Chapter 1

Equilibrium phases and thermodynamics of phase transitions

1.1 Equilibrium phases as minima of Gibbs free energy

Experimentally, any element or compound can be found, depending on the thermodynamic conditions in which it is, in different phases. When we say that a system is in a particular phase we mean that its physical properties (like density or magnetization) are uniform.

Equilibrium states are given by maxima of the entropy and minima of internal energy, or by minima of thermodynamics potentials such as A and G. Let us consider for example the Gibbs potential per particle of a fluid system

$$\frac{G}{N} \equiv g = g(T, P) \tag{1.1}$$

that depends on two intensive variables T and P and is not anymore a function of N because we have divided for N. Let us define α as the phase of a one-component system (say $\alpha = \text{gas}$ or liquid). Therefore, the thermodynamic properties are described by surfaces of function $g_{\alpha}(T,P)$ and for all equilibrium phase we have a surface on the space (T,P,g). For each value of T and P the thermodynamically stable phase is the one for which $g_{\alpha}(T,P)$ is minimum.

1.2 First order phase transition and phase coexistence

Let us suppose for example that the system can be found in two phases α and β (for example liquid and solid). Consider the surface g_{α} and g_{β} , we are looking for the lower one.

For given values of T and P the stable phase will be that with the lowest value of g: for example, if we have $g_{\alpha}(T,P) < g_{\beta}(T,P)$ then the system will be in phase α . Therefore there will be regions in (T,P) space were the most stable phase will be α and others in which it will be β . If we now plot the values of g as a function of T and T in (g,P,T) space for every phase of the system, we can determine the regions where the two phases will be the stable ones, namely we can determine the phase diagram of the system, as illustrated in Figure 1.1.

The very interesting region of this space (and the one on which we will focus our attention in this section) is the line where the surfaces of the two phases intersect: along this the two phases coexist, and when the system crosses it we say that it undergoes a *phase transition*. The coexistence line is the projection on the (T,P) plane of the intersection between different surfaces, so the *coexistence condition* is:

$$g_{\alpha}(T, P) = g_{\beta}(T, P) \tag{1.2}$$

To fix the ideas, let us choose a given value of pressure $P = P^*$ and study the behavior of $g(T, P^*)$ as a function of T when we go from solid to gas, as illustrated in Figure 1.2.

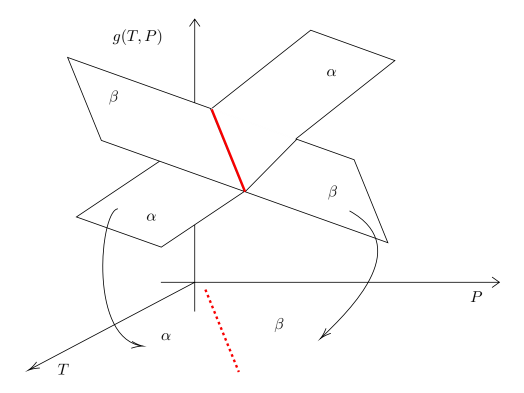


Figure 1.1: Phase diagram: stability of phases.

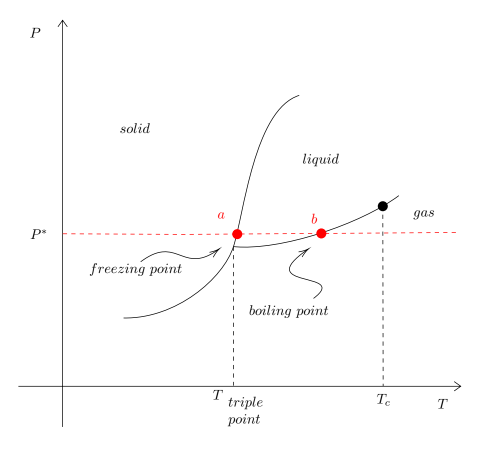


Figure 1.2: (T, P) projection.

The existence of a critical point has a very intriguing consequence: since the liquid-gas coexistence line ends in a point, this means that a liquid can continuously be transformed in a

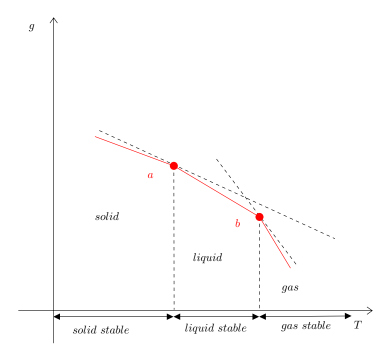


Figure 1.3: (g,T) projection at a fixed pression $P=P^*$. The red line is the coexistence line of the two phases α and β .

gas (or viceversa), and in such a way that the coexistence of liquid and gaseous phases is never encountered.

At the coexistence line, $g_{\text{solid}}(T_a, P^*) = g_{\text{liq}}(T_a)$ and $g_{\text{liq}}(T_b) = g_{\text{gas}}(T_b, P^*)$, as shown in Figure 1.3.

Note also that:

- At the coexistence points a and b of the two phases, one has $g_{\alpha}(T) = g_{\beta}(T)$.
- g(T) is a continuous function of T.
- Note that, $S = -\left(\frac{\partial G}{\partial T}\right)_V$ and $c_P = -T\left(\frac{\partial^2 G}{\partial T^2}\right)_P > 0$. This implies that g(T) is concave in T at fixed P.

How about its derivatives? Since P is fixed we can vary T and look for $s = -\left(\frac{\partial g}{\partial T}\right)_P$. As we cross different phases we have discontinuities, where ΔsT is called the *latent heat*. It is illustrated in Figure 1.4.

If there is a finite discontinuity in one, or more, of the *first* derivatives of the appropriate thermodynamic potential, the transition is called *first order transition*. In general, a phase transition is signaled by a singularity in a thermodynamic potential.

We can also fix the temperature $T=T^*$ and look at the variation of P, as shown in Figure 1.5. Note that, we have $v=\left(\frac{\partial g}{\partial P}\right)_T>0$:

$$\left(\frac{\partial^2 g}{\partial P^2}\right) = \left(\frac{\partial v}{\partial P}\right)_T = -vk_T < 0$$
(1.3)

so, also in this case we had a jump of the first order derivative of the thermodynamic potential g. It is illustrated in Figure 1.6.

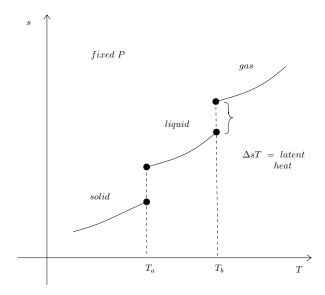


Figure 1.4: (s,T) projection.

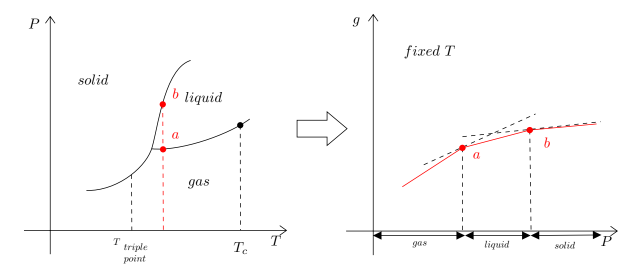


Figure 1.5: Left: (T, P) projection. Right: (g, P) projection at a fixed temperature $T = T^*$.

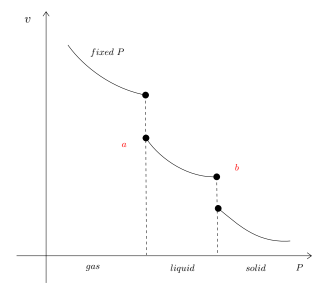


Figure 1.6: (v, P) projection.

1.2.1 Critical points

At the critical point (P_c, T_c) the system can pass from the liquid to the gas phase (and vice versa) in a continuous way

$$\Delta s = \Delta v = 0$$

Usually, critical points are end point of first order transition phases. Why there is no critical point between solid and liquid? The crossover between phases having the same symmetry define the Landau point. There is a break of symmetry, for instance we can think about the structure of the bravais lattice. Instead, from gas to liquid symmetries are not broken.

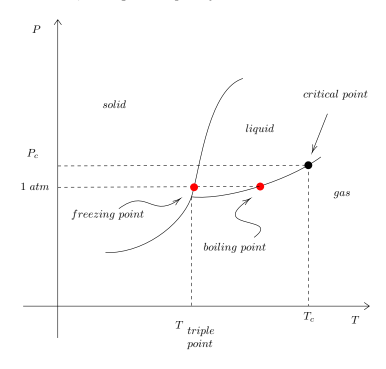


Figure 1.7: Phase diagram of a fluid. All the phase transition are first-order except at the critical point C. Beyond C it is possible to move continuously from liquid to a gas. The boundary between the solid and liquid phases is thought to be always first-oder and not to terminate in a critical point.

1.2.2 Ferromagnetic system

A similar behaviour can be encountered in magnetic systems. We can have a magnetization different from 0 even when the is no magnetic field. Supposing $P \leftrightarrow H, V \leftrightarrow M$, we have $(P,T) \leftrightarrow (H,T)$.

The magnetization M has a jump at H=0 for temperatures lower than the critical one; in this case since $M=-\frac{\partial F}{\partial H}$ we see that the first derivative of the free energy F with respect to H has a jump discontinuity. For instance, consider Figure 1.9. At the critical point the magnetization would pass through zero.

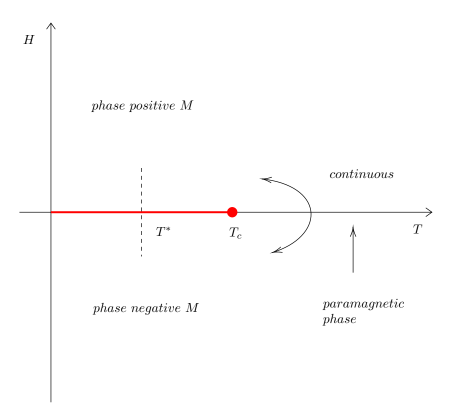


Figure 1.8: Phase diagram for a magnetic system in (T, H) space. A line of first-order transitions at zero field ends in a critical point at a temperature T_c .

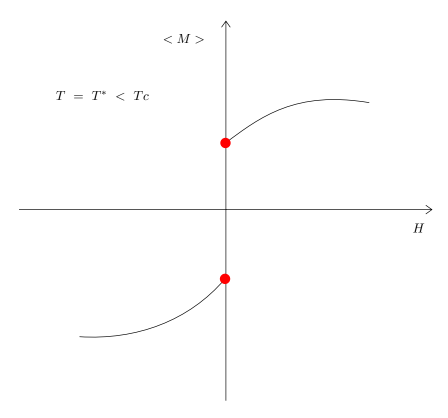


Figure 1.9: Plot of the Magnetization for $T = T^* < T_c$.

1.3 Second order phase transition

The transitions are classified in the first order transition and continuous transition. If the first derivatives are continuous, but second derivatives are discontinuous, or infinite, the transition will be described as higher order, continuous or critical. This is different from the previous situation, in which we had a jump for the first order derivative of a thermodynamic potential. Some examples are illustrated in Figure 1.10.

Let us suppose that

$$\left(\frac{\partial g}{\partial T}\right)_P = -s \tag{1.4a}$$

$$\left(\frac{\partial g}{\partial T} \right)_P = -s$$
 (1.4a)
$$\left(\frac{\partial g}{\partial P} \right)_T = v$$
 (1.4b)

are continuous. We suppose also that

$$\left(\frac{\partial^2 g}{\partial T \partial P}\right) = \left(\frac{\partial v}{\partial T}\right)_P = v\alpha_p \tag{1.5}$$

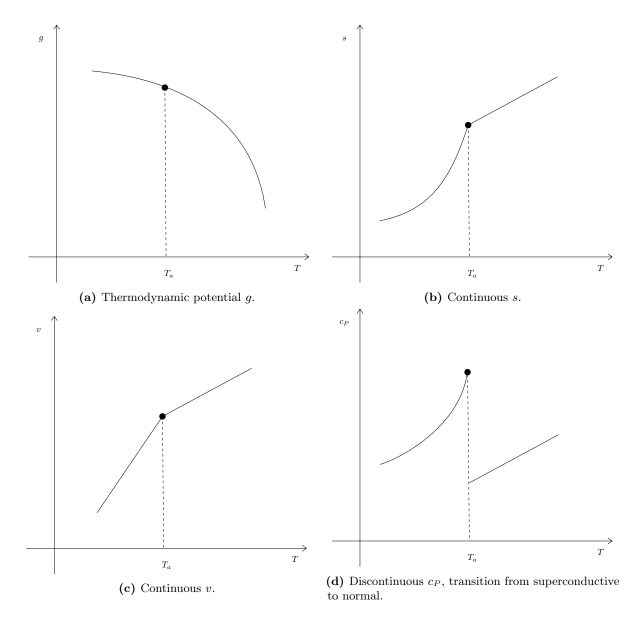


Figure 1.10: Example of a second order phase transition.

If we look for example at the specific heat c_P in Figure 1.10d, it represent the transition from superconducting.

The critical point is special because there is not a jump, so we can go continuously from gas to liquid. The response function when we plot this point shows that the specific heat diverges.

The *superfluid transition* is a transition where the second derivative of the thermodynamic potential diverges. There are many phase transitions that can be classified in different ways.

Remark. Note that at the coexistence line we increase V, but the pressure remains constant. At the coexistence line we see bubbles. It is the density that is changing locally, the bubbles becomes bigger and bigger and at the V_G , becomes a liquid.

1.3.1 Helmholtz free-energy

Consider A = A(T, V, N), here P is replaced by V which has the derivative discontinuous at the first order transition. Moreover, P > 0 implies $\partial A/\partial V < 0$ and

$$k_T = -\frac{1}{V} \left(\frac{\partial V}{\partial P} \right)_T = -\frac{1}{V} \left(\frac{\partial P}{\partial V} \right)_T = \frac{1}{V} \left(\frac{\partial^2 A}{\partial V^2} \right)_T > 0 \tag{1.6}$$

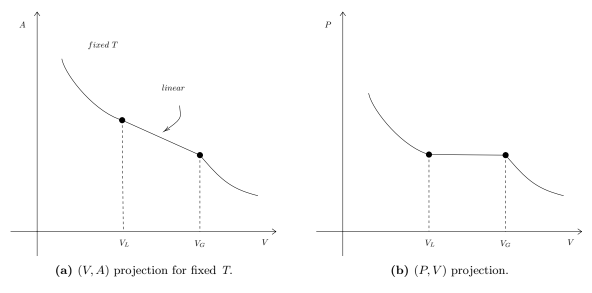


Figure 1.11: Helmholtz free-energy and phase transition.

so, A is an overall convex function of V. The behaviour of A when there is a first order phase transition is as in Figure 1.11a. The linear sector becomes an horizontal one in the $P = -(\partial A/\partial V)_T = P(V)$ curve (Figure 1.11b).

1.4 Thermodynamic of phase coexistence

1.4.1 Lever Rule

The lever rule is a rule used to determine the mole fraction of each phase of a binary equilibrium phase diagram. For instance, it can be used to determine the fraction of liquid and solid phases for a given binary composition and temperature that is between the liquid and solid line.

In an alloy or a mixture with two phases, α and β , which themselves contain two elements, A and B, the lever rule states that the mass fraction of the α phase is

$$w^{\alpha} = \frac{w_B - w_B^{\beta}}{w_B^{\alpha} - w_B^{\beta}} \tag{1.7}$$

where

- w_B^{α} : is the mass fraction of element B in the α phase.
- w_B^{β} : is the mass fraction of element B in the β phase.
- w_B : is the mass fraction of element B in the entire alloy or mixture.

Example 1

Consider Figure 1.12; at all points between A and B the system is a mixture of gas and liquid. Points D has global density $\rho_D = \rho_A + \rho_B$ and therefore $v_D = \frac{1}{\rho_D}, v_A = \frac{1}{\rho_A}, v_B = \frac{1}{\rho_B}$ which implies:

$$v_D = \frac{N_A}{N}v_A + \frac{N_B}{N}v_B = x_A v_A + x_B v_B$$

Since $x_A + x_B = 1$ we have $(x_A + x_B)v_D = x_Av_A + x_Bv_B$ and finally by rearranging, one finds the *Lever Rule*. It shows that the relative concentration of the liquid-gas mixture changes with V:

$$\frac{x_A}{x_B} = \frac{v_B - v_D}{v_D - v_A}$$

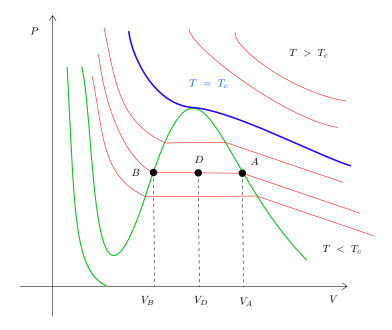


Figure 1.12: (V, P) projection. In the region between A and B the gas and the liquid phase coexist by keeping the pressure constant.

1.4.2 Phase coexistence (one component system)

Consider a (P, V, T) system as a mixture of two species (1, 2) at temperature T_1, T_2 , pressure P_1, P_2 and chemical potentials μ_1, μ_2 . The equilibrium condition is given by the maximum of the total entropy $S = S_1 + S_2$ and gives the conditions

$$T_1 = T_2, \quad P_1 = P_2, \quad \mu_1 = \mu_2$$
 (1.8)

this is the *coexistence condition* of the two phases.

In terms of the Gibbs potential G = U - TS + PV, where U is given by the Euler equation $U = TS - PV + \mu_1 N_1 + \mu_2 N_2$, the Gibbs per mole is

$$g_1(T, P) \equiv \frac{G_1}{N_1} = \mu_1$$
 (1.9a)

$$g_2(T, P) \equiv \frac{G_2}{N_2} = \mu_2$$
 (1.9b)

Therefore, on the coexistence line it should hold the relation

$$g_1(T, P) = g_2(T, P)$$
 (1.10)

1.4.3 Clausius-Clapeyron equation

The coexistence curves, as the one illustrated in Figure 1.13, are less arbitrary than is immediately evident; the slope $\mathrm{d}P\,/\,\mathrm{d}T$ of a coexistence curve is fully determined by the properties of the two coexisting phases.

The slope of a coexistence curve is of direct physical interest. Consider cubes of ice at equilibrium in a glass of water. Given the ambient pressure, the temperature of the mixed system is determined by the liquid-solid coexistence curve of water; if the temperature were not on the coexistence curve some ice would melt, or some liquid would freeze, until the temperature would again lie on the coexistence curve (or one phases would become depleted). If the ambient pressure were to decrease perhaps, by virtue of a change in altitude, then the temperature of the glass of water would appropriately adjust to a new point on the coexistence curve. If ΔP were the change in pressure, then the change in temperature would be $\Delta T = \Delta P/(\mathrm{d}P/\mathrm{d}T)_{coex}$, where the derivative in the denominator is the slope of the coexistence curve.

Remark. Ice skating presents another interesting example. The pressure applied to the ice directly beneath the blade of the skate shifts the ice across the solid-liquid coexistence curve, providing a lubricating film of liquid on which the skate slides. The possibility of ice skating depends on the negative slope of the liquid-solid coexistence curve of water.

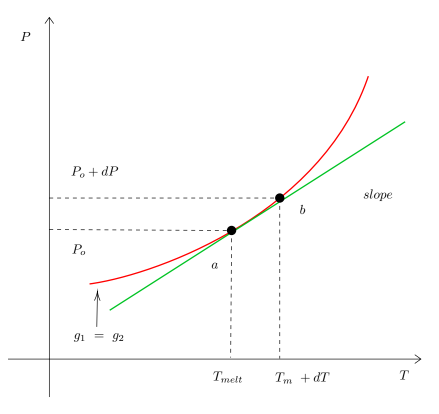


Figure 1.13: (T, P) projection. The coexistence line is represented in red, while in green the slope between the two points a and b.

Now, suppose to know the position on the coexistence line (for example the melt temperature T_m at the atmospheric pressure P_0). Is it possible to find other points on the curve? For example T_m at lower or higher pressure?

The answer is yes for small deviations of T and P from a. The idea is to compute the slope of the tangent of the coexistence curve, i.e. $(\mathrm{d}P/\mathrm{d}T)$. This is given by the Clausius-Clapeyron equation. Both at a and b the two phases 1 and 2 coexist. This means that at the coexistence line

$$\begin{cases}
g_1^{(a)} = g_2^{(a)} \\
g_1^{(b)} = g_2^{(b)}
\end{cases}$$
(1.11)

Hence, if a and b are very close:

$$\begin{cases}
dg_1 = g_1^{(b)} - g_1^{(a)} \\
dg_2 = g_2^{(b)} - g_2^{(a)}
\end{cases}$$
(1.12)

Therefore, the starting point for Clausius-Clapeyron is

$$\Rightarrow dg_1 = dg_2 \tag{1.13}$$

From the molar version of the Gibbs-Duhem relation, we have

$$\begin{cases} dg_1 = -s_1 dT + v_1 dP = d\mu_1 \\ dg_2 = -s_2 dT + v_2 dP = d\mu_2 \end{cases}$$
 (1.14)

taking the difference, one obtains

$$-(s_2 - s_1) dT + (v_2 - v_1) dP = 0$$

The slope is called **Clausius-Clapeyron equation**:

$$\left(\frac{\mathrm{d}P}{\mathrm{d}T}\right)_{coex} = \frac{(s_2 - s_1)}{(v_2 - v_1)} = \frac{\Delta s}{\Delta v} \tag{1.15}$$

Remark. Since $(dP/dT)_{coex}$ is finite, the equation explains why a first order transition is characterised by discontinuous changes in entropy and volume (or density). ΔS gives the latent heat L_{12} ¹:

$$L_{12} = T\Delta s \tag{1.16}$$

whence, the Clapeyron equation is

$$\frac{\mathrm{d}P}{\mathrm{d}T} = \frac{L_{12}}{T\Delta v} \tag{1.17}$$

1.4.4 Application of C-C equation to the liquid-gas coexistence line

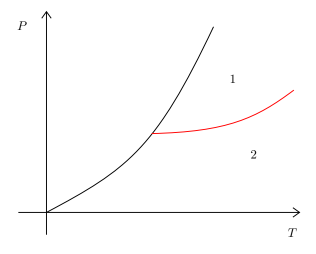


Figure 1.14: (T, P) projection. Region 1: liquid. Region 2: gas. The lines represent the combinations of pressures and temperatures at which two phases can exist in equilibrium.

Now, we go from gas (region 2) to liquid (region 1), we have:

$$\left(\frac{\mathrm{d}P}{\mathrm{d}T}\right)_{coex} = \frac{s_2 - s_1}{v_2 - v_1}$$

¹The latent heat of fusion is the quantity of heat required to melt one mole of solid.

The Clapeyron equation embodies the *Le Chatelier principle*². Consider a liquid-gas transition (the coexistence curves, from which the inequality sign is derived, are shown in Figure 1.14):

$$\left(\frac{\mathrm{d}P}{\mathrm{d}T}\right)_{coex} > 0 \quad \Rightarrow \frac{s_2 - s_1}{v_2 - v_1} > 0$$

and since $v_2 > v_1$, we have $s_2 > s_1$. The gas has more entropy as it should be. The slope of the phase curve is positive, then an increase in pressure at constant temperature tends to drive the system to the more dense (solid) phase, and an increase in temperature tends to drive the system to the more entropic (liquid) phase.

When going from a low-temperature phase to a high-temperature phase entropy always increases $\Delta S > 0$, because $c_P \equiv T(\partial S/\partial T)_P > 0$.

The sign of ΔV is more uncertain though. To see this point, let us consider the C-C equation at the solid-liquid (now solid is region 1 and liquid region 2) coexistence curve. At the melt temperature:

$$\left(\frac{\mathrm{d}P}{\mathrm{d}T}\right)_{coex} = \frac{\delta Q_{melt}}{T_{melt}\Delta v_{melt}}, \qquad \delta Q_{melt} = Q_{liq} - Q_{solid} > 0$$

In general, $\Delta v_m = v_{liq} - v_{solid} > 0$ which implies $(dP/dT)_{coex} > 0$. There are cases, however, where $\Delta v_m = v_{liq} - v_{solid} < 0$ because $\rho_{liq} > \rho_{solid}$ (for instance the H_20 , or also Silicon and Germanium). The paradigmatic example is the freezing of water where $v_{ice} > v_{liq}$ since ice is less dense than liquid water at the coxistence (0 < T < 4). This implies that dP/dT < 0.

Example 2: Melting point on Everest

Consider T = 237K and $P = P_0$. If we suppose that

$$\delta Q_m = 6.01kJ/mol, \quad \Delta v = -1.7cm^3/mol$$

we have

$$\frac{\mathrm{d}P}{\mathrm{d}T} = \frac{\delta Q_m}{T\Delta v} = \frac{6.0110^3 J/mol}{273 \cdot (-1.7cm^3/mol)} = -1.29 \cdot 10^4 J/m^3 = -1.29bar/K$$

$$\Rightarrow \Delta T = \frac{\Delta P}{(-1.29Pa/K)} = \frac{(P_0 - P_{\text{Everest}})}{(-1.29Pa/K)} = \frac{(1 - 0.36)atm}{(-1.29Pa/K)} = -0.5^{\circ}\text{C}$$
$$\Rightarrow T_m(\text{Everest}) = T_m(P_0) + 0.5^{\circ}\text{C}$$

Example 3: Boiling point on Everest

Let us consider

$$P_{\text{Everest}} = 0.36atm, \quad \rho(T = 100^{\circ}\text{C}) = 0.598kg/m^3, \quad L_{gl} = 2.257 \cdot 10^3 J/g$$

The density of the vapour (gas) is about 1000 less than water (liquid), it implies that:

$$\Delta V = V_g - V_l \approx V_g = \frac{1}{\rho_g}$$

We have

$$\frac{dP}{dT} = \frac{L_{ge}}{T\Delta V} = \frac{L_{ge}\rho_g}{T} = \frac{2.25 \cdot 10^3 J/g \cdot 0.593 kg/m^3}{373 K} = \frac{3.6}{K} \frac{10^3 J}{g} \frac{kg}{m^3} = 3.6 \cdot 10^3 \frac{Pa}{K}$$

$$\Rightarrow \Delta T \approx \Delta P/(3.610^3 Pa/K) = 18^{\circ} \text{C}$$

$$\Rightarrow T_0 - T_{\text{Everest}} = 18^{\circ} \text{C} \quad \Rightarrow T_{\text{Everest}} \approx 80^{\circ} \text{C}$$

²"When a settled system is disturbed, it will adjust to diminish the change that has been made to it".

1.5 Order parameter of a phase transition

An order parameter is a measure of the degree of order across the boundaries in a phase transition system. In particular, *order parameters* are macroscopic observable that are equal to zero above the critical temperature, and different from zero below:

$$O_p = \begin{cases} \neq 0 & T < T_c \\ = 0 & T \to T_c^- \end{cases}$$
 (1.18)

When a phase transition implies a breaking of a phase symmetry, the order parameter is related to this symmetry. Therefore, the order parameter reflects the symmetry of the system. Recall that, at T_c the system has a symmetry broken.

For instance, consider the densities of liquid and gas and the related order parameter of the gas-liquid transition $\Delta \rho = \rho_l - \rho_g$, that is $\neq 0$ for $T \neq T_c$ but $\to 0$ when $T \to T_c$ (see Figure 1.15).

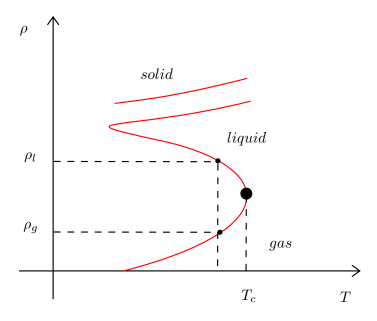


Figure 1.15: (T, ρ) projection of the (P, V, T) system, where $\rho = N/V$.

Remark. Note that $\rho = \frac{N}{V} = \frac{1}{v}$ hence either N or V varies.

In Figure 1.16 is shown the behaviour for a ferromagnetic system. We have

$$H = 0 \Rightarrow \begin{cases} M \neq 0 & T < T_c \\ M \to 0 & T \to T_c^- \end{cases}$$

Clearly $M \neq 0$ if $H \neq 0$. Recall that M is the order parameter of the paramagnetic-ferromagnetic phase transition.

It can be either a scalar $(\Delta \rho)$, a vector (\vec{M}) or even tensor $(Q_{\alpha\beta})$.

Variable conjugate to O_P

- Ferromagnetic system: $\vec{\mathbf{M}} \to \vec{\mathbf{H}}$ (magnetic field).
- Ferroelectric: $\vec{\mathbf{P}} \to \vec{\mathbf{E}}$ (electric field).
- Liquid crystals: $Q_{\alpha\beta} \to \vec{\mathbf{E}}, \vec{\mathbf{H}}$.
- Fluid: $V \to P$ (pressure), or $\rho \to \mu$.

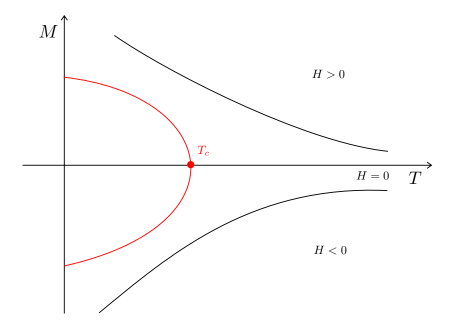


Figure 1.16: Magnetization of a ferromagnet. In red: zero-field magnetization. Below the critical temperature there is a spontaneous magnetization.

1.6 Classification of the phase transitions

1.6.1 Thermodynamic classification

Thermodynamically, one can distinguish two kinds of phase transitions:

- 1. Ones who develop latent heat.
- 2. Ones who do not develop latent heat. The entropy changes continuously at the transition.

1.6.2 Eherenfest classification

The *Eherenfest classification* is based on the behaviour of the derivatives of the thermodynamic potentials.

A phase transition is of order n if all the (n-1) derivatives are continuous and the n^{th} derivative displays a finite discontinuity.

Example 4

For instance, a first order transition in which $S = -(\partial G/\partial T)_P$ has finite discontinuity.

Remark. There are first order transitions where S is continuous (no latent heat), but ρ is discontinuous $(v = (\partial G/\partial P)_T)$.

Example 5

Second order transition. The specific heat displays a finite jump, see Figure 1.17c in the conductor-superconductor transition.

Another example is a second order transition but with divergence. Consider the fluid-superfluid transition (or λ transition) of the He₄ (Figure 1.17d).

Remark. λ transition: a second-order or higher-order transition, in which the heat capacity shows either a discontinuity (second-order) or a vertex (higher-order) at the transition temperature. It

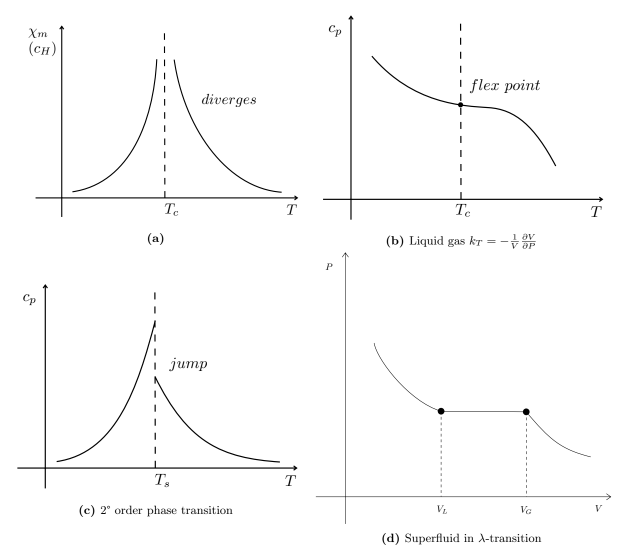


Figure 1.17: Plots of response functions.

is so named because the shape of the specific heat versus temperature curve resembles the Greek letter λ .

1.6.3 Modern classification

A phase transition is of the **first order** if exists a finite discontinuity in either one or more partial derivatives of the thermodynamic potentials. Instead, if the first derivatives are all continuous, but the second are either discontinuous, or infinite, one talks of **continuous transitions**. A critical point is a continuous transition.

1.7 Critical exponents

At the critical point response functions may diverge. How are these divergence? In general, when you are close to T_c , there are singularities. Now, we can ask, how the curve diverges? What is the behaviour close to the critical point? Power law, so which are the values of these critical exponents?

1.7.1 Divergence of the response functions at the critical point

While at the critical point the order parameter goes to zero continuously as $T \to T_c^-$, the response function may develop divergences.

Example 6

In a fluid system since at $T = T_c$ the curve P = P(V) develops an horizontal flex (Figure 1.18), we have $k_T = -\frac{1}{V} \left(\frac{\partial V}{\partial P} \right)_T \to \infty$. Similarly, in a magnetic since the curve is like Figure 1.16, we have $\chi_T = \left(\frac{\partial M}{\partial H} \right)_T \xrightarrow{T \to T_c} \infty$.

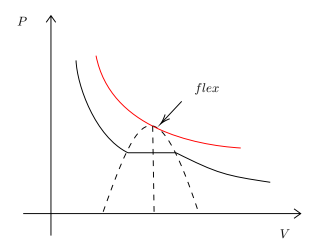


Figure 1.18: (V,T) projection.

1.7.2 Critical exponents definition

The notion of *critical exponent* describes the behaviour of the order parameter and the response functions in proximity of the critical point. In order to answer to these questions, let us define:

Definition 1: Critical Exponent, or Scale Exponent

Let us define the adimensional parameter measuring the distance from the critical point $t \equiv \frac{T - T_c}{T_c}$, the *critical exponent* λ associated to the function F(t) is defined as:

$$\lambda_{\pm} = \lim_{t \to 0^{\pm}} \frac{\ln |F(t)|}{\ln |t|} \tag{1.19}$$

We note that it behaves like a power law. One can also write the *power law*:

$$F(t) \stackrel{t \to 0^{\pm}}{\sim} |t|^{\lambda_{\pm}} \tag{1.20}$$

More generally, for $t \ll 1$:

$$F(t) = A|t|^{\lambda_{\pm}} (1 + bt^{\lambda_1} + \dots), \quad \lambda_1 > 0$$
 (1.21)

where all other terms are less important.

Definition 2: Thermodynamic Critical Exponents

- Exponent β : tells how the order parameter goes to zero. Consider Figure 1.19a, we have $|M|^{t\to 0^-}$ $(-t)^{\beta}$. No sense in going from above $(t\to 0^+)$ where it stays 0.
- Exponent γ_{\pm} (susceptibility): related to the response function. Consider Figure 1.19b, we have $\chi_T \stackrel{t \to 0^{\pm}}{\sim} |t|^{-\gamma_{\pm}}$. In principle, the value of γ can depend on the sign of t i.e. $\gamma^+ \neq \gamma^-$, but they are the same in reality and we have $\gamma^+ = \gamma^- = \gamma$.
- Exponent α_{\pm} : how specific heat diverges (second order derivative in respect of T). For instance see Figure 1.19c, we have $c_H \sim |t|^{-\alpha_{\pm}}$.
- Exponent δ : in this case one consider the isotherm $T = T_c$ and look for the behaviour of M at the critical point at small H (or viceversa). The result is $M \sim H^{1/\delta}$. In Figure 1.19d, $H \sim |M|^{\delta} \text{sign}(M)$.

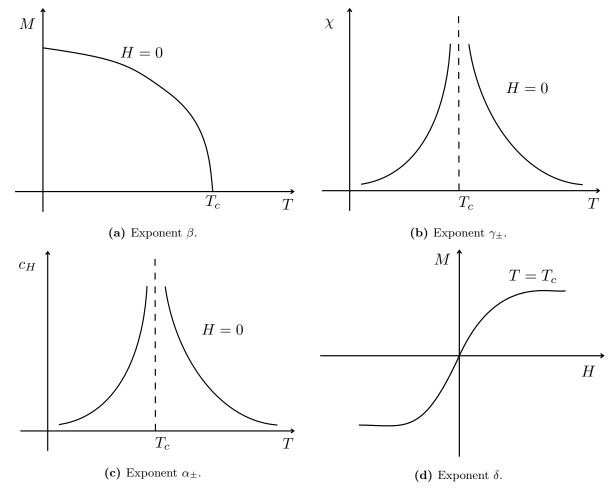


Figure 1.19

Remark. In compiling Table 1.1 and 1.2 we have made the as yet totally unjustified assumption that the critical exponent associated with a given thermodynamic variable is the same as $T \to T_c$ from above or below.

Critical exponents depend on very few parameters and are not system-specific: 1 - symmetries of the system 2 - dimension of the system 3 - range of interaction T_c instead does depend on specific chemistry, is not as universal as critical exponent.

| Zero-field specific heat | $C_H \sim t ^{-\alpha}$ |
|--------------------------------------|--|
| Zero-field magnetization | $M \sim (-t)^{\beta}$ |
| Zero-field isothermal susceptibility | $\chi_T \sim t ^{-\gamma}$ |
| Critical isotherm $(t=0)$ | $H \sim M ^{\delta} \mathrm{sign}(M)$ |
| Correlation length | $\xi \sim t ^{-\nu}$ |
| Pair correlation function at T_c | $G(ec{\mathbf{r}}) \sim rac{1}{r^{d-2+\eta}}$ |

Table 1.1: Definitions of the most commonly used critical exponents for a magnetic system [?].

| Specific heat at constant volume V_c | $C_V \sim t ^{-\alpha}$ |
|--|--|
| Liquid-gas density difference | $(ho_l- ho_g)\sim (-t)^{eta}$ |
| Isothermal compressibility | $k_T \sim t ^{-\gamma}$ |
| Critical isotherm $(t=0)$ | $P - P_c \sim \rho_l - \rho_g ^{\delta} \operatorname{sign}(\rho_l - \rho_g)$ |
| Correlation length | $\xi \sim t ^{- u}$ |
| Pair correlation function at T_c | $G(ec{\mathbf{r}}) \sim rac{1}{r^{d-2+\eta}}$ |

Table 1.2: Definitions of the most commonly used critical exponents for a fluid system [?].

1.7.3 Law of the corresponding states (Experimental estimates of order parameters)

The system displays correlation at very long distance, these goes to the size of the system when $T \to T_c$. We are talking about long range correlation. The *correlation function* is $\xi \sim t^{-\nu}$. For instance, consider a polymer as in Figure 1.20a.

Having defined the critical exponents, we need to justify why they are interesting and why they are more interesting than the critical temperature T_c itself. It turns out that, whereas T_c depends sensitively on the details of the interatomic interactions, the critical exponents are to a large degree *universal* depending only on a few fundamental parameters.

To summurize, the critical exponents are more interesting than T_c since their values do not depend on microscopic details, but only on few parameters such as the space dimension d and the symmetry of the system.

One of the first experimental evidence of this universality was given by the work of Guggenheim on the coexistence curves of g different fluids: A, Kn, χ_e , Ne, N_2 , CO_2 and O_2 . By plotting T/T_c versus ρ/ρ_c (Figure 1.20b) he found that all the data collapse on the same curve, i.e. different sets of data fit the same function. Moreover for $t \to 0$:

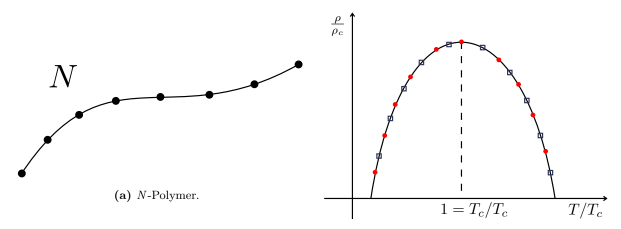
$$(\rho_l - \rho_c) \sim (-t)^{\beta}$$

and $\beta \sim 1/3 \approx 0.335$. Therefore, close to the critical point all the data lie on the same curve and hence can be described by the same exponent β . A further test of universality is to compare this value to that obtained for a phase transition in a completely different system with a scalar order parameter. For instance, if we do the same for a string ferromagnetic the result is $\beta = 1/3$ too.

Remark. The law of corresponding states gives a universal liquid-gas coexistence curve.

1.7.4 Thermodynamic inequalities between critical exponents

It is possible to obtain several rigorous inequalities between the critical exponents. The easiest to prove is due to Rushbrooke.



(b) Coexistence curve of different fluids plotted in reduced variables.

Figure 1.20

Rushbrocke inequality

It follows from the well known thermodynamic relation between the specific heats at constant field and constant magnetization. Remember the relation between response functions: For a fluid:

$$k_T(c_p - c_v) = Tv\alpha^2 = Tv\frac{1}{v^2} \left(\frac{\partial v}{\partial T}\right)_P^2 = T\frac{1}{v} \left(\frac{\partial v}{\partial T}\right)_P^2$$

For magnetic systems one has

$$\chi_T(c_H - c_M) = T \underbrace{\left(\frac{\partial M}{\partial T}\right)_H^2}_{\geq 0}$$

From thermodynamic stability we have $c_M \geq 0, c_H \geq 0, \chi_T \geq 0$. Hence, from the previous relation we have

$$c_H = \frac{T}{\chi_T} \left(\frac{\partial M}{\partial T}\right)_H^2 + \underbrace{c_M}_{\geq 0}$$

which implies

$$c_H \ge \frac{T}{\chi_T} \left(\frac{\partial M}{\partial T}\right)_H^2$$
 (1.22)

On the other hand, for $T \to T_c^ (t \to 0^-)$ and H = 0 (zero field) we have

$$\begin{cases} c_H \sim (-t)^{-\alpha} \\ \chi_T \sim (-t)^{-\gamma} \\ M \sim (-t)^{\beta} \end{cases}$$

that implies

$$\left(\frac{\partial M}{\partial T}\right)_{H=0} \sim (-t)^{\beta-1}$$

Since the inequality (1.22) is valid for all temperature T, it follows that can only be obeyed if

$$B(T_c - T)^{-\alpha} \ge B' T \frac{[(T_c - T)^{\beta - 1}]^2}{(T_c - T)^{-\gamma}}$$

with B, B' > 0. Take the limit $T \to T_c^-$, we have:

$$\lim_{T \to T_c^-} (T_c - T)^{2 - \alpha - 2\beta - \gamma} \ge \frac{B'T}{B} > 0$$

Since the left hand side must be strictly greater than zero, we have the RushBrook inequality:

$$\alpha + 2\beta + \gamma \ge 2 \tag{1.23}$$

Griffith inequality

The Griffith inequality is obtained from the convexity property (in T and V) of the Helmolds free energy and from $A \sim t^{2-\alpha}$ (as $t \to 0^-$ deduced from $c_P = t^{-\alpha}$):

$$\Rightarrow \alpha + \beta(1+\delta) \ge 2 \tag{1.24}$$

We have introduced two very new ideas, universality and inequalities between the critical exponents, which appear to hold as equalities.

In the intervening chapters, we look at models of systems which undergo phase transitions and how to calculate their critical exponents and other properties.