

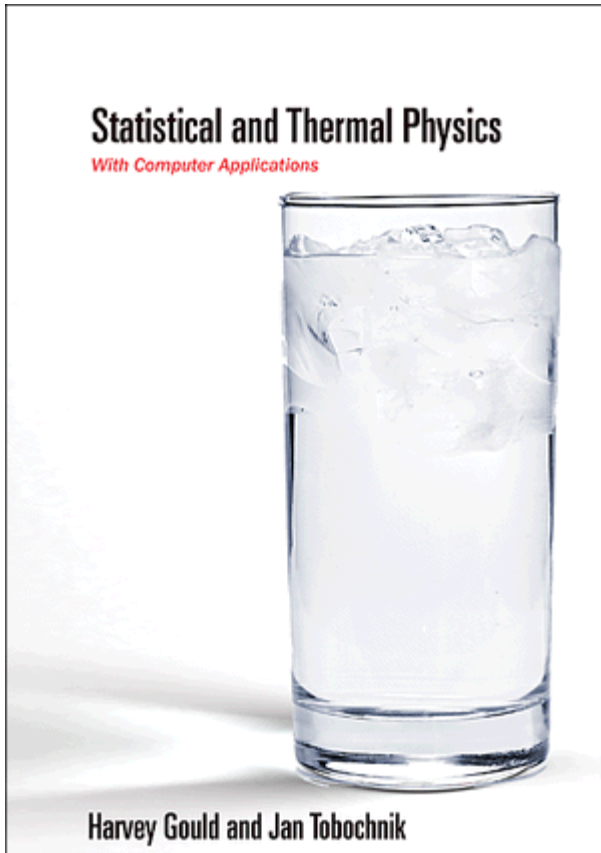
# **PHY 341/641**

# **Thermodynamics and Statistical Physics**

## **Lecture 2**

1. Continued discussion of microscopic models (Chapter 1)
  - a. Notion of equilibrium in statistical mechanics/thermodynamics
  - b. Macrostates/microstates
2. Introduction to thermodynamics (Chapter 2)
  - a. Definition of “the system”
  - b. Thermodynamic variables ( $T$ ,  $P$ ,  $V$ ,  $N$ , ...)
  - c. First law of thermodynamics

Review from last time --  
Chapter 1 – From Microscopic to Macrosopic Behavior



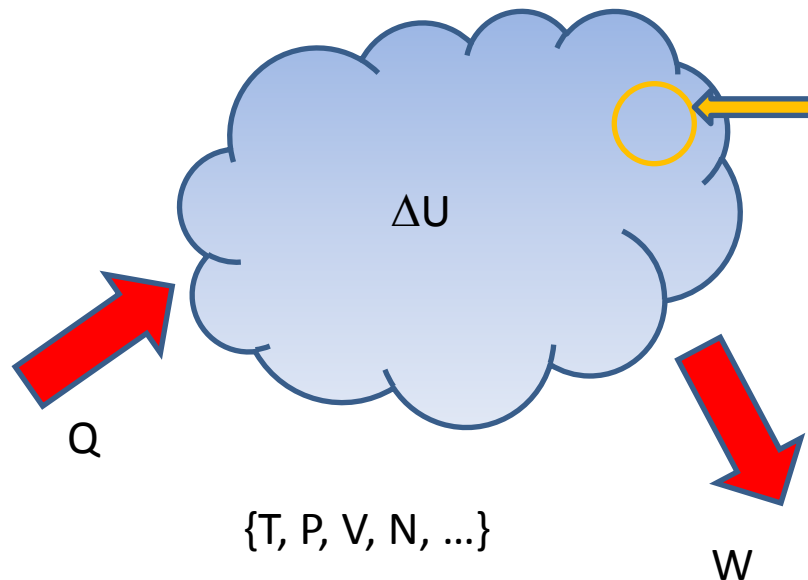
Assignment: Read Chapter 1 (quickly) during this week and checkout some of the corresponding simulations (HW 1 and HW 2) due Monday 1/23.

“The purpose of this introductory [material] is to whet your appetite... “ The chapter introduces a lot of the concepts that we will use (more carefully) throughout the course.

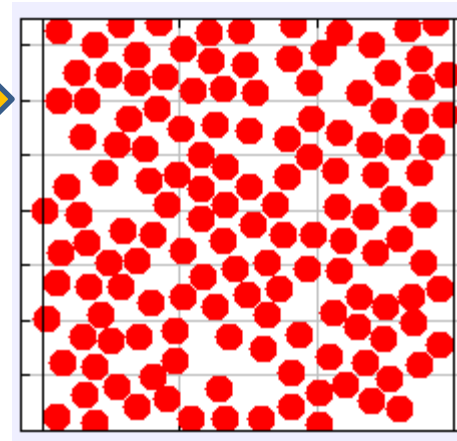
What will you learn?

# Energy Analysis

Macroscopic picture

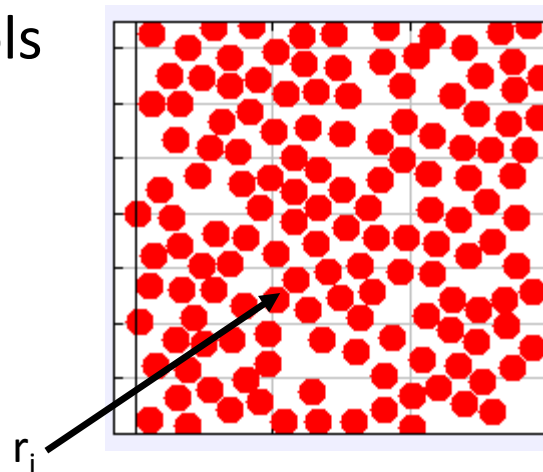


Microscopic picture



## Comment on simulation tools Molecular dynamics

$$m_i \frac{d^2 \mathbf{r}_i}{dt^2} = \mathbf{F}_i$$



$$\mathbf{F}_i = -\nabla_i \sum_{j \neq i} u_{pair}(|\mathbf{r}_i - \mathbf{r}_j|)$$

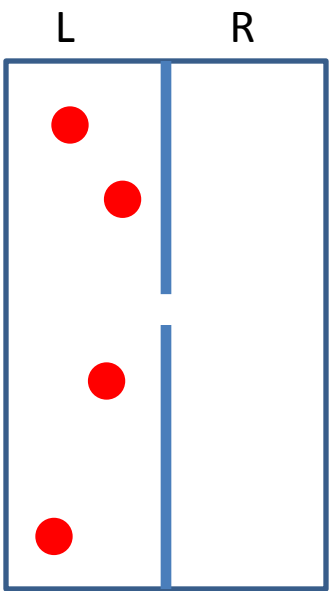
Example model pair potential (Lennard-Jones):

$$u_{LJ}(r) = 4\epsilon \left[ \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^6 \right]$$

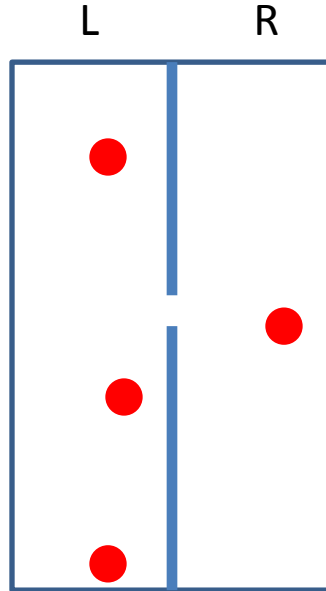
**For an “ideal gas”--**

$$m_i \frac{d^2 \mathbf{r}_i}{dt^2} = \mathbf{F}_i \equiv 0$$

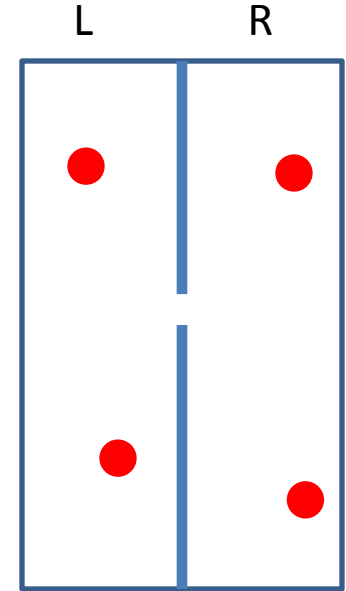
Modeling of a dilute gas of non-interacting particles:



$t = 0$



$t > 0$



$t \gg 0$

$$P_{L \rightarrow R} = \frac{n}{N}$$

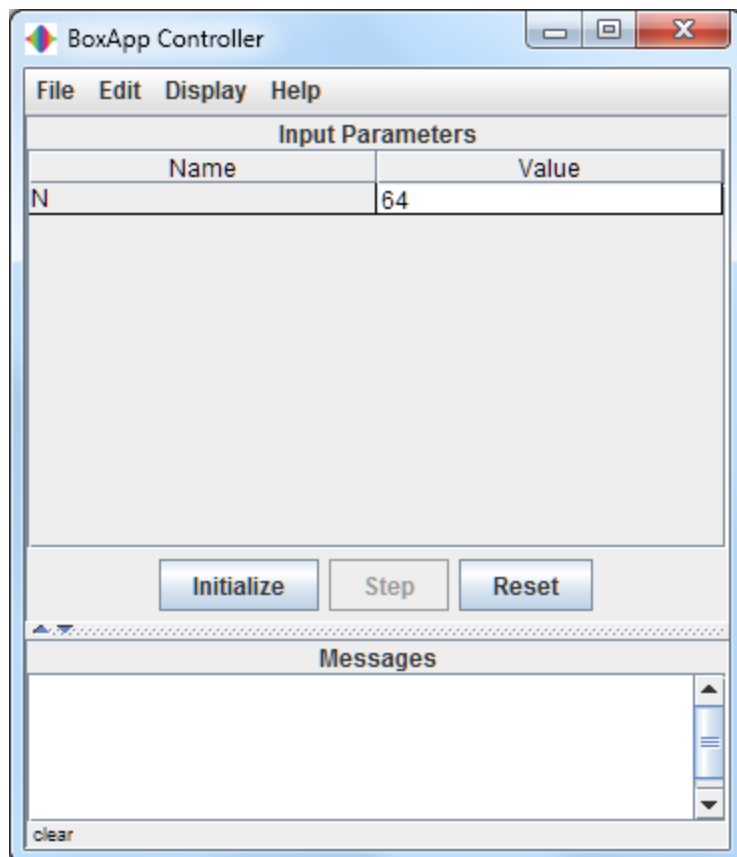
$$P_{R \rightarrow L} = \frac{N - n}{N}$$

## Enumeration of possibilities for N=4

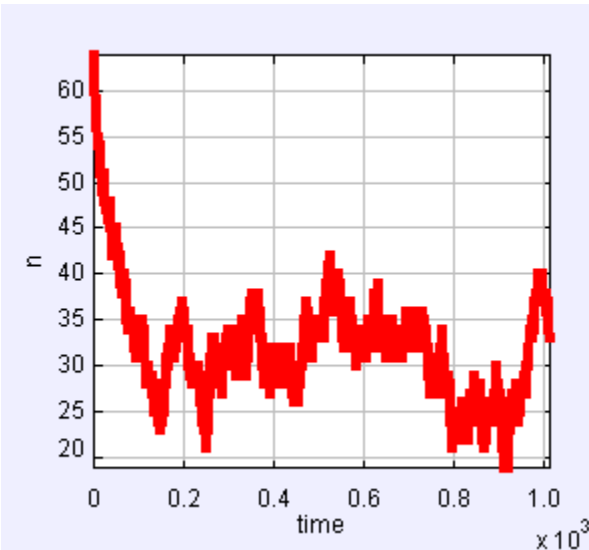
Microstates	n	$P_{L \rightarrow R}$
L L L L	4	1/16
L L L R		
L L R L	3	4/16
L R L L		
R L L L		
L L R R		
L R R L		
R R L L	2	6/16
L R L R		
R L R L		
R L L R		
R R R L		
R R L R	1	4/16
R L R R		
L R R R		
R R R R	0	1/16

Simulation available from: [http://www.compadre.org/STP/stp\\_ApproachToEquilibrium.jar](http://www.compadre.org/STP/stp_ApproachToEquilibrium.jar)

*Note: in order to easily control the simulation, you need to use:  
Display → Switch GUI*



$N=64, t \approx 1000$   $\langle n \rangle = 32.08, \langle n^2 \rangle = 1,067.13, \sigma^2 = 38.12, \sigma/\langle n \rangle = 0.19$

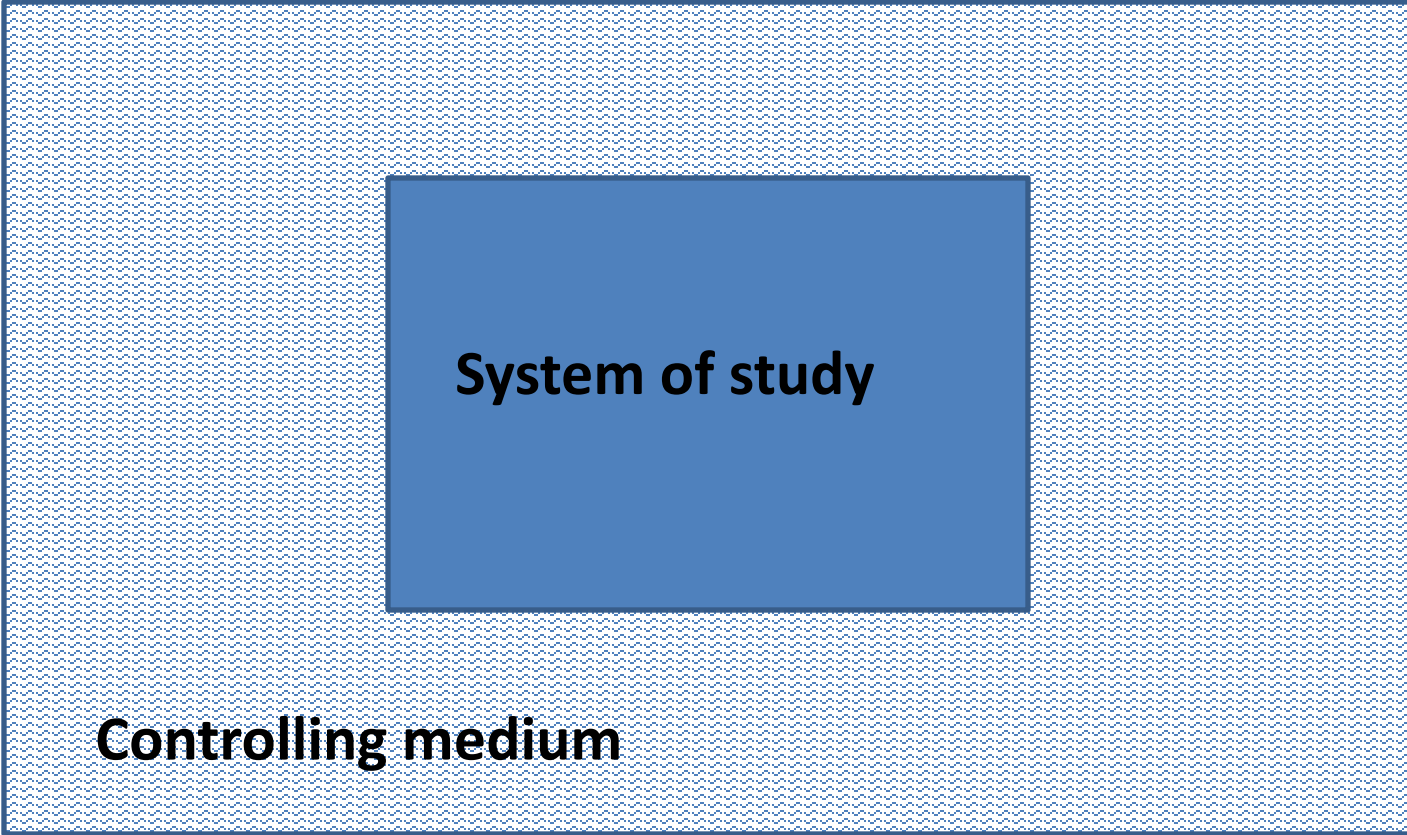


$N=256, t \approx 1000$   $\langle n \rangle = 149.57, \langle n^2 \rangle = 23,086.17, \sigma^2 = 714.56, \sigma/\langle n \rangle = 0.18$



# **Macroscopic viewpoint – thermodynamics**

(start reading Chapter 2)



**System of study**

**Controlling medium**

# Variables of thermodynamics

- **Temperature**
  - ❖  $T$
  - ❖ *Zeroth law of thermodynamics: Two systems in thermal equilibrium with a third system are in thermal equilibrium with each other.*
- **Pressure**
  - ❖  $P=F/A$
- **Volume**
  - ❖  $V$
- **Number of particles**
  - ❖  $N$

The relationships between these variables depends on the “equation of state” of the system.

## Examples of “Equations of State”

Ideal gas equation of state :

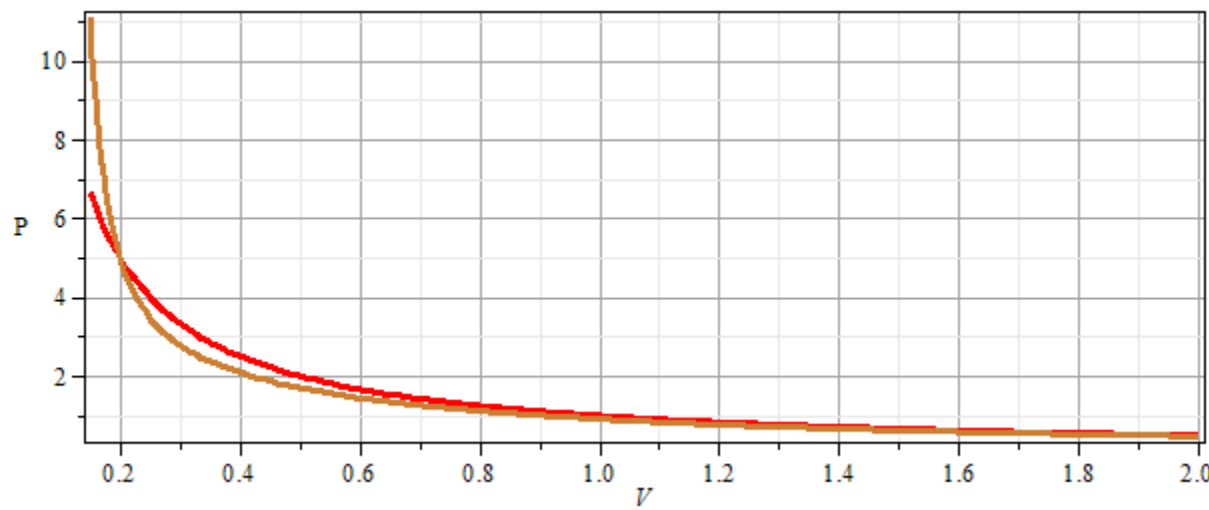
$$PV = Nk_B T \quad k_B = 1.38 \times 10^{-23} \text{ J/K}$$

van der Waals equation of state :

$$\left( P + \frac{N^2}{V^2} a \right) (V - Nb) = Nk_B T$$

## Comparison of

- Ideal gas equation
- van der Waals gas equation



In general the “state” of a system will depend on the thermodynamic variables. For example, the internal energy:

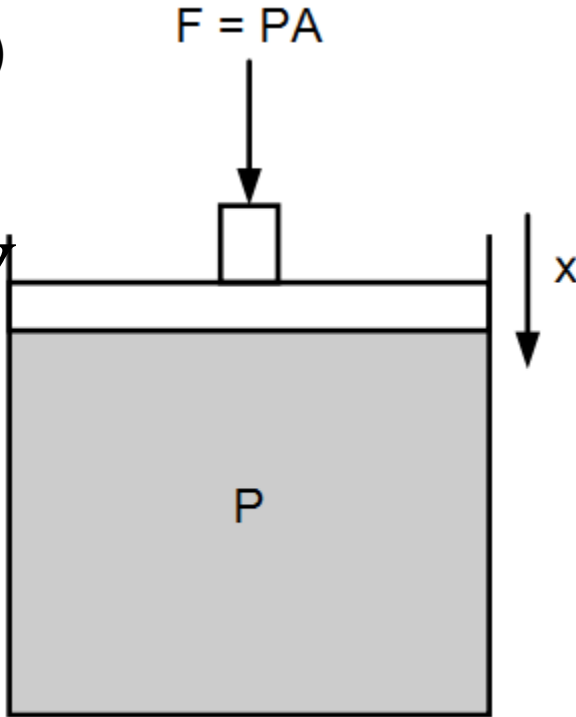
$$U = U(N, T, P, V)$$

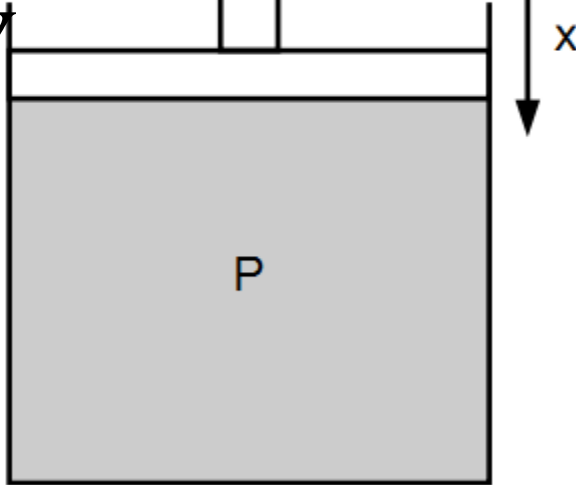
Thermodynamic “processes” → change the state of the system

$$U_1(N_1, T_1, P_1, V_1) \Rightarrow U_2(N_2, T_2, P_2, V_2)$$

Thermodynamic process -- work (performed ON the system)

$$F = PA$$


$$\downarrow$$



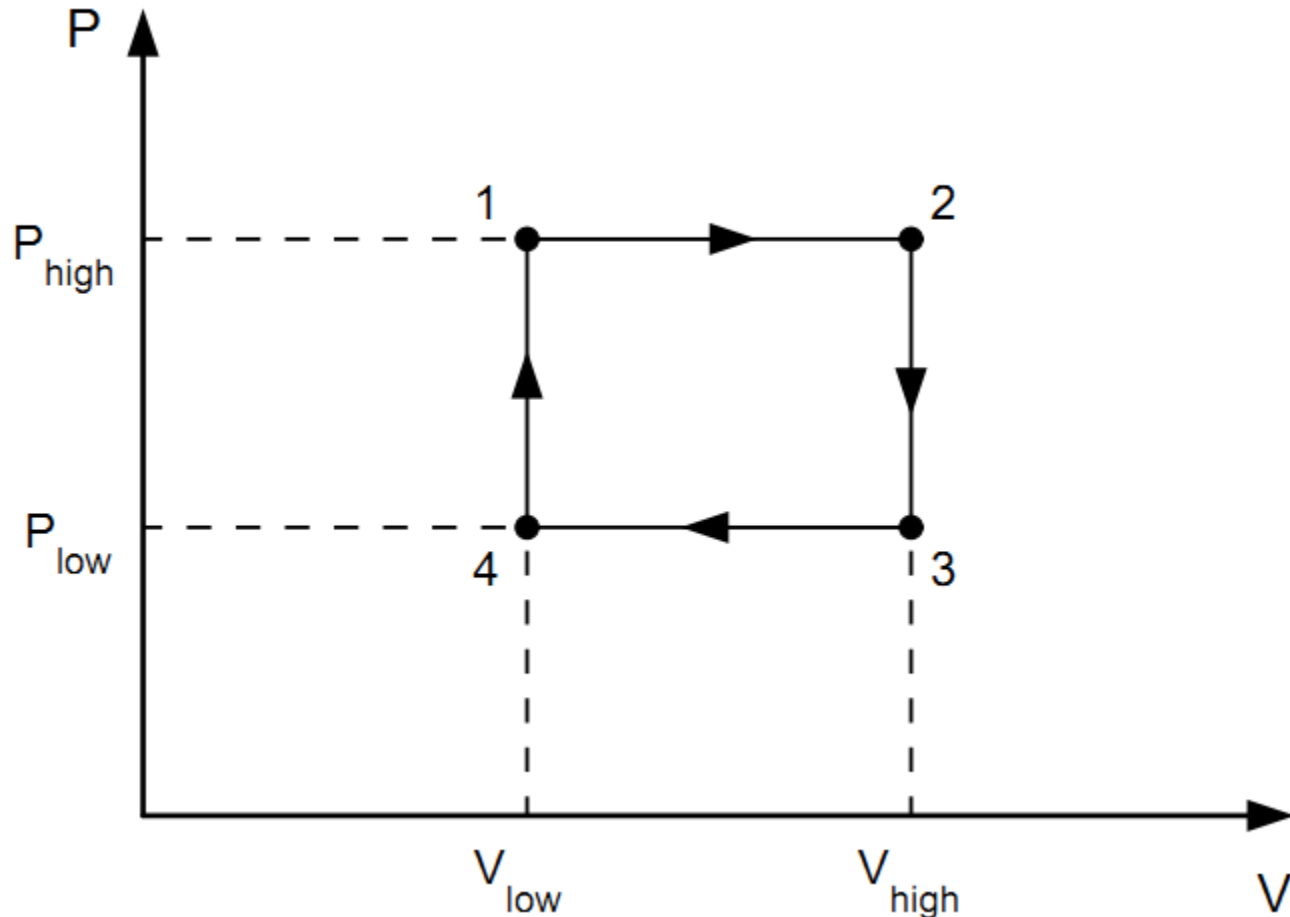
$$dW = -Fdx = -PAdx = -PdV$$

$$W_{1 \rightarrow 2} = - \int_{V_1}^{V_2} P(T, V) dV$$

For an ideal gas at constant T :

$$W_{1 \rightarrow 2} = - \int_{V_1}^{V_2} P(T, V) dV = - \int_{V_1}^{V_2} \frac{NkT}{V} dV = -NkT \ln \frac{V_2}{V_1}$$

Work performed during a cyclic process:



$$W_{net} = -(P_{high} - P_{low})(V_{high} - V_{low})$$

Thermodynamic process -- heat (added TO the system)

$Q$

For heat added TO the system:  $Q > 0$

For heat withdrawn FROM the system:  $Q < 0$

## First law of thermodynamics

$$U_2 - U_1 \equiv \Delta U = W + Q$$

Example: Adiabatic expansion  $V_1 \rightarrow V_2$  for an ideal gas system

$$U_2 - U_1 \equiv \Delta U = W + Q = W$$

$$PV = NkT$$

To be continued.....