

Physical Chemistry

Chapter 2 First Law of Thermodynamics

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First Law of Thermodynamics by Norhayati

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1.0 First Law of Thermodynamics

Aims

- To define the intensive and extensive state of function in thermal and not thermal
- To develop the calculation path for property change and estimate enthalpy and entropy values using equation state and generalized correlations
- To explain the conservation of the total energy of system plus surroundings for close and open systems
- To estimate heat, work, change of internal energy and change of enthalpy for various kinds of thermodynamic processes
- Expected Outcomes
 - Student should be able to define the intensive and extensive state of function in thermal and not thermal
 - Student should be able to develop the calculation path for property change and estimate enthalpy and entropy values using equation state and generalized correlations
 - Student should be able to explain the conservation of the total energy of system plus surroundings for close and open systems
 - Student should be able to estimate heat, work, change of internal energy and change of enthalpy for various kinds of thermodynamic processes



Subtopics

- 2.1 Introduction to First Law of Thermodynamics
- 2.2 Energy
- 2.3 Work
- 2.4 Heat transfer
- 2.5 Internal energy
- 2.6 Enthalpy
- 2.7 Measurement of Heat and Specific Heat
- 2.8 Thermochemistry



References

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2.1 First Law of Thermodynamics2.2 Energy

Terminology	Definition
2.1 First Law of Thermodynamics	Thermodynamics contributes the law of conservation of energy, which energy cannot be created or destroyed during a process; it can only change from one form to another
2.2 Energy	Energy is capacity to do work. Unit : J (1 J= 1 kgm ₂ s ⁻²) In thermodynamics analysis, forms of energy can be divided into two groups; macroscopic and microscopic energy. TWO form of energies: (a) Microscopic and(b) macroscopic
2.2 (a) Microscopic energy	 The forms of energy are those related to the molecular structure of a system and the molecular activity The sum of all the microscopic forms of energy is called the internal energy of a system and is denoted by U.
2.2 (b) Macroscopic energy	The macroscopic energy of a system is related to motion and the influence of some external effects such as gravity, magnetism, electricity and surface tension. Magnetism, electricity and surface tension are usually ignored. Consider 1- Kinetic energy 2- Potential energy

2.2 Energy

Terminology	Definition	
2.2 (b) Kinetic energy (Macroscopic energy)	The energy that a system possesses as a result of its motion is called kinetic energy (KE). $KE = m \frac{V^2}{2}$ (kJ)	
2.2 (b) Potential Energy <i>(Macroscopic energy)</i>	The energy that a system possesses as a result of its elevation in a gravitational field is called potential energy (PE) g is the gravitational acceleration, 9.81 m/s2 z is the elevation PE = mgz (kJ)	
Combination of energies	The total energy of a system consists of the kinetic, potential, and internal energies and is expressed as $E = U + KE + PE = U + m \frac{V^2}{2} + mgz (kJ)$	
Static energy	The total energy of a system, can be contained or stored in a system, and thus can be viewed as the static forms of energy.	
Dynamic energy	The forms of energy not stored in a system can be viewed as the dynamic forms. The only two modes of transfer energy interactions associated with a closed system are work and heat transfer.	

2.3 Work

Terminology	Definition	
2.3 Work	 Work is the energy transfer associated with a force acting through a distance or work is performed when the object moves some distance z due to the application of a force, F e.g. Chemical reaction to work Zn + 2HCI → ZnCl2 + H2 The gas produced drives out the piston and raise the weight. Work is done on a system is reported as positive quantity (energy is transferred to the system by doing work) 	
2.3(a) Expansion Work	 Maximum work is obtained when the external pressure is only infinitesimally less than the pressure of the gas in the system. System that remains in mechanical equilibrium with its surroundings at all stages of the expansion does maximum expansion work. When a system is in a state of mechanical equilibrium, an infinitesimal change in pressure results in opposite directions of change. In this case is said reversible process. 	

2.4 Heat Transfer

Terminology	Definition
2.4 Heat Transfer	 Energy can cross the boundary of a closed system in two distinct forms; <i>heat</i> and <i>work</i>. Heat – forms of energy that is transferred between 2 system by temperature difference. No heat transfer between two systems that are at the same temperature 3 modes of energy transfer : conduction, convection and radiation
(a) Exothermic	A process in a system that heats release to the surrounding
(b) Endothermic	A process in a system that is heated by the surrounding
(c) Diathermic	Walls that permit heating for heat transfer
(d) Adiabatic	Walls that do not permit heating for heat transfer





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2.5 Internal Energy

Definition

- Total energy of a system denoted as U
- KE + PE = U
- the change in internal energy of a system is

$$dU = dq + dw_{exp} + dw_e$$

 w_{exp} = expansion work

w_e = work in addition to the expansion work (e.g electrical work)

- If system kept at constant volume, w_{exp} = 0
- When the system release energy to surrounding, the internal energy of the system decrease, (-ve sign)
- Works done by the system
- Heat loses during heating process
- When the internal energy increase, (+ve sign)
- Work done on the system
- The system absorb heat from surrounding

- Enthalpy, H (J) is extensive property, while molar enthalpy, H_m (H/n; J/mol) is intensive property
- Enthalpy of a system is defined as

H = U + PV

• A change in enthalpy arises from a change in the internal energy and a change in the product PV.

 $\Delta H = \Delta U + \Delta (PV)$

• If the changes takes place at constant pressure, $\Delta H = \Delta U + P \Delta V$

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Molar enthalpy,

$$H_m = U_m + PV_m$$

• For perfect gas,

$$PV_m = RT$$

Where,

$$H_m = U_m + RT$$

Molar Enthalpy, $H_m = H/n$ Molar volume, $V_m = V/n$



• At constant pressure, no non-expansion work:

$$\Delta H = q_P$$

- Endothermic reaction; q > 0, $\Delta H > 0$
- Exothermic reaction; q < 0, $\Delta H < 0$



Three properties of enthalpy:
 Heat supplied at constant pressure

$$\Delta H = q_p$$

• Enthalpy change

$$\Delta H = H_f - H_i$$

Specific heat at constant pressure

$$C_p = \Delta H / \Delta T$$



2.7 Measurement of Heat and Specific Heat

• Specific Heats

$$C_{V} = \left(\frac{\partial u}{\partial T}\right)_{v} \quad and \quad C_{P} = \left(\frac{\partial h}{\partial T}\right)_{P}$$

 C_V = the change in internal energy with temperature at constant volume C_p= the change in enthalpy
with temperature at
constant pressure



2.7 Measurement of Heat and Specific Heat

Internal energy of a system rises as the temperature increased. The same as enthalpy, which also rises when temperature is increased.

Constant-volume heat capacity, Cv shows the temperature dependence of the internal energy at constant volume.

Constant-pressure heat capacity, Cp shows the enthalpy of a system changes as its temperature is raised at constant pressure.



2.7 Measurement of Heat and Specific Heat

• For a perfect gas,

$$C_{P,m} - C_{V,m} = R$$

 Molar heat capacity of a perfect gas is greater at constant pressure than at constant volume. Why?

At **constant volume**, all energy supplied as heat to the system remains inside.

At constant pressure, the energy supplied as heat escapes **back** to surrounding when the system expands and does work.



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- Thermochemistry is the study of heat change in chemical reactions.
- The application of enthalpy in chemistry.
- The thermochemistry processes contributing enthalpies are as follows:
 - 1. Phase changes
 - 2. Chemical changes (combustion, formation and reaction)

Enthalpy in phase changes

- Vaporization
- Fusion
- Freezing
- Sublimation



Enthalpy in phase changes;

- The conversion of one phase of a substance to another phase is called phase transition.
- Most phase transitions are accompanied by a change of enthalpy.
- The vaporization of a liquid is an endothermic process because heat must be supplied to bring about the change.
- Standard enthalpy of vaporization (+ve) is the energy that must be supplied as heat at constant pressure per mole of molecules that are vaporized under standard conditions (1 bar).

 $H_2O(l) \rightarrow H_2O(g) \qquad \Delta H^\circ = 40.7 \text{kJ/mol}$

Enthalpy in phase changes

• Standard enthalpy of fusion (+ve) is the change in molar enthalpy that accompanies fusion under standard conditions.

 $H_2O(s) \rightarrow H_2O(l) \qquad \Delta H^o = 6.01 kJ/mol$

• What is the **standard enthalpy of freezing**?



 Standard enthalpy of sublimation is the amount of energy required for direct conversion of a solid to a vapor.

$$\Delta H^{o}_{_{Sub}} = \Delta H^{o}_{_{Fus}} + \Delta H^{o}_{_{Vap}}$$

- The two enthalpies that are added together must be for the same temperature.
- Chemical changes
 - Combustion
 - Reaction
 - Formation



- Enthalpy of combustion
 - The standard enthalpy of combustion, ΔH^{o}_{c} is the change in standard enthalpy per mole of combustible substance.

 $CH_4(g) + 2O_2(g) \rightarrow CO_2(g) + H_2O(l)$ $\Delta H_c^o = -890 \, kJ$

- Commonly measured using bomb calorimeter (energy transfer as heat at constant volume).
- Gas phase

$$\Delta H_{c}^{o} = \Delta U_{c} + \Delta V_{gas} RT$$



	Symbol	Definition
Enthalpy of reaction,	$\Delta_r H$	The change in enthalpy that companies a chemical reaction Standard enthalpy of reaction is the value of the reaction enthalpy when all the reactants and products are in their standard state $\Delta H_{r}^{\theta} = \sum v H_{m}^{\theta} (\text{products}) - \sum v H_{m}^{\theta} (\text{reactants})$ $C + O_{2} \rightarrow CO_{2} \Delta H_{1} = -393.5 \text{ kJ}$ $CO + \frac{1}{2}O_{2} \rightarrow CO_{2} \Delta H_{2} = -282.9 \text{ kJ}$ $\Delta H_{rxn} = -110.62 \text{ kJ/mole}$

	Symbol	Definition
Enthalpy of formation,	$\Delta_f H$	Standard enthalpy of formation of a substance is the standard enthalpy (per mole of the substance) for it formation from its elements in their reference state. $\Delta H_{\rm r}^{\theta} = \sum v H_{\rm f}^{\theta} (\text{products}) - \sum v H_{\rm f}^{\theta} (\text{reactants})$ $H_2(g) + \frac{1}{2}O_2(g) \rightarrow H_2O(l) \Delta H^{\theta} = -286 kJ$



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Conclusions

- Conclusion
 - First Law of Thermodynamics is also known as the law of conversation of energy
 - State that energy can be neither created nor destroyed.
 - Energy can change forms, and total energy can flow from one place to another place
 - Provide a sound basis to study between different forms of energy and their interactions



Authors Information

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