# PHYSICS 7B: Physics for Scientists and Engineers

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### Chapter 0

## Syllabus

#### 0.1 Important Info

Syllabus schedule here

#### 0.1.1 Contacts + Sections

Instructor OH:

- 1. Robert Birgeneau; robertjb@berkeley.edu; Tu 2-3pm
- 2. Dr.Chen/Giles-Donovan OH are Th 2-3pm.

Lecture 1: MWF 11-12p @ Physics North 1

Head GSI: Yucehn Tang; ytang5@berkeley.edu; begin emails with [7B Lec.1]

2 Sections: DIS + LAB; Labs not every week and may be replaced by DIS (check syllabus)

#### 0.1.2 Materials

- 1. 7B Workbook by Hedeman (on bCourses)
- 2. Mastering Physics (Purchase access code)
- 3. Physics for scientists and engineers 5th edition by Giancoli (may come  $\mathbf{w}/$  mastering physics purchase)

#### 0.1.3 Assignments

Homework will be made available Friday 11:59 and due next Friday 11:59.

Must complete all 4 Labs - Sep 29; Oct 27; 11/17; 12/1.

Midterms - Pimentel 1 - 9/25 8-10pm + 10/30 8-10pm.

Final - TBD.

### Chapter 17

## Temperature, Thermal Expansion, and the Ideal Gas Law

#### 17.1 Atomic Theory of Matter

**Definition 1** (Unified Atomic Mass Units (u)). We define  $^{12}C$  to have exactly 12.0000 unified atomic mass units (u) such that  $1 u = 1.6605 \times 10^{-27}$  kg.

**Definition 2** (Elements, Compounds, Atoms, Molecules). *Elements* are substances that cannot be broken down into simpler substances by chemical means, *compounds* are substances made up of elements that can be broken down, *atoms* are the smallest pieces of an element, and *molecules* are the smallest pieces of compounds made up of atoms.

**Proposition 1.** To convert between Celsius and Fahrenheit, use:

$$T(^{\circ}C) = \frac{5}{9}[T(^{\circ}F) - 32] \quad T(^{\circ}F) = \frac{9}{5}[T(^{\circ}C) + 32]$$

**Remark.** Different materials do not expand precisely linearly over a wide temerpature rage. Thus, we standardize with the *constant-volume gas thermometer*. This thermometer consists of a hollow rigid bulb with a low-pressure gas connected by a thin tube to a mercury manometer. If the height of the mercury tube is adjusted to ensure the gas maintains a constant volume, the new height reached by the mercury is the temperature.

**Definition 3** (Freezing/Boiling Point). The *freezing point* of a substance is defined as that temperature at which the solid and liquid phases coexist

in equilibrium s.t. there is no net liquid or solid changing into the other one. The *boiling point* is defined analogously for liquid and gas. These temperatures vary with pressure so pressure is specified usually at 1 atm.

**Definition 4** (Thermal Equilibrium). Two objects are defined to be in *thermal equilibrium* is, when placed in contact, no net energy flows from to the other, and their temperatures don't change.

**Remark.** When two systems are in thermal equilibrium, their temperatures are (by definition) equal and no net energy is exchanged.

**Definition 5** (0th Law of Thermodynamics). Specifically experiments indicate that if two systems are in thermal equilibrium with a third system, then they are in thermal equilibrium with each other.

**Proposition 2** (Thermal Expansion). The thermal expansion in length, area, and volume of a material at a fixed pressure due to change in temperature  $\Delta T$  is approximately given by:

$$\Delta l = \alpha l_0 \Delta T$$
  $\Delta A = \gamma A_0 \Delta T$   $\Delta V = \beta V_0 \Delta T$ 

where  $\alpha, \gamma, \beta$  denote the material's coefficient of linear expansion with unit  $(C^{\circ})^{-1}$  and  $l_0, A_0, V_0$  denote its initial length, area, and volume.

**Proof.** At a certain temperature, a thin rod of length  $l_0$  at temperature  $T_0$  is heated by  $\Delta T$  to a temperature T. This causes it to expand by  $\Delta l$  to a length l, or l(T), that is dependent on the original length  $l_0$ , the temperature change  $\Delta T$ , and its coefficient of linear expansion  $\alpha$ . Thus,  $\Delta l = \alpha l_0 \Delta T$  so  $l = l_0 (1 + \alpha \Delta T)$ . For a thin plate with area  $A_0$ , this becomes  $A = A_0 + \Delta A = A_0 (1 + \alpha \Delta T)^2 = A_0 + 2\alpha A_0 \Delta T + \alpha^2 A_0 \Delta T^2$ . Thermal expansion for volume is similarly  $V = V_0 + \Delta V = V_0 (1 + \alpha \Delta T)^3 = V_0 + 3\alpha V_0 \Delta T + 3\alpha^2 V_0 \Delta T^2 + \alpha^3 V_0 \Delta T^3$ .

Because  $\alpha$  and (usually) $\Delta T$  are extremely small, we say  $\Delta A \simeq 2\alpha A_0 \Delta T$  and  $\Delta V \simeq 3\alpha V_0 \Delta T$  implying the coefficient for area  $\gamma \simeq 2\alpha$  and the coefficient for volume  $\beta \simeq 3\alpha$ .

**Note.** If water at  $0^{\circ}$ C is heated, its volume *decreases* until it reaches  $4^{\circ}$ C when it behaves normally and expands as the temperature increases.

This implies that above  $4^{\circ}$ C the surface water in a lake/river in contact with cold air sinks because it is denser bringing in warmer water from below causing convection to bring the whole body to the same temperature. As water cools below  $4^{\circ}$ C, however, its volume expands so its density decreases meaning the the surface water turning to ice is less dense than the water below so a layer of ice forms on top. The ice acts as an insulator, allowing life to exist under ice.

**Definition 6** (Thermal Stresses). If 2 ends of a material are rigidly fixed, temperature change can cause compressive or tensile stresses called *thermal stresses*.

If the beam tries to expand by  $\Delta \ell$  while rigid braces exert a force to hold the beam in place, compressing OR expanding it, the force required is  $\Delta \ell = \frac{1}{E} \frac{F}{A} \ell_0$  where E is Young's modulus for the material. Substituting the thermal expansion equation gives the stress to be  $\frac{F}{A} = \alpha E \Delta T$ .

**Definition 7** (State Variables). Quantities detectable by instruments are called *state variables*. For a gas in a container, they are pressure P, volume V, temperature T, and quantity of gas – mass m or equivalently moles.

**Definition 8** (Mole). One *mole*, abbreviated mol, is defined to be the number of atoms in exactly 12 g of  $^{12}$ C. This number  $N_A$  is called *Avogadro's number* and equal to  $6.022 \times 10^{23}$ .

**Definition 9** (Equilibrium States). When the state variables of a system are not chaning in time and equal throughout the system.

**Definition 10** (Absolute Zero). Absolute zero of temperature is -273.15 °C or 0 K on the Kelvin scale s.t.  $T(K) = T(^{\circ}C) + 273.15$ .

**Proposition 3** (Gas Laws). For a fixed quantity of gas, the following 'laws' are valid (as long as the pressure and density are not too great and the gas is not too close to condensating):

- 1. (Boyle's Law) [At constant temperature],  $P_1V_1 = P_2V_2$  and  $P \propto \frac{1}{V}$ .
- 2. (Charles's Law) [At constant pressure],  $\frac{V_1}{T_1} = \frac{V_2}{T_2}$  s.t.  $V \propto T$ .
- 3. (Gay-Lussac's Law) [At constant volume],  $\frac{P_1}{T_1}=\frac{P_2}{T_2}$  s.t.  $P\propto T.$

Together these imply  $PV \propto T$  so  $\frac{P_1V_1}{T_1} = \frac{P_2V_2}{T_2}$ .

**Definition 11** ((Equation of State for an) Ideal Gas Law). Of course, once mass is left unfixed, experiments show  $PV \propto mT$ . Specifically,

$$PV = nRT$$

where n is the number of moles of gas and R is the universal gas constant found to be roughly  $8.314 \frac{\text{J}}{\text{mol.K}} = 0.0821 \frac{\text{L.atm}}{\text{mol.K}} = 1.99 \frac{\text{calories}}{\text{mol K}}$ .

**Remark** (Important Remarks for Ideal Gas Law). Always give T in kelvins and P as absolute, rather than gauge, pressure. The equation is less 'ideal'

for gases at high pressure/density or near the boiling/condensation point. Recall  $1\,L=1\times 10^{-3}\,\rm m^3.$ 

**Definition 12** (Standard Temperature and Pressure). Standard temperature and pressure, abbreviated STP, implies  $T=273.15K(=0^{\circ} \text{ C})$  and  $P=1.00 \text{ atm} = 1.013 \times 10^{5} \text{ Nm}^{2} = 101.3 \text{ kPa}$ .

Note. 1.00 mol of gas at STP has  $V=22.4\,\mathrm{L}.$ 

**Remark** (Avogadro's Hypothesis). Amedeo Avogadro proposed equal volumes of gas at the same pressure and temperature contain equal numbers of molecules. In other words, he proposed R is the same for all gases.

**Definition 13** (Boltzmann constant). Because  $N_A$  is constant, we can write  $PV = \frac{N}{N_A}RT$  where N is the total number of molecules in a gas. Setting the  $Boltzmann\ constant\ k = R/N_A = 1.38 \times 10^{-23}\ \mathrm{J/K}$  gives PV = NkT.

**Definition 14** (Ideal Gas Temperature Scale). The *ideal gas temperature scale* is based on the property of an ideal gas that pressure is directly proportional to the absolute temperature. Real gases approach this ideal at very low density. This scale takes points P=0 at T=0 K and the *triple point* of water when its solid, liquid, and gas states coexist in equilibrium at P=4.58 torr and T=0.01 °C = 372.16 K.

**Definition 15** (Absolute/Kelvin Temperature). The absolute temperature T of a substance is determind by putting that substance in good thermal contact with a constant-volume gas thermometer s.t., at constant volume,  $T = (273.16 \,\mathrm{K}) \lim P_{tp} \to 0 (\frac{P}{P_{tp}})$ .

Here,  $P_{tp}$  denotes the pressure of the gas in the rigid bulb of the thermometer when placed in water at triple point and P is the pressure in the thermometer when it's in contact with the substance determining T.

### Chapter 18

## Kinetic Theory of Gases

# 18.1 The Ideal Gas Law and the Molecular Interpretation of Temperature

**Definition 16** (Kinetic Theory). The analysis of matter in terms of atoms in continuous random motion is called *kinetic theory*.

**Proposition 4** (Postulates of Kinetic Theory). Under these conditions describing an 'ideal gas', real gases follow the ideal gas law quite closely.

- 1. There are a large number, N, of molecules, each of mass m, moving in random directions with a variety of speeds.
- 2. The molecules are, on average, far apart from one another. I.e. their average separation is much greater than their diameter.
- 3. The molecules obey the laws of classical mechanics and only interact when they collide s.t. the potential energy relating to attractive forces between them is much weaker than the kinetic energy.
- 4. Collisions with another molecule or the wall of the vessel are perfectly elastic and of very short duration relative to time between collisions.

**Proposition 5** (Temperature to Average Kinetic Energy of Molecules). The average translational kinetic energy of molecules in random motion in an ideal gas is directly proportional to the absolute temperature of the gas. In other words,  $\overline{K} = \frac{1}{2}m\overline{v^2} = \frac{3}{2}kT$ .

**Proof.** Take a box of length  $\ell$  and ends of area A filled with an ideal gas. Focus on a single molecule of mass m's collision with one wall. Newton's 2nd and 3rd laws tell us a force  $F = \frac{dp}{dt}$  is exerted on the molecule. Because collisions are elastic, its velocity  $v_x$  is equal in magnitude so  $\Delta p = 2mv_x$ . If the molecule takes time  $\Delta t$  to travel  $2\ell$  to the other wall and back,

 $F = \frac{\Delta(mv)}{\Delta t} = \frac{2mv_x}{2\ell/v_x} = \frac{mv_x^2}{\ell}$ . Recall that although the particle may collide with the tops and sides of the container, its x component doesn't change and neither does its momentum.

We can average the force on a wall due to all the N molecules in the box by  $F=\frac{m}{\ell}(v_{x1}^2+v_{x2}^2+\cdots+v_{xN}^2)$ . Averaging,  $\overline{v_x^2}=\frac{v_{x1}^2+v_{x2}^2+\cdots+v_{xN}^2}{N}$  s.t.  $F=\frac{m}{\ell}N\overline{v_x^2}$ . Because  $v^2=v_x^2+v_y^2+v_z^2\simeq 3v_x^2$ , we let  $F=\frac{m}{\ell}\frac{N\overline{v^2}}{3}$ . Thus, the pressure on the wall is  $P=\frac{F}{A}=\frac{1}{3}\frac{Nm\overline{v^2}}{A\ell}=\frac{1}{3}\frac{Nm\overline{v^2}}{V}\Rightarrow PV=\frac{2}{3}N(\frac{1}{2}m\overline{v^2})$ . From the ideal gas law PV=NkT, such that  $\frac{3}{2}kT=\frac{1}{2}(m\overline{v^2})=\overline{KE}$ .  $\square$ 

**Definition 17** (Thermal Motion). This definition explains the relationship between temperature as a measure of motion of molecules such that random motion of a gas is sometimes called *thermal motion*.

**Definition 18** (Root-Mean-Square Speed  $v_{rms}$ ). To calculate how fast moleciles molecules move on average in an ideal gas, we can derive the *root-mean-square speed*  $v_{rms} = \sqrt{\overline{v^2}} = \sqrt{\frac{3kT}{m}}$ .

**Remark.** It's noteworthy that the average speed  $\overline{v} \neq v_{rms}$  necessarily. In fact, for an ideal gas, they differ by about 8%.

**Note.**  $\overline{KE} = \frac{3}{2}kT$  tells us as  $T \to 0$ ,  $\overline{KE} \to 0$ . However, modern quantum theory tells us this is not true and kinetic energy appraoches a small nonzero minimum.

#### 18.2 Distribution of Molecular Speeds

**Definition 19** (Maxwell Distribution of Speeds). In 1859, James Maxwell worked out a formula for the most probable distribution of speeds in a gas with N molecules, that is

$$f(v) = 4\pi N \left(\frac{m}{2\pi kT}\right)^{\frac{3}{2}} v^2 e^{-\frac{1}{2}\frac{mv^2}{kT}},$$

where f(v)dv represents the number of molecules that have speeds between v and v+dv such that  $\int_0^\infty f(v)dv=N$ .

**Definition 20** (Activation Energy). Two molecules may chemically react only if their kinetic energy is great enough to partially penetrate each other. This minimum energy is called *activation energy*. The rate of a chemical reaction is proportional to the number of molecules with energy greater than  $E_A$  such that rates increase rapidly with increased temperature.

**Example** (Determining  $\overline{v}$  and  $\overrightarrow{v_p}$ ). To find the average speed  $\overline{v}$ , we must integrate over the product of v and the number f(v)dv which have speed v and divide by N the number of molecules. Thus,  $\overline{v} = \frac{\int_0^\infty v f(v) dv}{N} = 4\pi (\frac{m}{2\pi kT})^{\frac{3}{2}} \int_0^\infty v^3 e^{-\frac{1}{2}\frac{mv^2}{kT}} dv = 4\pi (\frac{m}{2\pi kT})^{\frac{3}{2}} (\frac{2k^2T^2}{m}) = \sqrt{\frac{8}{\pi}\frac{kT}{m}}.$ 

To find the most probable speed, we need simply find when the slope is 0 such that  $\frac{df(v)}{dv}=4\pi N(\frac{m}{2\pi kT})^{\frac{3}{2}}(2ve^{-\frac{mv^2}{2kT}}-\frac{2mv^3}{2kT}e^{-\frac{mv^2}{2kT}})=0$ . Solving for v gives  $v_p=\sqrt{\frac{2kT}{m}}$ .

Note. In summary,

$$v_p \approx 1.41 \sqrt{\frac{kT}{m}}, \quad \overline{v} \approx 1.60 \sqrt{\frac{kT}{m}}, \quad v_{rms} \approx 1.73 \sqrt{\frac{kT}{m}}.$$

#### 18.3 Real Gases and Changes of Phase

**Remark.** At high pressure, the volume of a real gas is *less* than that predicted by the ideal gas law.

**Explanation.** This is because ideal gases assume the potential energy originating from attractive forces between molecules to be negligible relative to their kinetic energy. This energy pulls molecules closer together so the volume decreases. At lower temperatures, these forces result in *liquefaction* or *condensation*.

#### 18.4 Vapor Pressure and Humidity

**Definition 21** (Critical Point/Temperature). At a certain *critical temperature*, a gas will change to liquid phase if sufficient pressure is applied. This is also called the *critical point*.

**Definition 22** (Gas and Vapor). A substance in a gaseous state *below* its critical temperature is called a *vapor*; above the critical temperature it is called a *gas*.

**Remark** (Phase Diagram). A *PT Diagram* is often called a *phase diagram* because it compares the different phases of a substance.

**Definition 23** (Sublimation). Sublimation refers to the process wherein, at low pressures, a solid changes directly into the vapor phases without passing through the liquid phase.

#### 18.5 Partial Pressure and Humidity

Remark (Cooling Process). Molecules in a liquid roughly follow the Maxwell distribution. Molecules of high speeds in a liquid – i.e.  $E_A$  – may leave the liquid temporarily but be pulled back by attractive forces. Only those of highest speed *evaporate*, ultimately decreasing the average speed such that the absolute temperature is less. Kinetic theory, then, predicts evaporation is a *cooling process*.

**Definition 24** (Saturated Vapor Pressure). At equilibrium, an equal number of molecules enter the vapor above a liquid and enter the liquid. At this point, the pressure of the vapor is said to be *saturated*.

**Remark.** The concentration of particular molecules in the gas phase above the liquid will not affect the saturated vapor pressure, but will lengthen the amount of time to reach equilibrium due to collisions.

**Remark.** Increased temperature increases the saturated vapor pressure of a liquid until it equals the external pressure and *boiling* occurs. Bubbles which indicate a change from liquid to the gas phase are crushed if vapor pressure inside the bubbles is less than the external pressure.

**Definition 25** (Partial Pressure). Partial pressure is the pressure each gas would exert it it alone were present in a mixture. The relative humidity is defined to be the ratio of the partial pressure of water vapor to the saturated vapor pressure of water at a given temperature. If the partial pressure of water exceeds the saturated vapor pressure, the air is supersaturated. This excess water may appear as dew or clouds.

**Definition 26** (Dew Point). The *dew point* is when water is cooled s.t. the saturated vapor pressure of water equals its partial pressure.

**Note.** The pressure P of the amosphere as a function altitude y above sea level is, where  $P_0$  is 1.00 atm:

$$P = P_0 e^{-(\rho_0 g/P_0)y} = P_0 e^{-(1.25 \times 10^{-4} m^{-1})}$$

#### 18.6 van der Waals Equation of State

**Theorem 1** (van der Waals Equation of State). Given gas-dependent constants a and b,

$$(P + \frac{a}{(V/n)^2})(\frac{V}{n} - b) = RT.$$

**Proof.** To make the ideal gas law more accurate, we now resolve (1) the finite size of molecules in comparison to one another and the container and (2) forces between molecules may be greater than the size of molecules.

Let the molecules in a gas behave like hard spheres of radius r such that the volume molecules can move around in is less than the volume V of the container because the distance between molecules never shrinks below 2r. Where b is the unavailable volume per mole of gas, replace V in PV = nRT by (V - nb). This relation,  $P(\frac{V}{n} - b) = RT$  is called the *Clausius equation of state* and predicts ideal gases have less pressure than real gases. Next, molecules at the edge of the gas leaving towards a wall are slowed by a net attractive force pulling them back in, thus exerting less force and pressure on the wall. We say this pressure is proportionally reduced by the density squared, or  $(n/V)^2$ , for constant a s.t.  $P = \frac{RT}{(V/n)-b} - \frac{a}{(V/n)^2}$  or

$$(P + \frac{a}{(V/n)^2})(\frac{V}{n} - b) = RT.$$

Note that at low densities, van der Waals reduces to the ideal gas law.  $\qed$ 

#### 18.7 Mean Free Path

**Definition 27** (Mean Free Path). We define the *mean free path*,  $\ell_M$  to be the average distance a molecule travels between collisions such that

$$\ell_M = \frac{1}{4\pi\sqrt{2}r^2(N/V)}.$$

**Proof.** Suppose the molecules of a gas are hard spheres of radius r. Let the path of one molecule with mean speed  $\overline{v}$  be a cylinder of radius 2r such that if another molecule's center lies in the cylinder a collision will occur. We can assume for now the other molecules are not moving and the concentration of molecules is N/V. Then,  $V_cylinder*N/V$  is the number of collisions that will occur. Over a time  $\Delta t$ , the molecule travels  $\overline{v}\Delta t$  so the number of collisions is expected to be  $\pi(2r)^2\overline{v}\Delta t(N/V)$ . Thus, the average distance between collisions is

$$\ell_M = \frac{\overline{v}\Delta t}{\pi(2r)^2\overline{v}\Delta t(N/V)} = \frac{1}{4\pi r^2(N/V)}.$$

If the other molecules are moving, the number of collisions in  $\Delta t$  actually depends on the *relative* speed  $v_r e l \approx \sqrt{2} \overline{v}$  of the colliding molecules so

$$\ell_M = \frac{1}{4\sqrt{2}\pi r^2(N/V)}$$

. This of course loses meaning at low densities.

#### 18.8 Diffusion

**Definition 28** (Diffusion). In general, the *diffusing* substance moves from a region where its concentration is high to a region where its concentration is low.

Definition 29 (Fick's Law). Consider a tube of cross-sectional area A containg molecules of increasingly smaller concentration left-to-right. Take a small middle section of tube of length  $\Delta x$  with section 1 on the left and 2 on the right. 1 has more molecules causing greater pressure such that more will strike the boundary into the middle section than from 2. Let J be the rate of diffusion in numbehr of molecules/moles/kg per second. This is proportional, with diffusion constant D to the concentration gradient or difference in concentration per unit distance  $\frac{C_1-C_2}{\Delta x}$ . Thus,  $J=DA\frac{C_1-C_2}{\Delta x}=DA\frac{dC}{dx}$ . This is Fick's Law.

### Chapter 19

# Heat and the First Law of Thermodynamics

#### 19.1 Heat as Energy Transfer

**Remark.** An 18-th century model pictured heatflow as movement of a fluid substance called *caloric* which could never be detected. Today, heat, like work, represents a transfer of energy.

**Definition 30** (Calorie). A calorie (cal) is defined as the amount of heat necessary to raise the temperature of 1 gram of water by 1 Celsius degree. The more often used kilocalorie (kcal), or Calorie is 1000 calories. In the UK, heat is measured through British thermal units (Btu) defined as the heat needed to raise the temperature of 1 lb of water by 1 F°. Gas companies use the therm defined to be  $1 \times 10^5$  Btu.

Note (Mechanical Equivalent of Heat). The mechanical equivalent oheat is known as  $4.186\,\mathrm{J} = 1\,\mathrm{cal}$ . And,  $1\,\mathrm{Btu} = 0.252\,\mathrm{kcal} = 1056\,\mathrm{J}$ .

**Definition 31** (Heat). *Heat* is energy transferred from one object to another because of a difference in temperature.

**Definition 32** (Internal/Thermal Energy). The sum total of all the neergy of the molecules in an object is called its *internal/thermal energy*. e.g. 2 objects of equal temperature have more internal temperature than just 1.

For monatomic (one atom/molecule) gas, the internal energy

$$E_{int} = N(\frac{1}{2}\bar{v^2}) = \frac{3}{2}NkT = \frac{3}{2}nRT.$$

If the gas molecules contain more than one atom, the rotaional and vibrational energy of the molecules must also be taken into account.

**Remark.** Internal energy of real gases, as opposed to ideal gases, also depends on pressure and volume (due to atomic potential energy). Similarly, the internal energy of liquids and solids includes electric potential energy associated with chemical bonds.

#### 19.2 Specific Heat

**Definition 33** (Specific Heat). An amount of heat Q put into an object of mass m results, depending on its *specific heat* c, in a temperature change of  $\Delta T$  such that

$$Q = mc\Delta T$$
.

**Definition 34** (Closed/Open/Isolated Systems). A *closed system* is one for which no mass enters or leaves, but energy may be exchanged with the environment. Mass may enter or leave in an *open system*. If no energy AND no mass passes its boundaries, a system is *isolated*.

**Remark** (Energy Conservation for Heat Transfer). Within an isolated system, we can write the energy conservation equation for heat transfer as  $\Sigma Q = 0$ .

**Definition 35** (Calorimetry). Calorimetry is a technique which quantitatively measures heat exchange using a calorimeter that must be well insulated.

**Definition 36** (Bomb Calorimeter). A bomb calorimeter measures the thermal energy released when a substance burns to determine their Calorie content. In this calorimeter, a carefully weighed sample is placed with an excess of oxygen in the "bomb" that is then placed in the water of a calorimeter with a fine wire passing into the bomb that ignites the mixture.

#### 19.3 Latent Heat

**Definition 37** (Change of Phase). A certain amount of energy is required to change a material's phase.

**Definition 38** (Heat of Fusion/Vaporization). The heat required to change 1.0 kg of a substance from the solid to liquid state is called the *heat of fusion*, denoted by  $L_F$ . The heat required for the change from liquid to the vapor phase is called the *heat of vaporization*  $L_V$ . These are also called the *latent heats* of substances.

The heat involved in a change of phase is written as Q = mL where m is the mass of the substance.

**Remark.** These latent heats also refer to the amount of heat *released* by a substance when it changes from gas to liquid or liquid to solid.

Remark. At the melting point, the latent heat of fusion does not increase the average kinetic energy (and thus the temperature) of the molecules in a solid, but instead overcomes the potential energy from the forces between molecules. This is the same for vaporization. However, vaporization requires a greater average distance between molecules such that the heat of vaporization is far greater than the heat of fusion for a given substance.

#### 19.4 The First Law of Thermodynamics

**Definition 39** (The First Law of Thermodynamics). Because heat is defined to be the transfer of energy due to a difference in temperature, let work be the transfer of energy NOT due to a temperature difference such that, as a general statement of the law of conservation of energy  $\Delta E_{int} = Q - W$ .

**Note** (Sign Conventions). Because Q is the net heat added to the system while W is the net work done by the system, we say the following:

- 1. Heat added is +
- 2. Heat lost is -
- 3. Work on system is -
- 4. Work by system is +

**Remark.** A full summary of the first law of thermodynamics would include kinetic energy K and potential energy U such that

$$\Delta K + \Delta U + \Delta E_{int} = Q - W.$$

Definition 40 (Idealized Processes). An idealized process carried out at:

- 1. constant temperature ( $\Delta T = 0$ ) is called *isothermal*.
- 2. constant heat (Q=0) is called *adiabatic*. (This is the case for well-insulated systems or very rapid processes.)
- 3. constant pressure ( $\Delta P = 0$ ) is called *isobaric*.
- 4. constant volume ( $\Delta V=0$ ) is called isovolumetric (or isochoric).

**Definition 41** (Isotherms). Curves on a PV diagram at different temperatures are called *isotherms*.

**Definition 42** (Quasistatically). A process done slowly enough such that a series of equilibrium states are essentially maintained is called *quasistatically*.

**Definition 43** (Heat Reservoir). A body whose mass is so large that, ideally, its temperature does not change significantly when heat is exchanged, is called a *heat reservoir*.

**Example.** Suppose there is a gas of pressure P inside a cylindrical container of area A and length  $\ell$  fitted with a movable piston.

If our system is just the gas, as it exerts a force F = PA on the piston, the work done by the gas is  $dW = \mathbf{F} \cdot d\ell = PAd\ell = PdV$  where  $d\mathbf{l}$  is an infinitesimal displacement of the piston.

If the gas was compressed so  $d\ell$  would point into the gas, its volume would decrease and dV < 0. Thus, the work done by the gas would be negative s.t. positive work is being done on the gas. For a finite change in volume from  $V_A$  to  $V_B$ , the work W done by the gas will be  $W = \int dW = \int_{V_A}^{V_B} P dV$ .

For an quasistatic isothermal expansion of an ideal gas, P = nRT/V for constant T tells us  $W = nRT \ln \frac{V_B}{V_A}$ . This is the area under the curve between points A and B on a PV diagram.

We can replicate this same change of state from A to B for an ideal gas first isovolumetrically then isobarically. For an ideal gas undergoing an isovolumetric process where we lower the pressure from  $P_A \to P_B$ , we simply reduce the temperature by letting heat flow out  $(dV=0\Rightarrow W=0$  so no work is done). For an ideal gas undergoing an isobaric process taking the volume from  $V_A \to V_B$ , P is constant at  $P_B$  so  $W=P_B\Delta V=P_B(V_B-V_A)=\frac{nRT_B}{V_B}(V_B-V_A)=nRT_B(1-\frac{V_A}{V_B})$ . Thus, the total work from A to B in this case was  $\frac{nRT_B}{V_B}(V_B-V_A)$  which is quantitatively different from  $nRT \ln \frac{V_B}{V_A}$ .

**Definition 44** (Free Expansion). Free expansion is a type of adiabatic process when a gas is allowed to expand in volume without doing any work. Picture 2 well-insulated containers connected by a valve such taht one container has gas while the other is empty. When the valve is open, no heat flows or work is done because no objects are moved so  $\Delta E_{int} = Q = W = 0$ .  $E_{int}$  only depends on T so  $\Delta T = 0$  as well.

#### 19.5 Molar Specific Heats for Gases

**Remark.** Specific heats easily apply to solids and liquids. Gases depend much more on how the thermodynamic process is carried out however. Specific heats are given at constant volume  $c_V$  and pressure  $c_P$ .

**Definition 45** (Molar Specific Heats). Molar specific heats  $C_V$  and  $C_P$  denote the heat required to raise 1 mol of gas by 1 C° at constant volume and pressure respectively. Thus,

$$Q = nC_V \Delta T$$
 [volume constant]  
 $Q = nC_P \Delta T$  [pressure constant]

where  $C_V = Mc_V$  and  $C_P = Mc_P$  where M is the molecular mass of the gas (g/mol).

**Remark.** Imagine an ideal gas is slowly heated by  $\Delta T$ , first at constant volume, then constant pressure. In the constant volume process, no work is done since  $\Delta V = 0$ . Thus all the heat  $Q_V$  added goes towards increasing internal energy such that  $Q_V = \Delta E_{int}$ .

However, in the constant pressure process, work is done so the added heat  $Q_P$  increases the internal energy and does the work  $W = P\Delta V_P$  so  $\Delta E_{int} = Q_P - P\Delta V_P$ .

Because  $\Delta E_{int}$  is equal in both processes ( $\Delta T_V = \Delta T_P$ ),  $Q_P - Q_V = P\Delta V \Rightarrow nC_P\Delta T - nC_V\Delta T = P(\frac{nR\Delta T}{P})$  from the ideal gas law. This implies  $C_P - C_V = R$  which is very accurate to what's obtained experimentally.

**Remark.** Now, a process done at constant volume on a monatomic gas does no work so  $nC_V\Delta T = nC_V(T-0) = Q_V = \Delta E_{int} = N(\frac{1}{2}m\overline{v^2}) = \frac{3}{2}nRT$ . So  $C_V = \frac{3}{2}R$ .

**Definition 46** (Degrees of Freedom). Here, we denote *degrees of freedom* to mean the number of independent way molecules can possess energy.

**Example.** For instance, a diatomic molecule has 5 total degrees of freedom. 3 from translational energy (x,y,z) plus 2 from rotaional kinetic energy (not 3 because the axis along the line of the 2 molecules has such small inertia it is negligible).

**Definition 47** (Principle of Quipartition of Energy). Energy is shared equally among the active degrees of freedom and each active degree of a molecule has on average an energy equal to  $\frac{1}{2}kT$ .

Note. This makes sense as diatomic molecules have energy  $\frac{5}{2}nRT$  about 5/3 times monatomic molecules without degrees of freedom. Hence their  $C_V$  being 5/3 as much. Yet, at extreme low and high temperatures, this diverges such that at low temps, molecules only have translational kinetic energy (3/2). And at high temps, molecules also have vibration as 2 degree of freedom (as if from a spring) such that it has kinetic and potential energy (7/2).

**Note.** For solids, we maintain this same spring idea such that molecules have potential and kinetic energy to do with vibration in the x, y, z directions implying 6 degrees of freedom.

#### 19.6 Adiabatic Expansion of a Gas

**Remark.** Take the first law of thermodynamics in an adiabatic process  $(\Delta Q=0)$  for an ideal gas such that  $dE_{int}=dQ-dW=-dW=-PdV$ . For an ideal gas,  $\Delta E_{int}=nC_V\Delta T$  tells us  $nC_vdT=-PdV$ . Taking the differential of the ideal gas law gives PdV+VdP=nRdT so  $nC_V(\frac{PdV+VdP}{nR})+PdV=0$ . Rearranging and  $C_V+R=C_P$  gives  $C_PPdV+C_VVdP=0$  or  $\frac{C_P}{C_V}PdV+VdP=0$ . Defining  $\gamma=\frac{C_P}{C_V}$  and integrating finally gives  $\ln(P)+\gamma\ln(V)=$  constant which simplifies to

$$PV^{\gamma} = \text{constant}.$$

It's important to note the ideal gas law holds for an adiabatic expansion however PV is clearly not constant implying T must be nonconstant.

#### 19.7 Heat Transfer

**Definition 48** (Conduction). Heat *conduction* can be visualized via molecular collisions such that faster molecules at a heated end collide with slower-moving neighbors and transfer kinetic energy. In metals, collisions of free electrons are mainly responsible for conduction.

Remark (Thermal Conductivity). Take a uniform cylinder of cross-sectional area A and length  $\ell$  such that ends with temperatures  $T_1$  and  $T_2$  with heat flow Q over a time interval t gives  $\frac{Q}{t} = kA\frac{T_1 - T_2}{\ell}$  where k is the thermal conductivity constant characteristic of the material. The rate of heat flow (J/s) is directly proportional to  $(T_1 - T_2)/\ell$ .

**Definition 49** (Thermal Gradient). When k or A cannot be considered constant, we instead take the limit of an infinitesimally thin slab of thickness dx such that our equation becomes  $\frac{dQ}{dt} = -kA\frac{dT}{dx}$  where dT/dx is the temperature gradient. Here, the negative sign denotes that heat flow is in direction opposite to the temperature gradient.

**Definition 50** (Conductors/Insulators). Substances for which k is very large conduct heat rapidly and are called good thermal *conductors*. When k is small, the substances are called good thermal *insulators*.

**Definition 51** (Thermal Resistance). The insulating properties of bulding materials are usually specified by R-values or thermal resistance, defined as  $R = \frac{\ell}{k}$  for a thickness  $\ell$  of a material. Larger R values imply better insulation.

**Definition 52** (Convection). Though liquids and gases are poor conductors of heat, *Convection* is the rapid process whereby heat flows via the bulk movement of molecules over larger distances.

**Definition 53** (Forced/Natural Convection). For example, *Forced convection* occurs through a fan blowing air while *natural convection* occurs through hot air rising. Hot air or water (in convection currents) rise from buoyancy because heat causes them to expand, decreasing their relative density.

**Definition 54** (Radiation). Radiation is heat transferred over empty space through EM waves. IR wavelengths are mainly responsible for heating the Earth.

**Definition 55** (Stefan-Boltzmann equation). The rate at which energy leaves a radiating object is  $\frac{Q}{t} = \varepsilon \sigma A T^4$  where  $\sigma$  is the *Stefan-Boltzmann constant* with value  $5.67 \times 10^{-8} \, \text{W/m}^2 \cdot \text{K}^4$ . The factor  $\varepsilon$ , or the *emissivity*, falls between 0 and 1 and is characteristic of the surface of the radiating material.

**Remark.** A good absorber is *also* a good emitter. Black and dark objects are good emitters with  $\varepsilon \approx 1$ .

**Example.** Take an object of emissivity  $\varepsilon$  and area A at temperature  $T_1$  surrounded by an environment at temperature  $T_2$ . Because both objects radiate energy to each other so the object absorbs energy proportional to  $T_2^4$ , the *net* rate of radiant heat flow from the object is given by  $\frac{Q}{t} = \varepsilon \sigma A(T_1^4 - T_2^4)$ .

**Definition 56** (Solar Constant). About 1350 J of energy from the Sun strike Earth's atmospher per second per square meter at right angles to the Sun's rays. This number 1350 W/m² is called the solar constant. The atmosphere may absorb as much as 70% of this energy such that an object of emissivity  $\varepsilon$  with area A facing the Sun absorbs at a rate about  $\frac{Q}{t} = (1000 \, \text{W/m}^2) \varepsilon A \cos(\theta)$  where  $\theta$  is the angle between the Sun's rays and a line perpendicular to the area A. Seasons result from this "effective"

area" $\cos(\theta)$  capture.

**Definition 57** (Thermography). Diagnostic *thermography* uses a thermograph to scan the body and measure the intensity of infrared radiation to detect areas of high metabolic activity.

### Chapter 20

# Second Law of Thermodynamics

#### 20.1 Introduction

**Definition 58** (Second Law of Thermodynamics - Clausius Statement). Heat can flow spontaneously from a hot object to a cold object; heat will not flow spontaneously from a cold object to a hot object.

**Remark.** Energy could be conserved in a broken cup putting itself back together as it rises, but this is not observed to happen.

#### 20.2 Heat Engines

**Definition 59** (Heat Engine). A *heat engine* is any device that changes thermal energy into mechanical work.

**Note.** Sign convention *only* for heat engines:  $Q_H, Q_L, W > 0$ . where  $Q_H$  heat is input at a high temperature  $T_H$  into a system that then transforms it partly into work W and partly as exhausted as heat  $Q_L$  at a lower temperature  $T_L$ . By conservation of energy,  $Q_H = W + Q_L$ .

**Remark.** Mechanical energy can only be obtained from thermal energy when heat is allowed to flow from a high to low temperature. If the temperature were constant throughout, pressure on the intake and outtake would be equal such that the work done by and on the gas would be equal so net work would be done.

**Definition 60** (Operating Temperatures). The high and low temperatures,  $T_H, T_L$  are the *operating temperatures* of the engine.

**Definition 61** (Working Substance). The material, usually steam, that is heated and cooled is the *working substance*.

**Definition 62** (Efficiency e of any Heat Engine). The efficiency, e, of any heat engine can be defined as the ratio of work done, W to the heat input at the high termperature,  $Q_H$ , over a complete cycle:  $e := \frac{W}{Q_H}$ . Equivalently, This is  $e = \frac{Q_H - Q_L}{Q_H} = 1 - \frac{Q_L}{Q_H}$ .

**Definition 63** (Carnot Engine). An ideal engine, or *Carnot engine*, consists of 4 processes done in a cycle, 2 adiabatic and 2 isothermal done reversibly.

The isothermal processes, where heats  $Q_H$  and  $Q_L$  are transferred assume constant temperatures  $T_H$  and  $T_L$  such that the system is in contact with idealized *heat reservoirs* which are large enough to not fluctuate in temperature.

**Definition 64** (Reversible Processes). Processes done *reversibly* are considered to be carried out infinitely slowly so that the process is a series of equilibrium states so the whole process could be reversed with no change in the magnitude of work done or heat exchanged.

Remark. Real processes are always irreversible.

**Remark.** Suppose a Carnot engine uses an ideal gas. In the first isothermal process ab,  $W_{ab}=nRT_H\ln\frac{V_b}{V_a}$ . This is isothermal so  $E_{int}$  does not change and thus the heat,  $Q_H$  added equals the work done by the gas. So

$$Q_H = nRT_H \ln \frac{V_b}{V_a}.$$

Similarly, the heat lost in the isothermal process cd is

$$Q_L = nRT_L \ln \frac{V_c}{V_d}.$$

The paths bc, da are adiabatic so  $P_bV_b^{\gamma} = P_cV_c^{\gamma}$  and  $P_dV_d^{\gamma} = P_aV_a^{\gamma}$  where  $\gamma = \frac{C_P}{C_V}$ . From the ideal gas law,  $\frac{P_bV_b}{T_H} = \frac{P_cV_c}{T_L}$  and  $\frac{P_dV_d}{T_L} = \frac{P_aV_a}{T_H}$ . Together, this tells us  $T_HV_b^{\gamma-1} = T_LV_c^{\gamma-1}$  and  $T_LV_d^{\gamma-1} = T_HV_a^{\gamma-1}$ . This implies  $(\frac{V_b}{V_a})^{\gamma-1} = (\frac{V_c}{V_d})^{\gamma-1}$  so  $\frac{V_b}{V_a} = \frac{V_c}{V_d}$ . Substituting this into our original equations show

$$\frac{Q_L}{Q_H} = \frac{T_L}{T_H} \qquad \text{[Carnot Cycle]}$$

SO

$$e_{ideal} = 1 - \frac{Q_L}{Q_H} = 1 - \frac{T_L}{T_H}.$$

**Theorem 2** (Carnot's Theorem). For any ideal reversible engine with fixed input and exhaust temperatures  $T_H$  and  $T_L$ , the fundamental upper limit to the efficiency of any heat engine is  $e = 1 - \frac{T_L}{T_H}$ . In other words,

All reversible engines operating between the same two constant temperatures  $T_H$  and  $T_L$  have the same efficiency. Any irreversible engine operating between the same two fixed temperatures will have an efficiency less than this.

**Note.** Real engines that are well designed reach 60 to 80% of the Carnot efficiency.

**Definition 65** (Second Law of Thermodynamics - Kelvin-Planck Statement). No device is possible whose sole effect is to transform a given amount of heat completely into work.

**Definition 66** (Otto Cycle). The operation of an automobile internal combustion engine can be approximated by a reversible cycle called the Otto Cycle wherein the input and exhaust tempeartures are not constant. Paths ab, cd are adiabatic while bc, da are isovolumetric.

**Remark.** For an ideal gas as working substance, the efficiency of an Otto cycle engine is:

$$e = 1 - \left(\frac{V_a}{V_b}\right)^{1-\gamma},$$

where  $V_a/V_b$  is the compression ratio.

**Explanation.** Because the exchange takes place at constant volume,

$$Q_H = nC_V(T_c - T_b)$$
 and  $Q_L = nC_V(T_d - T_a)$ .

Thus,

$$e = 1 - \frac{Q_L}{Q_H} = 1 - \left[\frac{T_d - T_a}{T_c - T_b}\right]. \label{eq:epsilon}$$

From the adibatic processes,  $P_a V_a^{\gamma} = P_b V_b^{\gamma}$  and  $P_c V_c^{\gamma} = P_d V_d^{\gamma}$ . From ideal gas law then,  $T_a V_a^{\gamma-1} = T_b V_b^{\gamma-1}$  and  $T_c V_c^{\gamma-1} = T_d V_d^{\gamma-1}$ . This gives efficiency

$$e = 1 - \left\lceil \frac{(V_b/V_a)^{\gamma - 1}(T_c - T_b)}{T_c - T_b} \right\rceil = 1 - (\frac{V_b}{V_a})^{\gamma - 1} = 1 - (\frac{V_a}{V_b})^{1 - \gamma}.$$

#### 20.3 Refrigerators, Air Conditioners, and Heat Pumps

**Definition 67** (Perfect Refrigerator). A *perfect refrigerator* is one in which no work is required to take heat from the low-temperature region to the high-temperature region. This is impossible

**Remark.** Refrigerators, air conditioners, and heat pumps do the reverse of a heat engine, transferring heat out of a cool environment int oa warm environment. Each does work W to remove heat  $Q_L$  from inside and exhaust heat  $Q_H$  outside. This is usually done by an electric motor which compresses a fluid.

**Definition 68** (The Coefficient of Performance (COP)). The coefficient of performance (COP) of a refrigerator is defined as the ratio between the heat  $Q_L$  removed and the work W done so  $COP = \frac{Q_L}{W} = \frac{Q_L}{Q_H - Q_L}$ . For an ideal refrigerator,  $COP_{ideal} = \frac{T_H}{T_H - T_L}$ .

**Definition 69** (Heat Pump). A heat pump does work W to take heat  $Q_L$  from outside to deliver heat  $Q_H$  inside. Thus its COP is  $\frac{Q_H}{W}$ . Typical heat pumps have COP  $\approx 2.5$  to 3.

Remark (SEER Rating). Cooling devices typically have SEER (Seasonal Energy Efficiency Ratio) ratings defined as  $\frac{\text{(heat removed in Btu)}}{\text{(electrical input in watt-hours)}}$ .

**Definition 70** (Entropy). When an amount of heat Q is added to a system by a reversible process at constant temperature (K), the change in entropy S of the system is given by

$$\Delta S = \frac{Q}{T}.$$

If the temperature is not constant,  $dS = \frac{dQ}{T}$  so

$$\Delta S = \int_a^b dS = \int_a^b \frac{dQ}{dT}$$
 [reversible process].

**Definition 71** (State Variable). Entropy depends only on the states a and b of the system. Thus, it is a state variable.

Remark (Important Remark). The change in entropy is path-independent.

**Remark.** Any reversible cycle can be approximated as a series of Carnot cycles. More cycles leads to better approximations. For each cycle, we can write  $\Sigma \frac{Q}{T} = 0$  so that over the limit of infinitely many Carnot cycles,

we get, as an integral around a closed path:

$$\oint \frac{dQ}{T} = 0 \quad \text{[reversible paths]}.$$

Thus entropy for an irreversible can be calculated for a reversible process between the same two states.

**Note.** One form of the *third law of thermodynamics* states that as  $T \to 0$ ,  $S \to 0$ .

# 20.4 Entropy and the Second Law of Thermodynamics

**Remark.** Given an isolated system of two objects at temperatures  $T_H$  and  $T_L$ , the objects eventually come to some intermediate tempearture  $T_M$  so that the heat  $Q_H$  lost equals -Q and the heat  $Q_L$  gained by the colder object equal Q so the total change is  $\Delta S = \Delta S_H + \Delta S_L = -\frac{Q}{T_{HM}} + \frac{Q}{T_{LM}}$  where  $T_{HM}, T_{LM}$  are the intermediate temperatures.  $T_{HM} > T_{LM}$  at all times so  $\Delta S > 0$ . So the total change is positive. (While a reversible (ideal, impossible) process has  $\Delta S = 0$ .)

**Definition 72** (General Statement - Second Law of Thermodynamics). The entropy of an isolated system never increases. It either stays constant (reversible) or increases (irreversible processes).

The total entropy of any system plus that of its environment increases as a result of any natural process:  $\Delta S = \Delta S_{sys} + \Delta S_{env} > 0$ .

We can also say natural processes tend to move toward a state of greater disorder. Hence, entropy is called time's arrow.

#### 20.5 Order to Disorder

**Note.** The entropy of a system can be considered a measure of the disorder of the system.

**Example.** A stone falling is ordered macroscopically because each molecule is moving at the same velocity in the same direction. However, its fall turns into thermal energy which is more random and disordered.

#### 20.6 Unavailability of Energy; Heat Death

Remark. An important aspect of the 2nd Law of Thermodynamics is that in any natural processes, some energy becomes unavailable to do useful

work. As time goes on, energy is never lost, but is instead degraded.

**Definition 73** (Heat Death). The *heat death* of the universe is the prediction that all energy of the universe will degrade to thermal energy.

#### 20.7 Statistical Interpretation of Entropy

**Definition 74** (Micro/Macrostate). The *microstate* of a system describes the position and velocity of every particle (or molecule) while the *macrostate* of a system describes the properties of a system like temperature, pressure, number of moles, etc.

**Remark.** A basic assumption behind the statistical approach is that each microstate is equally probable. And a great number of microstates can equal the same macrostate.

**Remark.** The most probable state is the one with the greatest entropy or greatest disorder and randomness. In other words, it corresponds to the most *microstates*.

**Definition 75** (Thermodynamic Probability). In consistency with the Clausius statement that dS = dQ/dT, we can say the entropy of a system in a (macro)state is  $S = k \ln \mathcal{W}$  where  $\mathcal{W}$  is the number of corresponding microstates, or the *thermodynamic probability*, of the occurrence of some state.

**Definition 76** (Third Law of Thermodynamics). It is not possible to reach absolute zero in any finite number of processes.

**Remark.** From  $\frac{Q_L}{Q_H} = \frac{T_L}{T_H}$ , we can assign the value of the triple point of water  $T_{tp} = 273.16K$  to the thermodynamic scale such taht  $T = (273.16K) * (\frac{Q}{Q_{tp}})$  where  $Q, Q_{tp}$  are the magnitudes of the heats exchanged by a Carnot engine with such reservoirs. Thus, the thermodynamic scale is identical to the ideal gas scale as long as the ideal gas is valid.

**Note.** Because  $T_L$  of an engine is thus never 0, a 100% efficiency engine such that  $e=1-\frac{T_L}{T_H}=1$  is not possible.

**Note.** Mechanical energy is transformed to electric energy with a turbine and generator.

**Definition 77** (Thermal Pollution). The heat  $Q_L$  output from every heat engine is referred to as *thermal pollution* because the environment must absorb it.

### Chapter 21

# Electric Charge and Electric Field

# 21.1 Static Electricity, Electric Charge and Conservation

**Definition 78** (Law of Conservation of Electric Charge). The net amount of electric charge produced in any process is 0.

**Definition 79** (Free/Conduction Electrons). In a good (metal) conductor, some *free/conduction electrons* are bound very loosely and can move about freely inside.

When a positively charged object is brought close, these electrons move quickly toward it. And swiftly away from a negatively charged object.

There are fewer free electrons in a semiconductor, and almost none in an insulator.

#### 21.2 Induced Charge

**Definition 80** (Charging by Conduction). Take a positively charged object A and neutral object B. If A touches B, electrons from B will pass to A such that when A leaves, B will have a net positive charge. This is *charging by conduction*.

**Definition 81** (Charging by Induction). If a positively charged object simply approaches a neutral metal rod but doesn't touch it, a charge is *induced* at either end.

**Definition 82** (Grounded). Because the Earth is so large and can conduct, it easily accepts and gives up electrons causing any object connected to it to be *grounded*.

**Definition 83** (Electroscope). An *electroscope* is a device used to detect charge. A knob on the electroscope charges 1 or 2 gold leaves which are insulated with glass. The leaves separate when a positive object is brought close as electrons in the leaves go to the charge making the leaves positive so they repel each other.

If instead, the knob is charged by touching/conduction, the whole apparatus acquires a net charge which separates the leaves.

Note the electroscope doesn't tell you the sign of the charge unless you charged it first by conduction.

Modern sensitive electroscopes are called electrometers.

#### 21.3 Coulomb's Law

**Definition 84** (Coulomb's Law). The force between 2 charged particles is

$$F = -k \frac{Q_1 Q_2}{r^2} \quad \text{where}$$

 $k = 8.988 \times 10^9 \, \mathrm{Nm^2/C^2}$  and C is the coulomb.

In vector form, with unit vector  $\hat{r}_{21}$ , we can write this as

$$\vec{F}_{12} = k \frac{Q_1 Q_2}{r_{21}^2} \hat{r}_{21}.$$

**Definition 85** (Elementary Charge). The smallest charge observed in nature, the elementary charge, is denoted as e and equal to  $1.6022 \times 10^{-19}$  C.

Note this is positive so the charge on the electron is -e.

**Definition 86** (Quantized). Charge cannot be created or destroyed or split into partial charges (except for quarks) so electric charge is *quantized* such that it exists only in discrete amounts like 1e, 2e, 3e, etc.

**Remark** (Permittivity of Free Space). The constant k in Coulomb's Law is actually often written in terms of the *permittivity of free space* where  $k = \frac{1}{4\pi\varepsilon_0}$  and  $\varepsilon_0 = 8.85 \times 10^{-12} \, \mathrm{C}^2/\mathrm{Nm}^2$ .

Remark. These laws only apply for objects whose size are much smaller

than the distance between them. Ideally, like point charges. For finite-sized objects, charge may not be distributed uniformly and r values may not be clear.

**Definition 87** (Electrostatic/Coulomb Force). The *electrostatic* or *Coulomb* force is when one leectrically charged particle exerts a force on a second one. When charges are all at rest, this is the study of *electrostatics*.

**Definition 88** (Principle of Superposition). With several charges, the net force is the vector sum of the forces due to each charge. This sum relies on the principle of superposition and is based on experiment.

#### 21.4 Electric Field

**Definition 89** (Electric Field). The electric field  $\vec{E}$  defined at any point in space is defined as  $\vec{E} = \frac{\vec{F}}{q}$  as  $q \to 0$ . Here q is a test charge whose impact on the rest of the field is negligible.

E also magnitude  $k_{\pi^2}^Q$ .

Note. The superposition principle also applies to electric fields.

**Remark.** We can split a charge distribution into infinitesimal charges dQ which will act on a tiny point charge at a distance r such that  $dE = \frac{1}{4\pi\varepsilon_0} \frac{dQ}{r^2}$  which shows that  $\vec{E} = \int d\vec{E}$ .

**Example.** A thin ring of radius a holds a uniform total charge of +Q. Its center is x away from a point P. Thus, any point charge dQ along the ring is  $\sqrt{a^2+x^2}$  away from point P. Clearly, any electric field force perpendicular to the line of symmetry through the center of the ring will cancel out due to the ring's symmetry. So, the net force, and net field then, is only along and away from the line of symmetry. Thus,  $dE=k\frac{dQ}{a^2+x^2}\cos(\theta)=k\frac{dQ}{a^2+x^2}\frac{x}{\sqrt{a^2+x^2}}$ . Suppose a segment  $d\ell$  of the ring has point charge dQ with distribution  $\lambda$ . Thus  $E=\int dE=\frac{\lambda}{4\pi\varepsilon_0}\frac{x}{(x^2+a^2)^{\frac{3}{2}}}\int_0^{2\pi a}d\ell=\frac{1}{4\pi\varepsilon_0}\frac{\lambda x(2\pi a)}{(x^2+a^2)^{\frac{3}{2}}}$ .

This comes out to just

$$\frac{1}{4\pi\varepsilon} \frac{Qx}{(x^2 + a^2)^{\frac{3}{2}}}.$$

This makes sense as  $x \gg a$ , the field comes out to just the formula for if the ring were a point charge.

**Example.** Take a very long straight wire of uniformly distributed positive charge with charge per length  $\lambda$ . Take a point P a distance x from its

midpoint. Clearly,  $dE=\frac{1}{4\pi\varepsilon_0}\frac{dQ}{r^2}=\frac{1}{4\pi\varepsilon_0}\frac{\lambda dy}{x^2+y^2}$  where the wire lies along the y axis. Again, the y-component of the field will cancel from both sides so only  $\cos(\theta)$  will be summed giving  $E=E_x=\int dE\cos(\theta)=\frac{\lambda}{4\pi\varepsilon_0}\int \frac{\cos\theta dy}{x^2+y^2}$ . Note that we can write  $y=x\tan\theta$  so  $dy=xd\theta/\cos^2\theta$  giving  $\frac{1}{x^2+y^2}=\frac{\cos^2\theta}{x^2}$  so

$$E = \frac{\lambda}{4\pi\varepsilon_0 x} \int_{-\pi/2}^{\pi/2} \cos\theta d\theta = \frac{1}{2\pi\varepsilon_0} \frac{\lambda}{x}$$

**Example.** Take a thin circular disk of radius R with a uniformly distributed charge per unit area  $\sigma$ . Take a point P z above the disk's center. Everything not in the z-axis cancels out so  $dE = \frac{1}{4\pi\varepsilon_0} \frac{zdQ}{(z^2+r^2)^{\frac{3}{2}}}$  for each ring of radius r from the 1st example. Clearly, the ring has area  $(dr)(2\pi r)$  so  $dQ = \sigma 2\pi r dr$  giving  $dE = \frac{z\sigma r dr}{2\varepsilon_0(z^2+r^2)^{\frac{3}{2}}}$ . Now, integrating over all the rings from 0 to R gives

$$E = \frac{z\sigma}{2\varepsilon_0} \int_0^R \frac{rdr}{(z^2 + r^2)^{\frac{3}{2}}} = \frac{\sigma}{2\varepsilon_0} \left[ 1 - \frac{z}{(z^2 + R^2)^{\frac{1}{2}}} \right].$$

As  $z \ll R$ ,  $E = \frac{\sigma}{2\varepsilon_0}$  which approximates a point over an infinite plane of charge.

#### 21.5 Field Lines

**Definition 90** (Lines of Force). Electric field lines or *lines of force* are drawn to indicate the direction of force due to a given field on a positive test charge.

Lines starting on a positive charge terminate radially at negative charges.

**Remark.** The closer together lines are, the stronger the electric field is in that region. In fact, fields lines can be drawn so the number of lines crossing unit area perpendicular to  $\vec{E}$  is proportional to the magnitude of the electric field.

**Definition 91** (Electric Dipole). An electric field created by 2 equal charges of opposite sign is an *electric dipole*.

**Remark.** In the central reguon between two closely spaced, oppositely charged, parallel plates, the electric field has the same magnitude at all points  $E = \frac{\sigma}{\varepsilon_0} = \frac{Q}{\varepsilon_0 A}$ .

Note. We can equivalently make gravitational field diagrams.

#### 21.6 Electric Fields and Conductors

**Remark.** The electric field inside a conductor is 0 in the static situation.

Any net charge on a conductor distributes itself on the surface.

Remark. Imagein a neutral spherical metal shell with a positive charge fixed inside. There will be an induced charge on the inside of the shell surface. Electric fields from the positive charge will thus end at the shell inside. An induced charge will emerge at the surface of the shell given off new field lines radially. No field exists within the conducting shell itself.

**Remark.** The electric field is always perpendicular to the surface outside of a conductor.

However, the electric field outside a nonconductor does not necessarily make an angle of  $90^{\circ}$  to the surface.

**Remark.** Obviously, the force on a particle due to a field is  $\vec{F} = q\vec{E}$ .

#### 21.7 Electric Dipoles

**Definition 92** (Dipole Moment). Given an electric dipole formed by 2 equal charges Q of opposite sign separated by a distance  $\ell$ , the quantity  $p=Q\ell$  is called the *dipole moment*.  $\vec{p}$  can be considered as a vector pointing from the negative to positive charge.

**Definition 93** (Polar Molecules). Molecules with dipole moments are called *polar molecules*.

**Remark.** If a dipole of moment  $p=Q\ell$  is placed in a uniform electric field  $\vec{E}$  will result in a torque  $\tau=QE\frac{\ell}{2}\sin\theta+QE\frac{\ell}{2}\sin\theta=pE\sin\theta$ . That is,

$$\vec{\tau} = \vec{p} \times \vec{E}$$

will be exerted until the dipole is parralel to the field.

Thus, the work done on the dipole by the electric field to change the angle from  $\theta_1$  to  $\theta_2$  is  $W=\int_{\theta_1}^{\theta_2} \tau d\theta$ . In this case,  $\tau=-pE\sin\theta$  because its direction is opposite to the direction of increasing  $\theta$ . This gives  $W=pE(\cos\theta_2-\cos\theta_1)$ . If U=0 when  $\vec{p}\perp\vec{E}$ ,  $U=-W=-pE\cos\theta=-\vec{p}\cdot\vec{E}$ .

**Definition 94** (Hydrogen Bond). When  $H^+$  is involved, the weak bond it can make with a nearby negative charge is relatively strong and referred to as a *hydrogen bond*.