Lectures on Statistical Mechanics and Thermodynamics

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Part I Foundations of Statistical Mechanics

The object of Statistical Mechanics is to study systems with a large number of degrees of freedom by focusing solely on the macroscopic properties of the system.

Why, you may ask, are we only focusing on these macroscopic properties? Surely, given enough information we could model any system by following its individual members! This is not necessarily the case because the sheer volume of information necessary to do such a calculation is in fact an impediment to performing such a calculation. Let's illustrate this using a simple example of a system consisting of 20 particles in the ground state. Suppose, through divine intervention or otherwise, we have obtained the complete wave equation for each particle in the system. Before we begin to calculate any properties of this system we must first store this information.

As shown in figure 2, every particle wavefunction will be reasonably approximated by 10 points in each dimension. Taking the 3-dimensional case, we find that the amount of memory we would need would be $(10)^{3N}*8 \simeq 10^{61}$ bytes. We include the factor of 8 since each real number will be represented by 8 bytes. Assuming we store all of this data on terabyte drives that have a storage capacity of $\frac{10^{12}bytes}{0.4kgs}$, we find that the total mass of all of the drives required to store the information describing this 20 particle system to be on the order of 10^{49} kilograms, which is much much more than the mass of the Milky Way galaxy! Clearly, it is impossible to store this much information, much less manipulate it. Therefore, we employ statistics to deal with the system at a macroscopic level.

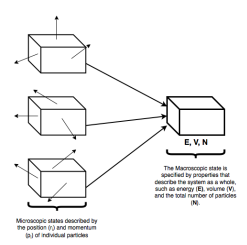


Figure 1: Multiple microscopic states make up a single macroscopic state of a system.

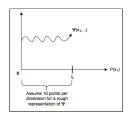


Figure 2: Wavefunction of our system in dimension x_1

1 Hamiltonian Dynamics

Consider a hypothetical system consisting of some number of particles which can be described by their individual positions, q_i and their momentums, p_i . We can indicate the paths these particles travel on a **Phase Space** diagram.

This diagram displays the evolution of the systems constituent particles as their positions and momentums change. The evolution of this system is governed by the Hamilton Equations: $\frac{dq_1}{dt} = \frac{\delta \mathcal{H}}{\delta p_i}$ and $\frac{dp_i}{dt} = -\frac{\delta \mathcal{H}}{\delta q_i}$ The evolution of p and q are clearly deterministic, therefore we can express these values as time-dependant functions. $q_i(t), p_i(t) \to q_i(t=0) = q_i^0$ and $p_i(t=0) = p_i^0$

There are a couple of tricks we can employ to idealize the measurement process and simplify manipulations of these functions.

2 Idealization of the Measurement Process

Let's start by defining some value \overline{f} , which is the result of measuring function $f(p_i, q_i)$.

$$\overline{f} = \lim_{T \to \infty} \frac{1}{T} \int_{\delta}^{\delta + T} dt f(q(t), p(t)) = \widetilde{f}(q_i^0, p_i^0)$$

T= duration of the measurement, $\delta=$ time the measurement starts, $\widetilde{f}=$ some function of the initial conditions.

This expression is an idealization of an arbitrary measurement. In reality, the time T is much larger than the microscopic scales in which the functions of q(t) and p(t) vary. For example, suppose the function f is the pressure on the wall of a gas containing vessel. The quantity f(q(t), p(t)) flucuates like the diagram below.

However, except in specialized cases, the measuring apparatus is not accurate enough the capture every tiny fluctuation of the force on the wall of the vessel. Instead, it measures an average of f(q(t), p(t)) over a finite period of time:

Taking the $\lim_{T\to\infty}$ is a mathematical idealization of this averaging out of function fluctuations. Returning to our analysis of the hypothetical measurement \overline{f} , we see that the function $\widetilde{f}(q_i^0,p_i^0)$ is not really a function of the initial condition. Rather, it is a function of the trajectory of the system evolution. This may not be immediately apparent, but can be illustrated by choosing another initial point on the trajectory of our function $\widetilde{f}=(q_i^0,p_i^0)$. As $T\to\infty$, we see that our new initial points $q_i^{0'}$ and $p_i^{0'}$ yield the same value of \widetilde{f} as q_i^0 and p_i^0 . i.e., $\widetilde{f}(q_i^0,p_i^0)=\widetilde{f}(q_i^{0'},p_i^{0'})$ This is because the difference between the two trajectories is neglible compared to the infinite extended trajectories as $T\to\infty$.

Now, it could be possible that two different trajectories give different values for \overline{f} .

However, it turns out that, for systems we are interested in, that it does not happen. These systems have the property that any individual trajectory will pass through almost **every** point in phase space. This is called the **Ergodic Property**.

A couple of important things to keep in mind. The assumption that the trajectory passes through every point in the phase space is contingent on the the assumption that these points have the same value of energy (and any other conserved quantities) as the initial point. After all, these values are conserved by the Hamiltonian Flow. The phrase "almost every point" also has a specific meaning, in that it is "almost every point" in the measure theory sense.

It is really difficult to prove that a realistic system has the ergodic property. Only a few systems have actually been proven rigorously to be ergodic (for example, the Sinai Stadium). Moving forward, we will operate on the assumption that the systems we are interested are, in fact, ergodic.

To expand on the concept of ergodicity, it is very easy to find systems that are not ergodic. Any conserved value restricts the trajectories to lie on a submanifold of the phase space. If the only conserved quantities are the ones resulting from the standard symmetries (i.e. energy, momeentum, etc.) we can question the validity of ergodicity within the restricted subspace with constant values of these quantities. There are systems with so many other conserved quantities that the subspace is one dimensional (integrable systems), thereby invalidating the ergodicity of the system. The Integrable/Ergodic classification is also non-binary, there are intermediate types of systems that exist between the two extremes, such as mixing systems.

3 Introduction to Thermodynamics

At its core level, many thermodynamic problems can be phrased as a comparison between two states of a system. The pre-evolutionary equilibrium state and the post-evolutionary equilibrium state.

Since every microstate of the system is equally probably, the final macroscopic state will be the one which corresponds to the largest number of microstates:

$$\Gamma(E_1',V_1',N_1',E_2',V_2',N_2') = \Gamma_1(E_1',V_1',N_1')\Gamma_2(E_2',V_2',N_2')$$

Where $\Gamma(E'_1, V'_1, N'_1, E'_2, V'_2, N'_2)$ is the number of states of the composite system, $\Gamma_1(E'_1, V'_1, N'_1)$ is the number of microstates of subsystem 1, and $\Gamma_2(E'_2, V'_2, N'_2)$ is the number of microstates of subsystem 2.

Let's look at the case where the wall between the two subsystems allows the exchange of energy but not of volume or particle number. In this case we find:

$$E_1 + E_2 = E_1' + E_2'$$

$$V_1 = V_1', V_2 = V_2'$$

$$N_1 = N_1', N_2 = N_2'$$

$$\Gamma(E_1', E_2') = \Gamma_1(E_1')\Gamma_2(E_2') \quad (1)$$

This implies that

$$\Gamma_{Max,E_{total}fixed} \Leftrightarrow \frac{1}{\Gamma_1} \frac{d\Gamma_1(E_1')}{dE_1'} - \frac{1}{\Gamma_2(E_2')} \frac{d\Gamma_2(E_2')}{dE_2'} = 0$$

So, knowing the total number of initial microstates of the two subsystems allows us to predict the final state of the overall system. This is summed up in a very clean and useful fashion in the definition of **entropy**:

$$S = k_B \ln(\Gamma)$$

Notice that entropy is extensive as long as the subsystems are large enough that the boundary effects are negligible.

How can Entropy grow?

Within the constraints of the problem, there is an apparent paradox in staying that the entropy is maximized. That's because, using Hamiltonian equations, we can show S is a constant.

Define $\rho(q, p, t = 0)$ to be an initial distribution of identical systems (this is called an **ensemble**.) As the Hamiltonian Flow evolves in time every element of the ensemble is carried with it and the distribution evolves to $\rho(q, p, t) = \rho(q(t), p(t))$.

Because the number of systems in the ensemble is fixed ρ obeys the continuity equation:

$$\frac{d\rho}{dt} + \nabla * (\rho \nabla) = 0$$

or

$$0 = \frac{\delta \rho}{\delta t} + \sum_{i=1}^{3N} \left[\frac{\delta(\rho \dot{q}_i)}{\delta q_i} + \frac{\delta \rho \dot{p}_i}{\delta p_i} \right]$$

$$= \frac{\delta \rho}{\delta t} +$$
(2)

This result is known as the **Liouville Theorem**. It implies that the density of elements on the ensemble does not change as it evolves along the Hamiltonian Flow (of course, it does change at any fixed position in phase space). As the number of systems in the ensemble is fixed, the Liouville Theorem implies that the volume of the phase space occupied by the ensemble is fixed.

This means that the entropy, being the log of the volume, cannot change either. So how can the entropy grow if the microscopic equations of motion show that it does not? A resolution to this paradox is to notice that, for systems in which statistical mechanics holds, the Hamiltonian Flow takes nice looking, civilized ensembles and turns them into convoluted shapes:

The average of smooth observables (i.e. the value of f(q,p)) does not distinguish between the average performed with the true $\rho(t)$ or the "coarse grained" $\tilde{\rho}(t)$, as shown below.

$$\overline{f} = \int dq dp f(q, p) \rho(q, p, t) \simeq \int dq dp f(q, p) \widetilde{\rho}(q, p, t)$$

For all practical purposes, the entropy grows so long as we only look at observables that are very smooth. Macroscopic observables don't care about the precise poition of the particles and tend to be smooth in phase space. Systems whose Hamiltonian Flow have this property are called "Mixing." It is easy to show that ergodic systems are mixing, but keep in mind that mixing systems are not necessarily ergodic.

Part II

Thermodynamics

From the general property of entropy we can derive a massive amount of information about a system. In fact, if the entropy as a function of extensive variables is known, everything else about the thermodynamic behavior of the system can be found! The definition of entropy is reiterated below:

$$S = k_B \ln(\Gamma)$$

We include the Boltzmann Constant, k_B , for historical reasons. However, if we judiciously choose our units (by measuring temperature in units of energy, as we should) we can set $k_B = 1$. A good thing to remember in this regard is that $10,000K \simeq 1eV$.

Since entropy is so important, let's review a few properties satisfied by the entropy:

- S as a function of the <u>extensive</u> variables contains all the information about a system that you need.
- S is itself extensive: $S(\lambda * E, \lambda * V, \lambda * N) = \lambda * S(E, V, N)$
- $\frac{\delta S}{\delta E} \geq 0$ We will eventually see that this implies $T \geq 0$
- S(E,V,N) is a concave function. We will eventually see that this implies stability:

$$S(E + \Delta * E, V + \Delta * V, N + \Delta * N) + S(E - \Delta * E, V - \Delta * V, N - \Delta * N) \leq 2S(E, V, N)$$

- S=0 at $\frac{1}{\frac{\delta S}{\delta E}}=0(T=0)$ This property depends on how we normalize the number of microstates, Γ . Quantum mechanics determines the right normalization that implies S(T=0)-0
- S is maximized in equilibrium within the constraints set by the physical setup. This is a direct consequence of the fact that all microstates are equally probable.

Inverting the entropy function S(E, V, N) allows us to obtain a function for the energy of the system in terms of entropy: E(S, V, N). If S, V, N are changed slowly (i.e. "adiabatically") so that the system stays in equilibrium at all intermediate states we find:

$$dE = \frac{\delta E}{\delta S}|_{V,N} dE + \frac{\delta S}{\delta V}|_{E,N} dV + \frac{\delta S}{\delta N}|_{E,V} dN$$

where

$$\frac{\delta S}{\delta E}|_{V,N} = \frac{1}{T}, \frac{\delta S}{\delta V}|_{E,N} = \frac{P}{T}, \frac{\delta S}{\delta N}|_{E,V} = \frac{\mu}{T}$$

To derive the first relation, we used $\frac{\delta E}{\delta S}|_{V,N}=\frac{1}{den}$ It's important to note that S is defined for equilibrium macroscopic states only. So dS or dE refer to changes slow enough such that the system continuously remains in equilibrium during its transition from its initial state to its final state. i.e., at any given time during the transition the system is in an equilibrium state. For instance, if the volume is changed suddenly and then the system is left to relax to equilibrium the change in entropy is not given by $\frac{p}{T}dV$ (it will, in fact, be larger).

From these relations we can derive the Equations of State:

	1
$T = \frac{\delta E}{\delta S} _{V,N} = T(S,V,N)$	$rac{1}{T} = rac{\delta S}{\delta E} _{V,N} = rac{1}{T(E,V,N)}$
$-P = \frac{\delta E}{\delta V} _{S,N} = P(S, V, N)$	$-\frac{P}{T} = \frac{\delta S}{\delta V} _{E,N} = -\frac{P(E,V,N)}{T(E,V,N)}$
$\mu = \frac{\delta E}{\delta N} _{S,V} = \mu(S, V, N)$	$\frac{\mu}{T} = \frac{\delta S}{\delta N} _{E,V} = \frac{\mu(E,V,N)}{T(E,V,N)}$

These equations of state are particularly important because they are frequently accessible to experiments. Notice that the same quantities as functions of other variables are not equations of state and may not contain the same information.

As mentioned earlier, all thermodynamical problems can be reduced to a transition between two states of a system:

Depending on the properties of the wall between the two subsystems, we

- Wall allows Energy exchange $\to T_1' = T_2'$
- Wall allows Volume exchange $\rightarrow V_1' = V_2'$
- Wall allows Particle exchange $\rightarrow \mu'_1 = \mu'_2$

Non-Relativistic, Non-Interacting, Monoatomic Gas

Let us compute the entropy, S, for one example:

$$\mathcal{H}(\searrow_i, \bowtie_i) = \sum_{i=1}^N \frac{\bowtie_i^2}{2m}, (0 < \searrow_i < L)$$

$$\Gamma(E,V,N) = \int \prod_{i=1}^{N} d \mathbf{1}_{i} \delta(E - sumseri = 1N \frac{l_{i}^{2}}{2m})$$

$$= [\int_{0}^{L} dx]^{3N} \int_{-\infty}^{\infty} dp_{ix} dp_{iy} dp_{iz} \dots dp_{nx} dp_{ny} dp_{nz} * 2m\delta(2mE - \sum_{i=1}^{3N} p_{i}^{2})_{S} = k_{B} \ln \Gamma = k_{B}[N \ln V + \frac{3N-1}{2} \ln E - V^{N} 2mS^{3N}(R = \sqrt{2mE})]$$

$$= 2mV^{N} \frac{2\pi^{\frac{3N}{2}}}{\Gamma(\frac{3N}{2})} (2mE)^{\frac{3N-1}{2}}$$

The above expression violates extensivity (2) and the Neust Postulate (5). The violation of the Neust Postulate is not surprising: the behavior of a gas at T=0 is dictated by quantum mechanics, not classical mechanics (the atoms wavelength is much larger than the separation between them). What is surprising is that quantum mechanics connects the calculation above even at large T. What is missing from the classical calculation above is that the gas is composed of identical particles (bosons or fermions, it doesn't matter at high T). So the correct way of counting states is:

$$S = k_B \ln \Gamma = k_B \int \prod_{i=1}^{N} \frac{dq_i dp_i}{N!h^N} \delta(E - \sum_{i=1}^{N} \frac{\iota_i^2}{2m})$$

The denominator factor of N! avoids double counting of microstates with the same particle position and the factor of h^N becomes important when considering the classical limit of quantum gases.

As the number of particles in the system becomes very large we find that the expression becomes

$$S = k_B * N \ln \frac{VE^{\frac{3}{2}}}{N^{\frac{5}{2}}}$$

From this entropy we can find the equations of state:

from this entropy we can find the equations of state.		
$\frac{1}{T} = \frac{\delta S}{\delta E} _{V,N} = \frac{3}{2} \frac{k_B * N}{E}$	$E = \frac{3}{2}k_B * N * T$	
$-\frac{P}{T} = \frac{\delta S}{\delta V} _{E,N} = -\frac{k_B * N}{V}$	$PV = k_B * NT$	
$\frac{\mu}{T} = \frac{\delta S}{\delta N} _{E,V} =$		
$k_B \ln \frac{VE^{\frac{3}{2}}}{N^{\frac{5}{2}}} - \frac{5}{2}k_B$		

Note that we found the Boyle-Mariott law without even discussing atoms colliding against walls or other kinetic theory considerations. There's a lot behind the assumption of ergodicity.

This leads us to....

5 Euler Relation

$$S(\lambda * E, \lambda * V, \lambda * N) = \lambda * S(E, V, N)$$

Taking the derivative of this expression in terms of λ yields

$$E()\frac{\delta S}{\delta E} = \frac{1}{T}) + V(\frac{\delta S}{\delta V} = -\frac{P}{T}) + N(\frac{\delta S}{\delta N} = \frac{\mu}{T}) = S$$

Which can be re-arranged to yield the following equations:

$$E = TS - PV + \mu * N$$

$$S = \frac{E}{T} - \frac{PV}{T} + \frac{\mu * N}{T}$$

5.1 Recovering the Entropy or Energy from an Equation of State

If three equations of state are available:

$$T(S, V, N) = \frac{\delta E}{\delta S}|_{V,N}$$
$$-P(S, V, N) = \frac{\delta E}{\delta V}|_{S,N}$$
$$\mu(S, V, N) = \frac{\delta E}{\delta N}|_{S,V}$$

$$\rightarrow E(S, V, N) = T(S, V, N) = T(S, V, N)S - P(S, V, N)V + \mu(S, V, N)N$$

If only two equations of state are available: $E(S, V, N) = NE(\frac{S}{N}, \frac{V}{N}, 1)$ The equality above is due to extensivity.

$$T = \frac{\delta E}{\delta S}|_{V,N} = T(S, V, N) = T(\frac{S}{N}, \frac{V}{N}, 1)$$

$$-P = \frac{\delta E}{\delta V}|_{S,N} = P(S,V,N) = P(\frac{S}{N},\frac{V}{N},1)$$

Note that there is no N!. T and P are intensive.

1

$$E(\frac{S}{V}, \frac{V}{N}, 1) = \int_{0}^{\frac{S}{N}} d(\frac{S}{N}) T(\frac{S}{N}, \frac{V}{N}, 1) + E(0, \frac{V}{N}, 1) = \int_{0}^{\frac{S}{N}} d(\frac{S}{N}) T(\frac{S}{N}, \frac{V}{N}, 1) - \int_{0}^{\frac{V}{N}} d(\frac{V}{N}) P(0, \frac{V}{N}, 1) + E(0, 0, 1) = \int_{0}^{\frac{S}{N}} d(\frac{S}{N}) T(\frac{S}{N}, \frac{V}{N}, 1) + E(0, 0, 1) = \int_{0}^{\frac{S}{N}} d(\frac{S}{N}) T(\frac{S}{N}, \frac{V}{N}, 1) + E(0, \frac{V}{N}, \frac{V}{N}, 1) = \int_{0}^{\frac{S}{N}} d(\frac{S}{N}) T(\frac{S}{N}, \frac{V}{N}, \frac{V}{N}$$

In the above expressions, $T(\frac{S}{N}, \frac{V}{N}, 1) = \frac{\delta E}{\delta S}|_{V,N}$ and $E(0,0,1) = E(\frac{S}{N}, \frac{V}{N}, 1)$. The latter equality can be found up to one constant independant of S,V, or N.

Now, if only one equation of state is available, you cannot recover the entropy or energy. There is simply not enough information in a single equation of state to extract these system properties.

5.2 EXAMPLE: Non-relativistic, Monoatomic, Non-interacting Gas

$$S = k_B * N \ln \frac{VE^{\frac{3}{2}}}{N^{\frac{5}{2}}}, E = \left(\frac{N^{\frac{5}{2}}}{V}e^{\frac{S}{k_B*N}}\right)^{\frac{2}{3}} = \frac{N^{\frac{5}{2}}}{V^{\frac{2}{3}}} * e^{\frac{2S}{3k_B*N}}$$

$$T = \frac{\delta E}{\delta S}|_{V,N} = \frac{2}{3k_B*N} \frac{N^{\frac{5}{3}}}{V^{\frac{2}{3}}} e^{\frac{2S}{3k_B*N}}$$

$$-P = \frac{\delta E}{\delta V}|_{S,N} = -\frac{2}{3} \frac{N^{\frac{5}{3}}}{V^{\frac{5}{3}}} e^{\frac{2S}{3k_B*N}}$$

$$E(\frac{S}{N}, \frac{V}{N}, 1) = \int_0^{\frac{S}{N}} d(\frac{S}{N}) \frac{2}{3k_B} (\frac{N}{V})^{\frac{2}{3}} e^{\frac{2S}{3k_B*N}} - \int_0^{\frac{V}{N}} d(\frac{V}{N}) \frac{2}{3} (\frac{N}{V})^{\frac{5}{3}} + E(0, 0, 1)$$

$$E(\frac{S}{N}, \frac{V}{N}, 1) = \frac{2}{3k_B} (\frac{N}{V})^{\frac{2}{3}} \frac{3k_B*N}{2} [e^{\frac{2S}{3k_B*N}} - 1] - \frac{2}{3} (\frac{-3}{2}) (\frac{N}{V})^{\frac{2}{3}}$$

$$E(\frac{S}{N}, \frac{V}{N}, 1) = \frac{N^{\frac{5}{3}}}{V^{\frac{1}{3}}} e^{\frac{2S}{3k_B*N}}$$

6 Minimum Energy Principle

Before formally stating the minimum energy principle, let's begin with an analogy. We know that a circle is the figure with the largest area for a given perimeter. Of course, it is also the the figure with the smallest perimeter for a given area. Even if a given circle was built by someone with a fixed length string trying to enclose the largest area it will still be true that the final circle minimizes the perimeter for the given area.

The same thing is true for the maximum entropy principle. After some constraints are changed the final state will be the one with the largest entropy for a given **total** energy (the constraint that the total energy is unchanged is always there). But the final equilibrium state can also be characterized as the state with the smallest energy for a fixed entropy:

Convince yourself that S as a function of total E and subsystem energy E_1^1 has the general shape depicted above. The minimum energy principle follows from the picture above.

Now let's see how the minimum energy principle works in our archetypal example :

$$E = E_1(S_1^1, V_1, N_1) + E_2(S_2', V_2, N_2)$$

Where $S_{2}' = S - S_{1}'$.

Therefore, we find that the minimum energy, E, for fixed entropy, S, is:

$$\frac{\delta E}{\delta S_1'} = 0 \to \frac{\delta E_1}{\delta S_1'} - \frac{\delta E_2}{\delta S_2'} = 0 \to T_1 = T_2$$

Which confirms what we expected: In its final state our example subsystem pair reaches an equilibrium temperature.

7 Efficiency of Engines

In our thermodynamic terminology, an 'engine' is a machine that extracts work from the the energy difference between two systems in contact. To clarify, the engine takes energy from the 'hot' source, uses some of that energy to perform work, and ejects the remaining energy into the 'cold' sink. In many cases, the 'cold' sink is the atmosphere: one example where this is true is a car engine, which throws off heat as it performs the work of moving the car.

In designing our thermodynamic engine, we would like it to perform this energy transfer over and over again in a closed cycle. I.e. we want the system to end up in the same macroscopic state as it started. For our purposes right now we will assume that all changes in the system caused by the energy transfers occur slowly (adiabatically) so, that at every point in the cycle the machine is in equilibrium and its macroscopic state is specified by E,V,N or S,V,N. It turns out, that for thermodynamic calculations it is better to use the variables T,S,N instead. In our engine we want to keep N fixed by having impermeable walls separating the machine from the rest of the Universe. This machine cycle can be represented by a closed loop in the TxS space:

Let us start by considering one specific cycle, the Carnot Cycle:

For a fixed N: dE = TdS - PdV so that

$$\Delta Q_1 = \int_{S_1}^{S_2} T dS = T_h(S_2 - S_1)$$
$$\Delta Q_2 = \int_{S_2}^{S_1} T dS = -T_c(S_2 - S_1)$$

Since after a full cycle the energy of the system goes back to what it was in the beginning we find:

$$\Delta Q_1 + \Delta Q_2 = \Delta W =$$

the area inside the cycles in the TxS plane.

In the expression above, $\Delta Q_1 + \Delta Q_2$ is the net heat absorbed by the system and ΔW is the work done by the system.

For practical reasons, we would like to get as much work as possible from a given amount of heat absorbed. The efficiency of the engine is defined as"

$$\epsilon = \frac{\Delta W}{\Delta Q_1} = \frac{(T_h - T_c)(S_2 - S_1)}{T_h(S_2 - S_1)} = 1 - \frac{T_c}{T_h} < 1$$

Now, one very important thing to note about this expression is that the total efficiency is *less than* not *less than or equal to* one because some of the energy is wasted and dispersed into the cold sink. This begs the question: is there a way that we could change the cycle to obtain a larger efficiency?

Going back to our TxS plane, we would want to maximize the area inside of the cycle (ΔW) and minimize the area below the top curve (ΔQ_1) . It's clear, for instance, that the cycle on the left can have its efficiency improved by changing it as shown below. We end up then with the Carnot cycle, which is the best that can be done. Thus, we have the universal result:

$$\epsilon \le 1 - \frac{T_c}{T_h}$$

Notice that we only used the fact that an ideal gas was used inside the engine to describe the different phases of the cycle in words. The main result, $E \leq 1 - \frac{T_c}{T}$ does not depend on that and is valid universally.

 $E \leq 1 - \frac{T_c}{T_h}$ does not depend on that and is valid universally. Finally, let us revisit the assumption of adiabaticity, namely, that all phases of the cycle were infinitely slow. In each phase of the cycle the machine foes from the one initial state A to a final state B exchanging a certain amount of heat ΔQ (with a source at temperature T) and performing some work ΔW (positive or negative). I'll assume the work is done on a reversible work source, namely, a system which can exchange work but not heat (a paper weight on top of a piston is an example). Similarly, heat is exchanged with a reversible heat source.

For given initial and final states A and B (and consequently, given changes in energy $E_B - E_A$ and entropy) the maximum amount of work will be delivered if ΔQ is a minimum as $E_B - E_A = -(\Delta W + \Delta Q)$. The process with the smallest ΔQ will correspond to the smallest possible increase of the entropy of the universe (system+heat source, the work source has no entropy). How much is the minimum amount of heat to be transferred to the heat source? As the total entropy of the universe must increase (otherwise the system would stay at A) we have

$$S_B - S_A + T_r \Delta Q \ge 0$$

Where T_r is the temperature of the heat source. That means that the minimum amount of heat "wasted" on the heat source happens when the total entropy of the universe is unchanged as is given by

$$\Delta Q = \frac{S_A - S_B}{T_r}$$

If more heat than $\frac{S_A - S_B}{T_r}$ is delivered to the heat source the universe's entropy will increase and less work will be delivered to the work source. A similar analysis applies when the system is absorbing heat or work. The main point is that **reversible processes**, those where the entropy of the universe does not change, lead to the maximum efficiency. Thus, our bound on the efficiency of an engine obtained assuming adiabaticity, is not violated by non-adiabatic processes. Notice that the adiabaticity condition means that the engine works really slowly. Practical Machines, which need to do their job fast, are not adiabatic and much less efficient than the theoretical limit would expect (roughly a factor of 3 less efficient).

8 Thermodynamic Stability

Previously we determined the equilibrium condition between two subsystems:

$$T_1 = T_2$$

for a diathermal wall

$$P_1 = P_2$$

for a moveable wall

$$\mu_1 = \mu_2$$

for a porous wall

using the maximum entropy principle. But we only imposed the $S = S_1 + S_2$ was a critical point: it could have been a maximum, a minimum, or a saddle point. We'll now see what the consequences are of demanding that the final equilibrium state is indeed a maximum.

Let's consider first the case of two subsystems separated by a diathermal wall. The condition that $S=S_1+S+2$ is a maximum implies:

$$\frac{dS}{dE_1'} = 0 \rightarrow \frac{\delta S_1}{\delta E_1} = \frac{\delta S_2}{\delta E_2}$$
$$\frac{d^2S}{dE_1^2} < 0 \rightarrow \frac{\delta^2 S_1}{\delta E_1^2} + \frac{\delta^2 S_2}{\delta E_2^2} < 0$$

The quantity $\frac{\delta^2 S}{\delta E^2}$ has a physical interpretation:

$$\begin{split} \frac{\delta^2 S}{\delta E^2}|_{V,N} &= \frac{\delta \frac{1}{T}}{\delta E}|_{V,N} = -\frac{1}{T^2} \frac{\delta T}{\delta E}|_{V,N} \\ &= -\frac{1}{T^2} \frac{1}{\frac{\delta E}{\delta T}|_{V,N}} = -\frac{1}{T^2 N} \frac{1}{c_v} \end{split}$$

Where c_v is the system's specific heat at a constant volume. From the above we see that the stability condition is then:

$$-\frac{1}{N_1T^2}\frac{1}{c_v^1}-\frac{1}{N_2T^2}\frac{1}{c_v^2}<0$$

or

$$\frac{1}{N_1 c_v^1} + \frac{1}{N_2 c_v^2} > 0$$

This is true because $T_1 = T_2$ in equilibrium. By taking one subsystem to be much larger we conclude that each one of c_v^1 and c_v^2 should be positive. This result is physically obvious. Suppose $c_v^1 < 0$. In this case, the system would get colder by absorbing heat. This would, of course, cause the system to absorb even more heat in a runaway cycle. Clearly, this is an unstable situation.

Next, let's consider the case where both E and V are exchanged between the subsystems. We have:

$$\begin{split} 0 &= \frac{\delta S}{\delta E_1^1} = \frac{\delta S_1}{\delta E_1} - \frac{\delta S_2}{\delta E_2} \\ 0 &= \frac{\delta S}{\delta V_1^1} = \frac{\delta S_1}{\delta V_1} - \frac{\delta S_2}{\delta V_2} \\ 0 &> d^2 S = (dE_1^1 dV_1^1) \left[\begin{smallmatrix} \frac{\delta^2 S_1}{\delta (E_1^1)^2} & \frac{\delta^2 S_1}{\delta (E_1^1)^2} & \frac{\delta^2 S_1}{\delta (E_1^1)^2} \\ \frac{\delta^2 S_1}{\delta (E_1^1)^2} & \frac{\delta^2 S_1}{\delta (V_1^1)^2} \end{smallmatrix} \right] \left[\frac{dE_1^1}{dV_1^1} \right] + 1 \leftrightarrow 2 \end{split}$$

We need that last term, 2, to be negative for any dE_1^1, dV_1^1 . For the second variation d^2S to be negative for any E_1^1, V_1^1 it's necessary that the eigenvalues of

$$\begin{bmatrix} \frac{\delta^2 S_1}{\delta (E_1^2)^2} & \frac{\delta^2 S_1}{\delta E_1^1 \delta V_1^1} \\ \frac{\delta^2 S_1}{\delta E_1^1 \delta V_1^1} & \frac{\delta^2 S_1}{\delta (V_1^1)^2} \end{bmatrix}$$

to be negative. So let's work out a general condition for this to be true.

$$\left| \begin{smallmatrix} a-\lambda & b \\ b & c-\lambda \end{smallmatrix} \right| = 0$$

where

$$\lambda^{2} - (a+c)\lambda + ac - b^{2} = 0$$

$$\lambda = \frac{a+c}{2} \pm \sqrt{\frac{a+c^{2}}{2} - (ac - b^{2})} < 0$$

$$a+c < 0\∾ - b^{2} > 0$$

$$a < 0\&c - \frac{b^{2}}{a} < 0$$

In this case, $a=\frac{\delta^2 S}{\delta E^2}|_{V,N}=\frac{\delta \frac{1}{T}}{\delta E}|_{V,N}=-\frac{1}{T^2}\frac{\delta T}{\delta E}|_{V,N}=-\frac{1}{T^2N}\frac{1}{c_v}<0$. Note that the last part of the expression is the same condition as before, namely that $c_v>0$.

$$0 > c - \frac{b^2}{a} = \frac{\delta^2 S}{\delta V^2} |_{E,N} - \left(\frac{\delta^2 S}{\delta E \delta V}\right)^2 \frac{1}{\frac{\delta^2 S}{\delta E^2} |_{V,N}}$$

Although this last expression is rather messy, we can rewrite in a more clear way with a transparent physical meaning.

$$\begin{split} \frac{\delta \frac{P}{T}}{\delta V}|_{T,} &= \nabla (\frac{P}{T}) * \bowtie \\ \frac{\delta \frac{P}{T}}{\delta V}|_{T,} &= \frac{\delta \frac{P}{T}}{\delta V}|_{E,} - \frac{\delta T}{\delta V}|_{E,} \frac{1}{\frac{\delta T}{\delta E}|_{V,}} \frac{\delta \frac{P}{T}}{\delta E}|_{V,} \end{split}$$

$$\frac{\delta \frac{P}{T}}{\delta V}|_{T,} = \frac{\delta^2 S}{\delta V^2} + T^2 \frac{\delta^2 S}{\delta E \delta V}|_E \frac{1}{-T^2 \frac{\delta^2 S}{\delta E^2}|_V} \frac{\delta^2 S}{\delta V \delta E}$$

So we find $\frac{\delta \frac{P}{T}}{\delta V}|_{T_{,}} = \frac{1}{T} \frac{\delta P}{\delta V}|_{T_{,}} < 0$. The $\frac{1}{T} \frac{\delta P}{\delta V}|_{T_{,}} < 0$ term can be rewritten to yield:

$$-\frac{1}{VT}[-\frac{1}{V}\frac{\delta V}{\delta P}|_{T,}]^{-1} = -\frac{1}{VT}\frac{1}{R_T}$$

We have defined a new quantity above, R_T . The quantity is the **Isothermal Compressibility** of the gas. The stability condition we defined earlier implies that the isothermal compressibility is positive, however, this result is not immediately obvious. A negative compressibility would mean that the system expands as the pressure increases. This behaviour would increase the pressure of the other system leading to a runaway expansion of our observed system.

Part III

Canonical Ensemble

For many situations, it is much easier to treat a system as if it is in contact with an infinite heat reservoir. This important property of this reservoir is that it is unaffected by any energy it absorbs due to its size. In order to derive an ensemble to describe this system, let's start with the microcanonical ensemble:

$$\rho(q,p) \Gamma(E - \mathcal{H}(q,p))$$

Where q, p are the coordinates of the subsystem only, Γ is the function which represents the number of microstates of the reservoir with energy $E - \mathcal{H}(q, p)$. The terms within the gamma function represent the total energy of the reservoir (E) and the subsystem energy (\mathcal{H}) . Continuing our derivation:

$$\rho(q,p) e^{\frac{1}{k_B}*S_R(E-\mathcal{H}(q,p))}$$

Where S_R is the reservoir's entropy.

$$\rho(q,p) \cong e^{\frac{1}{k_B}S_R(E) - \frac{1}{k_B}\mathcal{H}(q,p)\frac{\delta S_R}{\delta E}}$$

In the expression above, we see that $\frac{1}{k_B}S_R(E)$ is a constant independent of the coordinants (q,p), and the $\frac{\delta S_R}{\delta E} \to \frac{1}{T_R} = \frac{1}{T}$ term shows the equilibrium equivalence of the reservoir temperature (T_R) and the subsystem temperature (T). Thus:

$$\rho(q,p) e^{-\frac{\mathcal{H}(q,p)}{k_B T}}$$

Therefore, we see that $\rho(q,p)$ is independent of the reservoir's properties. This expression illustrates what is known as the **Canonical Ensemble**.

$$\rho(q, p) = \frac{e^{-\beta \mathcal{H}(q, p)}}{Z}$$

$$Z = \int dq dp e^{-\beta \mathcal{H}(q, p)}$$

$$\beta = \frac{1}{k_B T}$$

9 Equivalence with the Microcanonical Ensemble

To see the equivalence of these two ensembles, let's start with the partition function:

$$Z = \int dq dp e^{-\beta \mathcal{H}(q,p)} = \int dE \Gamma(E) e^{-\beta E}$$

Where $dE\Gamma(E)$ is the number of microstates with energy between E and E+dE. Therefore

$$Z = \int dE e^{\frac{S(E)}{k_B} - \beta E}$$

As we see in the above expression, S(E) is the microcanonical ensemble and the entire exponent of e is an extensive quantity which is N very concentrated around the maximum of the integrand.

$$Z \cong \Delta e^{-\beta(E-TS(E))}|_{E=E*}$$

The energy constraint insists that the energy minimizes the value E-TS(E). This value has a particular significance and therefore has a specific name...

10 Helmholtz Free Energy and the Partition Function

As stated before, the quantity $E - TS(E, V, N)|_{E=E*}$ has a specific name: **Helmholtz Free Energy**, and is denoted by the function F(T, V, N). Before we further discuss the importance of this quantity, let's first explore the significance of the normalization factor Z in the Canonical Ensemble. Z also has a specific name, the **Partition Function**. It is extremely useful in its own right:

$$\frac{\delta \ln Z}{\delta \beta} = \frac{1}{T} \frac{\delta Z}{\delta \beta} = \frac{1}{Z} \int dq dp (-\mathcal{H}) e^{-\beta \mathcal{H}(q,p)} = <-E>$$

Where $\langle -E \rangle$ is the average of the energy ion the canonical ensemble.

$$\begin{split} \frac{\delta^2 \ln Z}{\delta \beta^2} &= \frac{\delta}{\delta \beta} \frac{1}{Z} \frac{\delta Z}{\delta \beta} = -\frac{1}{Z^2} (\frac{\delta Z}{\delta \beta})^2 + \frac{1}{Z} \frac{\delta^2 Z}{\delta \beta^2} \\ \frac{\delta^2 \ln Z}{\delta \beta^2} &= - < E >^2 + \frac{1}{Z} \int dq dp \mathcal{H}^2(q,p) e^{-\beta \mathcal{H}(q,p)} \\ \frac{\delta^2 \ln Z}{\delta \beta^2} &= - < E >^2 + < E^2 > \\ \frac{\delta^2 \ln Z}{\delta \beta^2} &= < (E - < E >)^2 > \end{split}$$

However, keep in mind that

$$\frac{\sqrt{<(E-)^2}}{} \frac{\sqrt{N}}{N} \frac{1}{\sqrt{N}} \to_{N>>1} 0$$

Where $\sqrt{\langle (E-\langle E \rangle)^2}$ is the fluctuation in energy and the $\langle E \rangle$ is the average energy.

This shows that in the thermodynamic limit $(N \to \infty)$ the energy does not fluctuate and only the microstates with an energy equal to the average value contributes significantly when computing averages using the canonical ensemble. In other words, the canonical ensemble is equivalent to the microcanonical ensemble.

There are two specific situations where the canonical ensemble is used:

- The system is coupled to a heat reservoir
- The system is isolated but it is easier to compute averages in the canonical ensemble. This ends up being a very common situation

Previously we have argued that the averages of observables computed in the canonical ensemble are the same (in the Thermodynamic Limit) as the microcanonical ensemble. Now, we would like to demonstrate that the information contained in S(E,V,N) (or, equivalently, E(S,V,N)) is also contained in the Helmholtz Free Energy, F(T,V,N)(notice the arguments T,V,N; **NOT** S,V,N!). To do this, we'll have to utilize a mathematical algorithm known as the Legendre Transform.

10.1 Legendre Transforms

Suppose f(x) is not known but f(p), where $p = \frac{df}{dx}$, is known. Can we recover the function f(x) from f(p)? **NO!** This is clearly illustrated by the figure below:

As an analytical example, let's take a look at a couple of functions and see if we can perform this recovery:

$$f(x) = x^2, p = \frac{df}{dx} = 2x \to f(p) = (\frac{p}{2})^2$$

$$f(x) = (x - a)^2, p = \frac{df}{dx} - 2(x - a) \to f(p) = (\frac{p}{2})^2$$

From two different starting functions we extracted two different values of p, however, f(p) in both cases is the same. Clearly we cannot recover a unique initial function from the information contained solely in f(p).

Now, for single variable functions this constant shift is not a big deal. But for functions of several variables we end up seeing shifts and alterations that are functions of other variables, which greatly complicates the situation.

Even though we cannot recover the initial function from the f(p) values shown above, is it possible to construct some f(p) where it is possible to reconstruct the initial function? It turns out the answer is yes! A function of $p = \frac{df}{dx}$ containing all of the information needed to reconstruct f(x) is the intercept of the tangent with the vertical axis:

$$\rightarrow p = \frac{f-f}{x} \rightarrow \tilde{f}(p) = px - f(x)|_{xsuchthat \frac{f}{x} = p}$$

is the Legendre Transform of E(S, V, N) in relation to the variable S. Indeed, the Legendre Transform of E is:

$$\begin{split} E(S,V,N) \to \tilde{E}(T,V,N) = TS - E(S,V,N)|_{S=S*suchthat \frac{\delta E}{\delta S} = T} \\ T = \frac{\delta E}{\delta S} \end{split}$$

From the definition of the Helmholtz Free Energy:

$$F(T,V,N) = E - TS|_{E=E*,suchthat0 = \frac{\delta}{\delta E}(E-TS) = 1 - T\frac{\delta S}{\delta E}|_{E=E*}}$$

But the value S^* for which $\frac{\delta E}{\delta S}|_{S^*} = T$ corresponds to $E^* = E(S^*, V, N)$, so

$$F(T, V, N) = -\tilde{E}(E, V, N)$$

Notice that:

$$\begin{split} \frac{\delta F}{\delta T}|_{V,N} &= \frac{\delta}{\delta t} [(E(S,V,N) - TS)|_{Ssuchthat} \frac{\delta E}{\delta S}|_{S*,=T}]|_{V,N} \\ \frac{\delta F}{\delta T}|_{V,N} &= \frac{\delta E}{\delta S}|_{V,N} \frac{\delta S*}{\delta T}|_{V,N} - S^* - T \frac{\delta S^*}{\delta T}|_{V,N} \end{split}$$

(Remember that $\frac{\delta E}{\delta S}|_{V,N} = T!$)

$$\frac{\delta F}{\delta T}|_{V,N} = -S^*$$

For expedience, from now on we will be dropping the * superscript.

$$\begin{split} &\frac{\delta F}{\delta V}|_{T,N} = \frac{\delta}{\delta V} [(E(S,V,N) - TS)|_{Ssuchthat} \frac{\delta E}{\delta S}|_{S^*,}]|_{T,N} \\ &= \frac{\delta E}{\delta V}|_{S,N} + \frac{\delta E}{\delta S}|_{V,N} \frac{\delta S^*}{\delta V} - T \frac{\delta S^*}{\delta V} = \frac{\delta E}{\delta V}|_{S,N} = -P \end{split}$$

$$\frac{\delta F}{\delta N}|_{T,V} = \frac{\delta E}{\delta N}|_{S,V} + \frac{\delta E}{\delta S}|_{V,N} \frac{\delta S^*}{\delta N} - T \frac{\delta S^*}{\delta N} = \frac{\delta E}{\delta N}|_{V,N} = \mu$$

Thus, $dF = -SdT - PdV + \mu dN$ Also, as a consequence of Euler's relations, $E = TS - PV + \mu N \rightarrow F = -PV + \mu N$

10.1.1 EXAMPLE: Classical, Ideal, Non-Relativistic, Monoatomic

$$\mathcal{H}(\mathbf{x}_{i}, \mathbf{x}_{i}) = \sum_{i=1}^{N} \frac{\mathbf{x}_{i}^{2}}{2m}$$

$$Z = \int \prod_{i=1}^{N} \frac{d^{3} \mathbf{x}_{i} d^{3} \mathbf{x}_{i}}{N!} e^{-\beta \sum_{i=1}^{N} \frac{\mathbf{x}_{i}^{2}}{2m}}$$

$$= \left[\int d^{3} \mathbf{x} \int d^{3} \mathbf{x} e^{-\beta \frac{\mathbf{x}_{i}^{2}}{2m}} \right]^{3} \frac{1}{N!}$$

(Remember that $\int d^3 \setminus$ is simply V)

$$= V^{N} \left[\int_{-\infty}^{\infty} dp e^{-\frac{p^{2}}{2mk_{B}T}} \right]^{3N} \frac{1}{N!}$$
$$= V^{N} \left(\frac{2mk_{B}T}{\pi} \right)^{\frac{3N}{2}} \frac{1}{N!}$$

Therefore, in the N >> 1 limit,

$$F \cong -k_B T \ln Z = -Nk_B T \left[\ln \left(V \left(\frac{2mk_B T}{\pi} \right)^{\frac{3}{2}} \right) - \ln N \right]$$
$$F \cong -Nk_B T \ln \frac{V}{N} \left(\frac{2mk_B T}{\pi} \right)^{\frac{3}{2}}$$

This shows that F is extensive, which it should be. Now, we take the Legendre Transform of the Helmholtz Free Energy to find E(S, V, N) from F(T, V, N):

$$\begin{split} \frac{\delta F}{\delta T} &= -S = -Nk_B \ln \frac{V}{N} (\frac{2mk_BT}{\pi})^{\frac{3}{2}} \frac{-3Nk_B}{2} \\ &\rightarrow k_BT = \frac{\pi}{2m} \frac{N^{\frac{2}{3}}}{V^{\frac{2}{3}}} e^{\frac{2}{3} \frac{S}{Nk_B} - 1} \\ E &= F + TS = -Nk_BT \ln \frac{V}{N} (\frac{2mk_BT}{\pi})^{\frac{3}{2}} + \frac{\pi}{2m} (\frac{N}{V})^{\frac{2}{3}} e^{\frac{2S}{3Nk_B} - 1} S \\ E &= \frac{3}{2} \frac{\pi}{2m} \frac{N^{\frac{5}{3}}}{V^{\frac{2}{3}}} e^{\frac{2S}{3Nk_B} - 1} \end{split}$$

Which is the same value that we calculated before using the microcanonical ensemble.

10.1.2 EXAMPLE: Classical model of a Paramagnet

$$\mathcal{H}(\theta_i, L_i) = \sum_{i=1}^{N} \frac{L_i^2}{2I} - H\mu \cos \theta_i$$

$$Z = \int \prod_{i=1}^{N} d\theta_i dL_i e^{-\beta \sum_{i=1}^{N} \left[\frac{L_i^2}{2I} - H\mu \cos \theta_i\right]}$$

$$= \left[\int_0^{2\pi} d\theta e^{\beta H\mu \cos \theta}\right]^N \left[\int_{-\infty}^{\infty} dL e^{-\frac{\beta L^2}{2I}}\right]^N$$

Here we see that $[\int_0^{2\pi} d\theta e^{\beta H\mu\cos\theta}]^N$ is simply the I_0 Bessel Function, specifically $I_0(\beta H\mu)$. The second term, $[\int_{-\infty}^{\infty} dL e^{-\frac{\beta L^2}{2I}}]^N$, is approximately the value of \sqrt{T} . Therefore,

$$Z [I_0(\beta H\mu)\sqrt{T}]^N$$

Thus we can easily express the Helmholtz Free Energy of the system as

$$F = -k_B T \ln Z = -k_B T \ln \sqrt{T} I_0(\beta H \mu)$$

Now, let's derive the value of an observable. For example, the average magnetic moment per site in our system: < M >

$$\frac{1}{N} \langle M \rangle = \langle \frac{1}{N} \sum_{i=1}^{N} \mu \cos \theta_i \rangle = \frac{1}{Z} \int d\theta_i dL_i \frac{\mu}{N} \sum \cos \theta_i e^{-\beta \mathcal{H}}$$

$$= \frac{1}{\beta N} \frac{\delta}{\delta H} \ln Z = -\frac{\delta F}{\delta H}$$

$$= \frac{I_1(\frac{H\mu}{k_B T})}{I_0(\frac{H\mu}{k_B T})}$$

The above expression yields two different values based on the limit of $H\mu$.

$$H\mu << k_B T \rightarrow \frac{H\mu T}{2}$$

 $H\mu >> k_B T \rightarrow sign(H)$

10.1.3 EXAMPLE: Classical Harmonic Oscillators and the Equipartition Theorem

Systems of linked harmonic oscillators are extremely important in modelling many physical phenomena. Let's examine such a system using the equipartition theorem:

In this case, the Hamiltonian that describes the system is: $\coprod_{\rangle}, \sqrt{\rangle} = \sum_{i=1}^{N} \left[\frac{p_i^2}{2m} + \frac{m\omega^2}{2}q_i^2\right]$

Therefore, we can use this to write the system's partition function:

$$Z = \int \prod_{i=1}^{N} dq_i dp_i e^{-\beta \sum_{i=1}^{N} \left[\frac{p_i^2}{2m} + \frac{m\omega^2}{2} q_i^2 \right]}$$

$$= \left[\int_{-\infty}^{\infty} dq e^{-\frac{\beta m\omega^2}{2}q^2} \right]^N \left[\int_{-\infty}^{\infty} dp e^{-\frac{\beta p^2}{2m}} \right]^N$$

The first term is approximately $\frac{1}{\sqrt{m\omega^2 k_B T}}$ and the second term is approximately $\sqrt{2mk_B T}$.

Many of these variables cancel or reduce, leaving us with

$$Z \left(\frac{k_B T}{\omega}\right)^N = (\beta \omega)^{-N}$$

Solving for the average energy yields:

$$= -\frac{1}{Z}\frac{\delta Z}{\delta\beta} = -\frac{\delta \ln Z}{\delta\beta} = \frac{N}{\beta} = Nk_BT$$

This result is important, as it shows in our classical model that the average energy of any one harmonic oscillator at temperature T is k_BT , entirely independent of ω . Oscillators with a large ω (stiff spring constant) will oscillate less, the oscillators with a small ω (loose spring constant) will oscillate more, but all will have the same average energy:

If we had worked this example for free particles in one dimension instead of oscillators we would have found that $\langle E \rangle = \frac{1}{2}k_BT$ (see appendix). The fact that many physical systems can be written in terms of coupled harmonic oscillators (normal modes) implies the **Equipartition Theorem**. This theorem is extremely powerful and is very useful in solving classical systems, so it is stated separately below:

Equipartition Theorem: Each quadratic term in the Hamiltonian contributes $\frac{k_BT}{2}$ to the system's average energy (< E>).

10.1.4 EXAMPLE: Classical, Non-relativistic, Diatomic Gas

Assuming the vibrational mode can be treated as classical, we have:

$$\mathcal{H} = \sum_{i=1}^{N} \frac{\underline{\mathbf{j}}}{2m} + \frac{p_i^2}{2m} + \frac{m\omega^2}{2} x_i^2$$

Disecting this hamiltonian, we see that the first term is a momentum vector. This represents the center of mass momentum, so it contributes 3 quatratic terms per molecule. The second and third terms together represent the molecule's vibrational energy, where x_i is the distance between the atoms in the molecule. The vibrational energy contributes 2 quadractic terms per molecule. Since we know this information, it is now trivial to construct an expression for the average energy using the Equipartition Theorem:

$$\frac{1}{N} < E > = (3+2)\frac{1}{2}k_BT = \frac{5}{2}k_BT$$

11 Quantum Statistical Mechanics

So far we have been assuming that the systems we have been working with behaved Classically at the microscopic level. Now let's rework our previous formalism to describe systems which exist within the realm of Quantum Mechanics. There are two ways to accomplish this: We can trust analogies and guess the form of most expressions or we can generalize the concept of the ensemble for quantum systems. The second option is much more powerful, however, the first is quicker and will be enough for the majority of the systems we will be dealing with.

11.		
	Classical	Quantum
Observables:	Function in Phase	Operator in
	Space $\to f(q, p)$	Hilbert Space
		$ ightarrow \hat{f}$
Averages in the	$\frac{1}{Z}\int dqdpe^{-\beta H}f(q,p)$), $ZTx[e^{-\beta \hat{H}}\hat{f}],Z = Tr[e^{-\beta \hat{H}}]$
Canonical	$\int dq dp e^{-\beta H}$	$Tr[e^{-\beta \hat{H}}]$
Ensemble:		, ,
Partition	Z =	$Z = \sum_{n} <$
Function:	$\int dE \Gamma(E) e^{-\beta E}$	$Z = \sum_{n} < n e^{-\beta \hat{H}} n > = \sum_{n} e^{-\beta E_{n}} \cong \int dE\Gamma(E)e^{-\beta E}$
		$\sum_{n} e^{-\dot{\beta}E_n} \cong$
		$\int \overline{dE} \Gamma(E) e^{-\beta E}$

A couple of notes regarding the quantum partition function: n is are the eigenstates of \hat{H} and E_n are the eigenvalues. $dE\Gamma(E)$ is the number of quantum states between E and E+dE. The integral over these states approximates the partition function only for very dense spectra.

11.1 The Density Matrix

Although this concept won't actually be used in this course, it is a basic thing that any good physicist worth their salt should know. Suppose we know the quantum state of a system at a given time, for example

$$|\Psi>$$

We can compute the expectation value of an observable (represented by operator \hat{A} in this case) by

$$\langle A \rangle = \langle \Psi | \hat{A} | \Psi \rangle$$

Where $\langle A \rangle$ is the average over results obtained by many identically prepared systems. An interesting observation of this statement is that we can get different results from identical measurements on identically prepared systems.

This uncertainty is due to the nature of Quantum Mechanics. In particular, if a system is prepared in the state $|\Psi\rangle = \alpha |\Psi_1\rangle + \beta |\Psi_2\rangle$, $\langle A\rangle$ is given by:

$$< A> = <\Psi |\hat{A}|\Psi> = |\alpha|^2 <\Psi_1 |\hat{A}|\Psi_1> + |\beta|^2 <\Psi_2 |\hat{A}|\Psi_2> + \alpha^*\beta <\Psi_1 |\hat{A}|\Psi_2> + \alpha\beta^* <\Psi_2 |\hat{A}|\Psi_1> + \alpha^*\beta <\Psi_1 |\hat{A}|\Psi_2> + \alpha\beta^* <\Psi_2 |\hat{A}|\Psi_1> + \alpha^*\beta <\Psi_1 |\hat{A}|\Psi_2> + \alpha\beta^* <\Psi_2 |\hat{A}|\Psi_1> + \alpha\beta^*\beta <\Psi_1 |\Psi_1> + \alpha\beta^*\beta <\Psi_1 |\Psi_1> + \alpha\beta^*\beta <\Psi_1 |\Psi_1> + \alpha\beta^*\beta <\Psi_1 |\Psi_1> + \alpha\beta^*\beta <\Psi_$$

 $\alpha^*\beta < \Psi_1|\hat{A}|\Psi_2> +\alpha\beta^* < \Psi_2|\hat{A}|\Psi_1>$ are the interference terms, so they may have either sign. Now, suppose that eh state $|\Psi>$ is not known for sure, not because of any uncertainty imposed by quantum mechanics but just because of practical considerations. Let us state that there is a probability p_1 that the system is in state $|\Psi_1>$ and probability p_2 that it is in state $|\Psi_2>$. The actual wavefunction is either $|\Psi_1>$ or $|\Psi_2>$. Therefore, we expect that the expectation value of observable A is:

$$< A> = p_1 < \Psi_1 |\hat{A}|\Psi_1> + p_2 < \Psi_2 |\hat{A}|\Psi_2> = Tr[(p_1|\Psi_1> < \Psi_1| + p_2|\Psi_2> < \Psi_2|)\hat{A}] = Tr[\hat{\rho}\hat{A}]$$

The operator $\hat{\rho}$ is called the **Density Matrix** and it is the quantum analogue of the phase space distribution $\rho(q_i, p_i)$. Of course, this definition generalizes to the case where the classical uncertainty extends to several states:

$$\hat{\rho} = \sum_{n} p_n |n\rangle \langle n|$$

From this definition a few properties directly follow:

- $Tr\hat{\rho}$ (since $\sum_{n} p_n = 1$)
- $\hat{\rho}^{\dagger} = \hat{\rho}$ (ρ is hermitian, since the analogue of $\rho(q_i, p_i)$ must be real)

If $\hat{\rho} = |\Psi> < \Psi|$, that is, if we know the quantum state of a system with 100% confidence, we say that the system is in a *pure* state. Otherwise we say the system is in a *mixed* state. For a pure state $\hat{\rho}^2 = |\Psi> < \Psi|\Psi> < \Psi| = |\Psi> < \Psi| = \hat{\rho}$, so $\hat{\rho}^2 = \hat{\rho} \leftrightarrow \hat{\rho}$ is a pure state.

 $|\Psi><\Psi|=\hat{\rho},$ so $\hat{\rho}^2=\hat{\rho}\leftrightarrow\hat{\rho}$ is a pure state. Notice the difference between $|\Psi>=\frac{1}{\sqrt{2}}|\Psi_1>+\frac{1}{\sqrt{2}}|\Psi_2>$ and $\hat{\rho}=\frac{1}{2}|\Psi_1><\Psi_1|+\frac{1}{2}|\Psi_2><\Psi_2|$. Both could be described as a 50-50 combination of $|\Psi_1>$ and $|\Psi_2>$ but they represent completely different physical situations.

11.2 EXAMPLE: Quantum Harmonic Oscillator

We begin, as always, with the Hamiltonian.

$$H = \frac{\hat{p}^2}{2m} + \frac{m\omega^2}{2}\hat{x}^2 \to E_n = (n + \frac{1}{2})\hbar\omega, |n>, n = 0, 1, \dots$$

$$Z = \sum_{n_1=0}^{\infty} \sum_{n_2=0}^{\infty} \sum_{n_3=0}^{\infty} \dots \sum_{n_N=0}^{\infty} e^{\beta \hbar \omega [(n_1 + \frac{1}{2}) + (n_2 + \frac{1}{2}) + \dots + (n_N + \frac{1}{2})]}$$

$$Z = \left[\sum_{n=0}^{\infty} e^{-\beta\hbar\omega(n+\frac{1}{2})}\right]^{N}$$

$$Z = \left[e^{-\frac{\beta\hbar\omega}{2}} \frac{1}{1 - e^{-\beta\hbar\omega}}\right]^{N}$$

$$Z = \left[\frac{1}{e^{\frac{\beta\hbar\omega}{2}} - e^{-\frac{\beta\hbar\omega}{2}}}\right]^{N}$$

$$Z = \left[\frac{1}{2\sinh\frac{\beta\hbar\omega}{2}}\right]^{N}$$

$$Z = \left[\frac{1}{2\sinh\frac{\beta\hbar\omega}{2}}\right]^{N}$$

$$\frac{1}{N} < E > = -\frac{1}{ZN} \frac{\delta Z}{\delta\beta} = -\frac{1}{N} \frac{\delta \ln Z}{\delta\beta} = \frac{1}{N} \frac{\delta N \ln 2 \sinh\frac{\beta\hbar\omega}{2}}{\delta\beta}$$

$$= \frac{\cosh\frac{\beta\hbar\omega}{2}}{\sinh\frac{\beta\hbar\omega}{2}} = \frac{\hbar\omega}{2} \coth\frac{\beta\hbar\omega}{2}$$

Similar to the classical situation, this expression again yields two different solutions based on the limit of k_BT .

$$k_B T >> \hbar \omega \to \frac{\hbar \omega}{2} + k_B T$$

 $k_B T << \hbar \omega \to \frac{\hbar \omega}{2} + \hbar \omega e^{-\frac{\hbar \omega}{k_B T}}$

Note that the first result is the classical result k_BT modified by the addition of zero-point energy $(\frac{\hbar\omega}{2})$. The second result includes the same zero-point energy, but also includes "Boltzmann Supressed" thermal energy. This arises because in the $k_BT \ll \hbar\omega$ limit the high frequency modes are "frozen". This runs contrary to the classical equipartition theory.

11.3 **EXAMPLE: Specific Heat of Molecular Gases**

Molecules have vibrational and rotational degrees of freedom in addition to the standard translational ones. For instance, for H_2 :

$$\mathcal{H} = \sum_{i=1}^{N} \frac{\mathbb{P}_i^2}{2M} + \frac{\mathbb{P}_i^2}{2m_{red}} + m_{red} \frac{\omega^2}{2} x_i^2 + \frac{\mathbb{L}_i^2}{2I}$$

Breaking this down by terms, we have: $\frac{\mathbb{P}_i^2}{2M}$, which is the center of mass kinetic energy (M=2m); $\frac{\mathbb{P}_i^2}{2m_{red}}$, which is the vibrational kinetic energy $(\frac{1}{m_{red}}=1)$ $\frac{1}{m} + \frac{1}{m}$); $m_{red} \frac{\omega^2}{2} x_i^2$, which is the vibrational potential energy (x_i is the separation between atoms); and $\frac{\mathbb{L}_i^2}{2I}$ which is the rotational energy. Classically, we express the partition function thusly: $Z = \int \prod_{i=1}^N d^3 \mathbb{R}_i d^3 \mathbb{P}_i dp_i dx_i d^2 \mathbb{L}_i e^{-\beta \mathcal{H}}$

$$Z = \int \prod_{i=1}^{N} d^3 \mathbb{R}_i d^3 \mathbb{P}_i dp_i dx_i d^2 \mathbb{L}_i e^{-\beta \mathcal{H}_i}$$

Note that there is no rotation around the molecule's axis. This is because such rotation would result in a state that is indistinguishable from the initial state and therefore have no effect on our system.

If all of these degrees of freedom are classical,

Part IV

Quantum Gases

Previously, using the grand canonical ensemble we found the grand canonical potential of a quantum ideal gas of identical particles:

$$\Omega(T, V, \mu) = \pm g k_B T V \int \frac{d^3 k}{(2\pi)^3} \ln(1 \mp e^{-\beta(\frac{\hbar^2 k^2}{2m} - \mu)})$$

In this expression, we nicely sum up the main components of the canonical potential for both bosonic and fermionic gases. To clarify, bosons behave according to the upper operators and fermions utilize the lower operators (i.e. for bosons there is a plus sign outside the log and a minus sign inside, and the converse is true for fermions). We also rely on g, which is the degeneracy of single particle (orbital) states, 2s+1 for particles of spin s. Within the log, the value $\frac{\hbar^2 k^2}{2m}$ is the energy E_k . This is the energy of one particle with momentum of $\hbar k$.

Now,
$$N = -\frac{\delta\Omega}{\delta\mu}$$
 since $-\frac{\delta\Omega}{\delta\mu} = \frac{\delta}{\delta\mu} \frac{1}{\beta} \ln Tr[e^{-\beta(\hat{H} - \mu\hat{N})}]$
$$= \frac{1}{\beta} \frac{Tr[\beta e^{-\beta(\hat{H} - \mu\hat{N})}\hat{N}]}{Tr[e^{-\beta(\hat{H} - \mu\hat{N})}]} = Tr[\hat{\rho}\hat{N}] = \langle N \rangle$$

Using this information we can find the particle density:

$$\frac{N}{V} = -\frac{\delta\Omega}{\delta\mu} = \mp k_B T g \int \frac{d^3k}{(2\pi^3)} \frac{\mp \beta e^{-\beta(E_k - \mu)}}{1 \mp e^{-\beta(E_k - \mu)}}$$
$$= g \int \frac{d^3k}{(2\pi^3)} \frac{1}{e^{\beta(E_k - \mu)} \mp 1}$$

Similarly,

$$S = -\frac{\delta\Omega}{\delta T} = \mp k_B g V \int \frac{d^3k}{2\pi^3} \ln 1 \mp e^{-\beta\epsilon_k}$$