

### Physics 1

Voronkov Vladimir Vasilyevich

#### Lecture 6

- Heat flow and the first law of thermodynamics.
- Kind of thermodynamic process.
   Adiabatic processes.

#### Heat

- When the temperature of a thermal system in contact with a neighboring system changes, we say that there has been a **heat flow** into or out of the system.
- An energy unit related to thermal processes is the calorie (cal), which is defined as the amount of energy transfer necessary to raise the temperature of 1 gram of water by 1 degree (from 14.5°C to 15.5°C).

# Mechanical equivalent of heat

Mechanical energy is not conserserved in the presence of nonconservative forces. It transforms into internal energy. For example, friction produces heating

1 cal = 4.186 J

# Specific heat capacity

The heat capacity C of a particular sample of a substance is defined as the amount of energy needed to raise the temperature of that sample by 1 °C.

$$C=Q/\Delta T$$

 The specific heat capacity c of a substance is the heat capacity per unit mass.

$$c=C/m=Q/(m\Delta T)$$

Specific heat is essentially a measure of how thermally insensitive a substance is to the addition of energy. The greater a material's specific heat, the more energy must be added to a given mass of the material to cause a particular temperature change.

# Energy transfer and specific heat capacity

From this definition, we can relate the energy Q transferred between a sample of mass m and specific heat capacity c of a material and its surroundings to a temperature change △T as

 $Q=mc \Delta T$ 

# Specific Heats of Some Substances at 25°C and Atmospheric Pressure

	Specific	heat c		Specific	heat c
Substance	J/kg·°C	cal/g·°C	Substance	J/kg·°C	cal/g·°C
Elemental solids Aluminum Beryllium Cadmium Copper Germanium Gold Iron Lead Silicon Silver	900 1 830 230 387 322 129 448 128 703 234	0.215 0.436 0.055 0.092 4 0.077 0.030 8 0.107 0.030 5 0.168 0.056	Other solids  Brass Glass Ice (-5°C) Marble Wood  Liquids Alcohol (ethyl) Mercury Water (15°C)  Gas  Steam (100°C)	380 837 2 090 860 1 700 2 400 140 4 186	0.092 0.200 0.50 0.21 0.41 0.58 0.033 1.00

# Dependence of specific heat capacity on temperature

Specific heat varies with temperature. For example, the specific heat of water varies by only about 1% from 0 c °C to 100 °C at atmospheric pressure. Usually such variations are negligible.

# Dependence of specific heat capacity on volume and pressure

Measured values of specific heats are found to depend on the conditions of the experiment. In general, measurements made in a constant pressure process are different from those made in a constant volume process. For solids and liquids, the difference between the two values is usually no greater than a few percent and is often neglected.

#### Phase transition

- It can be that transfer of energy does not result in a change in emperature. This is the case when the physical characteristics of the substance change from one form to another; such a change is called a phase change. Two common phase changes:
  - melting: from solid to liquid
  - boiling: from liquid to gas
  - change in the crystalline structure of a solid
- All such phase changes involve a change in internal energy but no change in temperature.
- The increase in internal energy in boiling, for example, is represented by the breaking of bonds between molecules in the liquid state; this bond breaking allows the molecules to move farther apart in the gaseous state, with a corresponding increase in intermolecular potential energy.

#### Latent heat

Quantitative measure of phase transition is latent heat L:

$$Q=\pm mL$$

- Latent heat of fusion  $L_f$  is the term used when the phase change is from solid to liquid,
- Latent heat of vaporization  $L_{\nu}$  is the term used when the phase change is from liquid to gas (the liquid "vaporizes vaporizes").

#### Latent Heats of Fusion and Vaporization

Substance	Melting Point (°C)	Latent Heat of Fusion (J/kg)	Boiling Point (°C)	Latent Heat of Vaporization ( J/kg)
Helium	-269.65	$5.23 \times 10^{3}$	-268.93	$2.09 \times 10^{4}$
Nitrogen	-209.97	$2.55 \times 10^{4}$	-195.81	$2.01 \times 10^{5}$
Oxygen	-218.79	$1.38 \times 10^{4}$	-182.97	$2.13 \times 10^{5}$
Ethyl alcohol	- 114	$1.04 \times 10^{5}$	78	$8.54 \times 10^{5}$
Water	0.00	$3.33 \times 10^{5}$	100.00	$2.26 \times 10^{6}$
Sulfur	119	$3.81 \times 10^{4}$	444.60	$3.26 \times 10^{5}$
Lead	327.3	$2.45 \times 10^{4}$	1 750	$8.70 \times 10^{5}$
Aluminum	660	$3.97 \times 10^{5}$	2 450	$1.14 \times 10^{7}$
Silver	960.80	$8.82 \times 10^{4}$	2 193	$2.33 \times 10^{6}$
Gold	1 063.00	$6.44 \times 10^4$	2 660	$1.58 \times 10^{6}$
Copper	1 083	$1.34 \times 10^{5}$	1 187	$5.06 \times 10^{6}$

# State variables - Thermodynamic process - Thermal equilibrium

We describe the state of a system using such variables as pressure, volume, temperature, and internal energy. These quantities are called state variables. Macroscopic state of a system can be specified only if the system is in thermal equilibrium. When we regard a thermodynamic process we imply that all its state variables change quasi-statically, that is, slowly enough to allow the system to remain essentially in thermal equilibrium at all times.

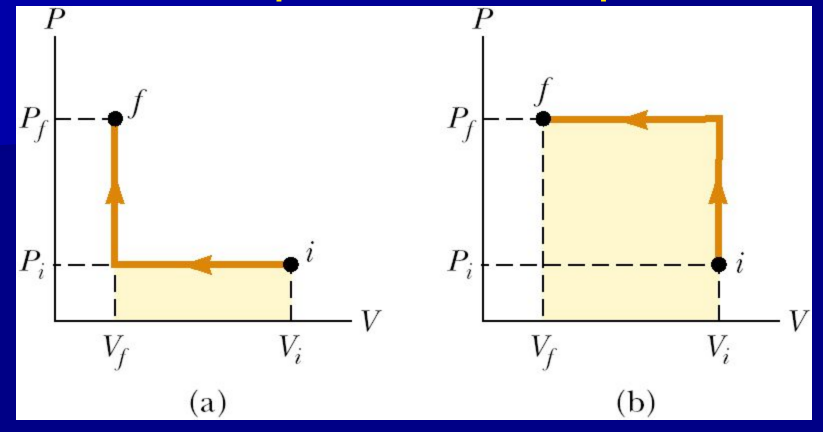
# Work and heat in thermodynamic process

The total work done by the gas as its volume changes from  $V_i$  to  $V_f$  is

$$W=\int_{V_i}^{V_f}\!PdV$$

The work done by a gas in a quasi-static process equals the area under the curve on a PV diagram, evaluated between the initial and final states. It depends on the path between the initial and final states.

#### Work depends on the path:



- (a):  $W_a = P_i(V_f V_i)$ (b):  $W_b = P_f(V_f - V_i)$ 1)  $W_i < W_i$  as  $P_i < V_i$
- 1) W<sub>a</sub> < W<sub>b</sub> as P<sub>f</sub> < P<sub>i</sub>
  2) W<sub>a</sub> < W<sub>b</sub> as the coloured area in (b) case is large then the area in (a) case

# Two ways of energy transfer

There exist two ways in which energy can be transferred between a system and its surroundings:

- One way is work done by the system, which requires that there be a macroscopic displacement of the point of application of a force.
- The other is heat, which occurs on a molecular level whenever a temperature difference exists across the boundary of the system.

Both mechanisms result in a change in the internal energy of the system and therefore usually result in measurable changes in the macroscopic variables of the system, such as the pressure, temperature, and volume of a gas.

### The First Law of Thermodynamics

The change in internal energy  $\Delta U$  of the system is equal to the heat Q put into a system minus the work W done by the system.

$$\Delta U = Q - W$$

Note: here W is with the minus sign as the work is done by the system. The first law of thermodynamics is a special case of the law of conservation of energy that encompasses changes in internal energy and energy transfer by heat and work. It provides a connection between the microscopic and macroscopic approaches.

#### Ideal Gas Processes

- Here W is work done by the system,
- ΔQ heat flow into the system.
- Isobaric (constant pressure):

$$W=P\Delta V$$
$$dQ = C_p dT$$

Isochoric (constant volume):

$$\Delta W = 0$$

$$\Delta Q = \Delta U$$

$$dQ = C_{V}dT$$

 $C_p$ ,  $C_V$  are specific heat capacities,  $C_p = C_V + nR$ , n is the number of moles.

Isothermal (constant temperature):

$$\Delta U = 0$$
  
 $\Delta Q = \Delta W$ 

Adiabatic (no heat flow, Q=0):

$$\Delta W = -\Delta U$$

The curve of adiabatic process is described by formula:

$$PV^{\gamma} = const$$

$$TV^{\gamma-1} = const$$

$$\gamma = C_P/C_V$$

# Polytropic processes

$$PV^{\gamma} = const, \gamma = const.$$

- Isobaric  $\gamma=0$
- Isotermic  $\gamma$ =1
- Adiabatic  $\gamma = C_p/C_V$
- Isochoric γ=∞

### Cyclic Processes

If a nonisolated system is performing a cyclic process, the change in the internal energy must be zero. Therefore the energy Q added to the system must equal the negative of the work W done by the system during the cycle:

$$\Delta U = 0,$$

$$Q = W$$

On a *PV* diagram, a cyclic process appears as a closed curve. In a cyclic process, the net work done by the system per cycle, equals the area enclosed by the path representing the process on a PV diagram.

