# Thermodynamics

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## 1 Zeroth law

Zeroth law

temperature scales

## 2 Definitions

- 2.1 Ideal gas state equation
- 2.2 Systems and processes

- 3 Isothermal expansion
- 4 First law

## 5 Heat capacities

include derivations

### 6 Enthalpy

### 6.1 State function enthalpy

We define enthalpy as:

$$H = U + PV$$
.

This state function greatly simplifies our  $C_P$  definition:

$$C_P = \left(\frac{\mathrm{d}Q}{\mathrm{d}T}\right)_P = \left(\frac{\partial H}{\partial T}\right)_P$$

and its differential is:

$$dH = dU + VdP + PdV.$$

- 6.2 Latent heats
- 6.3 Constant pressure expansion
- 6.4 Filling empty tank
- 6.5 Chemical reactions

## 7 Heat engines

include hurricanes application

### 8 Entropy

### 8.1 State function entropy

**Entropy** is defined as:

$$S = \frac{Q}{T},$$

where Q is <u>heat from reservoir</u> and T the reservoir temperature.

Entropy has units  $JK^{-1}$ , and if we have a **reversible cyclic** heat engine:

$$\mathrm{d}S = \frac{\mathrm{d}Q}{T}.$$

This is the differential form of entropy. Applying the first law under again **reversible** conditions gives:

$$dU = TdS - PdV$$

The entropy change of a system is:

$$\Delta S = \int_{(1)}^{(2)} \frac{dU}{T} + \int_{(1)}^{(2)} P dV$$
$$= \int_{(1)}^{(2)} \frac{dQ}{T},$$

for a reversible path.

### 8.2 Entropy of ideal gases

Let's now assume an ideal, monoatomic gas. Then its heat capacity becomes:

$$C_V = \frac{\mathrm{d}U}{\mathrm{d}T}.$$

So  $C_V dT = T dS - P dV$  and  $C_V = \frac{3}{2} nR$ . Combining these:

$$dS = nR \left( \frac{3}{2} \frac{dT}{T} + \frac{dV}{V} \right)$$

which can be discretised to find  $\Delta S$ . Rearranging we get  $T dS = C_V dT + P dV$ . Integrating and using identity  $C_P - C_V = nR$  we get:

$$\Delta S = nc_V \ln \left(\frac{P_2}{P_1}\right) + nc_P \ln \left(\frac{V_2}{V_1}\right)$$

for here n is moles and the heat capacities are in molar quantities.

### 8.3 Entropy of mixing

Consider a box with separator, containing  $n_A$  moles at  $V_A$  of gas A on one side and  $n_B$  moles at  $V_B$  of gas B on the other. We set  $\Delta T = 0$ ,  $\Delta P = 0$  and proceed to  $\underline{\text{mix}}$  gases A and B.

The entropy of mixing is:

$$\begin{split} \Delta S &= \Delta S_A + \Delta S_B \\ &= n_i R \ln \left( \frac{V_f}{V_i} \right) \\ &= n_A R \ln \left( \frac{V_A + V_B}{V_A} \right) + n_B R \ln \left( \frac{V_A + V_B}{V_B} \right), \end{split}$$

Since P and T are fixed, the ideal state equation PV = nRT implies that:

$$\frac{V_A + V_B}{V_A} = \frac{n_A + n_B}{n_A}$$

and so we define **inverse mole fractions** as:

$$x_A = \frac{n_A}{n_A + n_B}$$

and

$$x_B = \frac{n_B}{n_A + n_B}.$$

After substituting and dividing through by  $n_A + n_B$  we find the **molar specific** entropy of mixing:

$$\Delta s_{mix} = -R(x_A \ln x_A + x_B \ln x_B).$$

#### 8.4 Measuring entropy

Practically,  $C_P$  is much easier to measure than  $C_V$ . Recall definition:

$$C_P = \frac{\mathrm{d}Q_p}{\mathrm{d}T} = \left(\frac{\mathrm{d}H}{\mathrm{d}T}\right)_p$$

where state function H is the enthalpy. Since H = U + PV:

$$dH = TdS + VdP$$

and dividing by dT:

$$\frac{\mathrm{d}H}{\mathrm{d}T} = T\frac{\mathrm{d}S}{\mathrm{d}T} + V\frac{\mathrm{d}P}{\mathrm{d}T}.$$

If  $\Delta P = 0$  then:

$$C_P(T) = T \left(\frac{\mathrm{d}S}{\mathrm{d}T}\right)_P.$$

Rearranging and integrating gives us the change in entropy:

$$\Delta S = \int_{(1)}^{(2)} \frac{C_P(T)}{T} dT.$$

## 9 Second law

### 9.1 Reversibility

### 9.2 Second law

include spontaneous process sections 12 and 13

## 10 Helmholtz and Gibbs free energy

include chemical reactions and colloidal particles sections 14 and 15

### 11 Maxwell relations

#### 11.1 Derivation

So from previous sections we have the following four differential relations<sup>1</sup>:

$$dU = TdS - PdV,$$

$$dH = TdS + VdP,$$

$$dF = -SdT - PdV,$$

$$dG = -SdT + VdP.$$

For equation of state f(P, V, T) = 0, the **Maxwell relations** are the following:

$$\begin{split} \left(\frac{\partial T}{\partial V}\right)_S &= -\left(\frac{\partial P}{\partial S}\right)_V, \\ \left(\frac{\partial T}{\partial P}\right)_S &= \left(\frac{\partial V}{\partial S}\right)_P, \\ \left(\frac{\partial S}{\partial V}\right)_T &= \left(\frac{\partial P}{\partial T}\right)_V, \\ -\left(\frac{\partial S}{\partial P}\right)_T &= \left(\frac{\partial V}{\partial T}\right)_P. \end{split}$$

Notice how each equation relates entropy S to measurable quantities.

These relations are derived by taking **total differentials** and then applying **Clairaut's theorem**. We are only going to do the first relation:

$$\begin{split} \mathrm{d}U &= T\mathrm{d}S - P\mathrm{d}V \\ &= \left(\frac{\partial U}{\partial S}\right)_V \mathrm{d}S + \left(\frac{\partial U}{\partial V}\right)_S \mathrm{d}V \end{split}$$

Therefore we have that:

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

and

$$P = -\left(\frac{\partial U}{\partial V}\right)_S.$$

Taking partials again and equating gives us the first relation.

 $<sup>^1\</sup>mathrm{Here}\ U$  is the internal energy, H is enthalpy, F is the Helmholtz free energy and G is the Gibbs free energy.

### 11.2 Applications

We can define **isobaric expansivity**  $\beta$  with units  $K^{-1}$  as:

$$\beta = \frac{1}{V} \left( \frac{\partial V}{\partial T} \right)_P$$

and isothermal compressibility  $\kappa_T$  with units  $Pa^{-1}$  as:

$$\kappa_T = -\frac{1}{V} \left( \frac{\partial V}{\partial P} \right)_T.$$

These may be thought of as volume change per physical unit change.

The Maxwell relations can be used to find entropy. For example:

$$\Delta S = \int_{(1)}^{(2)} dS$$
$$= -\int_{(1)}^{(2)} \left(\frac{\partial V}{\partial T}\right)_{P} dP.$$

#### 11.2.1 Ideal gas internal energy

Now we can show why U = U(T) if we have an ideal gas. Beginning with:

$$dU = TdS - PdV$$

and dividing through by dV gives:

$$\begin{split} \left(\frac{\partial U}{\partial V}\right)_T &= T \left(\frac{\partial S}{\partial V}\right)_T - P \\ &= T \left(\frac{\partial P}{\partial T}\right)_V - P \end{split}$$

via a Maxwell relation. Finally our ideal gas assumption implies that:

$$P = \frac{nRT}{V}$$

and substituting this into the previous expression results in:

$$\left(\frac{\partial U}{\partial V}\right)_T = 0$$

which means that U = U(T).

#### 11.2.2 Difference in heat capacities

We begin with the following expression:

$$C_{P} = C_{V} + \left(P + \left(\frac{\partial U}{\partial V}\right)_{T}\right) \left(\frac{\partial V}{\partial T}\right)_{P}$$

$$= C_{V} + T \left(\frac{\partial P}{\partial T}\right)_{V} \left(\frac{\partial V}{\partial T}\right)_{P}$$

$$= C_{V} + T \left(\frac{\partial P}{\partial T}\right)_{V} \beta V.$$

$$\therefore C_{P} - C_{V} = \beta T V \left(\frac{\partial P}{\partial T}\right)_{V}$$

Then using the following identities:

$$\begin{split} \left(\frac{\partial P}{\partial T}\right)_{V} \left(\frac{\partial V}{\partial P}\right)_{T} \left(\frac{\partial T}{\partial V}\right)_{P} &= -1 \\ \frac{1}{\left(\frac{\partial V}{\partial P}\right)_{T}} &= \left(\frac{\partial P}{\partial V}\right)_{T} \end{split}$$

we get that the difference in heat capacities is:

$$C_P - C_V = \frac{\beta^2}{\kappa_T} TV$$

where  $\beta$  is the **isobaric expansivity** and  $\kappa_T$  is the **isothermal compressibility**. Furthermore it is clear that  $C_P - C_V > 0$  for all cases.

#### 11.2.3 Liquefying gases

## 12 Phase transitions

### 12.1 PT diagrams

phase transition diagrams (PT) supercritical points phase boundaries triple point clausius clapeyron equation

### 12.2 Van der Waals state equation

The Van der Waals equation is a modification of the ideal gas equation:

$$\left(P + \frac{an^2}{V^2}\right)\left(V - nb\right) = nRT$$

and defines the volume V occupied by n moles of gas at pressure P and temperature T. The <u>molar</u> version of this is the following:

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

where we define the <u>molar volume</u> as  $v = \frac{V}{n}$ .

#### 12.2.1 Helmholtz and Gibbs free energies

Since we have that dF = -SdT - PdV at fixed temperatures:

$$\left(\frac{\partial F}{\partial V}\right)_T = -P$$

and integrating the molar version of this:

$$f = -\int P \, dv$$
$$= \tau(T) - RT \ln(v - b) - \frac{a}{v}$$

where we have substituted the Van der Waals equation.

For the Gibbs free energy because f = u - TS we then have that:

$$g = u + Pv - TS$$

$$= f + Pv$$

$$= \tau(T) - RT \ln(v - b) - \frac{a}{v} + Pv$$

## 12.3 PV diagrams

## 13 Chemical potentials

## 14 Third law