The energy of the gas

The energy of the gas U is the sum of the Kinetic and potential energies

U = U (T, V; N)

As before we will regard N as being fixed,

And the dependence on volume simply reflects

the change in density. Specifically the energy

per volume takes the form:

U = f(T, n) f = some function

If I double the system size double the number of particles at fixed temperature, the density remains the same and energy per volume is the same. At small density we expand in density

 $U = n f(T) + n^2 f(T) + n^3 f(T) + ...$

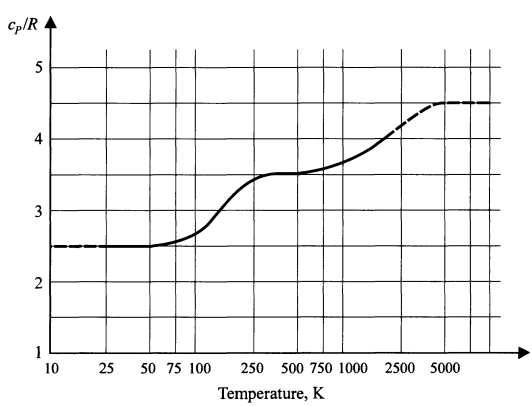
So the low density (ideal gas) expansion takes the form with n = N/V

U(T,V) = N(f(T) + Nf(T) + ...)

· First lets take first term. This is an ideal gas	
U(T,V)= Nfo(+) Ideal gas only	
For a mono-atomic gas, $U = N \stackrel{?}{=} k_B T$; for a diatomic gas $U = N \stackrel{?}{=} k_B T$, specifying $f(T)$.	
· What about more complicated gasses, such as H2O	?
stretch mode 7 H	
It can rotate the H's can vibrate. All of that complexity is incorporated in f(T). The molecules are indepent of each other but the system still needs to decide how to partition energy	
· We note that for an ideal gas	
And The molecules are so far apart they don't talk to each other	
$C_V = \left(\frac{\partial U}{\partial T}\right) = N f'(T)$ Ideal gas only	
We will want to figure out how to coloulate	

fo(T), which is directly related to measured Cp+Cv
· For non ideal gasses: non-ideal gas
DU + 0 and this reflects the potential energy between the
Wibice Wes,
$\frac{\partial U}{\partial V} \approx -\frac{(N)^2 f_1(T)}{V} \Rightarrow \text{for dilute gasses}. This term clearly involves}$
pairs of particles and Scales as the (density) ² , we won't calculate reflecting the interactions $f_{i}(T)$ in this course.
we work calculate reflecting the interactions $f_i(T)$ in this course.
Finally note that the curve Cp is directly related to fott). For one mole R
$C_{V}^{lmol} = N_{A} k_{B} \left(\frac{1}{k_{B}} \frac{df_{o}}{dT} \right) = R \left[\frac{1}{k_{B}} \frac{df_{o}}{dT} \right]$
$C_p^{lmol} = C_v^{lmol} + R = R \left[\frac{1}{K_B} \frac{df_0}{d\tau} + 1 \right]$ this what is plotted
on Cp/R graph. So we see that measurements about specific heats
are directly telling us about U (or fo(T)). (Reconsider what Cp for Hydrogen is telling you looking
at the Slide) below

Specific Heat of H_2



The specific heats reflect the internal energy of the system

$$U = N f_0(T)$$

We have for one mole

$$C_V = (\partial U/\partial T)_V = R\left(\frac{1}{k_B}\frac{df_0}{dT}\right)$$

and
$$C_p = C_V + R$$