Chapter 5: The First Law of Thermodynamics: Closed Systems

The first law of thermodynamics can be simply stated as follows: during an interaction between a system and its surroundings, the amount of energy gained by the system must be exactly equal to the amount of energy lost by the surroundings.

A closed system can exchange energy with its surroundings through *heat* and *work* transfer. In other words, work and heat are the forms that energy can be transferred across the system boundary.

Based on kinetic theory, heat is defined as the energy associated with the random motions of atoms and molecules.

Heat Transfer

Heat is defined as the form of energy that is transferred between two systems by virtue of a *temperature difference*.

Note: there **cannot** be any heat transfer between two systems that are at the **same temperature**.

Note: It is the thermal (internal) energy that can be stored in a system. Heat is a form of energy in transition and as a result can only be identified at the system boundary.

Heat has energy units kJ (or BTU). Rate of heat transfer is the amount of heat transferred per unit time.

Heat is a directional (or vector) quantity; thus, it has magnitude, direction and point of action.

Notation:

- -Q(kJ) amount of heat transfer
- $-Q^{\circ}(kW)$ rate of heat transfer (power)
- -q (kJ/kg) heat transfer per unit mass
- $-q^{\circ}(kW/kg)$ power per unit mass

<u>Sign convention</u>: Heat Transfer *to a system is positive*, and heat transfer *from a system is negative*. It means any heat transfer that increases the energy of a system is positive, and heat transfer that decreases the energy of a system is negative.



Fig. 5-1: Sign convention: positive if to the system, negative if from the system.

Modes of Heat Transfer

Heat can be transferred in three different modes *conduction, convection*, and *radiation*. All modes of heat transfer require the existence of a temperature difference.

<u>Conduction</u>: is the transfer of energy from the more energetic particles to the adjacent less energetic particles as a result of interactions between particles.

In solids, conduction is due to the combination of *vibrations* of the molecules in a *lattice* and the energy transport by *free electrons*. Conduction will be discussed in more details in Ch 10 & 11.

<u>Convection</u>: is the mode of energy transfer between a solid surface and the adjacent liquid or gas which is in motion, and it involves the combined effects of conduction and fluid motion (*advection*).

Convection is called *forced* if the fluid is forced to flow by external means such as a fan or a pump. It is called *free or natural* if the fluid motion is caused by buoyancy forces that are induced by density differences due to the temperature variation in a fluid. Convection will be discussed in more details in Ch 12, 13 & 14.

<u>Radiation</u>: is the energy emitted by matter in the form of electromagnetic waves (or photons) as a result of the changes in the electronic configurations of the atoms or molecules. Radiation will be discussed in more details in Ch 15.

Work

Work is the energy interaction between a system and its surroundings. More specifically, work is the energy transfer associated with *force acting through a distance*.

Notation:

- W (kJ) amount of work transfer
- W° (kW) power
- w (kJ/kg) work per unit mass
- w° (kW/kg) power per unit mass

<u>Sign convention</u>: work *done by* a system is *positive*, and the *work done on* a system is *negative*.



Fig. 5-2: Sign convention for heat and work.

Similarities between work and heat transfer:

- Both are recognized at the boundaries of the system as they cross them (boundary phenomena).
- Systems posses energy, but not heat or work (transfer phenomena).
- Both are associated with a process, not a state. Heat or work has no meaning at a state.
- Both are path functions, their magnitudes depend on the path followed during a process as well as the end states.

<u>Path functions</u>: have **inexact differentials** designated by symbol δ . Properties, on the other hand, are <u>point functions</u> which depend on the state only (not on how a system reaches that state), and they have **exact differentials**.

$$\int_{1}^{2} dV = V_{2} - V_{1} = \Delta V \quad \text{(Point function)}$$

$$\int_{1}^{2} \delta W = W_{12} \quad \text{(Path function, not } \Delta W \text{ nor } W_{2} - W_{1}\text{)}$$

Electrical Work

The work that is done on a system by *electrons*. When N coulombs of electrons move through a potential difference V, the electrical work done is:

 $W_e = VN \quad (kJ)$

Which can be explained in the rate form as

 $W_{e}^{\bullet} = VI$ (*kW*)

Example 5-1: Electrical work

A well-insulated electrical oven is being heated through its heating element. Determine whether it is work or heat interaction. Consider two systems: a) the entire oven (including the heater), and b) only the air in the oven (without the heater) see Fig 5-3.

Solution:

The energy content of the oven is increased during this process.

- a) The energy transfer to the oven is not caused by a temperature difference between the oven and air. Instead, it is caused by electrical energy crossing the system boundary and thus: this is a work transfer process.
- b) This time, the system boundary includes the outer surface of the heater and will not cut through it. Therefore, no electrons will be crossing the system boundary. Instead, the energy transfer is a result of a temperature difference between the electrical heater and air, thus: this is a he at transfer process.



Fig. 5-3: Schematic for example 5-1.

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Mechanical work

There are several ways of doing work, each in some way related to a force acting through a distance.

$$W = F.s \qquad (kJ)$$

If the force is not constant, we need to integrate:

$$W = \int_{1}^{2} F.ds \qquad (kJ)$$

There are two requirements for a work interaction:

- there must be a force acting on the boundary
- the boundary must move

Therefore, the displacement of the boundary without any force to oppose or drive this motion (such as expansion of a gas into evacuated space) is not a work interaction, W=0.

Also, if there are no displacements of the boundary, even if an acting force exists, there will be no work transfer W = 0 (such as increasing gas pressure in a rigid tank).

Moving Boundary Work

The expansion and compression work is often called moving boundary work, or simply *boundary work*.

We analyze the moving boundary work for a quasi-equilibrium process, see Chapter 3. Consider the gas enclosed in a piston-cylinder at initial P and V. If the piston is allowed to move a distance *ds* in a quasi-equilibrium manner, the differential work is:

$$\delta W_{b} = F.ds = PAds = PdV$$

The quasi-equilibrium expansion process is shown in Fig. 5-4. On this diagram, the differential area dA under the process curve in P-V diagram is equal to PdV, which is the differential work.

Note: a gas can follow several different paths from state 1 to 2, and each path will have a different area underneath it (work is path dependent).

The net work or cycle work is shown in Fig. 5-5. In a cycle, the net change for any properties (point functions or exact differentials) is zero. However, the net work and heat transfer depend on the cycle path.

 $\Delta U = \Delta P = \Delta T = \Delta$ (any property) = 0



Fig. 5-4: the area under P-V diagram represents the boundary work.



Fig. 5-5: network done during a cycle.

Polytropic Process

During expansion and compression processes of real gases, pressure and volume are often related by $PV^n=C$, where *n* and *C* are constants. the moving work for a polytropic process can be found:

$$W_{polytopic} = \int_{1}^{2} P dV = \int_{1}^{2} CV^{-n} dV = \frac{P_2 V_2 - P_1 V_1}{1 - n}$$

Since $P_1V_1^n = P_2V_2^n = C$. For an ideal gas (*PV*= *mRT*) it becomes:

$$W_{polytropic} = \frac{mR(T_2 - T_1)}{1 - n}, \quad n \neq 1 \quad (kJ)$$

The special case n =1 is the *isothermal* expansion $P_1V_1 = P_2V_2 = mRT_0 = C$, which can be found from:

$$W_{b,isothermal} = \int_{1}^{2} P dV = \int_{1}^{2} \frac{C}{V} dV = P_{1}V_{1} \ln\left(\frac{V_{2}}{V_{1}}\right), \quad n = 1 \qquad (kJ)$$

Since for an ideal gas, $PV=mRT_0$ at constant temperature T_0 , or P=C/V.

Example 5-2: Polytropic work

A gas in piston-cylinder assembly undergoes a polytropic expansion. The initial pressure is 3 bar, the initial volume is 0.1 m³, and the final volume is 0.2 m³. Determine the work for the process, in kJ, if a) n=1.5, b) n=1.0, and c) n=0.

Solution:

Assume that i) the gas is a closed system, ii) the moving boundary is only work mode, and iii) the expansion is polytropic.

a) n =1.5

$$W = \int_{V_1}^{V_2} P dV = \frac{P_2 V_2 - P_1 V_1}{1 - n}$$

We need P₂ that can be found from $P_1V_1^n = P_2V_2^n$:

$$P_{2} = P_{1} \left(\frac{V_{1}}{V_{2}} \right)^{n} = (3bar) \left(\frac{0.1}{0.2} \right)^{1.5} = 1.06 \ bar$$
$$W = \left(\frac{(1.06bar)(0.2m^{3}) - (3)(0.1)}{1 - 1.5} \right) \left(\frac{10^{5} N / m^{2}}{1 \ bar} \right) \left(\frac{1 \ kJ}{10^{3} \ N.m} \right) = 17.6 \ kJ$$

b) n =1, the pressure volume relationship is PV = constant. The work is:

$$W = \int_{1}^{2} P dV = P_{1}V_{1} \ln\left(\frac{V_{2}}{V_{1}}\right)$$
$$W = (3bar)(0.1 m^{3})\left[\frac{10^{5} N/m^{2}}{1 bar}\right]\left(\frac{1 kJ}{10^{3} N.m}\right) \ln\left(\frac{0.2}{0.1}\right) = 20.79 kJ$$

c) For n = 0, the pressure-volume relation reduces to P=constant (*isobaric process*) and the integral become W= P (V_2 - V_1).

Substituting values and converting units as above, W=30 kJ.

Spring work

For linear elastic springs, the displacement x is proportional to the force applied:

$$F = k_s x$$

where k_s is the spring constant and has the unit kN/m. The displacement x is measured from the undisturbed position of the spring. The spring work is:

$$W_{spring} = \frac{1}{2}k_s \left(x_2^2 - x_1^2\right)$$
 (kJ)

Note: the work done on a spring equals the energy stored in the spring.

Non-mechanical forms of work

Non-mechanical forms of work can be treated in a similar manner to mechanical work. Specify a *generalized force* F acting in the direction of a *generalized displacement* x, the work transfer associated with the displacement dx is:

$$\delta W = F.dx$$

Example 5-3: Mechanical work

Calculate the work transfer in the following process:



Fig. 5-6: Schematic P-V diagram for Example 3-3.

Solution:

Process 1-2 is an expansion $(V_2 > V_1)$ and the system is doing work $(W_{12} > 0)$, thus: $W_{12} = P_1 (V_2 - V_1) + [0.5(P_1 + P_2) - P_1] (V_2 - V_1)$ $= (V_2 - V_1) (P_1 + P_2) / 2$

Process 2 - 3 is an isometric process (constant volume $V_3 = V_2$), so $W_{23} = 0$

Process 3 - 1 is a compression ($V_3 > V_1$), work is done on the system, ($W_{31} < 0$)

$$\begin{split} W_{31} &= - P_1 (V_2 - V_1) \\ W_{cycle} &= W_{net} = W_{12} + W_{23} + W_{31} = (V_2 - V_1) (P_2 - P_1) / 2 \\ \text{Note that in a cycle } \Delta U &= \Delta P = \Delta T = \Delta (\text{any property}) = 0 \end{split}$$

First Law of Thermodynamics

First law, or the *conservation of energy principle*, states that energy can be neither created nor destroyed; it can only *change forms*.

The first law <u>cannot be proved mathematically</u>, it is based on *experimental* observations, i.e., there are no process in the nature that violates the first law.

The first law for a closed system or a fixed mass may be expressed as:

net energy transfer to (or from) = net increase (or decrease) in the the system as heat and work total energy of the system

$$Q - W = \Delta E (kJ)$$

where

Q = net heat transfer (= $\Sigma Q_{in} - \Sigma Q_{out}$)

W= net work done in all forms $(=\Sigma W_{in} - \Sigma W_{out})$

 ΔE = net change in total energy (= $E_2 - E_1$)

The change in total energy of a system during a process can be expressed as the sum of the changes in its internal, kinetic, and potential energies:

$$\Delta E = \Delta U + \Delta KE + \Delta PE \qquad (kJ)$$
$$\Delta U = m (u_2 - u_1)$$
$$\Delta KE = \frac{1}{2} m (V_2^2 - V_1^2)$$
$$\Delta PE = mg (z_2 - z_1)$$

Note: for stationary systems $\Delta PE=\Delta KE=0$, the first law reduces to

$$Q - W = \Delta U$$

The first law can be written on a unit-mass basis:

$$q - w = \Delta e$$
 (kJ/kg)

or in differential form:

$$\delta Q - \delta W = dU$$
 (kJ)
 $\delta q - \delta W = du$ (kJ/kg)

or in the rate form:

$$Q^{\circ} - W^{\circ} = dE / dt$$
 (kW)

For a cyclic process, the initial and final states are identical, thus $\Delta E=0$. The first law becomes:

$$Q - W = 0$$
 (kJ)

Note: from the *first law* point of view, there is no difference between *heat transfer* and *work*, they are both *energy interactions*. But from the *second law* point of view, heat and work are *very different*.

Example 5-4: Fist law

Air is contained in a vertical piston-cylinder assembly fitted with an electrical resistor. The atmospheric pressure is 100 *kPa* and piston has a mass of 50 *kg* and a face area of 0.1 m². Electric current passes through the resistor, and the volume of air slowly increases by 0.045 m³. The mass of the air is 0.3 kg and its specific energy increases by 42.2 kJ/kg. Assume the assembly (including the piston) is insulated and neglect the friction between the cylinder and piston, g = 9.8 m/s². Determine the heat transfer from the resistor to air for a system consisting a) the air alone, b) the air and the piston.



Fig. 5-7: Schematic for problem 5-4.

Assumptions:

- > two closed systems are under consideration, as shown in schematic.
- > the only heat transfer is from the resistor to the air. $\Delta KE = \Delta PE = 0$ (for air)
- > the internal energy is of the piston is not affected by the heat transfer.

a) taking the air as the system,

$$(\Delta KE + \Delta PE + \Delta U)_{air} = Q - W$$

 $Q = W + \Delta U_{air}$

For this system work is done at the bottom of the piston. The work done by the system is (at constant pressure):

$$W = \int_{V_1}^{V_2} P dV = P(V_2 - V_1)$$

The pressure acting on the air can be found from:

$$PA_{piston} = m_{piston} g + P_{atm} A_{piston}$$

$$P = \frac{m_{\text{piston}}g}{A_{\text{piston}}} + P_{atm}$$
$$P = \frac{(50kg)(9.81m/s^2)}{(0.1m^2)} \left(\frac{1Pa}{1N/m^2}\right) \left(\frac{1kPa}{1000Pa}\right) + 100kPa = 104.91 \, kPa$$

Thus, the work is

With $\Delta U_{air} = m_{air} \Delta u_{air}$, the heat transfer is

Q = W + $m_{air} \Delta u_{air}$ = 4.721 kJ + (0.3 kg)(42.2 kJ/kg) = 17.38 kJ

b) system consisting the air and the piston. The first law becomes:

$$(\Delta KE + \Delta PE + \Delta U)_{air} + (\Delta KE + \Delta PE + \Delta U)_{piston} = Q - W$$

where $(\Delta KE = \Delta PE)_{air} = 0$ and $(\Delta KE = \Delta U)_{piston} = 0$. Thus, it simplifies to:

$$(\Delta U)_{air} + (\Delta PE)_{piston} = Q - W$$

For this system, work is done at the top of the piston and pressure is the atmospheric pressure. The work becomes

$$W = P_{atm} \Delta V = (100 \text{ kPa})(0.045 \text{ m}^3) = 4.5 \text{ kJ}$$

The elevation change required to evaluate the potential energy change of the piston can be found from the volume change:

$$\Delta z = \Delta V / A_{piston} = 0.045 \text{ m}^3 / 0.1 \text{ m}^2 = 0.45 \text{ m}$$

$$(\Delta PE)_{piston} = m_{piston} g \Delta z = (50 \text{ kg})(9.81 \text{ m/s}^2)(0.45 \text{ m}) = 220.73 \text{ J} = 0.221 \text{ kJ}$$

$$Q = W + (\Delta PE)_{piston} + m_{air} \Delta u_{air}$$

Note that the heat transfer is identical in both systems.

Specific Heats

The specific heat is defined as the energy required to raise the temperature of a unit mass of a substance by one degree. There are two kinds of specific heats:

- specific heat at constant volume, Cv (the energy required when the volume is maintained constant)
- specific heat at constant pressure, Cp (the energy required when the pressure is maintained constant)

The specific heat at constant pressure Cp is **always higher than** Cv because at constant pressure the system is allowed to expand and energy for this expansion must also be supplied to the system.

Let's consider a stationary closed system undergoing a constant-volume process ($w_b = 0$). Applying the first law in the differential form:

at constant volume (no work) and by using the definition of C_{ν} , one can write:

$$C_{v}dT = du$$

or
$$C_{v} = \left(\frac{\partial u}{\partial T}\right)_{v}$$

Similarly, an expression for the specific heat at constant pressure *Cp* can be found. From the first law, for a constant pressure process ($w_b + \Delta u = \Delta h$). It yields:

$$C_p = \left(\frac{\partial h}{\partial T}\right)_p$$

- specific heats (both C_v and C_p) are properties and therefore independent of the type of processes.
- C_v is related to the changes in *internal energy u*, and C_p to the changes in *enthalpy*, *h*.

It would be more appropriate to define: C_v is the change in specific internal energy per unit change in temperature at constant volume. C_p is the change in specific enthalpy per unit change in temperature at constant pressure.

Specific heats for ideal gases

It has been shown mathematically and experimentally that the internal energy is a function of temperature only.

$$u = u(T)$$

Using the definition of enthalpy (h = u + Pv) and the ideal gas equation of state (Pv = RT), we have:

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$$h = u + RT$$

Since *R* is a constant and u is a function of *T* only:

$$h = h(T)$$

Therefore, at a given temperature, u, h, C_v and C_p of an ideal gas will have fixed values regardless of the specific volume or pressure. For an ideal gas, we have:

$$du = C_v(T) dT$$
$$dh = C_v(T) dT$$

The changes in internal energy or enthalpy for an ideal gas during a process are determined by integrating:

$$\Delta u = u_2 - u_1 = \int_{1}^{2} C_{\nu}(T) dT \qquad (kJ / kg)$$
$$\Delta h = h_2 - h_1 = \int_{1}^{2} C_{\nu}(T) dT \qquad (kJ / kg)$$

As low pressures, all real gases approach ideal-gas behavior, and therefore their specific heats depend on temperature only. The specific heats of real gases at low pressures are *called ideal-gas specific heats* (or *zero-pressure specific heats*) and are often denoted by C_{p0} and C_{v0} . To carry out the above integrations, we need to know $C_v(T)$ and $C_p(T)$. These are available from a variety of sources:

- Table A-2a: for various materials at a fixed temperature of T = 300 K
- > Table A-2b: various gases over a range of temperatures $250 \le T \le 1000 \text{ K}$
- ➤ Table A-2c: various common gases in the form of a third order polynomial For an ideal gas, we can write:

$$RT = h(T) - u(T)$$
$$R = \frac{dh}{dT} - \frac{du}{dT}$$
$$R = C_p - C_v$$

The ratio of specific heats is called the *specific heat ratio* $k = C_p/C_v$:

- > varies with temperature, but this variation is very mild.
- ➢ for monatomic gases, its value is essentially constant at 1.67.
- Many diatomic gases, including air, have a specific heat ratio of about 1.4 at room temperature.

Specific heats for solids and liquids

A substance whose specific volume (or density) is constant is called *incompressible substance*. The specific volumes of solids and liquids (which can be assumed as incompressible substances) essentially remain constant during a process.

The constant volume assumption means that the volume work (boundary work) is negligible compared with other forms of energy. As a result, it can be shown that the constant-volume and constant-pressure specific heats are identical for incompressible substances:

$$C_p = C_v = C$$

Specific heats of incompressible substances are only a function of temperature,

$$C = C(T)$$

The change of internal energy between state 1 and 2 can be obtained by integration:

$$\Delta u = u_2 - u_1 = \int_{1}^{2} C(T) dT \qquad (kJ / kg)$$

For small temperature intervals, a C at averaged temperature can be used and treated as a constant, yielding:

$$\Delta u \approx C_{ave} \left(T_2 - T_1 \right)$$

The enthalpy change of incompressible substance can be determined from the definition of enthalpy (h = u + Pv)

$$h_2 - h_1 = (u_2 - u_1) + v(P_2 - P_1)$$
$$\Delta h = \Delta u + v \Delta P \quad (kJ/kg)$$

The term $v\Delta P$ is often small and can be neglected, so $\Delta h = \Delta u$.

Example 5-5: Specific heat and first law

Two tanks are connected by a valve. One tank contains 2 kg of CO_2 at 77°C and 0.7 bar. The other tank has 8 kg of the same gas at 27°C and 1.2 bar. The valve is opened and gases are allowed to mix while receiving energy by heat transfer from the surroundings. The final equilibrium temperature is 42°C. Using ideal gas model, determine a) the final equilibrium pressure b) the heat transfer for the process.

Assumptions:

- > the total amount of CO_2 remains constant (closed system).
- ➢ ideal gas with constant Cv.
- > The initial and final states in the tanks are equilibrium. No work transfer.

The final pressure can be found from ideal gas equation of state:

$$P_{f} = \frac{m_{t}RT_{f}}{V_{1} + V_{2}} = \frac{(m_{1} + m_{2})RT_{f}}{V_{1} + V_{2}}$$

For tank 1 and 2, we can write: $V_1 = m_1 R T_1 / P_1$ and $V_2 = m_2 R T_2 / P_2$. Thus, the final pressure, Pf becomes:

$$\begin{split} P_{f} &= \frac{\left(m_{1} + m_{2}\right)RT_{f}}{\left(\frac{m_{1}RT_{1}}{P_{1}}\right)\left(\frac{m_{2}RT_{2}}{P_{2}}\right)} = \frac{\left(m_{1} + m_{2}\right)T_{f}}{\left(\frac{m_{1}T_{1}}{P_{1}}\right)\left(\frac{m_{2}T_{2}}{P_{2}}\right)}\\ P_{f} &= \frac{\left(10kg\right)(315K)}{\frac{(2kg)(350K)}{0.7bar} + \frac{(8kg)(300K)}{1.2bar}} = 1.05bar \end{split}$$

b) the heat transfer can be found from an energy balance:

$$\Delta U = Q - W$$

With W = 0,

 $Q = U_f - U_i$

where initial internal energy is: $U_i = m_1 u(T_1) + m_2 u(T_2)$

The final internal energy is: $U_f = (m_1 + m_2) u(T_f)$

The energy balance becomes:

$$Q = m_1 [u(T_f) - u(T_1)] + m_2 [u(T_f) - u(T_2)]$$

Since the specific heat C_v is constant

$$Q = m_1 C_v [T_f - T_1] + m_2 C_v [T_f - T_2]$$
$$Q = (2kg) \left(0.745 \frac{kJ}{kg.K} \right) (315K - 350K) + (8kg) \left(0.745 \frac{kJ}{kg.K} \right) (315K - 300K) = 37.25kJ$$

The plus sign indicates that the heat transfer is *into the system*.

Chapter 6: The First Law: Control Volumes

The first law is discussed for closed systems in Chapter 5. In this Chapter, we extend the conservation of energy to systems that involve mass flow across their boundaries, *control volumes*.

Any arbitrary region in space can be selected as *control volume*. There are no concrete rules for the selection of control volumes. The boundary of control volume is called a *control surface*.

Conservation of Mass

Like energy, mass is a conserved property, and it cannot be created or destroyed. Mass and energy can be converted to each other according to Einstein's formula: $E = mc^2$, where *c* is the speed of light. However, except for nuclear reactions, the conservation of mass principle holds for all processes.

For a control volume undergoing a process, the conservation of mass can be stated as:



Fig. 6-1: Conservation of mass principle for a CV.

The conservation of mass can also be expressed in the rate form:

$$\sum m^{\bullet}_{i} - \sum m^{\bullet}_{e} = dm_{CV} / dt$$

The amount of mass flowing through a cross section per unit time is called the mass flow rate and is denoted by m° . The mass flow rate through a differential area dA is:

$$dm^{\circ} = \rho V_n dA$$

where V is the velocity component normal to dA. Thus, the mass flow rate for the entire cross-section is obtained by:

$$m^{\bullet} = \int_{A} \rho V_n dA$$
 (kg/s)

Assuming one-dimensional flow, a uniform (averaged or bulk) velocity can be defined:

$$m^{\circ} = \rho V A$$
 (kg/s)

where V (m/s) is the fluid velocity normal to the cross sectional area. The volume of the fluid flowing through a cross-section per unit time is called the volumetric flow, V $^{\circ}$:

$$V^{\bullet} = \int_{A} V_n dA = VA \qquad (\mathrm{m}^3/\mathrm{s})$$

The mass and volume flow rate are related by: $m^{\circ}=\rho V^{\circ}=V^{\circ}/v$.

Conservation of Energy

For control volumes, an additional mechanism can change the energy of a system: mass flow in and out of the control volume. Therefore, the conservation of energy for a control volume undergoing a process can be expressed as

total energy crossing	+	total energy of	-	total energy	=	net change
boundary as heat		mass entering		of mass		in energy of
and work		CV		leaving CV		CV

$$Q - W + \sum E_{in,mass} + \sum E_{out,mass} = \Delta E_{CV}$$

This equation is applicable to *any* control volume undergoing *any* process. This equation can also be expressed in rate form:



Fig. 6-2: Energy content of CV can be changed by mass flow in/out and heat and work interactions.

<u>Work flow:</u> is the energy that required to push fluid into or out of a control volume. Consider an imaginary piston (that push the fluid to CV) where the fluid pressure is P and the cross sectional area is A. The force acting on the piston is F = PA.



Fig. 6-3: schematic for flow work.

The work done in pushing the fluid is:

$$W_{flow} = F.s = PA.s = PV$$
 (kJ)

or in a unit basis,

$$w_{flow} = W_{flow} / m = Pv$$
 (kJ/kg)

Note that the flow work is expressed in terms of properties.

The flow work can also be written as a rate equation.

The fluid entering or leaving a control volume possesses an additional form of energy (flow energy Pv). Therefore, the total energy of a flowing fluid on a unit-mass basis (denoted by θ) becomes:

$$\theta = Pv + e = Pv + (u + ke + pe)$$
 (kJ/kg)

Recall that enthalpy is defined as: h = u + Pv. Therefore, the above equation becomes:

$$\theta = h + ke + pe = h + V^2 / 2 + gz$$
 (kJ/kg)

The property θ is called **methalpy**. By using enthalpy instead of internal energy, the energy associated with *flow work into/out of control volume is automatically taken care of*. This is the main reason that enthalpy is defined!

Steady-State Flow Process

A process during which a fluid flows through a control volume steadily is called *steady-state process*. A large number of devices such as turbines, compressors, and nozzles operates under the same conditions for a long time and can be modeled (classified) as steady-flow devices.

The term *steady* implies no change with time. The term *uniform* implies no change with location over a specified region.

A steady flow is characterized by the following:

1- No properties within the CV change with time. Thus, volume, mass, and energy of CV remains constant. *As a result, the boundary work is zero*. Also, total mass entering the CV must be equal to total mass leaving CV.

2- No properties change at the boundary of the CV with time. It means that the mass flow rate and the properties of the fluid at an opening must remain constant during a steady flow.

3- The heat and mass interactions between the CV and its surroundings do not change with time.

Using the above observation, the conservation of energy principle for a general steady-flow system with multiple inlets and exits can be written as:

$$Q^{\bullet} - W^{\bullet} = \sum m_e^{\bullet} \left(h_e + \frac{V_e^2}{2} + gz_e \right) - \sum m_i^{\bullet} \left(h_i + \frac{V_i^2}{2} + gz_i \right)$$
$$Q^{\bullet} - W^{\bullet} = \sum m_e^{\bullet} \theta_e - \sum m_i^{\bullet} \theta_i$$

Nozzles and Diffusers

A *nozzle* is a device that *increases the velocity* of a fluid at the expense of pressure. A *diffuser* is a device that *increases the pressure* of a fluid by slowing it down.

The cross sectional area of a nozzle decreases in the flow direction for subsonic flows and increase for supersonic flows.



Fig. 6-4: Schematic of nozzle

For a nozzle, common assumptions and idealizations are:

Q° = 0, no time for heat transfer, due to high velocity

 $W^{\circ} = 0$, nozzles include no shaft or electric resistance wires

 $\Delta PE = 0$, no change in fluid elevation.

Mass equation for nozzles becomes:

$$m_{1}^{\circ} = m_{2}^{\circ}$$
 or $\rho_{1}A_{1}V_{1} = \rho_{2}A_{2}V_{2}$

Energy balance:

$$Q^{\circ} - W^{\circ} = m^{\circ}_{1} \theta_{1} - m^{\circ}_{2} \theta_{2}$$

(h + V²/2)_{at inlet} = (h + V²/2)_{at exit}

Diffusers are exactly the same device as nozzles; the only difference is the direction of the flow. Thus, all equations derived for nozzles hold for diffusers.



Fig. 6-5: Schematic for diffuser.

Example 6-1: Nozzle

Steam enters a converging-diverging nozzle operating at steady state with $P_1 = 0.05$ MPa, $T_1 = 400$ °C and a velocity of 10 m/s. The steam flows through the nozzle with negligible heat transfer and no significant change in potential energy. At the exit, $P_2 = 0.01$ MPa, and the velocity is 665 m/s. The mass flow rate is 2 kg/s. Determine the exit area of the nozzle, in m².

Assumptions:

- 1. Steady state operation of the nozzle
- 2. Work and heat transfer are negligible, $Q^\circ = W^\circ = 0$.
- 3. Change in potential energy from inlet to exit is negligible, $\Delta PE = 0$.

The exit area can be calculated from the mass flow rate m° :

$$A2 = m^{\circ}v_2 / V_2$$

We need the specific volume at state 2. So, state 2 must be found. The pressure at the exit is given; to fix the state 2 another property is required. From the energy equation enthalpy can be found:

$$h_2 + \frac{V_2^2}{2} = h_1 + \frac{V_1^2}{2} \implies h_2 = h_1 + \frac{1}{2} (V_1^2 - V_2^2)$$

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From Table A-6, h_1 = 3278.9 kJ/kg and velocities are given.

$$h_2 = 3278.9 \ kJ \ / \ kg \ + \left[\frac{10^2 - 665^2}{2}\right] \left(\frac{m}{s}\right)^2 \left[\frac{1 \ N}{1 \ kg \ m \ / \ s^2}\right] \left[\frac{1 \ kJ}{10^3 \ N \ m}\right] = 3057.84 \ kJ \ / \ kg$$

From Table A-6, with P₂ = 0.01 MPa, and h_2 = 3057.84 kJ/kg, (using interpolation) one finds T₂ = 300 °C (approximately) and v₂ = 26.445 m³/kg.

The exit area is:



Fig. 6-6: Converging-diverging nozzle.

Turbines and Compressors

In steam, gas, or hydroelectric power plants, the device that derives the electric generator is turbine. The work of turbine is positive since it is done by the fluid.

Compressors, pumps, and fans are devices used to increase the pressure of the fluid. Work is supplied to these devices, thus the work term is negative.

Common assumptions for turbines and compressors:

Q° = 0.

 $\Delta PE = \Delta KE = 0.$

Example 6-2: Turbine

Steam enters a turbine at steady state with a mass flow rate of 4600 kg/h. The turbine develops a power output of 1000 kW. At the inlet the pressure is 0.05 MPa, the temperature is 400 °C, and the velocity is10 m/s. At the exit, the pressure is 10 kPa, the quality is 0.9, and the velocity is 50 m/s. Calculate the rate of heat transfer between the turbine and surroundings, in kW.

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Assumptions:

Steady-state operation. The change in potential energy is negligible. The energy balance for the turbine is:

$$Q_{CV}^{\bullet} - W_{CV}^{\bullet} = m^{\bullet} \left(h_2 + \frac{V_2^2}{2} + gz_2 \right) - m^{\bullet} \left(h_1 + \frac{V_1^2}{2} + gz_1 \right)$$
$$Q_{CV}^{\bullet} = W_{CV}^{\bullet} + m^{\bullet} \left[\left(h_2 - h_1 \right) + \frac{1}{2} \left(V_2^2 - V_1^2 \right) \right]$$



Fig. 6-7: Schematic for turbine.

Using Table A-6, $h_1 = 3278.9 \text{ kJ/kg}$, using Table A-5 with x = 0.9,

 $h_2 = h_f + x_2 h_{fg} = 191.83 \text{ kJ/kg} + 0.9(2392.8 \text{ kJ/kg}) = 2345.35 \text{ kJ/kg}$ Therefore,

 $h_2 - h_1 = 2345.35 - 3278.9 = -933.55 \text{ kJ/kg}$

The specific kinetic energy change is:

$$\frac{1}{2} \left(V_2^2 - V_1^2 \right) = 0.5 \left(50^2 - 10^2 \right) \left(\frac{m}{s} \right)^2 \frac{1 N}{1 \text{ kg.m/s}^2} \frac{1 \text{ kJ}}{10^3 \text{ N.m}} = 1.2 \text{ kJ / kg}$$

 Q°_{CV} = (1000 kW) + 4600 kg/h (-831.8 + 1.2 kJ/kg) (1h / 3600 s) (1kW / 1 kJ/s) = -61.3 kW

Note that change in specific kinetic energy is very small. Also Q°_{CV} is small compared to W°_{CV} . The negative sign in Q°_{CV} means that heat transfer is from CV to surroundings.

Throttling Valves

Any kind of flow restricting devices that causes a significant pressure drop in the fluid is called throttling valve.



Fig. 6-8: Three kinds of throttling valves.

The pressure drop in fluid is often accompanied with a temperature drop in fluids. The magnitude of this temperature is governed by a property called the *Joule-Thomson coefficient*.

Common assumptions for throttling valves:

Q° = 0

W° = 0

 $\Delta PE = \Delta KE = 0.$

The conservation of energy becomes:

$$h_2 = h_1$$

 $u_1 + P_1 v_1 = u_2 + P_2 v_2$

The flow energy increases during the process ($P_2v_2 > P_1v_1$), and it is done at the expense of the internal energy. Thus, internal energy decreases, which is usually accompanied by a drop in temperature.

For ideal gases h = h(T) and since h remains constant the temperature has to remain constant too.

Mixture Chambers (Direct Contact Heat Exchangers)

Mixture chambers are used to mix two streams of fluids. Based on mass principle, the sum of incoming flow equals to sum of leaving fluid.



Fig. 6-9: T-elbow is a mixing chamber.

Common assumptions for mixing chamber:

 $Q^\circ = 0$

 $W^{\circ} = 0$

 $\Delta PE = \Delta KE = 0.$

The conservation of energy becomes:

 $\Sigma(m^{\circ}h)_{inlet} = \Sigma(m^{\circ}h)_{outlet}$

Heat Exchangers

Heat exchangers are devices where two moving fluid streams exchange heat without mixing.

Common assumptions for heat exchangers:

W° = 0

 $\Delta PE = \Delta KE = 0.$

Q° could be different depending on the selected CV, see example 6-3.

The mass and energy balance become:



Fig. 6-10: Schematic for heat exchanger.

Example 6-3: Heat exchanger

Engine oil is to be cooled by water in a condenser. The engine oil enters the condenser with a mass flow rate of 6 kg/min at 1 MPa and 70°C and leaves at 35°C. The cooling water enters at 300 kPa and 15°C and leaves at 25°C. Neglecting any pressure drops; determine a) the mass flow rate of the cooling water required, and b) the heat transfer rate from the engine oil to water.



We choose the entire heat exchanger as our control volume, thus work transfer and heat transfer to the surroundings will be zero. From mass balance:

$$m_1^{\bullet} = m_2^{\bullet} = m_W^{\bullet}$$
 and $m_3^{\bullet} = m_4^{\bullet} = m_{Oil}^{\bullet}$

The conservation of energy equation is:

$$Q^{\bullet} - W^{\bullet} = \sum m_{e}^{\bullet} \left(h_{e} + \frac{V_{e}^{2}}{2} + gz_{e} \right) - \sum m_{i}^{\bullet} \left(h_{i} + \frac{V_{i}^{2}}{2} + gz_{i} \right)$$

$$\sum m_{e}^{\bullet} h_{e} = \sum m_{e}^{\bullet} h_{e}$$

$$m_{W}^{\bullet} h_{1} + m_{Oil}^{\bullet} h_{3} = m_{W}^{\bullet} h_{2} + m_{Oil}^{\bullet} h_{4}$$

$$m_{W}^{\bullet} = \frac{h_{3} - h_{4}}{h_{2} - h_{1}} m_{Oil}^{\bullet}$$

Assuming constant specific heat for both the oil and water at their average temperature,

$$m_{W}^{\bullet} = \frac{C_{P,Oil}(T_{3} - T_{4})}{C_{P,Water}(T_{2} - T_{1})} = \frac{\left(2.016\frac{kJ}{kg.C}\right)(70 - 35)}{\left(4.18\frac{kJ}{kg.C}\right)(25 - 15)}(6 \ kg \ / \min) = 10.1 \ kg \ / \min$$

b) To determine the heat transfer from the oil to water, choose the following CV. The energy equation becomes:

$$Q_{Oil}^{\bullet} - W^{\bullet} = m_{Oil}^{\bullet} (\Delta h) = m_{Oil}^{\bullet} C_{P,Oil} (T_4 - T_3)$$
$$Q_{Oil}^{\bullet} = (6 \ kg \ / \min) \left(2.016 \frac{kJ}{kg.C} \right) (35 - 70) = -423.36 \ kJ \ / \ kg$$



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