

17

Soil Temperature and Heat Flow in Soil

17.1 TEMPERATURE

Temperature is a measure of the thermal state of a body with respect to its ability to transfer heat. It is also defined as the measure of intensity or potential energy or heat. Temperature is the driving force for heat flow as pressure head is for water flow. Temperature is measured in three scales: Celsius ($^{\circ}\text{C}$), Fahrenheit ($^{\circ}\text{F}$), and Kelvin (K). The conversion from one scale to another is given in Table 17.1.

17.2 THE DEVELOPMENT OF THERMOMETER AND TEMPERATURE SCALES

One of the first attempts to make a standard temperature scale occurred about 170 AD, when Galen proposed a standard neutral temperature made up of equal quantities of boiling water and ice with four degrees of heat and cold on either side of this temperature, respectively. The earliest device used to measure the temperature was known as a “thermoscope” and consisted of a glass bulb having a long tube, which extended downward into a container of colored water. Before filling the liquid, some of the air in the bulb was

TABLE 17.1 Mathematical Expressions and Relations for Temperature Scales

Temperature scales	Mathematical formula
Celsius ($^{\circ}\text{C}$)	$^{\circ}\text{C}=(5/9)*(^{\circ}\text{F}-32)$
Fahrenheit ($^{\circ}\text{F}$)	$^{\circ}\text{F}=(9/5)*^{\circ}\text{C}+32$
Kelvin (K)	$\text{K}=^{\circ}\text{C}+273.15$

removed, causing the liquid to rise into the tube. As the remaining air in the bulb was heated or cooled, the level of the liquid in the tube would vary reflecting the change in

the air temperature. An engraved scale on the tube allowed for a quantitative measure of the temperature fluctuations.

The first sealed thermometer using liquid rather than air as the thermometric medium was developed for Ferdinand II in 1641. The thermometer was a sealed alcohol-in-glass device, with 50 “degree” marks on its stem, but no “fixed point” was used to zero the scale. This device was referred to as a “spirit” thermometer. Robert Hook (1664) used a red dye in the alcohol and the scale needed only one fixed point, for which every degree represented an equal increment of volume equivalent to about 1/500 part of the volume of the thermometer liquid, which was the freezing point of water. Hook demonstrated that a standard scale could be established for thermometers of a variety of sizes. Hook’s original thermometer was known as the standard of Gresham College, and was used by the Royal Society until 1709.

Ole Roemer of Copenhagen, Denmark, developed the thermometer scale in 1702 based upon two fixed points: snow (or crushed ice) and the boiling point of water, and recorded the daily temperatures at Copenhagen in 1708 and 1709. Gabriel Fahrenheit, an instrument maker in Amsterdam, The Netherlands, was the first to use mercury as the thermometric liquid in 1724. Mercury’s thermal expansion is large and uniform, and does not stick to the glass, and remains a liquid over a wide range of temperatures. The silvery appearance also makes it easy to read. Fahrenheit measured the boiling and freezing points of water to be 212 and 32, respectively, and designated temperatures in degrees Fahrenheit (°F).

In 1745, Carolus Linnaeus of Uppsala, Sweden, described the freezing point of water as zero, and the boiling point as 100, making it a “centigrade” (one hundred steps) scale. Anders Celsius (1701–1744) used the reverse scale in which 100 represented the freezing point and zero the boiling point of water, still, with 100 degrees between the two defining points. In 1948 use of the centigrade scale was dropped in favor of a new scale using degrees Celsius (°C). A degree Celsius equals the same temperature change as a degree on the ideal-gas scale. An “ideal gas” is one whose physical behavior is accurately described by the ideal-gas equation*. On the Celsius scale, the boiling point of water at standard atmospheric pressure is 99.975°C in contrast to the 100 degrees defined by the centigrade scale.

In 1780, J.A.C.Charles, a French physician, showed that for the same increase in temperature, all gases exhibited the same increase in volume. Because the expansion coefficient of gases is about the same, it is possible to establish a temperature scale based on a single fixed point rather than the two fixed-point scales, such as the Fahrenheit and Celsius scales. This brings us back to a thermometer that uses a gas as the thermometric medium.

P.Chappuis in 1887 conducted extensive studies of gas thermometers with constant pressure or with constant volume using hydrogen (H₂), nitrogen (N₂), and carbon dioxide (CO₂) as the thermometric medium. Based on his results, the Comité International des Poids et Mesures adopted the constant-volume hydrogen scale based on fixed points at the ice point (0°C) and the steam point (100°C) as the practical scale for international meteorology.

17.3 MEASUREMENT OF TEMPERATURE

The temperature of a substance (such as soil) is generally measured indirectly by measuring a property that responds to changes in its heat content. Some of these instruments are the liquid-in-glass thermometer, electric resistance thermometer, bimetallic thermometer, thermocouple, and remote-sensing thermometer (www.temperatures.com; Childs et al., 2000; Scott, 2000).

17.3.1 Liquid-in-Glass Thermometer

The liquid-in-glass thermometer is placed in close contact with soil or any substance, the conduction of heat between thermometer and its surrounding soil causes the change in volume of liquid in the glass thermometer (Childs et al., 2000; Scott, 2000). The traditional liquid-in-glass thermometer consists of a reservoir and capillary tubes and is based on the design proposed by Daniel Fahrenheit in 1714. This type of thermometer is used very commonly in the field and is sufficient for a reliable measurement of soil temperature provided good contact between reservoir and soil is

* $PV=nRT$, where P is the pressure (atm), V is the volume (m^3), n is number of moles, T is temperature (K), and R universal gas constant.

ensured. The accuracy of these devices ranges from ± 0.01 to $\pm 4^\circ\text{C}$. For the measurement of maximum and minimum temperatures, alcohol, toluene, or mercury is used as thermometric liquid. Since mercury vapors are toxic to human (ATSDR, 1999), cheaper resistance devices giving a digital readout have replaced mercury-in-glass thermometers.

17.3.2 Electric Resistance Thermometer

Electric methods are mostly based on the thermoelectric effect of temperature or change in resistance of a metal with a change in temperature. The motion of free electrons and atomic lattice vibrations are temperature dependent, which makes it possible to relate the resistance of a conductor to temperature. Resistance thermometers consist of a thin platinum or nickel wire, which is spiraled on a cylinder. The resistance measured using a bridge circuit generally increases by 0.4–0.5% per $^\circ\text{C}$ rise in temperature. A semiconductor known as a thermistor is a special type of resistance thermometer whose resistance decreases exponentially with an increase in temperature as follows (Scott, 2000):

$$R = B \exp\left(\frac{a}{T}\right) \quad (17.1)$$

where a and B are constants and T is the absolute temperature. The advantage of electric thermometers is that they can be easily used for continuous and rapid temperature measurements and can be highly accurate (Childs et al., 2000). However, these thermometers need to be frequently calibrated during use (Scott, 2000).

17.3.3 Bimetallic Thermometer

Bimetallic thermometers have two metals strips, which are joined together. These strips have different thermal expansion coefficients. The strips are also connected to a pointer. When temperature changes, the metal strips get deformed, which moves the pointer on a temperature scale. These thermometers are commonly used in thermographs and their accuracy is few tenths of a °C. An advantage of these devices is that they do not require a power supply (Childs et al., 2000; Scott, 2000).

17.3.4 Thermoelectric Thermometer

Sir William Siemens, in 1871, proposed a thermometer whose thermometric medium is a metallic conductor, whose resistance changes with temperature. The element platinum does not oxidize at high temperatures and has a relatively uniform change in resistance with temperature over a large range. The platinum resistance thermometer is now widely used as a thermoelectric thermometer and covers the temperature range from about -260°C to 1235°C . It defines the international temperature scale between the triple point of hydrogen (H_2), 13.8023 K, and freezing point of silver, 1234.93 K, within an accuracy of ± 0.002 K. Errors associated with platinum resistance thermometers are self heating, oxidation, corrosion, and strain of sensing element (Childs et al., 2000). If accuracy is less critical, a cheaper form of resistance thermometer known as a “thermistor” can be used. This utilizes a semiconductor (e.g., mixtures of oxides of nickel, magnesium, iron, copper, cobalt, manganese, titanium, etc.) in place of platinum. The accuracy of these devices for commercial application is $\pm 1^{\circ}\text{C}$.

17.3.5 Thermocouple

Thermocouple, the most widely used soil temperature measurement instrument, is made up of two wires of different metals (commonly copper-constantan, iron-constantan, or chromel-constantan) welded together at two places with the welds kept at different temperatures. The temperature difference causes a roughly proportional electric potential difference between the welds and current flows through the circuit formed by two wires. This effect is known as the thermoelectric effect. For measurement of soil temperature, one of the welds is kept at reference temperature while the other is kept in contact with soil. The compensation method measures the thermoelectric potential difference, and a galvanometer, the thermoelectric current between welds. Thermocouples are less economic, robust, and capable of monitoring temperatures between -270 and 3000°C . The sensitivity and speed of these devices is sufficient for many applications but are less accurate than resistance temperature devices (Childs et al., 2000; Scott, 2000).

17.3.6 Remote Sensing Thermometer

Temperature measurement devices based on thermal radiation monitoring can measure temperatures from 50 to 6000 K (Childs et al., 2000). Infrared thermometry is the most popular methods of estimating the temperature of the surfaces of soil, plant leaves, and crop canopies. According to the Stefan–Boltzmann equation the infrared radiations emitted by the surface are expressed as follows [see also Eqs. (17.15) and (17.16)]

$$R_l = e\sigma T^4 \quad (17.2)$$

where R_l is the long wave radiation, e is emissivity, which is close to 1 for most soil and plant surfaces, σ is Stefan–Boltzmann constant ($5.675 \times 10^{-8} \text{Wm}^{-2}\text{K}^{-4}$), and T is absolute temperature (Scott, 2000). An infrared measurement system comprises a source, a medium through which heat energy is transferred (e.g., gas), and a measurement device (e.g., optical system, a detector, a control and analysis system).

17.4 TEMPERATURE AS A THERMODYNAMIC PROPERTY (see also Chapter 14)

Experiments with gas thermometers have shown that there is very little difference in the temperature scale for different gases. Thus, it is possible to set up a temperature scale that is independent of the thermometric medium if it is a gas at low pressure. In this case, all gases behave like an “ideal gas” and have a very simple relation between their pressure, volume, and temperature:

$$PV = (\text{constant})T \quad (17.3)$$

where P is partial pressure of gas, V is volume of gas, T is temperature (also known as thermodynamic temperature), which is defined as the fundamental temperature and whose unit is the Kelvin (K), named after Lord Kelvin. Note that there is a naturally defined zero on this scale that is the point at which the pressure of an ideal gas is zero, making the temperature zero. With this as one point on the scale, only one other fixed point needs to be defined. In 1933, the International Committee of Weights and Measures adopted this fixed point as the triple point of water, the temperature at which water, ice, and water vapor coexist in equilibrium; its value is set as 273.16 K.

17.4.1 Entropy

Entropy (S_e) is a thermodynamic quantity, which is a measure of the degree of disorder within any system. The greater the degree of disorder, the higher the S_e . For an increase in disorder, S_e is positive and has the units of joules per degree K per mole. The entropy has a standard that is fixed by the third law of thermodynamics (see the following section).

17.4.2 Enthalpy

Enthalpy (H) is a thermodynamic state function, generally measured in kilojoules per mole. In chemical reactions the enthalpy change (ΔH) is related to changes in the free energy (ΔG) and entropy (ΔS_e) by the Gibbs equation:

$$\Delta G = \Delta H - T\Delta S_e \quad (17.4)$$

The enthalpy of an element has an internationally defined value at 298.15 K and 101.32 kPa and its entropy is zero at 0K and 101.32 kPa. The temperature that is most often used for recording thermodynamic data is 298.15 K, and by international convention the enthalpy of a pure element at 298.15 K and standard pressure is zero.

17.5 HEAT AND THERMODYNAMICS

Heat is the kinetic energy of random thermal motion of soil particles. Prior to the nineteenth century, it was believed that the sense of how hot or cold an object felt was determined by how much “heat” it contained. Heat was envisioned as a liquid that flowed from a hotter to a colder object, this weightless fluid was called caloric, and no distinction was made between heat and temperature. Black was the first to distinguish between the quantity (caloric) and the intensity (temperature) of heat. Joule (1847) conclusively showed that heat was a form of energy.

The zeroth law of thermodynamics states that if two bodies (e.g., masses of soil; A and B) are at the same temperature, and a third body C has the same temperature as body B , then the temperature of body C is equal to the temperature of body A .

$$\text{Temperature } A=B=C$$

(17.5)

The first law of thermodynamics is the conservation of energy and it states, “When heat is transformed into any other form of energy, or when other forms of energy are transformed into heat, the total amount of energy (heat plus other forms) in the system (plus surrounding) remains constant.” To express it another way, the law states, “It is in no way possible either by mechanical, thermal, chemical, or other means, to obtain a perpetual motion machine; i.e., one that creates its own energy.” At the same time, it is not possible to construct a cyclic machine that does nothing but withdraw heat energy and converts it into mechanical energy. No cyclic machine can convert heat energy wholly into other forms of energy, because efficiency of a cyclic machine can never be 100%. In the simplest form, the first law states “energy can neither be created nor destroyed.” It can change from one form to another, for example, electricity to heat, heat that can boil water and make steam, hot steam that can push a piston (mechanical energy) or turn a turbine that makes electricity, which can be changed into light (in a light bulb) or can change to sound in an audio speaker system, and so forth. If the total energy of a system is E , then between any two equilibrium states (E_1 for system 1 and E_2 for 2), the change in internal energy is equal to the difference of heat transfer (Q) into a system and work done (W) by the system

$$E_2 - E_1 = Q - W$$

(17.6)

A process, which does not involve heat transfer, is known as an adiabatic process. The second law of thermodynamics implies that there is an irreversibility of certain processes—that of converting all heat into mechanical energy. The law states that “there exists useful state variable called entropy (S_e) and the change in entropy is equal to the

heat transfer divided by the temperature.” For a given physical process, the entropy of the system and the environment will remain constant if the process can be reversed.

$$\Delta S_e = \frac{\Delta Q}{T} \quad (17.7)$$

If we denote the initial and final states of the system by “*i*” and “*f*”, then for a reversible system the change in entropy is zero (i.e., $S_{ef}=S_{ei}$) and for an irreversible system the entropy will increase (i.e., $S_{ef} > S_{ei}$).

An example of a reversible process is ideally (no boundary layer losses) forcing a flow through a constricted pipe. As the flow moves through the constriction, the pressure, temperature and velocity would change, but these variables would return to their original values downstream of the constriction. The state of the gas would return to its original conditions and the change of entropy of the system would be zero. The second law states that if the physical process is irreversible, the entropy of the system and the environment must increase and the final entropy must be greater than the initial entropy. An example of an irreversible process is when a warm soil is kept in contact with a cold one and after some time both achieve the same equilibrium temperature. If we then separate two soils, they do not naturally return to their original (different) temperatures. The process of bringing them to the same temperature is irreversible.

The third law of thermodynamics was formulated by Walter Nernst and is also known as the Nernst heat theorem. The law states, “at absolute zero, all bodies have the same entropy.” In other words, a body at absolute zero could exist in only one possible state, which possesses a definite energy, called the zero-point energy. This state is defined as having zero entropy, which is the entropy of a pure perfect crystal at 0K. At 0K, the atoms in a pure perfect crystal are aligned perfectly and do not move. Moreover, there is no entropy of mixing since the crystal is pure. For a mixed crystal containing the atomic or molecular species *A* and *B*, there are many possible arrangements of *A* and *B* and there is, therefore, entropy associated with the arrangement of the atoms/molecules.

17.5.1 Heat Capacity

The amount of temperature change in a body in response to heat adsorption or release is known as heat capacity. There are two types of heat capacities. The gravimetric heat capacity (C_g) is “the amount of heat energy required to raise the temperature of 1 kg of a substance by 1 K.” The volumetric heat capacity (C_v) is “the amount of heat required to raise the temperature of 1 m³ of a substance by 1 K.” The units of C_g and C_v in SI system are $\text{Jkg}^{-1}\text{K}^{-1}$ and $\text{Jm}^{-3}\text{K}^{-1}$, respectively. These two heat capacities are related by soil bulk density (ρ_b) as follows

$$C_g \rho_b = C_v \quad (17.8)$$

The specific heat of a substance is the ratio of the heat capacity of substance and water, and is dimensionless (see also the section on heat capacity of soils).

17.5.2 Blackbody

A blackbody is assumed to satisfy the ideal conditions, such as, (i) absorbs all incident radiation regardless of wavelength and direction, (ii) for a prescribed temperature and wavelength, no surface can emit more energy than a blackbody, and (iii) radiation emitted by a blackbody is a function of wavelength and is independent of direction. A blackbody is also known as a diffuse emitter.

17.6 FACTORS AFFECTING INSOLATION AT THE SOIL SURFACE

Radiations received at the soil surface are affected by a number of physical factors, which include vegetation, albedo, exposure, distribution of land and water, etc.

17.6.1 Vegetation

Vegetation cover buffers the soil beneath against sudden fluctuations in temperature. Bare soil is unprotected from the direct rays of the Sun and gets warm during the day and loses heat to atmosphere during the night. However, a good vegetative cover intercepts significant amount of solar radiation and prevents soil from getting warmer in summer. During winter or cold seasons, it prevents soil from losing heat as well, thereby reducing the daily variation of soil temperature as well as frost penetration and depth of freezing. The vegetation alters the soil energy balance in a number of ways, which include (i) altering albedo, (ii) insulating soil surface to prevent heat exchange, (iii) reducing depth of penetration of solar radiation, and (iv) increasing the removal of latent heat by evapotranspiration. Application of mulches on soil surface also alters the heat exchange in bare soil. The light colored mulches transmit short wave thermal energy to soil but prevent the loss of long wave thermal radiation and keep the soil warm by producing a green house effect.

17.6.2 Albedo

The fraction of all incoming solar radiations reflected back into space at the crop or soil surface is known as albedo. Albedo depends upon the nature of soil surface, angle of sunlight, and latitude. The albedo increases significantly with the distance from the equator. Water surfaces generally reflect 10% of the incoming radiations and therefore have lower albedo as compared to crop or soil surface. The albedo for canopy surfaces ranges from 5% for forest canopies to 25% for nonequatorial crops at full ground cover (Jury et al., 1991). The color of the soil is an important factor and affects the amount of reflection, for example, a light-colored soil has higher albedo than a dark-colored soil. Similarly a dry soil has higher albedo than a wet soil. The value of albedo for some soils and crops is presented in Table 17.2.

17.6.3 Latitude

The angle at which the Sun's rays meet the earth influences the amount of radiation received per unit area because of two reasons: (i) the albedo is high because of the angle, and (ii) radiation is subject to higher scattering reflection and adsorption, since they move through more atmospheres. Albedo is the highest in polar areas, decreases slightly in the middle latitudes, and is the lowest in tropical regions.

17.7 SOIL TEMPERATURE

Soil temperature is one of the most important factors affecting plant growth. Until soil reaches a certain critical temperature neither seeds

TABLE 17.2 Albedo from Soil, Forest, and Crops

Cover/surface	Albedo (%)	Reference
Light sand (Dry)	30–60	Geiger (1965)
Serozem (Dry)	25–30	Chudnovskii (1966)
Serozem (Wet)	10–12	Chudnovskii (1966)
Chernozem (Dry)	14	Chudnovskii (1966)
Chernozem (Wet)	8	Chudnovskii (1966)
Clay (Dry)	23	Chudnovskii (1966)
Clay (Wet)	16	Chudnovskii (1966)
Forest	5–20	Geiger (1965)
Corn (New York)	23.5	Chang (1968)
Sugar cane (Hawaii)	5–18	Chang (1968)
Pineapple (Hawaii)	5–8	Chang (1968)
Potato (Russia)	15–25	Chang (1968)

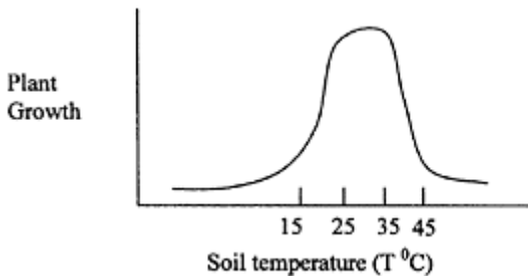


FIGURE 17.1 A graph of plant growth with soil temperature.

germinate nor plants have a normal growth because it affects root and shoot growth and availability of water and nutrients (Fig. 17.1). The optimum range of soil temperature for plant growth is between 20 and 30°C. The rate of plant growth declines drastically when temperature is less than 20°C (suboptimal) and above 35°C (supraoptimal) (Figs. 11.2–11 A). Further, all soil processes are temperature dependent. Consequently, the thermal regime of soil strongly influences the edaphic environment. The release of soil nutrients for root uptake is also dependent upon soil temperature regime. The biological processes (such as respiration by plants) are temperature dependent. Respiration rate (R_T) at a temperature (T) is expressed as follows:

$$R_T = R_0 Q_{10}^{(T-T_0)/10} \quad (17.9)$$

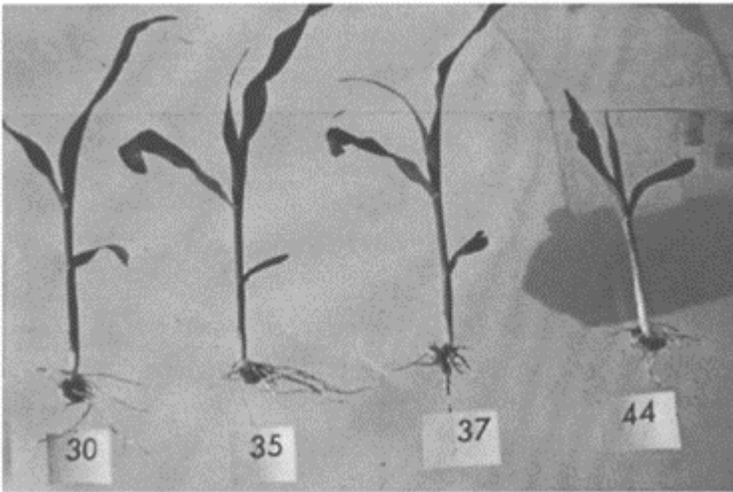


FIGURE 17.2 Corn seedling growth in relation to constant soil temperature maintained from 30 to 44° C in the root zone. (From Lal, 1972, greenhouse experiments.)

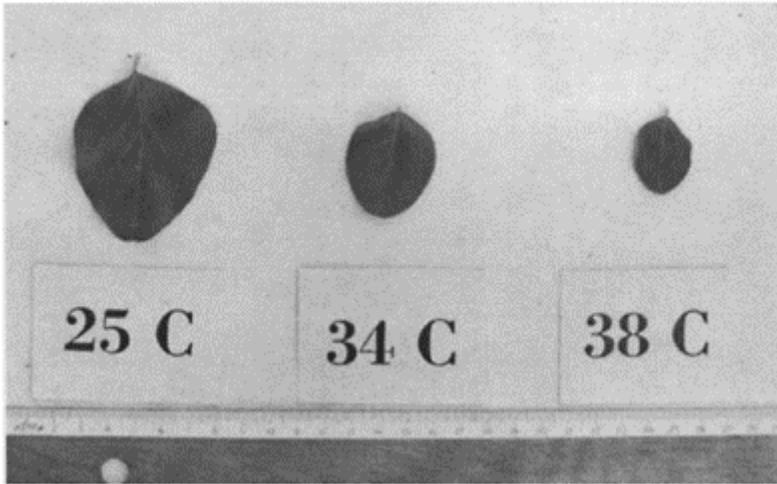


FIGURE 17.3 Soybean leaflets of seedlings grown at constant soil temperature of 25 to 38°C. (From Lal, 1972, greenhouse experiments.)

where R_0 is rate at reference temperature T_0 , Q_{10} is the factor which relates respiration to each 10°C change in temperature. The reaction rate (K_{rea}) in soil is mostly described by the Arrhenius equation as follows:

$$K_{rea} = A \exp\left(-\frac{E}{RT}\right) \quad (17.10)$$

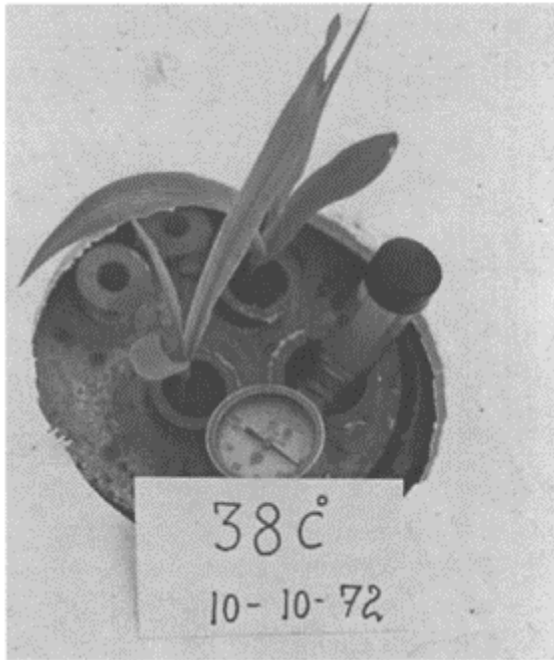


FIGURE 17.4 Chlorotic symptoms of nutrient imbalance in corn seedlings grown at constant soil temperature of 38°C. (From Lal, 1972, greenhouse experiments.)

where A is the preexponential factor, E is activation energy (J), R is gas constant ($8.314 \text{ Jmol}^{-1}\text{K}^{-1}$), and T is absolute temperature. A plot of $\log K_{\text{rea}}$ vs. $1/T$ provides the values of empirical constants E (as the slope) and A (as the intercept).

Soil-water movement, soil-water availability, evaporation, and aeration are also governed by soil temperature. Heat stored near the soil surface has a strong influence on evaporation from soil. A drier soil warms up relatively more quickly and cools down faster than wetter soil because heat capacity of water is several times more than that of soil. Soil temperature also influences the properties of water, such as surface tension, and to a lesser degree, viscosity and density (Table 17.3). Hence, soil-water characteristic curves and hydraulic conductivity functions are also temperature dependent. Bouyoucos (1915) was among the first to observe water movement caused by the soil temperature gradient. He imposed temperature gradients across soil columns, which were at different water contents and contained different soil materials. He found that the difference in water between two halves of column was dependent on both soil material and initial temperature.

Soil temperature varies as a result of radiant, thermal, and latent heat energy exchange processes, which take place primarily through the soil

TABLE 17.3 Density and Viscosity of Water at Various Temperatures

Temperature (°C)	Density (g cm ⁻³)	Viscosity (cp)
0	0.99987	1.787
3.98	1.00	1.568
5	0.9999	1.519
10	0.9997	1.307
20	0.9982	1.002
30	0.9957	0.7975
50	0.988	0.5468
80	0.971	0.3547
100	0.9584	0.2818

Source: Adapted from Handbook of Chemistry and Physics, 1988–89.

surface. Soil characteristics, which govern temperature regime, include bulk density, degree of wetness, soil heat capacity, and thermal sources and sinks present in soil matrix.

17.8 SOIL TEMPERATURE REGIMES

Soil temperature continuously varies in response to the changing meteorological regimes acting upon the soil atmosphere interface. The meteorological regimes are characterized by periodic succession of days and nights and winters and summers. Daytime heating is by short-wave radiation from the sun and sky, whereas nighttime cooling is from long-wave radiation emitted by soil. The temperature regimes of soil surface have two cyclical periods, namely diurnal and annual cycles.

17.8.1 The Diurnal Cycle

The variations in soil temperature owing to daytime heating and night-time cooling are known as diurnal variations. In the morning before sunrise, the minimum temperature of soil is the lowest at surface and increases with increase in depth. Similarly, the temperature continues to rise in the lower layers even after the top layer starts to cool down. However, the amplitude of the diurnal wave continues to decrease with soil depth (Fig. 17.5). The amplitude of the surface temperature fluctuation is the range from maximum or minimum to the average temperature (Fig. 17.5).

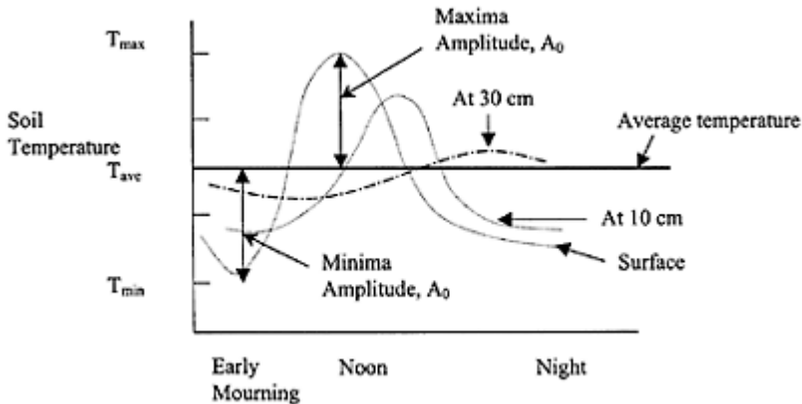


FIGURE 17.5 Schematic of diurnal variations in temperature measured at different depths.

17.8.2 The Annual Cycle

The annual variations in soil temperature result from the variations in short-wave radiation throughout the year. As one goes farther away from the equator, the annual variations in soil temperatures become significant. The summer months in June and July in the Northern Hemisphere represent the peak of global radiations and temperatures, whereas winter months have effects similar to nocturnal daily temperatures. During summer months, the soil temperature at surface is less than that of deeper layers (Fig. 17.6) (Smith, 1932).

The diurnal and seasonal variations of heat can be mathematically represented by assuming that soil temperature oscillates as a pure harmonic (sinusoidal) function of time around an average temperature. Let us also assume that average temperature of soil for all depths is the same. Assuming starting temperature as 0°C , the temperature at the soil surface and at any time t [$T(0, t)$] can be expressed as

$$T(0, t) = \bar{T} + A_0 \sin \omega t \quad (17.11)$$

where A_0 is the amplitude of surface temperature fluctuation, ω is the angular or radial frequency (1 radian = 57.3 degrees), which is 2π times the actual frequency, \bar{T} is the average temperature and t is time. Assuming that at infinite depth ($z = \infty$), the temperature is constant and equal to \bar{T} (Fig. 17.5). Therefore, temperature at any depth [$T(z, t)$]

$$T(z, t) = \bar{T} + A_z \sin(\omega t + \phi(t)) \quad (17.12)$$

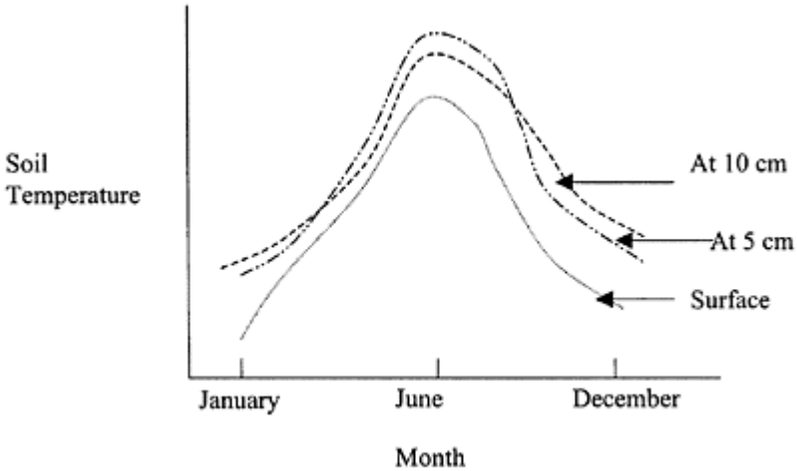


FIGURE 17.6 Schematic of annual variations in temperature measured at different depths in the Northern Hemisphere.

where A_z the amplitude at depth z and $\phi(z)$ are the functions of z , but not time. Incorporating the characteristic depth, also known as damping depth (d), which is defined as the depth at which temperature amplitude decreases to the fraction $1/e * A_0$, or $1/2.718 * A_0$, or $0.37 * A_0$, provides the following equation

$$T(z, t) = \bar{T} + \frac{A_0 \sin(\omega t + \phi(t))}{e^{z/d}} \quad (17.13)$$

The damping depth is also related to the thermal properties of the soil and the frequency of temperature fluctuation by the following relationship

$$d = \left(\frac{2k_t}{C_v \omega} \right)^{1/2} = \left(\frac{2D_T}{\omega} \right) \quad (17.14)$$

where C_v is volumetric heat capacity and D_T is thermal diffusivity of soil.

17.8.3 Soil Temperature Classes

Based upon the mean annual soil temperature, soil temperature regime is expressed in six categories (SSSA, 1987) namely pergelic, cryic, frigid, mesic, thermic, and hyperthermic (Table 17.4). In pergelic soils, mean annual temperature is lower, whereas in cryic soils, it is higher than 0°C . If mean annual temperature is lower than 8°C , the soils are known as frigid,

TABLE 17.4 Classes of Soil Temperature
According to Taxonomy of Soils

Class	Mean annual temperature	Remarks
Pergelic	$<0^{\circ}\text{C}$	Permafrost is present
Cryic	$0^{\circ}\text{C} < T < 8^{\circ}\text{C}$	Mean $(T_{\text{summer}} - T_{\text{winter}}) = 5^{\circ}\text{C}$ at depth 0.5 m
Frigid	$< 8^{\circ}\text{C}$	Mean $(T_{\text{summer}} - T_{\text{winter}}) > 5^{\circ}\text{C}$ at depth 0.5 m
Mesic	$8^{\circ}\text{C} < T < 15^{\circ}\text{C}$	Mean $(T_{\text{summer}} - T_{\text{winter}}) > 5^{\circ}\text{C}$ at depth 0.5 m
Thermic	$15^{\circ}\text{C} < T < 22^{\circ}\text{C}$	Mean $(T_{\text{summer}} - T_{\text{winter}}) > 5^{\circ}\text{C}$ at depth 0.5 m
Hyperthermic	$> 22^{\circ}\text{C}$	Mean $(T_{\text{summer}} - T_{\text{winter}}) > 5^{\circ}\text{C}$ at depth 0.5 m

Source: Modified from SSSA, 1987; Scott, 2000.

otherwise as mesic, provided mean annual temperature is below 15°C . For isofrigid, isomesic, isothermic, and isohyperthermic soils, the temperature differs by less than 5°C .

17.9 HEAT TRANSFER IN SOIL

There are three principle heat transport processes: radiation, conduction, and convection.

17.9.1 Radiation

Radiation is the process of heat transfer in which the emission of energy is expressed in the form of electromagnetic waves. The energy of the radiation field can also be transmitted through a vacuum since it does not require a carrier. The energy travels as discrete packets called quanta or photons, whose energy content depends on their wavelength or frequencies. According to the Stefan–Boltzmann law, the total energy emitted by a body, J_i , integrated over all wavelengths is proportional to the fourth power of the absolute temperature of the body, T , and can be expressed as below:

$$J_i = \sigma AT^4 \quad (17.15)$$

where A is the cross-sectional area of body (m^2), and σ is the Stefan–Boltzmann constant and in SI units is expressed as $5.675 \times 10^{-8} \text{ Wm}^{-2}\text{K}^4$. Eq. (17.15) gives the maximum energy flux that can leave a surface area A at any absolute temperature T . The ratio of radiant energy emitted by soil and maximum amount of radiant energy emitted (ϵ_s/ϵ_b) is known as emissivity coefficient, which equals one for a perfect emitter. Normally a blackbody transmits the maximum and is known as a perfect emitter. Soils emit much less radiant energy. The ϵ varies as a function of the wavelength of radiation and serves as a correction factor or indicator of the efficiency of natural resources. Equation (17.15) can be modified to Eq. (17.16):

$$J_f = \varepsilon \sigma A T^4 \quad (17.16)$$

The T also determines the wavelength distribution of the emitted energy and is inversely proportional to maximum radiation intensity, λ_m , micrometers (μm), which is also known as Wien's law.

$$\lambda_m = 2900/T \quad (17.17)$$

Assuming the temperature of the soil as 300 K, the radiations emitted by soil surface [Eq. (17.15)] have peak intensity at about 10 μm [refer to Eq. (17.17)] and its wavelength distribution is over the range of 3–50 μm , which falls in the infrared region. The Sun is a blackbody and has a temperature of 6000 K. The radiation emitted by the Sun has a peak intensity of about 500 nm [2900/6000]. The Sun's radiation includes a visible light range from 400 to 700 nm [400–425—Violet; 425–490—blue; 490–575—green; 575–585—yellow; 585–650—orange; 650–700—red; and invisible light range from 100–400—ultraviolet and 700–1400 nm—infrared (WHO, 1979), where 1nm=10⁻⁹m]. Planck's law describes the actual intensity distribution as a function of the wavelength, λ , and temperature T as follows:

$$E_\lambda = \frac{C_1}{\lambda^5} (e^{C_2/\lambda T} - 1) \quad (17.18)$$

where E_λ is the energy emitted for a given wavelength or range and C_1 and C_2 are constants. In general, the incoming solar radiations are referred to as short-wave radiations and the spectrum emitted by Earth comprises long-wave radiation. Most of the solar radiation reaching Earth's atmosphere is dissipated before it strikes the soil surface. The dissipation occurs partially as a result of the reflection of radiation by clouds, absorption by water vapor, oxygen, carbon dioxide*, and ozone, and diffusion by molecules and particles in air. Solar radiation reaching Earth's

* Greenhouse gases allow incoming solar radiation to pass through Earth's atmosphere, but prevent most of the outgoing infrared radiation from the surface and lower atmosphere from escaping into outer space. The greenhouse effect is the rise in temperature that Earth experiences because certain gases in the atmosphere (water vapor, carbon dioxide, nitrous oxide, methane, halogenated fluorocarbons, ozone, perfluorinated carbons, and hydrofluorocarbons) trap energy from the Sun. Without these gases, heat would escape back into space and Earth's average temperature would be about 33°C colder. Because of how they warm Earth, these gases are referred to as greenhouse gases (<http://www.epa.gov/globalwarming>).

surface is partly direct and partly in the form of scattered beams. After striking the crop or canopy, a fraction of incoming radiation is reflected back to the atmosphere, which is known as albedo (α). The thermal radiations are also transmitted from soil surface into the atmosphere Dearth, and onto soil surface from clouds R_{sky} , therefore net radiation (R_N) is:

$$R_N = (1 - \alpha)R_s + R_{in} \quad (17.19)$$

where R_s is global solar radiation (sum of direct and scattered beam) and R_{nt} is net long-wave thermal radiation ($R_{sky} - R_{earth}$). The R_N varies significantly with climate, latitude, and surface cover. For a known value of emissivity, both R_{sky} and R_{earth} can be calculated by the Stefan–Boltzmann equation (17.15).

17.9.2 Conduction

Conduction is the primary heat transfer mechanism in soil and refers to the propagation of heat within a soil or another body by molecular motion. It is the transfer of translational, rotational, and vibrational energy from molecule to molecule. The process of heat conduction is analogous to diffusion and both try to equilibrate, or even out, mixer's distribution of molecular kinetic energy. The heat flow by conduction in soil takes place from warmer locations towards the cooler regions. Fourier law explains the heat flow by conduction and macroscopically one-dimensional conduction of heat energy through a soil section is described as follows

$$q_h = -k_T A \left(\frac{\partial T}{\partial z} \right) \quad (17.20)$$

where q_h is the heat flux (Js^{-1}), k_T is proportionality constant or thermal conductivity ($\text{Jm}^{-1}\text{s}^{-1}\text{K}^{-1}$), A is the area of cross section (m^2), T is the temperature in $^\circ\text{K}$, and $\partial T/\partial z$ is the temperature gradient in degrees per unit length and the slope of the temperature-distance curve. The negative sign in Eq. (17.20) indicates that heat transfer occurs in the direction of decreasing temperature. Thermal conductivity of solids (Table 17.5) varies from $1 \text{ J m}^{-1} \text{ s}^{-1} \text{ K}^{-1}$ to $100 \text{ J m}^{-1} \text{ s}^{-1} \text{ K}^{-1}$. For liquids and gases, it ranges from $0.01 \text{ Jm}^{-1}\text{s}^{-1}\text{K}^{-1}$ to $1.0 \text{ Jm}^{-1}\text{s}^{-1}\text{K}^{-1}$ and $0.001 \text{ J m}^{-1} \text{ s}^{-1} \text{ K}^{-1}$ to $0.1 \text{ J m}^{-1} \text{ s}^{-1} \text{ K}^{-1}$, respectively. The ratio of q_h and A is also known as heat flux density ($\text{Jm}^{-2}\text{s}^{-1}$).

17.9.3 Convection

The transfer of heat energy in a convection process involves the movement of a heat-carrying mass. Infiltration of warm water into an initially cold soil

TABLE 17.5 Thermal Conductivity of Certain Metals (at 25°C)

Metal	Thermal conductivity ($\text{cal cm}^{-1}\text{s}^{-1}\text{C}^{-1}$)
Aluminium	9.56
Copper	16.25
Gold	2.44
Iron	2.24
Platinum	0.75
Silver	7.05

Tungstun

4.88

Source: Adapted from Handbook of Chemistry and Physics, 1988–89.

results in heat transfer by the process of convection. Newton's first law of cooling can be used to calculate energy fluxes in and out of the system

$$q_v = C_v A v (T_s - T_0) \text{ or } L * E \quad (17.21)$$

where q_v is the heat flux of convection (Js^{-1} or W), C_v is the volumetric heat capacity ($\text{Jm}^{-3}\text{K}^{-1}$), v is the velocity of fluid (m s^{-1}), T_s is the temperature of the soil in contact with fluid (K), T_0 is the temperature of the fluid far away from the surface (K), L is latent heat of vaporization, and E is evaporation rate. The convection phenomenon is probably more important in the atmosphere, where there is a consistent circulation of warm and cold air and heat exchange. In soils, the heat convection phenomenon is less important in general, however, during infiltration and redistribution of water in the soil profile, which is cooler than the incoming water, convective heat energy transport becomes important.

17.10 OTHER PROCESSES OF HEAT EVOLUTION AND TEMPERATURE IN SOIL

17.10.1 Condensation

The conversion of water vapor to a liquid state is known as condensation. Condensation is an exothermic process, and the heat energy released in condensation warms the soil surface. A similar phenomenon is observed when liquid water freezes. About 600 cal g^{-1} of heat energy is released when water vapor condenses, whereas 80 cal g^{-1} of heat (of fusion) is taken up when soil freezes. The six-phase changes that water can undergo and the heat gained and lost is given in Table 17.6.

TABLE 17.6 The Six Phase Changes That Water Can Undergo and the Heat Gained and Lost

Process	From	To	Heat gained/lost (calg^{-1})
Condensation	Vapor	Liquid	600
Evaporation	Liquid	Vapor	-600
Freezing	Liquid	Ice	80
Melting	Ice	Liquid	-80
Deposition	Vapor	Ice	680
Sublimation	Ice	Vapor	-680

Source: Modified from <http://www.usatoday.com/weather/wlatentl.htm>.

17.10.2 Microbial Processes and Heat Evolution

Diverse communities of organisms are present in soil. Variations of microbial population, distribution and activity are a function of depth and type of soil including structure, texture, and water status (Misthustin, 1956). Metabolism is defined as the sum of all chemical reactions occurring within a living organism. Catabolic reactions are exergonic or energy-releasing reactions, which break down more complex molecules, usually by hydrolysis, into simpler components (e.g., chemical processes of digestion). Anabolic reactions are endergonic or energy requiring and build more complex molecules, usually by condensation, from subunit components. The energy for anabolic reactions is provided by catabolic reactions. Microorganisms decompose the organic matter present in the soils and heat energy is released. An example of an exergonic reaction is the fermentation of alcohol as follows:



17.10.3 Chemical Reactions and Heat Evolution

Oxidation and reduction reactions (REDOX reactions) are always coupled in biological systems. Oxidation reactions are exothermic (or exergonic) and release energy, whereas reduction reactions are endothermic (or endergonic) and harness energy. Chemical reactions may be viewed in terms of the amount of energy required by the reaction at various stages. A convenient way to do this is with an energy hill diagram. In these diagrams, the total amount of energy, both kinetic and potential in the chemicals involved in the reaction, is plotted as a function of time. Conversion of sulfur (S) to sulfur dioxide (SO₂) and to sulfuric acid (H₂SO₄) are examples of exothermic processes.



The above chemical reaction releases about 178 Kcal/mole of energy (Bohn et al., 1934). Chemical reactions are known as *exothermic* when the chemical products of the reaction have less energy than the starting materials. Another example of an exothermic reaction is burning wood or a burning match. Wood is mainly cellulose and has a lot of chemical energy. The products of burning (e.g., CO₂ and H₂O) have much less energy because the net balance of the energy is converted into light and heat.

Chemical reactions that lead to products having more energy at the end than at the beginning are called *endothermic*. Endothermic reactions typically involve the synthesis of complex molecules from simple ones. Examples of endothermic reactions are cells making proteins from amino acids and photosynthesis in plant cells. In photosynthesis, CO₂ and H₂O, which are the starting materials for photosynthesis, have less energy than

the final product, i.e., carbohydrates
(<http://old.jccc.net/~pdeccl/metabolism/energyhill.html>).

17.11 ENERGY BALANCE OF SOIL

Net radiation is the sum of all incoming minus all outgoing radiation on Earth's surface. Steady state one-dimensional heat energy balance at the soil surface or crop canopy can be written as

$$\begin{aligned} &\text{Net heat energy arriving at surface} \\ &- \text{net heat energy leaving surface} = 0 \end{aligned} \quad (17.26)$$

Equation (17.26) disregards the transient energy changes due to heating or cooling of soil surface and lateral heat energy inputs. Heat transfer

* Except in a greenhouse, where net heat energy leaving the surface is smaller, therefore, Eq. (17.26) is not satisfied.

from the soil surface takes place as (i) convective heat flux (H_c), (ii) soil heat flux (J_H) and (iii) latent heat flux (L^*E). H_c represents the transport of warm air from the soil surface to the atmosphere vertically above it. The J_H represents the vertical transport of heat into the soil, and L^*E denotes evaporation and subsequent transport of water vapor from the soil surface (L is the latent heat of vaporization and E is the evaporation rate). The net radiation received by the soil surface is transformed into heat, which warms soil and air and vaporizes water. Therefore, under steady state conditions the heat balance equation can be written as

$$R_N = H_c + J_H + L^*E \quad (17.27)$$

Combining Eqs. (17.19) and (17.27) provides the total surface energy balance as follows

$$(1-\alpha)R_s + R_{nt} - (H_c + J_H + L^*E) = 0 \quad (17.28)$$

17.12 HEAT CAPACITY OF SOIL

The heat capacity of soil as defined in Sec. 17.5.1 is the amount energy required to change the temperature of a body by 1°C by heat adsorption or release. The relationship between volumetric and gravimetric heat capacities for a dry soil is described in Eq. (17.8), which is rewritten below:

$$C_v = \rho_b * C_g \quad (17.8)$$

and for a wet soil

$$C_v = \rho'_b * C_g = \rho_b(1 + w) * C_g = (\rho_b + \theta) * C_g \quad (17.29)$$

where ρ'_b is wet bulk density, w is gravimetric water content, and θ is volumetric water content. The C_v is dependent on composition of solid phase, which constitutes mineral and organic matter, bulk density, and water content of soil. The total C_v is calculated by summing the heat capacities of various constituents, weighted according to their volumes (de Vries, 1975)

$$C_v = \sum(f_{si}C_{si} + f_wC_w + f_aC_a) \quad (17.30)$$

TABLE 17.7 Thermal Conductivity and Heat Capacity of Gases (Oxygen at 25°C, 1 atm), Water (26.7°C), Quartz (37.8°C), and Sandstone (100°C)

Gas	Thermal conductivity (k_T) (10^{-6} cal $\text{cm}^{-1} \text{s}^{-1} \text{ } ^\circ\text{C}^{-1}$)	Heat capacity (C_g) (cal $\text{g}^{-1} \text{ } ^\circ\text{C}^{-1}$)
Air	62.2	0.25
Carbon dioxide	39.67	–
Oxygen	63.64	0.219
Water	42.57	0.998
Quartz (C-axis)	6.4	0.18
Sandstone	3.82	0.26
Iron		

Source: Adapted from Handbook of Chemistry and Physics, 1988–89.

where f is the volumetric fraction of each constituent phase, subscript s , w , a , and f stand for solid, water, air, and number of components in a given phase. The gravimetric heat capacity is the ratio of volumetric heat capacity and particle density (Table 17.7). The heat capacity of loam, sandy loam, sandy clay loam, a clay loam, and clay soil is given in Table 17.8. The heat capacity for each of the components solid, water, and air is the product of their particle density and specific heat or heat capacity per unit mass, i.e., $C_{si} = \rho_{si}C_{mi}$, $C_w = \rho_wC_{mw}$, and $C_a = \rho_aC_{ma}$. In general, the contribution of air is almost negligible because of the very small density and is ignored. The solid phase is divided into two components: mineral (m) and organic matter (o). Eq. (17.30) can be rewritten as

$$C_v = \sum(f_mC_m + f_oC_o + f_wC_w) + f_aC_a + f_wC_w \quad (17.31)$$

17.13 THERMAL CONDUCTIVITY

The thermal conductivity (k_T) is defined as the quantity of heat transferred through a unit cross-sectional area in unit time under a unit temperature gradient. The K_T of soil depends upon the volumetric proportions of the solid, liquid, and gaseous phase of the soil medium. The other factors, which influence K_T , are the size and arrangement of solid particles, and interfacial contact between solid and liquid phases. The K_T values of some materials is presented in Tables 17.7 and 17.9, which show that air has a much lower K_T than water and solid, therefore, high air content reduces the thermal contact between soil

TABLE 17.8 Heat Capacity of Some Nigerian Soils

Soil texture	Heat capacity ($\text{calg}^{-1} \text{ } ^\circ\text{C}^{-1}$)
Sandy loam	0.322
Sandy clay loam	0.350
Loam	0.279
Clay loam	0.224
Clay	0.248

Source: Modified from Ghuman and Lal, 1985.

TABLE 17.9 Range and Averages of Thermal Diffusivity and Thermal Conductivity of Soil Particles

Soil type	Moisture state	Thermal diffusivity $\times 10^{-3}$ (cm^2s^{-1})		Thermal conductivity $\times 10^{-3}$ ($\text{calcm}^{-1}\text{s}^{-1}\text{ } ^\circ\text{C}^{-1}$)	
		Range	Average	Range	Average
Sand	Dry	3.5–1.5	2.23	0.55–0.37	0.42
	Wet	12.6–4.4	8.0	4.35–3.7	4.02
Clay	Dry	1.8–1.2	1.5	0.37–0.17	0.26
	Wet	11–3.2	5.97	3.5–1.4	2.69

Source: From Geiger, 1965; Nakshabandi and Kohnke, 1965; and van Duin, 1963.

particles and reduces the K_T of soil. On the other hand, an increase in bulk density of soil lowers the porosity and improves the thermal contact between soil particles and increases K_T and D_T (cm^2s^{-1}). The increase in water content of the soil also improves the thermal contact between soil particles and increases K_T as well as D_T .

17.14 HEAT FLOW IN SOILS

Fourier (1822) analyzed the heat conduction in solids and developed a mathematical relationship, which is analogous to the diffusion equation by Pick (see Chapters 16 and 18), conduction of fluid flow in porous media by Darcy (refer to Chapters 12 and 13), and conduction of electricity by Ohm (Table 17.10). The first law of heat conduction is known as Fourier's law and under steady state condition in one-dimension, the heat flux density (q_h , $\text{Jm}^{-2}\text{s}^{-1}$) equation and the heat energy balance equation for

TABLE 17.10 The Analogous Laws of Water, Air, Heat, and Electricity Flow

Process	Law	Equation
Water movement	Darcy's	$q = -K \frac{\partial H}{\partial x}$
Air movement	Fick's	$q_x = -D \frac{\partial C}{\partial x}$
Heat movement	Fourier's	$h = -k \frac{\partial T}{\partial z}$
Electric flow	Ohm's	$a = -k \frac{\partial \phi}{\partial x}$

homogeneous soils is expressed as follows:

$$q_h = -K_T \frac{\partial T}{\partial z} \quad (17.32)$$

$$\frac{\partial H}{\partial t} = -\frac{\partial q_h}{\partial z} \quad (17.33)$$

where H is the volumetric heat content (Jm^{-3}), and volumetric heat capacity of soil (C_v) is equal to dH/dT . The early developments of heat conduction dealt with dry media, i.e., solids only (Carslaw and Jaeger, 1959). The theory was later expanded to soils containing water. At first the heat flow was studied for homogeneous soils with constant C_v and K_T .

Combining these two equations [(17.32) and (17.33)], which ignores the existence of a sink or source term, results in general heat conduction equation. These equations can be solved numerically for nonhomogeneous soil profiles by assuming that heat transfer takes place by conduction only. The addition of water complicates the process as water may evaporate and condense. As the heat energy inside a soil matrix is transported by convection (by flowing water, air, and latent heat), conduction, and radiation. The first two are the most important heat transport process through soil. The convection can be represented as given by Eq. (17.21). The expression for net flux of heat through soil can be written as

$$q_h = -k_T \frac{dT}{dz} + L * E_v \quad (17.34)$$

where E_v is the water vapor mass flux ($\text{gcm}^{-2} \text{s}^{-1}$). The temperature gradient across a moist soil results in movement of water both as liquid and gas along with heat, therefore, the effective value of K_T exhibits a temporal variation and therefore cannot be measured in the soil directly (de Vries, 1958). If D_v is the thermal vapor diffusivity then water vapor flux in soil in one-dimension flow where relative humidity is above unity can be written as

$$E_v = -D_v \frac{dT}{dz} \quad (17.35)$$

Transferring Eq. (17.35) into Eq. (17.34) results in

$$q_h = -(K_T + D_v L) \frac{dT}{dz} \cong -K_{Te} \frac{dT}{dz} \quad (17.36)$$

where K_{Te} is the effective thermal conductivity of the porous medium, which includes the effects of conduction and convection of latent heat.

17.15 HEAT CONSERVATION EQUATION

The heat conservation equation for a small cubic soil matrix (Fig. 17.7) can be derived by accounting for the amount of heat energy entering a system, leaving a system, and change in the heat energy of a system. Mathematically a heat balanced equation can be written as follows (Jury et al., 1991)

$$\begin{aligned} &\text{The amount of heat energy entering soil matrix} \\ &= \text{amount of heat leaving the soil matrix} \\ &+ \text{increase in heat energy of the soil matrix} \\ &+ \text{loss of heat energy from the soil matrix.} \end{aligned} \quad (17.37)$$

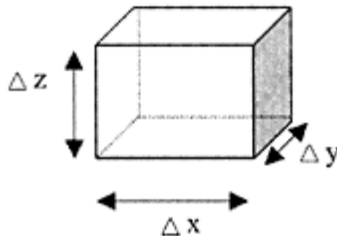


FIGURE 17.7 Schematic of a soil matrix for heat conservation equation.

Assuming the heat flow is in z -direction, the one-dimensional heat flow in vertical direction during time interval Δt through the soil matrix (Jury et al., 1991; Scott, 2000)

$$= h(x, y, z, t + \frac{1}{2} \Delta t) \Delta x \Delta y \Delta t \quad (17.38)$$

where h is the heat flux at the average time $t + (\Delta t/2)$ and Δx and Δy are the cross-sectional area of the soil matrix. The amount of heat flowing out of the matrix for the same time interval

$$= h(x, y, z + \Delta z, t + \frac{1}{2} \Delta t) \Delta x \Delta y \Delta t \quad (17.39)$$

The net change in H within the soil matrix

$$[H(x, y, z + \frac{1}{2} \Delta z, t + \frac{1}{2} \Delta t) - H(x, y, z + \frac{1}{2} \Delta z, t)] \Delta x \Delta y \Delta z \quad (17.40)$$

$$= \Delta H \Delta x \Delta y \Delta z$$

where H is at the middle or $z + (\Delta z/2)$. If the amount of heat lost from the soil matrix per unit system is r_s , the total heat loss

$$= r_s \Delta x \Delta y \Delta z \Delta t \quad (17.41)$$

Transferring Eqs. (17.38), (17.39), (17.40), and (17.41) into Eq. (17.37), and after rearranging and assuming that ΔZ and $\Delta t \rightarrow 0$, the resultant equation is known as the differential form of heat conservation equation (Jury et al., 1991; Scott, 2000)

$$\frac{\partial H}{\partial t} + \frac{\partial h}{\partial z} + r_r = 0 \quad (17.42)$$

The H can also be expressed as a function of volumetric heat capacity of soil C_v and temperatures

$$H = C_v (T - T_{ref}) \quad (17.43)$$

where T_{ref} is the reference temperature at which $H=0$. Transferring Eq. (17.43) into (17.42), and assuming C_v a constant, we obtain

$$C_v \frac{\partial T}{\partial t} = \frac{\partial}{\partial z} \left(K_T \frac{\partial T}{\partial z} \right) \quad (17.44)$$

if k_t is assumed to be independent of z then following heat flow equation is obtained.

$$\frac{\partial T}{\partial t} = D_T \frac{\partial^2 T}{\partial z^2} \quad (17.45)$$

where $D_T=K_T/C_V$, and is known as soil thermal diffusivity (L^2T^{-1} or m^2s^{-1}). D_T can be expressed in three different ways for soil water diffusivity, diffusion in air, and soil thermal diffusivity as follows:

Definition	Equation	Unit
Soil water diffusivity	$D_\theta=K_\theta(d\Phi_w/d\theta)$	$cm^2 s^{-1}$ or $L^2 T^{-1}$
Diffusion coefficient in air	$D_s=D_0*0.66*f_a$	$cm^2 s^{-1}$ or $L^2 T^{-1}$ (see Chapter 18)
Soil thermal diffusivity	$D_T=K_T/C_V$	$cm^2 s^{-1}$ or $L^2 T^{-1}$

17.16 MEASUREMENT OF THERMAL CONDUCTIVITY OF SOIL

Similar to hydraulic conductivity (K_s), thermal conductivity (K_T) can be measured using the steady state method or transient methods. Let us assume a soil column of thickness L placed between two glass plates of thickness d . The outer surface of each of these two plates is at constant temperature. The temperature is measured at several different positions as shown in Fig. 17.8. Let us assume the datum at point A, i.e., $z=0$ at A_1 and that heat flux through soil and glass plates is equal, glass and soil are in good contact, and lateral movement of heat is negligible. If the temperature measured at the plate at A is T_0 , then according to Fourier’s law, the heat flux across the plate and soil column can be written as

$$q_h = -K_T \frac{(T_2 - T_0)}{d} \tag{17.46}$$

$$q_h = -K_{Te} \frac{(T_3 - T_2)}{L} \tag{17.47}$$

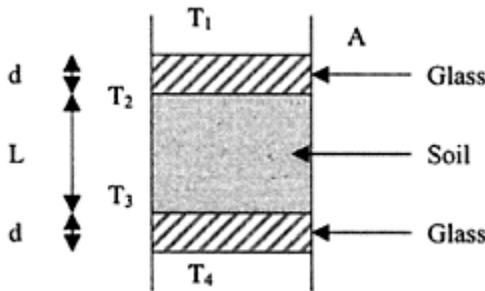


FIGURE 17.8 Apparatus for measurement of thermal conductivity under steady state.

where K_{Te} is the effective thermal conductivity of porous media. Equating these two equations [(17.46) and (17.47)] and rearranging them gives the expression for effective thermal conductivity as follows

$$K_{Te} = - \frac{K_T L (T_2 - T_0)}{d(T_3 - T_2)} \quad (17.48)$$

The steady state methods are useful for measurement of K_T in the laboratory. The basic drawback of this method is that a nonuniform profile within the column is created due to the redistribution of water under the influence of a steady state temperature gradient (Jury and Miller, 1974). The transient method for the measurement of K_T can be used in situ and does not have the drawback of the steady state method (De Vries and Peck, 1968). The method consists of a thin metal wire, which serves as a heat source. The wire and the sensors for temperature measurement (for example thermocouples) are kept inside a cylindrical tube, which is inserted into the soil. The flow of heat takes place radially from the wire and the temperature of the thermocouple probe in contact with the soil is given by the following equation (Carslaw and Jaeger, 1959)

$$T = T_0 = \frac{q_h}{4\pi k_T} [d + \ln(t + t_0)] \quad (17.49)$$

where T_0 is the temperature at $t=0$, $T-T_0$ is rise in temperature, q is heat flowing per unit time and length of wire, d is a constant, which depends on the location of thermocouple, and t_0 is a correction constant, which depends upon the dimensions of the probe. The equation (17.49) can be rearranged for $t \gg t_0$ and the K_T can be calculated by the following relationship

$$K_T = \frac{q_h}{4\pi m} \quad (17.50)$$

where m is the slope measured by plotting T versus $\ln(t)$. The q_h is calculated from the current (I) applied to the wire and the measured resistance per unit length of wire.

17.17 MANAGEMENT OF SOIL TEMPERATURE

Soil temperature can be managed in a number of ways, which includes mulching, tillage, irrigation, drainage, cover crop or shading, and application of dark or light powder. The management options depend whether the temperature of soil needs to be increased or reduced. The duff layer, which is the thatch of plant material on the surface, reduces the frequency of freeze-thaw cycles in the seed zone, maintains aggregates, and prevents crusting. Different types of mulches are used to either lower or raise the soil temperature, depending upon the need. Light-colored mulches (e.g., chopped straw, plastic mulches) reflect a portion of incoming solar radiation and reduce the amount of radiant flux reaching the soil surface, thus lowering the soil temperature (Fig. 17.9). On the other hand, application of thick and dark mulches (e.g., charcoal or bitumen) enhance the soil temperature. Dark plastic mulch absorbs most of the radiant solar energy but transmits

very little to the soil, keeping the soil cool (Fig. 17.10). On the other hand, transparent plastic mulch transmits short-wave (visible



FIGURE 17.9 A light-colored mulch (aluminium foil) decreases soil temperature. (Lal, 1975, field experiments.)



FIGURE 17.10 A dark-colored plastic mulch decreases soil temperature. (Field experiments, IITA, Ibadan, Nigeria, 1975.)

light) radiation to the soil surface while preventing infrared (long-wave) radiation, creating a greenhouse effect and warming the soil (Fig. 17.11) (Lal, 1979). Soil temperature (in °C) at 5 cm depth under a maize crop one week after planting and under black plastic, clear plastic, straw mulch, ridges, bare flat, and aluminum foil shows that the soil temperature fluctuation is the minimum for straw mulch and the maximum for ridges (Fig. 17.12). The soil temperature for bare flat treatment under different crops was in the order cassava > soybean > maize=cowpea (Fig. 17.13) (Lal, 1979).

Ridge tillage increases surface soil temperature by increasing the area exposed to radiation and decreasing soil moisture (Fig. 17.14). The surface 5 cm, or the seed zone, of no-till soils, may warm more slowly in spring and cool more slowly in autumn than in cultivated soils. Below 5 cm, in the root zone, no-till soils may be warmer and wetter from fall through spring. The amplitude of temperature variation at the soil surface is greater in plowed than no-till soils. However, because of the lower thermal conductivity of plow-till soil, the amplitude decreases more rapidly in plow-till than in no-till soils (van Duin, 1956). The maximum and minimum temperatures at the surface of a plow-till soil are also much higher than a no-till soil. Soil temperature in early spring is significantly affected by tillage methods (Table 17.11) (Fausey and Lal, 1989). Similar to no-till, mulching with crop residue decreases the maximum soil temperature and increases the minimum soil temperature (Fig. 17.15).

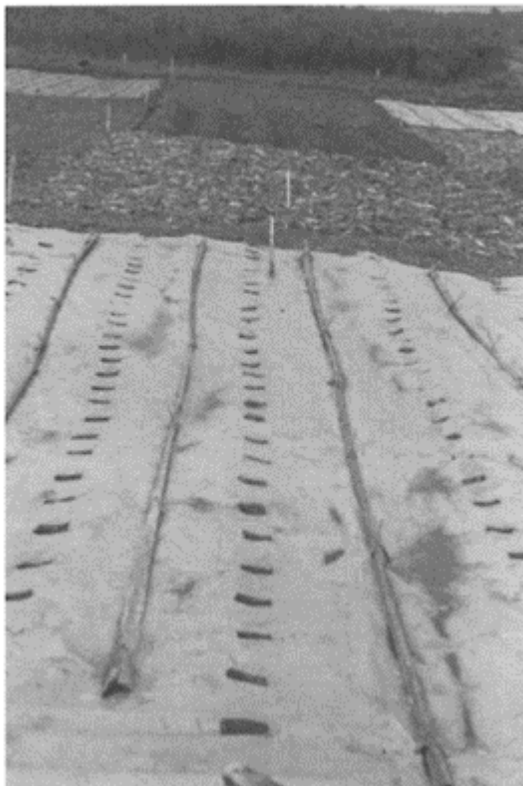


FIGURE 17.11 A clear plastic creates a greenhouse effect and increases soil temperature. (Field experiments, IITA, Ibadan, Nigeria, 1975.)

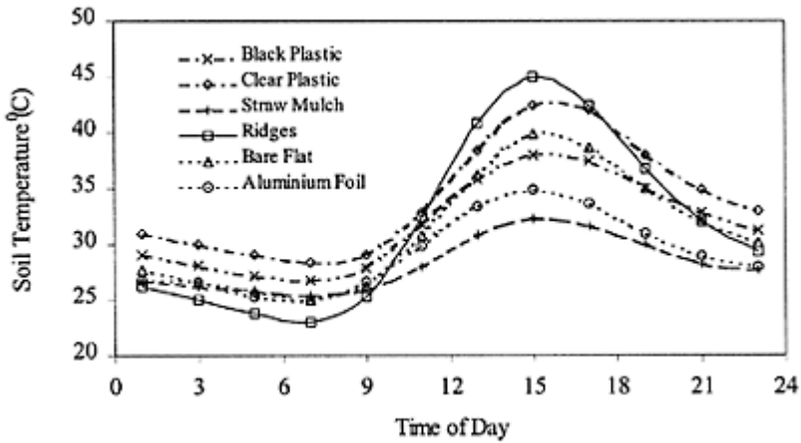


FIGURE 17.12 Soil temperature at a 5 cm depth under different mulches one week after planting crops during the first growing season in 1977 in Ibadan, Nigeria. (Redrawn from Lal, 1979.)

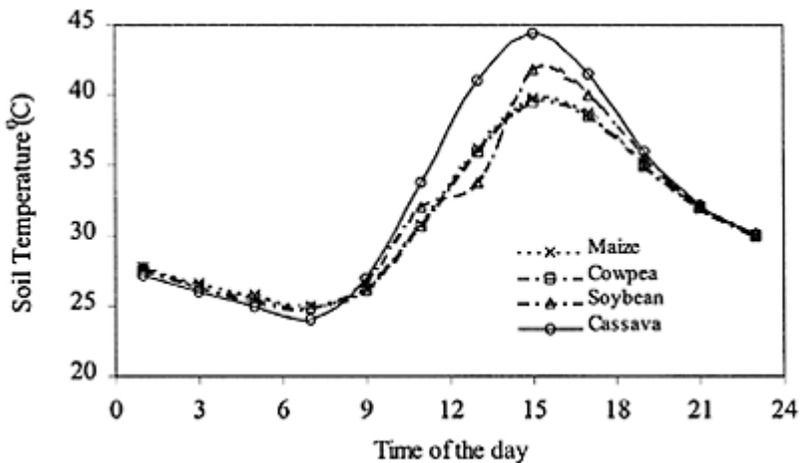


FIGURE 17.13 Soil temperature at a 5 cm depth under different crops for bare flat seedbed preparation during the first growing season in 1977 in Ibadan, Nigeria. (Redrawn from Lal, 1979.)



FIGURE 17.14 Ridge tillage may decrease the minimum and increase the maximum soil temperature.

Irrigation with cold water during the summer results in bringing down the temperature of surface soil (Fig. 17.16). Similarly, drainage has a strong influence on soil temperature. During spring, wet soils are cold at the soil surface, because of the increase in K_T of soil, which results in the conduction of heat in a downward direction reducing the temperature of the surface of

TABLE 17.11 The Tillage and Drainage Effects on Mean Daily Maximum Soil Temperature in °C

Tillage	Distance from drain (m)		
	0	9	27
No-till	6.2	6.1	6
Ridge-till	5.4	5.3	5.1
Plow-till	6.2	5.8	5.5
Beds	6	5.7	5.7

Source: Modified from Fausey and Lal, 1989.



FIGURE 17.15 Mulching with crop residue decreases the maximum and increases the minimum soil temperature.

soil. The high water content also increases heat capacity of soil, thus reducing the temperature of the surface of soil. The evaporation of water from wet soil also consumes the energy, which results in a reduction of temperature of the soil surface. Therefore, removal of excess water by surface or subsurface drainage increases soil aeration, which in turn warms the soil surface and improves seed germination and root growth. Building large mounds in a poorly drained (hydromorphic soil) increases soil temperature (Fig. 17.17). The effects of tillage and drainage on soil temperature are presented in Table 17.11, which show that as distance from drain increases, the mean daily temperature of soil reduces for all the treatments namely, no-till, ridge-till, plow-till, and beds.



FIGURE 17.16 Irrigation lowers the soil temperature in summer and raises it in winter.



FIGURE 17.17 Farmers in West Africa construct large mounds in hydromorphic soils to create well-aerated root zones and raise soil temperature. (Field experiments, IITA, Ibadan, Nigeria.)

Example 17.1

If the Stefan–Boltzmann constant (σ) is $5.67 \times 10^{-8} \text{Wm}^{-2}\text{K}^{-4}$ and emissivity (ε) is 0.94, calculate the long-wave thermal radiation energy flux R_{earth} for a temperature 300 K.

Solution

The energy flux density can be calculated as

$$\begin{aligned} R_{\text{earth}} &= \varepsilon \sigma T^4 = 0.94 * 5.67 \times 10^{-8} * (300)^4 \\ &= 431.713 \text{Wm}^{-2} = 0.61 \text{calcm}^{-2}\text{min}^{-1} \end{aligned}$$

Example 17.2

A soil column contains 40 cm of dry sand over 20 cm of dry loam soil. Both ends are attached to a constant temperature bath with the top maintained at 25°C and the bottom at 4°C. If the thermal conductivity of sand (K_{Ts}) is $0.5 \text{meal}^{-1}\text{s}^{-1}\text{°C}^{-1}$ and that of loam (K_{Tl}) is $0.25 \text{meal}^{-1}\text{s}^{-1}\text{°C}^{-1}$, calculate the steady state heat flux through the two layers and the temperature at the sand-loam interface.

Solution

The equivalent thermal conductivity of the sand-loam system (k_{eq}) for the total thickness of the sand-loam system (i.e., 60 cm) can be calculated as below

$$\begin{aligned} \frac{60}{K_{eq}} &= \frac{40}{K_{Ts}} + \frac{20}{K_{Tl}} \Rightarrow K_{Teq} = \frac{60}{(40/0.5) + (20/0.25)} = \frac{60}{(80 + 80)} \\ &= 0.375 \text{meal}^{-1}\text{s}^{-1}\text{°C}^{-1} \end{aligned}$$

The heat flux equation across the entire soil column of sand and loam will provide the steady state heat flux across column as follows

$$q_h = -\frac{K_{Teq}}{(40 + 20)}(25 - 4) = -\frac{0.375 * 21}{60} = -0.131 \text{calcm}^{-2}\text{s}^{-1}$$

The temperature across the sand-loam interface (T) can be calculated as

$$q_h = -\frac{K_{Tl}}{20}(T - 4) \Leftrightarrow -0.131 = -\frac{0.25 * (T - 4)}{20} \Leftrightarrow T = 14.48^\circ\text{C}$$

Example 17.3

If the bulk density of a soil 1.45gcm^{-3} and is at two different volumetric water contents: (i) 0.50 and (ii) 0.25, what will be the ratio of volumetric heat capacity, C_v , for these two situations?

Solution

The C_v can be calculated from Eq. (17.26).

For $\theta=0.50$, $C_{v1}=(1.45+0.5) C_g$

And for $\theta=0.25$, $C_{v2}=(1.45+0.25) C_g$

$$\frac{C_{v1}}{C_{v2}} = \frac{1.95}{1.70} = 1.15$$

Example 17.4

If the particle density of a soil is $2.65 \times 10^3 \text{ kg m}^{-3}$, and bulk density is $1.45 \times 10^3 \text{ kg m}^{-3}$, assuming the soil is water-saturated, calculate the volumetric heat capacity (C_v) of the soil if volumetric organic matter content is 15% of solid mass. Assume volumetric heat capacity of mineral, organic matter, and water as 2×10^6 , 2.5×10^6 , and $4.2 \times 10^6 \text{ J m}^{-3} \text{ deg}$, respectively.

Solution

The total porosity of the soil can be calculated as

$$f_t = \frac{(\rho_s - \rho_b)}{\rho_s} = \frac{2.65 * 10^3 - 1.45 * 10^3}{2.65 * 10^3} = 0.453,$$

since soil is saturated volumetric fraction of water is equal to porosity

Therefore volumetric fraction of solids is $1 - 0.453 = 0.547$

Organic matter fraction $= 0.547 * 0.1 = 0.0547$

Mineral matter fraction $= 0.547 * 0.9 = 0.492$

Therefore volumetric heat capacity (C_v) can be calculated as

$$C_v = f_m C_m + f_0 C_0 + f_w C_w$$

$$C_v = 0.492 * 2 * 10^6 + 0.0547 * 2.5 * 10^6 + 0.453 * 4.2 * 10^6$$

$$= 4.25 * 10^6 \text{ J m}^{-3} \text{ degree}$$

Example 17.5

For a temperature difference of 20°C across a 30-cm thick soil sample, calculate the one-dimensional thermal flux and total heat transfer under steady state condition. Assume thermal conductivity of soil as $1.6 \text{ J m}^{-1} \text{ s}^{-1} \text{ }^\circ\text{C}^{-1}$.

Solution

The heat flux across a soil column is expressed as

$$q_h = -K_T \frac{(T_2 - T_0)}{L} = \frac{1.6 * 20}{0.3} = 106.67 \text{ J m}^{-2} \text{ s}^{-1}$$

$$\text{Total heat transfer} = q_h * t = 106.67 * 3600 = 3.84 * 10^5 \text{ J m}^{-2}$$

Example 17.6

Assuming that the diurnal temperature wave is symmetrical and mean temperature is equal throughout the soil profile with surface temperature equal to mean temperature at 6 A.M. and 6 P.M., calculate the temperatures at noon and midnight for depths 10 cm and 25 cm. Assume daily maximum and minimum soil surface temperature 36°C and 8°C, respectively, and damping depth as 10 cm.

Solution

The temperature T at any depth z and time t can be calculated as follows

$$T(z, t) = T_{\text{ave}} + \frac{A_0[\sin(\omega t - (z/d))]}{e^{z/d}}$$

where A_0 is minimum value above mean, ω is the radial frequency ($2\pi/24$), z is depth, and d is the damping depth. The average temperature $T_{\text{ave}}=(38+8)/2=23^\circ$

Temperature above mean $A_0=38-23=15^\circ$

At soil surface $z=0$

Temperature 6 h after mean temperature, i.e., noon temperature

$$T(0, 6) = 23 + \frac{15 * [\sin((\pi/2) - 0)]}{e^0} = 23 + 15 = 38^\circ\text{C}$$

At midnight

$$T(0, 18) = 23 + \frac{15 * [\sin((3\pi/2) - 0)]}{e^0} = 23 - 15 = 8^\circ\text{C}$$

At depth 10 cm

$$T(10, 6) = 23 + \frac{15 * [\sin((\pi/2) - 0.1/0.1)]}{e^{0.1/0.1}} = 26^\circ\text{C}$$

and at midnight $T(10, 18)=22^\circ\text{C}$

PROBLEMS

1. If the particle density of a soil is $2.65 \times 10^3 \text{ kgm}^{-3}$, and bulk density is $1.45 \times 10^3 \text{ kgm}^{-3}$, assuming the soil is (a) dry and (b) volumetric water content is 30%, calculate the volumetric heat capacity (C_v) of the soil if volumetric organic matter content is 8% of solid mass. Assume volumetric heat capacity of mineral, organic matter, and water as 2×10^6 , 2.5×10^6 , and $4.2 \times 10^6 \text{ Jm}^{-3} \text{ deg}$, respectively.

2. Calculate one-dimensional thermal flux and total heat transfer under steady state condition for a temperature difference of 10°C across a 25-cm thick soil sample. Assume thermal conductivity of soil as $1.6 \text{ J m}^{-1} \text{ s}^{-1} \text{ }^\circ\text{C}^{-1}$

3. How much heat is required to change 20 kg of ice at -8°C to steam at 100°C ?

4. A soil column contains 35 cm of dry sand over 15 cm of dry loam soil. Both ends are attached to a constant temperature bath with top maintained at 28°C and bottom at 8°C . If the thermal conductivity of sand (k_s) is $0.48 \text{ meal}^{-1} \text{ s}^{-1} \text{ }^{\circ}\text{C}^{-1}$ and that of loam (k_l) is $0.23 \text{ meal}^{-1} \text{ s}^{-1} \text{ }^{\circ}\text{C}^{-1}$, calculate the steady state heat flux through the two layers and the temperature at the sand-loam interface.

5. Compute the amount of heat required to raise the temperature of a unit area of soil ($\rho_b=1.25 \text{ gcm}^{-3}$, $w=0.2 \text{ gg}^{-1}$) from an initial temperature of 10°C to 20°C to a depth of 50 cm.

6. Calculate the direction and quantity of heat per unit that will flow in one day, when soil temperature at the surface is 30°C and at 5 cm depth is 25°C . Assume thermal conductivity $=3 \times 10^{-3} \text{ cal cm}^{-1} \text{ s}^{-1} \text{ }^{\circ}\text{C}^{-1}$.

7. Why is soil temperature more important than air temperature?

REFERENCES

- ATSDR (1999). Mercury CAS#7439-97-6. Agency for Toxic Substance and Disease Registry, U.S. Dept. of Health and Human Services 1600 Clifton Road NE, Atlanta GA30333 (<http://www.atsdr.cdc.gov/>).
- Bohn H.L., McNeal B.L., and O'Connor G.A. (1934). *Soil Chemistry*. John Wiley & Sons, New York, p 52.
- Carlsaw H.S. and Jaeger J.C. (1959). *Conduction of Heat in Solids*. Oxford University Press, London.
- Chang J.H. (1968). Microclimate of sugar cane. *Hawaiian Planter's Rec.* 56:195-225.
- Childs P.R.N., Greenwood J.R. and Long C.A. (2000). Review of temperature measurement. *Review of Scientific Instrument.* 71(8):2959-2978.
- Chudnovskii A.F. (1966). Plants and light. I. Radiant energy. Pp1-51. In: *Fundamentals of Agrophysics*, Isreal Prog. For Scientific Translations, Jerusalem,
- de Vries D.A. (1958). Simultaneous transfer of heat and moisture in porous media. *Trans. Am. Geophys. Un.* 39:909-916.
- de Vries D.A. (1975). Heat transfer in soils. In: de Vries, D.A. and Afgan N.H., eds. *Heat and Mass Transfer in Biosphere*. Scripta Book Co., Washington DC, p 5-28.
- de Vries D.A. and Peck A.J. (1968). On the cylindrical probe method of measuring thermal conductivity with special reference to soils. *Aust. J. Phys.* 11:255-271.
- Fausey N.R. and Lal R. (1989). Drainage-Tillage effects on Crosby-kokomo soil association in Ohio II. Soil temperature regime and infiltrability. *Soil Technology* 2:371-383, Cremlingen.
- Geiger R. (1965). *The climate near the ground*. Harvard University Press, Cambridge, MA.
- Ghuman B.S. and Lal R. (1985). Thermal conductivity, thermal diffusivity, and thermal capacity of some Nigerian soils. *Soil Science* 139(1):74-80.
- Jury W.A. and Miller E.E. (1974). Measurement of transport coefficients for coupled flow of heat and moisture in a medium sand. *Soil Sci. Soc. Am. Proc.* 38:551-557.
- Jury W.A., Gardner W.R. and Gardner W.H. (1991). *Soil Physics*. John Wiley and Sons, USA.
- Lal R. (1979). Soil and micro-climate considerations for developing tillage systems in tropics. In Lal R., (ed.) *Soil Tillage and Crop Production*. International Institute of Tropical Agriculture, Ibadan, Nigeria, Proceedings Series number 2, p 52.

- Nakshabandi G.A. and Kohnke H. (1965). Thermal conductivity and diffusivity of soils as related to moisture tension and other physical properties.
- Scott H.D. (2000). Soil physics agricultural and environmental applications. Iowa State University Press, Ames, IA.
- Soil Science Society of America (1987). Glossary of soil science terms. Madison, WI.
- van Duin R.H.A. (1963). The influence of soil management on the temperature wave near the surface. Technical Bulletin 29, Institute of Land and Water Management Research, Wageningen.
- Weast R.C. et al., eds. (1989). Handbook of Chemistry and Physics. Boca Raton, Florida, CRC Press, 69th edition.
- WHO (1979). Environmental health criteria 14: Ultraviolet radiation. WHO, Geneva, Switzerland. 110pp.