_H \tag{189}$$

and similar to the calculation of l, we have:

$$h = T \left(\frac{\partial S}{\partial H}\right)_T = \mu_0 T \left(\frac{\partial M}{\partial T}\right)_H \tag{190}$$

We can obtain the relationship between C_M and C_H that is similar between C_V and C_p . Substructing Eq.181 and Eq.188, we obtain:

$$(C_H - C_M) dT = -T\mu_0 \left[\left(\frac{\partial H}{\partial T} \right)_M dM + \left(\frac{\partial M}{\partial T} \right)_H dH \right]$$
(191)

since M, T and H are all function of state, we can express the relation:

$$dT = \left(\frac{\partial M}{\partial T}\right)_H dM + \left(\frac{\partial T}{\partial H}\right)_M dH \tag{192}$$

So that we have the result:

$$C_H - C_M = -T\mu_0 \left(\frac{\partial H}{\partial T}\right)_M \left(\frac{\partial M}{\partial T}\right)_H = T\mu_0 \left(\frac{\partial M}{\partial T}\right)_H^2 \left(\frac{\partial H}{\partial M}\right)_T$$
(193)

where we used the relation:

$$\left(\frac{\partial H}{\partial T}\right)_{M} = -\left(\frac{\partial M}{\partial T}\right)_{H} \left(\frac{\partial H}{\partial M}\right)_{T} \tag{194}$$

For system with fixed energy, we maximize the entropy to find the equilibrium ¹². However, for system in contact with the reservior, we need to minimize the free energy, defined by:

$$F = U - TS = -\frac{1}{\beta} \ln Z \tag{195}$$

$$dF = \mu_0 H dM - S dT \tag{196}$$

which gives the Maxwell relationship:

$$\mu_0 \left(\frac{\partial H}{\partial T} \right)_M = -\left(\frac{\partial S}{\partial M} \right)_T \tag{197}$$

When the system is fixed at temperature T and external field H, The Gibbs potential should be minimized:

$$dG = -\mu_0 M dH - S dT \tag{198}$$

with the corresponding Maxwell relationship:

$$\mu_0 \left(\frac{\partial M}{\partial T} \right)_H = -\left(\frac{\partial S}{\partial H} \right)_T \tag{199}$$

Finally, if the number of particle in the system is not fixed: we define extra thermodynamic potentials: The particle number N is given by:

$$\Omega_F = F - \mu N \quad d\Omega_F = \mu_0 H dM - S dT - N d\mu$$

$$\Omega_G = G - \mu N \quad d\Omega_G = \mu_0 M dH - S dT - N d\mu$$

$$N = -\left(\frac{\partial\Omega_F}{\partial\mu}\right)_{T,M} = -\left(\frac{\partial\Omega_G}{\partial\mu}\right)_{T,M} \tag{200}$$

 $^{^{12}}$ That is to say, for microcanonical system, the equilibrium is found at states that correspond to the most number of microstates

Appendix A. Intensive and extensive properties (previous version)

11.1 Heat

We provide a definition of heat as The thermal energy in transit, denoted as Q^{13} . We define the heat capacity C of an object as the amount of heat that is needed to increase its temperature:

$$C = \frac{dQ}{dT} \tag{201}$$

thus C has the unit J/K. The specific heat is defined to be the heat capacity per unit mass, having the unit $J/(kg \cdot K)$

11.2 Entropy

We define entropy S for an isolated macroscopic system of N particles in volume V and energy E to be:

$$S(E, N, V, x) = k_B \ln \Omega(E, N, V, x)$$
(202)

where $\Omega(E, N, V, x)$ is the number of accessible states at a given value of E, N, V, and x is some constraints which influence the number of accessible states. As a non-equilibrium isolated system allow to relax to equilibrium, entropy will increase monotonically and eventually maximize at equilibrium.

Temperature consider an isolated system with two subsystem in weak contact but heat is allowed to follw between the two subsystem. The total number of accessible states (configurations) are given by the product of the number of configurations of the two subsystem. Entropy will be additive. Consider the energy of one of the subsystem E_1 as the constrain for the configurations.

$$\Omega(E, E_1) = \Omega_1(E_1)\Omega_2(E_2) \tag{203}$$

$$S(E, E_1) = S_1(E_1)S_2(E_2) \tag{204}$$

Relaxation process will increase entropy by changing E_1 , towards a macroscopic that correspond to more configurations. At equilibrium (heat no longer exchange), we have:

$$\frac{\partial S}{\partial E_1} = 0 \Rightarrow \frac{\partial S_1(E_1)}{\partial E_1} \bigg|_{N_1, V_1} = \frac{\partial S_2(E_2)}{\partial E_2} \bigg|_{N_2, V_2} = \frac{1}{T}$$
(205)

where the final equality gives the definition of temperature, thus if two subsystem reaches equilibrium in terms of energy flow, their temperature will be equal. This process is irreversible and thus define a arrow of time.

Chemical potential now, we fix only volume V of each subsystem and allow both energy and particles to exchange, then:

$$\frac{\partial S_1(N_1)}{\partial N_1}\Big|_{E_1,V_1} = \frac{\partial S_2(N_2)}{\partial N_2}\Big|_{E_2,V_2} = -\frac{\mu}{T}$$
 (206)

the last equality defines the chemical potential μ .

Pressure finally, we allow the volume of the system to exchange, and we can similar define pressure:

$$\frac{\partial S_1(V_1)}{\partial V_1}\Big|_{E_1, N_1} = \frac{\partial S_2(V_2)}{\partial V_2}\Big|_{E_2, N_2} = \frac{P}{T}$$
 (207)

Thus, we can see that if we set an initial system not in equilibrium, the two subsystem will start to exchange energy, particles and volume until T, P and μ become the same for the two subsystem.

We can separate the macroscopic properties of a system into two different catagory:

- Extensive properties that will increase proportional to the system size, such as N, E, V
- Intensive properties that will be same for any of the subsystem, such as P, μ, T

 $^{^{13}}$ Blundell, p14

Appendix B. Thermodynamic potentials(previous version)

It follows from the definition of T, P and μ that

$$\frac{1}{T} = \left(\frac{\partial S}{\partial E}\right)_{N,V}; \ \frac{\mu}{T} = \left(\frac{\partial S}{\partial N}\right)_{E,V}; \ \frac{P}{T} = \left(\frac{\partial S}{\partial V}\right)_{E,N}$$
 (208)

we can organize:

$$dS = \frac{1}{T}dE - \frac{\mu}{T}dN + \frac{P}{T}dV \tag{209}$$

$$dE = TdS + \mu dN - PdV \tag{210}$$

leading to:

$$T = \left(\frac{\partial E}{\partial S}\right)_{NV}; \ \mu = \left(\frac{\partial E}{\partial N}\right)_{SV}; \ P = -\left(\frac{\partial E}{\partial V}\right)_{SN}$$
 (211)

For the Eq.210, we can integrate the subsystems parts by parts into the whole system (S, N and V are extensive parts), since T, μ and P will be the same for all subsystems, the integration gives:

$$E = TS + \mu N - PV \tag{212}$$

Now, let's consider the canonical ensemble. The energy of the system is no longer fixed, as opposed to microcanonical ensemble in which all the microstates in the ensemble have fixed energy. However, for each energy, it is associated with a probability and thus we only consider the average energy, which we define as **internal energy**:

$$U \equiv \langle E \rangle = \sum_{j} P_{j} E_{j} = \frac{\sum_{j} E_{j} e^{-\beta E_{j}}}{\sum_{j} e^{-\beta E_{j}}} = -\frac{1}{Q_{N}} \frac{\partial Q_{N}}{\partial \beta}$$
 (213)

The internal energy is now the macroscopic observable instead of energy, as in the microcanonical case. We wish to establish an relationship between the internal energy and the free energy $Q_N = \exp(-\beta A)$

Let's write

$$Q_N = \sum_{j} e^{-\beta E_j} = \sum_{E_j} \Omega(E_j) e^{-\beta E_j}$$
 (214)

with $\Omega(E_j)$ gives the number of microstates that have energy E_j . With the definition of entropy $S(E) = k_B \ln \Omega(E)$, we have:

$$Q_N = \sum_{E_j} e^{-\beta(E_j - S(E_j)T)} \simeq e^{-\beta(E' - S(E')T)}$$
(215)

with E' be the value that minimize the function E - S(E)T. If the fluctuation is small, we can consider only the term $e^{-\beta(E'-S(E')T)}$ in the summation is not ignorable, which gives the approximation. The requirement that E' minimize function E - S(E)T is:

$$\frac{\partial (E - S(E)T)}{\partial E} \bigg|_{E'} = 1 - T \frac{\partial S(E)}{\partial E} \bigg|_{E'} = 0 \tag{216}$$

therefore, E' is the energy in which:

$$\left. \frac{\partial S(E)}{\partial E} \right|_{E'} = \frac{1}{T}$$
 (217)

that is to say, E' is the energy where the system's temperature defined by Eq.205 is equal to the temperature of the heat bath, which is the equilibrium condition. This motivate us to equal $E' = \langle E \rangle = U$ leading to the result that

$$A = U - TS(U) \tag{218}$$

U minimize U - TS(U) at equilibrium, therefore, the equilibrium condition of a canonical potential is the minimization of free energy A.

With Eq.218, Eq.210 and $U = \langle E \rangle$, we have

$$dA(N, V, T) = dU - TdS - SdT = -SdT - pdV + \mu dN$$
(219)

and

$$S = -\left(\frac{\partial A}{\partial T}\right)_{N,V}; \ \mu = \left(\frac{\partial A}{\partial N}\right)_{T,V}; \ P = -\left(\frac{\partial A}{\partial V}\right)_{T,N} \tag{220}$$

For Grand canonical ensemble, we can apply similar method as in the canonical ensemble, but this time adding the particle number as a summation variable in the calculation of partition function, we can obtain:

$$\Omega(\mu, T, V) = U - TS - \mu \bar{N} \tag{221}$$

with \bar{N} being the average particle number. The equilibrium condition of a grand canonical ensemble is then the minimization of Ω . We also have the following relationship:

$$d\Omega(\mu, T, V) = -SdT - Nd\mu - PdV \tag{222}$$

and

$$S = -\left(\frac{\partial\Omega}{\partial T}\right)_{\mu,V}; \ N = -\left(\frac{\partial\Omega}{\partial\mu}\right)_{T,V}; \ P = -\left(\frac{\partial\Omega}{\partial V}\right)_{T,\mu}$$
 (223)

11.3 Gibbs expression of entropy

For the canonical ensemble, we have:

$$P_i = \frac{1}{Q}e^{-\beta E_i}$$

$$A = -k_B T \ln Q = U - TS$$

This lead to the relationship:

$$S = -k_B \sum_{i} P_i \ln P_i \tag{224}$$

This is the Gibbs' form of entropy in terms of probability.