SRMVALLIAMMAI ENGINEERING COLLEGE

(An Autonomous Institution)

SRM Nagar, Kattankulathur- 603 203

DEPARTMENT OF CIVIL ENGINEERING



IV SEMESTER 1902405 SOIL SCIENCE LABORATORY

Regulation – 2019

Academic Year 2021 - 22

Prepared by

Mr. M.MOGANRAJ, Assistant Professor(O.G.)/ AGRI
Ms. K.THARINI, Assistant Professor(O.G.)/ CIVIL

1902405

SOIL SCIENCE LABORATORY

LTPC 0021

OBJECTIVE:

- Students will be able to identify the types of rocks and minerals.
- Students should be able to verify various quality aspects of soil and water studied in theory by performing experiments in lab.
- To provide hands on experience on the textural analysis of soil.
- Students should be able to determine the organic carbon and gypsumrequirements.
- Students should be able to perform in situ tests on soil samples

LIST OF EXPERIMENTS

- 1. Identification of rocks and minerals
- 2. Collection and processing of soil samples
- 3. Determination of soil moisture, EC and pH
- 4. Field density determination by Core Cutter and Sand Replacement method
- 5. Specific gravity determination by Pycnometer
- 6. Textural analysis of soil by International Pipette method
- 7. Grain size analysis by using Mechanical shaker
- 8. Determination of Organic carbon
- 9. Estimation of Gypsum requirements

TOTAL: 30 PERIODS

OUTCOMES:

- 1. Learn about the identification of minerals referred under theory.
- 2. Knowledge of field investigations including collection of soil sampling, Processing and storage through observation of soil.
- 3. Understanding of the physical properties of soil and different laboratory methods
- 4. Be able to perform test to identify the organic carbon and application of soil amendments through chemical analysis of soil.
- 5. Students know the techniques to determine various physical and chemical properties of soil that are applicable for agriculture and irrigation by conducting appropriate tests.

REFERENCES:

- 1. Punmia, B.C., "Soil Mechanics and Foundation Engineering", Laxmi Publishers, New Delhi, 2007.
- 2. Laboratory Manual, Centre for Water Resources, Anna University, Chennai, 2012.

TABLE OF CONTENTS

EXP.NO.	EXPERIMENT NAME	PAGE.NO.
1.	Identification of rocks and minerals	
2.	Collection and processing of soil samples	
3.(i)	Determination of soil moisture	
3.(ii)	Determination of ElectricalConductivityof soil	
3.(iii)	Determination of soil pH	
4.(i)	Field density determination by Core Cutter method	
4.(ii)	Field density determination by Sand Replacement method	
5.	Specific gravity determination by Pycnometer	
6.	Textural analysis of soil by International Pipette method	
7.	Grain size analysis by using Mechanical shaker	
8.	Determination of Organic carbon	
9.	Estimation of Gypsum requirements	

Ex. no: 1 IDENTIFICATION OF ROCKS AND MINERALS

Date:

AIM:

To identify various types of rock and mineral samples.

INTRODUCTION

Soils don't simply *exist* on the landscape; rather they grow, develop, erode, and slowly transform into other soils through time—a group of processes known collectively as *pedogenesis*. Pedogenesis begins with *parent material*, the stuff from which soils form. Although a small minority of soils develop in organic deposits, most derived from inorganic parent materials acted upon by *additions*, *losses*, *translocations*, and *transformations*

A major type of transformation is **weathering of geologic parent materials**. All mineral soils develop from the rocks and minerals of the earth's crust. Through weathering, minerals and rocks physically **disintegrate into smaller particle sizes**, and **chemically decompose into altered chemical and mineralogical products**. However, the original rock material commonly is transported or reworked by some mechanism before a soil is formed. Variations in the history of the initial material lead to a **range of types of soil parent materials** (Table 1). Additionally, these **weathering processes release elements**, including plant and animal nutrients, and convert primary minerals into clays.

Table 1. "Summary of transported parent materials" by Colby J. Moorberg

Mode of transportation	Mode of deposition	Name of parent material
Gravity	Gravity	Colluvium
	Stream	Alluvium
Water	Lake	Lacustrine
	Ocean	Marine
1	Ice	Till, moraine
ice	Meltwater	Outwash (alluvium, lacustrine)
Wind	Wind	Loess, dunes

These parent materials continue to weather over time based on a function of soil forming factors and processes. Another major consideration for weathering potential of soil parent materials is hillslope position. Hillslope position references where on the landscape a soil is located (Figure 1). Hillslope positions essentially delegate the degree of slope and interaction with the

groundwater table for the soils of a specific position location. These relationships between soil and slope, or the groundwater table, greatly influence weathering processes over time, increasing both physical and chemical weathering, depending on hillslope position.

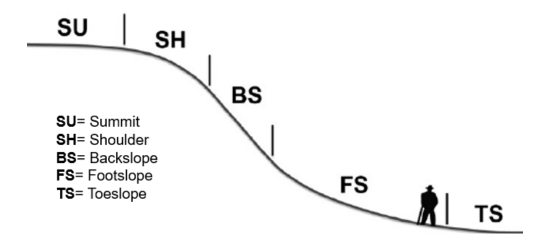


Figure 1. "Hillslope position across a landscape" Positions higher up and with a higher slope will have better drainage and less interaction with the groundwater table, while those positions that are lower lying on the landscape have a lower slope and decreased drainage. Wysocki et al., 2000 and Schoeneberger et al., 2012.

MINERALS

In geology and soil science, a mineral is a naturally occurring, inorganic, crystalline solid having a definite chemical composition and predictable physical properties. Of the six criteria included in this definition, three (naturally occurring, inorganic, and solid) need little elaboration; the remaining three are explained in succeeding paragraphs.

Crystallinity of Minerals

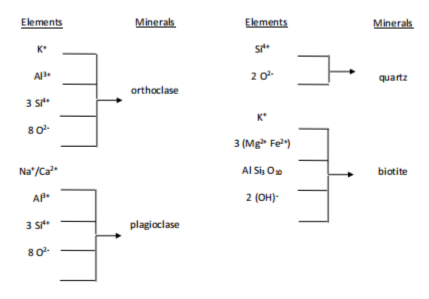
Crystalline means that a mineral's atoms are arranged in an orderly and repeatable manner; they are not random. The atomic arrangement, which varies among mineral groups, contributes to several of the mineral's physical properties, including hardness and cleavage, which are explained under "Physical Properties of Minerals." In contrast, a substance having randomly arranged atoms is amorphous, or noncrystalline, and as such is not considered to be a mineral. A good example of amorphous material is the rock obsidian, a form of volcanic glass.

Chemical Composition of Minerals

Minerals are made up of one or more elements, which give rise to definite chemical compositions (Figure 2). Although chemical formulas are precisely fixed for some minerals (e.g., quartz is SiO₂ and orthoclase is KAlSi₃O₈), they can vary within a narrow range for other minerals (e.g., plagioclase ranges from NaAlSi₃O₈ to CaAl₂Si₂O₈ because Na⁺ and Ca²⁺, which substitute for each other, occur together in varying proportions in the same mineral). Also, no

matter how precise the formula, irregularities occur in nature, allowing impurities to sneak in. Note the elements and minerals in Figure 2. What do the four minerals have in common?

Figure 2. Elemental composition of four common minerals important in soil formation.



On a mass basis, nearly three-fourths of Earth's crust is made up of only two elements: oxygen and silicon; and on a volume basis, these two elements constitute more than 90%! The next six most common elements bring the totals to about 98.5 percent by mass, and to nearly 100 percent by volume (Table 2). Obviously, very little room is left for the remaining 100 or so elements, including most plant nutrients, found in the crust. How is it then that soils, nearly all of which derive ultimately from crustal weathering, can supply enough nutrient elements for nearly all terrestrial life, including three-fourths of humanity's food supply? Your work in this laboratory exercise should help you begin to understand and appreciate the answer to this question.

Table 2. The eight most common elements in the Earth's crust.

Element	Ionic Formula	Mass Percent	Volume Percent
Oxygen	O ²⁻	46.60	91.97
Silicon	Si^{4+}	27.72	0.80
Aluminum	Al^{3+}	8.13	0.77
Iron	$Fe^{2+}\& Fe^{3+}$	5.00	0.68
Calcium	Ca^{2+}	3.63	1.48

Sodium	Na ⁺	2.83	1.60
Potassium	K^+	2.59	2.14
Magnesium	Mg^{2+}	2.09	0.56

Mineral Classification Based on Chemical Composition

Most minerals are grouped into seven classes based on their fundamental anionic unit (Table 3). (A few other classes exist, but these can be ignored for this laboratory). Of these, the silicates are by far the most common; nonetheless weathering products of others can be equally vital.

Table 3. Classification and composition of selected minerals important in soil science.

Mineral Class	Example	Chemical Composition (specimen #)	Comments
Sulfides	1) pyrite	FeS ₂ (19)	readily oxidized in moist
	2) sphalerite	ZnS (17)	soils
Oxides	hematite	Fe ₂ O ₃ (15)	readily oxidized, producing
			red colors
Halides	1) halite	NaCl (18)	rock salt; sometimes
			found in desert soils
	2) sylvite	KCl	
Carbonates	1) calcite	CaCO ₃ (13)	1) calcareous; reacts to
			acid; weathers easily in moist soils
	2) dolomite	CaMg(CO ₃)2	2) calcareous; less reactive
			than calcite
Sulfates	gypsum	CaSO ₄ 2H ₂ O (20)	a common agricultural
			amendment; sometimes
			found in desert soils
Phosphates	apatite	Ca ₅ (PO ₄)3(F,Cl,OH) (8)	the only significant source
			of phosphorus

Silicates	olivine	(Mg, Fe)2SiO4	most easily weathered of
			the silicates
	pyroxene:		susceptible to oxidation
	augite	(Ca, Na)(Mg, Fe, Al)(Si, Al)2O ₆ (12)	and hydration
	amphibole: hornblende		susceptible to oxidation
		(Ca,Na)2-3	and hydration, but more
		(Mg,Fe,Al)5Si ₆ (Si,Al)2O ₂₂ (OH)2 (11)	resistant than augite
	micas:		1) soft, but strongly resists
	1) muscovite	1) KAl ₂ (AlSi ₃ O ₁₀)(OH)2 (6)	chemical weathering
	2) biotite	2) K(Mg, Fe)3(AlSi ₃ O1 ₀)(OH)2 (7)	2) soft; weathers more
			easily than muscovite
	feldspars:		most common minerals
	1)	WAIS: 0 (0)	1)hard, resistant 2)hard,
	orthoclase	$KAlSi_3O_8(9)$	weathers
	2)	$Na(AlSi_3O_8)\square Ca(Al_2Si_2O_8)$ (10)	
	plagioclase		
	quartz	SiO ₂ (4)	hard and most resistant to
			weathering

Physical Properties of Minerals

<u>Color:</u> Minerals exist in nearly every color, and although color is easy to determine, it might not be reliable for identifying a mineral. For example, pyrite is nearly always brassy yellow, but quartz can range across several hues, values, and chromas from colorless to white, pink, purple, and gray. In some minerals, color can be strongly influenced by impurities.

<u>Luster</u>: independent of a mineral's color and determined by the nature of light reflected from a mineral's surface. Most lusters fall into two main groups: metallic and nonmetallic. Metallic lusters are typical of minerals that strongly absorb light, and as a result are opaque, even in very thin pieces. These minerals tend to look like metals, even though their surfaces might range from shiny to dull. Nonmetallic lusters are seen in minerals that allow light to pass through thinly cut slices. Surfaces of nonmetallic minerals might be brilliant like diamond, glassy like quartz, waxy like serpentine, or earthy like clay.

<u>Cleavage and fracture</u>: Cleavage is a mineral's tendency to break along certain predictable directions when the mineral is struck by a hammer. The resulting cleavage surfaces might be smooth and distinct, or they might be rough and indistinct, depending on the type of mineral. Different minerals also cleave along different directions and at different angles, depending on the type and orientation of atomic bonding. Although the number of possible cleavage directions is 1, 2, 3, 4, or 6, we will focus only on minerals that have 1 (e.g., biotite), 2 (e.g., orthoclase), and 3 (e.g., calcite) directions (Figure 3). Some minerals, when hit by a hammer, do not break along predictable planes: they simply shatter in any direction. These minerals are said to fracture, rather than cleave. Quartz is a good example of a mineral that exhibits fracture.

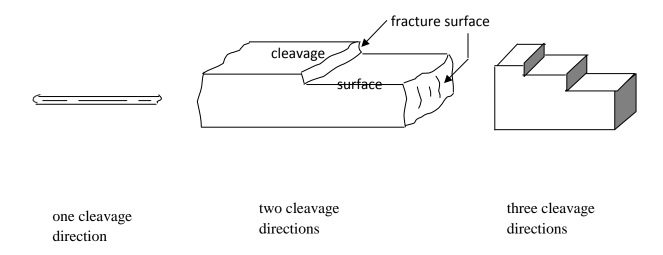


Figure 3. Examples of mineral cleavage.

<u>Hardness</u>: Hardness is determined by the resistance of a mineral's surface to scratching by various instruments of known hardness. It is judged by the Mohs hardness scale, which uses certain reference minerals ranked from 1 (soft) to 10 (hard): (see next page)

Mohs Hardness Scale

- 1. Talc (easily "scratched" by rubbing between fingers, giving a soapyfeel)
- 2. Gypsum (easily scratched by afingernail)
- 3. Calcite (same hardness as apenny)
- 4. Fluorite
- 5. Apatite (a bit softer than a pocket knifeblade)
- 6. Orthoclase (harder than a knife blade, but slightly softer thanglass)
- 7. Quartz (harder than both a knife blade andglass)
- 8. Topaz
- 9. Corundum
- 10. Diamond

(Soil minerals are very rarely harder than 7)

Essential elements for plants and animals

Essential elements are those that organisms need to grow and complete their life cycles. For plants, sixteen elements are universally recognized as essential; nonetheless research conducted in the late 20th century suggests that the number is eighteen or perhaps even greater. We will consider that plants require eighteen elements (Table 4), although you should understand that many plants, depending on their species, take up more than they seem to need.

A mnemonic phrase can help you memorize the 18 elements:

C HOPKNS CaFe Mg B MnCuZnClCoMo Ni

Translated, this reads "C Hopkns café, managed by "mine" cousins Clyde, Como, and Nicky."

Note that of the eight most common elements in Earth's crust, only five, oxygen, iron, calcium, potassium, and magnesium, are among the 18 essential elements (Table 4); and the first of these, oxygen, is supplied by O₂ in the atmosphere, **not** by weathering of minerals and rocks. Of the thirteen remaining essential elements, hydrogen comes from water, carbon comes from carbon dioxide (CO₂) in the atmosphere, and nitrogen is maintained by cycling of organic matter. (Nitrogen first is made available to plants by fixation from the atmosphere.) The remaining ten elements, phosphorus, sulfur, boron, manganese, copper, zinc, chlorine, molybdenum, nickel, and cobalt derive from weathering of minerals and rocks.

Table 4. Essential elements for plants.

Macronutrients			Micronutrients		
Element	Atomic	Plant Available	Element	Atomic	Plant Available
	Symbol	Form		Symbol	Form
carbon	С	CO ₂	iron	Fe	Fe ²⁺ , Fe ³⁺
hydrogen	Н	H ⁺	manganese	Mn	Mn^{2+}
oxygen	O	O_2 , H_2O	boron	В	H_3BO_3
nitrogen	N	, NH ₄ , NO ₃	zinc	Zn	Zn^{2+}
phosphorus	P	- 2- H ₂ PO ₄ , HPO ₄	copper	Cu	Cu^{2+}
potassium	K	K^+	chlorine	Cl	Cl
calcium	Ca	Ca^{2+}	cobalt	Co	CO^{2+}
magnesium	Mg	Mg^{2+}	molybdenum	Mo	2- MoO ₄
sulfur	S	2- SO ₄	nickel	Ni	Ni^{2+}

In addition to the 18 elements required by plants, animals also require arsenic (As), chromium (Cr), fluorine (F), iodine (I), lithium (Li), sodium (Na), selenium (Se), silicon (Si), and tin (Sn). Herbivorous animals obtain these elements by eating plants, which take up the elements, even though the plant might not need them. Carnivorous animals, in turn, obtain the elements by preying on other animals.

Regardless of whether the elements are required by plants or animals, all except carbon, hydrogen, oxygen, and nitrogen, are made available initially by weathering of minerals. The elements gradually become concentrated in the biosphere through biogeochemical cycles, including the carbon and nitrogen cycles, which will be covered in a later laboratory.

ROCKS

The most common rocks are consolidated assemblages of minerals, but a few (e.g., obsidian, pumice, and coal) comprise non-crystalline (hence, nonmineral) materials. Despite the great number of possible compositions and arrangements, rocks can be divided into three categories based on their mode of formation: igneous, sedimentary, and metamorphic.

Igneous Rocks

<u>Intrusive and Extrusive</u>: Igneous rocks form when molten material called magma cools and solidifies. Although the magma originates tens of kilometers below Earth's surface, it can solidify at any depth from its depth of origin to the surface. Magma that remains at depth while cooling and solidifying forms igneous intrusive rocks. Because of extremely slow cooling, minerals can grow to macroscopic size, and can be readily identified in a hand specimen.

Instead of remaining deep throughout the cooling process, some magmas erupt to the surface, where they cool and solidify rapidly. The resulting igneous extrusive rocks can be glassy (i.e., amorphous) or microcrystalline, although many contain a smattering of macrocrystals that formed before eruption. In erupting, the magma might ooze toward the surface as lava, or it might explode violently into the atmosphere, forming pumice and volcanic ash.

<u>Felsic and Mafic</u>: In addition to having distinctively different crystal sizes, igneous rocks also vary in mineralogical composition because of chemical differences among magmas. Some rocks are dominated by quartz and orthoclase (potassium-rich feldspar) because they derived from magma that was rich in silica and potassium. These rocks, which tend to be light colored, are called felsic. Other igneous rocks have little or no quartz or orthoclase, but instead are characterized by calcic plagioclase (calcium-rich feldspar) and ferromagnesian minerals (e.g., augite and hornblende). These are called mafic and are dark colored. Other rocks of intermediate compositions can be found between these extremes.

Classification of Igneous Rocks

The textural trend caused by varying rates of cooling, combined with the mineralogical trend resulting from chemical differences in magma provide the basis for classifying igneous rocks (Table 5). Selected elements also combine to form minerals, which then combine to form a common igneous intrusive rock (Table 6).

Accessory minerals, which can be significant, also vary from felsic to mafic. Of the micas, muscovite is felsic, and biotite is more mafic. As a result, muscovite is found in felsic rocks, including true granite, whereas biotite can occur in rocks ranging from felsic to moderately mafic. An important rock of intermediate composition is granodiorite, which has a characteristic salt-and-pepper appearance derived from light colored quartz and feldspars, especially sodic plagioclase, combined with dark colored biotite. Olivine, a strongly mafic mineral, and augite, which is slightly less mafic, are common constituents of gabbro andbasalt.

Table 5. Classification of igneous rocks

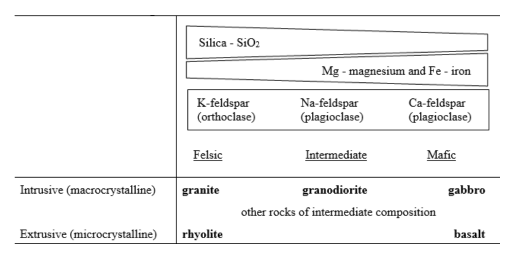
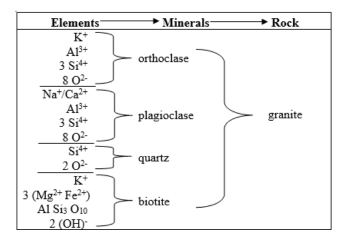


Table 6. An example of elements combining to form minerals and minerals combining to form an igneous rock.



Sedimentary Rocks

Sedimentary rocks, which make up the bulk of Earth's continental crust, can be either clastic (i.e., fragmental) or nonclastic (i.e., nonfragmental). See Table 7 for examples.

<u>Clastic rock</u>: consist of fragments of previously existing rocks (igneous, sedimentary, or metamorphic) that have been transported to a new location (often an ocean bottom), where they were deposited and buried layer upon layer, and cemented together. The deposits can be several thousand meters thick!

The fragments, or clasts, can vary in size to include clay, silt, sand, gravel, and cobbles.

<u>Nonclastic rock:</u> form by chemical precipitation of dissolved salts and by biochemical precipitation of organically derived compounds in water, most commonly on the floors of shallow seas. Organisms that generate nonclastic rocks include bacteria, algae, diatoms, corals, and mollusks. The most common nonclastic sedimentary rocks are limestone and dolomite (also called dolomitic limestone). Others include rock gypsum, chert, and coal.

Although the two major categories are recognized, most sedimentary rocks are a combination of clastic and nonclastic, nonetheless they usually are dominantly one or the other. For example, the sand grains in sandstone are clastic, but the cement that holds them together can be nonclastic.

Table 7. Classification of sedimentary rocks.

Clasti c		Nonclastic	
Dominant Constituent	Sedimentary Rock	Dominant Mineral	Sedimentary Rock
clay	claystone	calcite (CaCO3)	limestone
silt	siltstone	dolomite (CaMg)(CO3)2	dolomite
mud (silt + clay)	mudstone	gypsum (CaSO4•2H2O)	gypsum
thinly layered mud	shale	microcrystalline quartz (SiO2)	chert
sand (≥ 50%): mostly quartz, feldspars, micas sand: 50-85% sand with remainder mud and possibly some pebbles sand: > 85% sand with remainder mud and possibly pebbles	sandstones wacke (dirty sandstone) arenite (clean sandstone)		
gravel (pebbles) with sand and mud; quartz, orthoclase, quartzite	conglomerate		

Metamorphic Rocks

Metamorphic rocks form primarily by the application of extremely high heat and pressure to some previously existing rock. The original rock may be igneous, sedimentary, or even metamorphic, but sedimentary rocks are most susceptible to metamorphosis. Metamorphic rocks can be either foliated or nonfoliated. Foliated rocks consist of minerals that are aligned or oriented to yield a somewhat stratified

or layered appearance, whereas the minerals in nonfoliated rocks are about equally arranged in all directions, giving them a more uniformly massive appearance (Table 8).

Table 8. Classification of metamorphic rocks.

Foliated					
Metamorphic Rock	Common Original Rock	Appearance/Comments			
slate	shale	strongly stratified (layered), flat; often with very			
		fine quartz and mica; cleaves into sheets, but denser than shale			
schist	shale or other slate	wavy foliation; often with micas, quartz, feldspars			
gneiss	granitic rocks	minerals oriented to give more faint stratification			
(pronounced "nice")		than schist; often contains the same minerals as its granitic counterpart			
	Nonfe	pliated			
marble	limestone	calcite cleavage usually visible; fizzes to release			
		CO2 when tested with HCl			
quartzite	quartz arenite	hard; very resistant to weathering			
serpentinite	peridotite (an ultramafic igneous rock)	greenish assemblage of serpentine minerals; high magnesium; occurs intermittently along west coast of North America, and smaller areas of the East Coast			

WEATHERING

Weathering breaks rocks and minerals down, resulting in the production of soils. Weathering occurs via two processes: physical disintegration (breaking into smaller pieces) and chemical decomposition (changing the nature of the minerals themselves).

Physical disintegration causes rock masses to split apart or to abrade and wear away from the larger rock surface. Freezing and thawing, abrasion by particles suspended in wind or water, and grinding caused by glaciers result in disintegration. Repeated actions of these processes cause stresses, which eventually break and erode rock surfaces. Physical weathering usually precedes or occurs in concert

with the chemical decomposition process. With a decrease in the size of individual particles, the total number of corners and edges and the total surface area of the rock mass greatly increase.

Chemical decomposition

Chemical decomposition progresses most rapidly when more surface area is available for chemical reactions to occur. Warm and wet environments enhance chemical decomposition. The smaller the particles, the greater the surface area of a given volume of rock (the greater the total exposed corners and edges) and the greater the rate of chemical decomposition. The total number of corners exposed to weathering processes is the major factor in the determination of weathering rates. This is because a corner has three surfaces and three edges for chemical attack. Chemical weathering is a combination of five major processes: carbonation, hydration, hydrolysis, oxidation-reduction (redox), and solution.

<u>Hydrolysis and particle size:</u> Hydrolysis is a chemical decomposition process involving the splitting of water molecules during a reaction. A hydrolysis reaction can be easily identified by looking for water on the left side of a chemical equation and the absence of water (but the formation of H⁺ and/or OH⁻) on the right side of the equation. All minerals weather by hydrolysis. The hydrolysis reaction is particularly evident with silicate minerals (e. g. feldspars and micas). Hydrolysis is the primary reaction resulting in the release of most plant nutrient elements from minerals. The mineral orthoclase will weather by hydrolysis.

$$KAlSi_3O_8 + H_2O \square HAlSi_3O_8 + K + OH^-$$

 $Orthoclase + Water \square Al$ -silicate mineral + Potassium ion + Hydroxide ion

The potassium cation (K⁺) released by this hydrolysis reaction is soluble in water and can be adsorbed by the soil clay minerals and organic colloids, used by plants, or be removed (leached) by the drainage water. The aluminum silicate compound may crystallize into a clay mineral (e. g. kaolinite). This natural chemical weathering process occurs at a very slow but continual rate in all rocks, parent materials, and soils. The smaller the original mineral, the faster the hydrolysis reaction will occur. The hydroxide ion (OH⁻) will increase the pH. When a glacier grinds rocks with glacial ice (water), the grinding is a physical disintegration process. However, the resulting hydrolysis reaction of the fine particles hydrolyzing with the water causes the glacial till to have a high pH (abrasion pH). Initially, the pH of the glacial till will be 9 or 10, but as the glacial till absorbs carbon dioxide from the air, the pH will drop.

<u>Carbonation and Solution:</u> The metabolic activities of plant roots and microorganisms produce an abundance of CO_2 within soil pores. This CO_2 reacts with soil water to produce most of the carbonic acid found in soils.

$$CO_2 + H_2O \square H_2CO_3$$

Carbon Dioxide + Water \square Carbonic acid

The carbonic acid in the soil water continually reacts with minerals as the water percolates through soils and rocks. Carbonic acid, a weak acid, is the most common acid found in soils. Carbonic acid can ionize into acidic hydrogen ions (H⁺) and bicarbonate (HCO):

$$H_2CO_3 \square H^+ + HCQ^-$$

Carbonic acid \square Acidic hydrogen ion + Bicarbonate ion

At pH < 6, this reaction tends to go more to the left than to the right, indicating carbonic acid is a weak acid and does not produce very many acidic hydrogen ions. Carbonic acid is effective in accelerating the chemical decomposition processes of minerals because nature continually provides rainwater and carbon dioxide. Thus, rainwater drives this reaction to the right. As the rainwater moves through the soil, it carries soluble bicarbonate ions deeper into the soil profile and deposits the acidic hydrogen ions on the soil surfaces. The law of mass action allows the soil to eventually become acidic with time of weathering as rainfall drives the above reaction to the right.

As a result of a glacier grinding rocks, the hydrolysis reaction results in the glacial till containing soluble Ca²⁺ ions and OH ions (which causes a high pH, greater than 8.3). As the glacial till is exposed to carbonic acid, the glacial till undergoes two carbonation reactions. The first reaction process is quite rapid, while the second reaction is slow and requires time for completion. The first carbonation reaction is

$$Ca^{2+} + 2OH + H_2CO_3 \square CaCO_3 + 2H_2O$$

 $Soluble\ calcium + Soluble\ hydroxide + Carbonic\ acid\ \Box\ Low\ soluble\ calcite + Water$

The second reaction is:

$$CaCO_3 + H_2CO_3 \square Ca(HCO_3)_2$$

 $Calcite + Carbonic \ acid \ \square \ Soluble \ calcium \ bicarbonate$

The calcium bicarbonate dissolves slowly and releases soluble calcium ions (Ca²⁺) for plant uptake or possible leaching downward through the soil when excessive rainfall or irrigation occurs. The loss of the calcite minerals (the chief mineral in limestone) eventually leads to the development of acidic soils, carbonate hardpans, and the formation of huge underground caverns (e. g. Mammoth Cave and Carlsbad Caverns). The cave one walks through is the hole remaining after the soluble calcium bicarbonate has left the cavity where the carbonic acid dissolved the limestone rock.

The soluble calcium and bicarbonate move downward and enter the ground water, which eventually emerges as water in streams flowing to the ocean. The presence of the calcium bicarbonate in the ground water is the major reason limestone ground water is termed "hard water".

Eventually, the second reaction has two major effects in a high rainfall environment. First, the surface soil will become acidic and the pH will decrease due to increased rainfall and CO_2 production (H_2CO_3) . Second, soluble calcium is made available to plants. However, further rainfall will leach the soluble calcium bicarbonate downward.

Eventually, the soil will become depleted of calcium (Ca^{2+}) . The same phenomenon will be true for magnesium (Mg^{2+}) , potassium (K^+) , and sodium (Na^+) cations released to the soil by the hydrolysis of various minerals. This process results in a loss of overall plant nutrition and soil fertility.

<u>Reduction-oxidation (redox) reactions</u>: involve the transfer of electrons from one reactant to another. In every redox reaction, some substance becomes oxidized and some other substance becomes reduced by the transfer of the electron. The substance losing the electron is oxidized and the component accepting the electrons becomes reduced. Commonly, but not always, oxygen serves as the oxidizing agent.

Oxygen combines with an element in a mineral, such as iron (Fe²⁺), causing the iron to give up some of its electrons, which are accepted by oxygen. The iron becomes oxidized while the oxygen becomes reduced. These electron transfers constitute a form of chemical weathering, which disrupts the mineral's atomic structure, releasing several elements, and leading to the development of new compounds including clay minerals.

Although redox reactions take place continuously in soils, their reaction rates can be very slow and unnoticeable. If a soil is moist and well aerated, iron tends to become oxidized (Fe³⁺, loses electrons to oxygen), but if air (oxygen gas) is absent, as occurs with water-saturated soil, the iron becomes reduced

(Fe²⁺, gains electrons). The extent of either oxidation or reduction in soils is manifested in the color of the soil. Oxidized iron (Fe³⁺) occurs where oxygen is present in well-drained soils and produces red, yellow, or brown colors in soils. Reduced iron (Fe²⁺) exists in flooded or very poorly drained soils and produces black, gray, olive, and blue colors. Alternating periods of oxidation and reduction, caused by fluctuating water levels in a soil, result in mottling (blotches of red and yellow mixed with gray colors) to occur. Continuous strongly reducing conditions caused by long-term water-logging forms gleyed conditions. Gleyed soils have dark gray to blue to black soil colors.

$$Fe^{2+} \overset{o_2}{\longleftrightarrow} Fe^{3+}$$

Reduced (gray soil colors) \square Oxidized (red and yellow soil colors)

ACTIVITY 1: MINERALS

You will need to study the mineral specimens provided to fill in the blanks in the table below. Use the following guidelines for completing the table:

Color: Describe what you see. Munsell color designations are not necessary.

Luster: Write in either "M" for metallic or "N" for nonmetallic.

<u>Hardness:</u>Soft (S) minerals can be scratched by a fingernail. Medium (M) minerals can be scratched by a penny. Hard (H) minerals cannot be scratched by a penny and can scratch glass. <u>Cleavage:</u>Note whether present (1, 2, or 3 directions) or absent (0 directions).

<u>Plant-available nutrients:</u> Refer to the list of essential elements and to Tables 3 and 4. Write in the chemical symbol of nutrients that are <u>released and used by plants</u> by weathering of minerals.

Mineral	Color	Luster (M or N)	Hardness (S, M, or H)	Cleavage (yes/no)	Plant-available nutrients from Minerals (Table 4) (atomic symbol & charge)
pyrite					
calcite					
gypsum					
hornblende					
muscovite					
plagioclase					
orthoclase					
quartz					

1)) Which of the minerals in Table 3 is most resistant tow	veathering?
2)	2) Of the eight most common elements in the earth's cruare released to plants by the weathering of minerals?	st, which ones are essential elements that
3)	Name the source of each of the following plant-essen C H O	tialelements: N

ACTIVITY 2: ROCKS

Read the background information presented in this exercise, and study the minerals and rocks provided to fill in the blanks in the Properties of Rocks Table. Give the class as igneous intrusive (Ii), igneous extrusive (Ie), sedimentary (S), or metamorphic (M). List the crystal or grain size as microscopic (Mi) if the crystals or grains are too small to identify without a microscope, or macroscopic (Ma) if the crystals or grains are recognizable without a microscope.

Exercise: Fill in the properties of the following rocks provided in the rock and mineral kits. Ii=igneous intrusive, Ie=Igneous extrusive, S=sedimentary, M=Metamorphic, Mi=microcrystalline, Ma=macrocrystalline.

Rock	Class	Colo	Crystal or grain	Principal
	(Ii, Ie, S, M)	r	size (Mi, Ma)	Constituents or Minerals
granite				
basalt				
sandstone				
schist				
limestone				

chert				
conglomerate				
Rock	Class	Colo	Crystal or grain	Principal
	(Ii, Ie, S, M)	r	size (Mi, Ma)	Constituents or Minerals
marble				
gneiss				
shale				
ACTIVITY 3: V	WEATHERING			
Part 1				
	-	· ·	bench. Each beaker has a roughile backer #2 contains a se	
			while beaker #2 contains a so water beaker. After the reac	-
neasure the pH	of the solution. Rec	ord initial and fi	inal pH of the solution.	
<u>InitialpH:</u>			<u>FinalpH</u> :	
1) What are	e your predictions f	or what will hap	ppen when the liquids aread	ded?

2)	List	yourobser	vations
_,		jourobber	receive

3) Explain the chemical process youobserved:

Part 2

Your instructor has set out pictures detailing redoximorphic features displaying Oxidation and Reduction reactions. Observe each photo and detail your observations in the section below.							
1)	Observe the picture on display for gleyed soils (Image 1). Describe the appearance of the soil.	gleyed					
2)	Observe the colors of the mottled soil on display (Image 2). Describe the appearance of the mottledsoil.	he					
3)	What are the physical (i.e. color) and chemical (i.e. form of Fe) properties of a gleyed/reduced soil?						
4)	What environmental condition causes soils to becomegleyed?						
5)	In the field, what soil topographic or slope position (i.e. summit, backslope, or toeslope) we most conducive to the formation of reducing conditions or gleyed soils? Why?	vould be					

6) Most soils do NOT have gleyed horizons. If a gleyed horizon exists, which soil horizon is most commonly gleyed/reduced in a soil (Assume this soil has an impermeable layer at 100cm)?

7) Explain why this horizon (from Q # 6) would be most commonly reduced orgleyed.

8) Soil may have a mottled condition associated with a fluctuating water table. What are the physical and chemical properties of a mottled soil? What happens when the water table rises and falls in the mottled soilzone?

9)	Well oxidized soils are associated with what soil drainage condition? (See page 1-11 in the Red Book)
toeslop	field, what soil topographic or slope position (i.e. summit, backslope, or pe) would be most conducive to the formation of strongly oxidizing ions in a soil? Why?
10) For most soils, which is likely to have the most oxides, and most
10	likely to have an oxidized horizon?Why?
	AN GO
TIVITY 4:	ROCKS TO PAREN <mark>T MATERIAL</mark>

DEVELOPMENT OF ROCKS INTO PARENT MATERIAL THAT BECOMES SOIL

As mentioned above, inorganic parent materials are typically derived from a variety of geological species. These parent materials are typically identified by their mode of transportation (i.e. colluvium transported by gravity, or loess transported by wind), however, it is important to recognize how specific rock types physically weather in order to be able to identify parent materials properly in the field.

Observe the trays of parent materials and soil for the granite and sandstone rocks on display. The first tray contains the fresh rocks before being weathered, the second tray contains the parent material of the soil formed from this rock, and the third tray contains the soil formed from the parent material derived from this rock type.

1) List the main minerals in each type ofrock.

Sandstone:			
		nges (particle size and color) have occurred in weathered rock tosoil?	
most sus		veathered rock, or soil (circle one)—is ner physical and chemical	
fraction filling in	(rock, weathered the table below	rochloric acid (HCl) (a few drops) on each d rock, and soil). Indicate which materials fizz by with a "Y" for yes and an "N" for no.	
Rock Type	Rock (R)	Weathered rock parent material (Cr)	Soil
Granite			
Sandstone			
<u>.</u>			

COLLECTION AND PROCESSING OF SOIL SAMPLES



Exp. no.: 3.i. Date:

DETERMINATION OF SOIL MOISTURE

OBJECTIVE

Determine the natural content of the given soil sample.

NEED AND SCOPE OF THE EXPERIMENT

In almost all soil tests natural moisture content of the soil is to be determined. The knowledge of the natural moisture content is essential in all studies of soil mechanics. To sight a few, natural moisture content is used in determining the bearing capacity and settlement. The natural moisture content will give an idea of the state of soil in the field.

DEFINITION

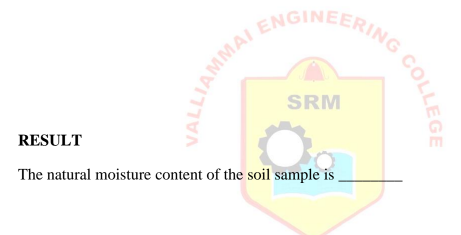
The natural water content also called the natural moisture content is the ratio of the weight of water to the weight of the solids in a given mass of soil. This ratio is usually expressed as percentage.

APPARATUS REQUIRED

- 1. Non-corrodible air-tight container.
- 2. Electric oven, maintain the temperature between 1050 C to 1100 C.
- 3. Desiccator.
- 4. Balance of sufficient sensitivity.

PROCEDURE

- 1. Clean the container with lid dry it and weigh it (W1).
- 2. Take a specimen of the sample in the container and weigh with lid (W2).
- 3. Keep the container in the oven with lid removed. Dry the specimen to constant weight maintaining the temperature between 1050 C to 1100 C for a period varying with the type of soil but usually 16 to 24 hours.
- 4. Record the final constant weight (W3) of the container with dried soil sample. Peat and other organic soils are to be dried at lower temperature (say 600) possibly for a longer period. Certain soils contain gypsum which on heating loses its water if crystallization. If it is suspected that gypsum is present in the soil sample used for moisture content determination it shall be dried at not more than 800 C and possibly for a longer time.



OBSERVATIONS:

S.No.	Description	Trial 1	Trial 2	Trial 3
1.	Weight of container with lid, W ₁ (gm)			
2.	Weight of container with lid + wet soil, W_2 (gm)			
3.	Weight of container with lid + dry soil, W_3 (gm)			
4.	Moisture content, W (%)			



Exp.No.: 3.ii. Date:

DETERMINATION OF ELECTRICAL CONDUCTIVITY OF SOIL



Exp.No.: 4.ii. Date:

FIELD DENSITY DETERMINATION BY SAND REPLACEMENT METHOD

OBJECTIVE

Determine the in situ density of natural or compacted soils using sand pouring cylinders.

NEED AND SCOPE

The in situ density of natural soil is needed for the determination of bearing capacity of soils, for the purpose of stability analysis of slopes, for the determination of pressures on underlying strata for the calculation of settlement and the design of underground structures. It is very quality control test, where compaction is required, in the cases like embankment and pavement construction.

APPARATUS REQUIRED

- 1. Sand pouring cylinder, mounted above a pouring cone and separated by a shutter cover plate.
- 2. Tools for excavating holes; suitable tools such as scraper tool to make a level surface.
- 3. Cylindrical calibrating container fitted with a flange surrounding the open end.
- 4. Balance to weigh unto an accuracy of 1g.
- 5. Metal containers to collect excavated soil.
- 6. Metal tray.
- 7. Glass plate.
- 8. Clean, uniformly graded natural sand passing through 1.00 mm I.S.sieve and retained on the 600micron I.S.sieve. It shall be free from organic matter and shall have been oven dried and exposed to atmospheric humidity.
- 9. Suitable non-corrodible airtight containers.
- 10. Thermostatically controlled oven with interior on non-corroding material to maintain the temperature between 1050 C to 1100 C.
- 11. A dessicator with any desiccating agent other than sulphuric acid.

THEORY

By conducting this test it is possible to determine the field density of the soil. The moisture content is likely to vary from time and hence the field density also. So it is required to report the test result in terms of dry density. The relationship that can be established between the dry density with known moisture content is as follows:

$$\gamma_d = \gamma_0 / [1 + w]$$

PROCEDURE

Calibration of the Cylinder:

1. Fill the sand pouring cylinder with clean sand so that the level of the sand in the cylinder is within about 10 mm from the top. Find out the initial weight of the cylinder plus sand (W1) and this weight should be maintained constant throughout the test for which the calibration is used.

2. Allow the sand of volume equal to that of the calibrating container to run out of the cylinder by opening the shutter, close the shutter and place the cylinder on the glass sand takes place in the cylinder close the shutter and remove the cylinder carefully. Weigh the sand collected on the glass plate. Its weight(W2) gives the weight of sand filling the cone portion of the sand pouring cylinder. Repeat this step at least three times and take the mean weight (W2) Put the sand back into the sand pouring cylinder to have the same initial constant weight (W1)

Determination of Bulk Density of Soil:

- 3. Determine the volume (V) of the container be filling it with water to the brim. Check this volume by calculating from the measured internal dimensions of the container.
- 4. Place the sand poring cylinder centrally on yhe of the calibrating container making sure that constant weight (W1) is maintained. Open the shutter and permit the sand to run into the container. When no further movement of sand is seen close the shutter, remove the pouring cylinder and find its weight (W3).

Determination of Dry Density of Soil In Place:

- 5. Approximately 60 sqcm of area of soil to be tested should be trimmed down to a level surface, approximately of the size of the container. Keep the metal tray on the level surface and excavate a circular hole of volume equal to that of the calibrating container. Collect all the excavated soil in the tray and find out the weight of the excavated soil (Ww). Remove the tray, and place the sand pouring cylinder filled to constant weight so that the base of the cylinder covers the hole concentrically. Open the shutter and permit the sand to run into the hole. Close the shutter when no further movement of the sand is seen. Remove the cylinder and determine its weight (W3).
- 6. Keep a representative sample of the excavated sample of the soil for water content determination.

RESULT:

Bulk density of insitu soil, $\gamma_b =$

Dry density of insitu soil, γ_d =

Water content of insitu soil w =

OBSERVATIONS AND CALCULATIONS:

S.No.	Calibration	1	2	3
1.	Weight of sand in cone (of pouring cylinder) W ₂ gm	ERING		
2.	Volume of calibrating container (V)	9		
	in cc SRN	n i	TI	
3.	Weight of sand + cylinder before		Q H	
	pouring W ₁ gm			
4.	Weight of sand + cylinder after			
	pouring W ₃ gm			
5.	Weight of sand to fill calibrating			
	Containers			
	$Wa = (W_1 - W_3 - W_2) gm$			
6.	Bulk density of sand $\gamma s = W_a / V$			
	gm/cc			

S.No.	Measurement of Soil Density	1	2	3
1.	Weight of wet soil from hole Wwgm			
2.	Weight of sand + cylinder before			
	pouring W ₁ gm			

3.	Weight of sand + cylinder after
	pouring W ₄ gm
4.	Weight of sand in hole $W_b = (W_1 - W_2 -$
	W ₄) gm
5.	Bulk density $\gamma_b = (W_w/W_b) \gamma_s (gm/cc)$
	Water content determination:
6.	Container number
7.	Weight of wet soil
8.	Weight of dry soil
9.	Moisture content, w (%)
10.	Dry density $\gamma_d = \gamma_b / [1+w]$ (gm/cc)



Exp.No: 5

SPECIFIC GRAVITY DETERMINATION BY PYCNOMETER

Aim

To determine the specific gravity of soil solids by Pycnometer bottle method.

Apparatus Required:

- 1. Pycnometer of about 1 litre capacity
- 2. Balance accurate to 1 g, glass rod, de-aired distilled water etc.

Theory:

Specific gravity of soil solids is defined as the weight of soil solids to weight of equal volume of water. In effect, it tells how much heavier (or lighter) the material is than water. This test method covers the determination of the specific gravity of soil solids that pass 4.75 mm sieve.

Equation for specific gravity,

 $G = (W_2 - W_1) / ((W_2 - W_1) - (W_3 - W_4))$

Where, W_1 = weight of Pycnometer in grams.

W₂= weight of Pycnometer + dry soil in grams.

W₃= weight of Pycnometer + soil+ water grams.

 W_4 = weight of Pycnometer + water grams.

PROCEDURE:

- 1. Clean and dry the pycnometer and weigh it along with the conical cap (W1 in gm).
- 2. Select about 300 gm of dry soil free of clods and put the same into the pycnometre. Weigh it (W2 in g) with cap and washer.
- 3. Fill the pycnometer with de-aired water up-to half its height and stir the mix with a glass rod. Add more water and stir it. Fit the screw cap and fill the pycnometer flush with the hole in the conical cap and take the weight (W3 in g).
- 4.Remove all the contents from the pycnometer, clean it thoroughly and fill it with distilled water. Weigh it (W4 in g).
- 5. Now use the above equation for determining G.

6. Repeat the same process for additional tests.



RESULT

Average value of Specific Gravity is _____

OBSERVATION:

S.No.	Description	Trial 1	Trial 2	Trial 3
1.	Weight of Pycnometer bottle, W ₁ (gm)			
2.	Weight of Pycnometer + dry soil, W ₂ (gm)			
3.	Weight of Pycnometer + soil + water, W ₃ (gm)			
4.	Weight of Pycnometer bottle, W_4 (gm)	GINER		
5.	Specific gravity, G	KING		
6.	Average, $G = (G_1 + G_2 + G_3) / 3$		0	•

EXP. NO. DETERMINATION OF PARTICLE SIZE DISTRIBUTION

BY SIEVING (GRAIN SIZE ANALYSIS)

Aim: To determine the particle size distribution by sieving (Grain size analysis) and to determine effective size and uniformity coefficient of the soil.

This test is performed to determine the percentage of different grain sizes

contained within a soil. The mechanical or sieve analysis is performed to determine the distribution of the coarser, larger-sized particles. The distribution of different grain sizes affects the engineering properties of soil. Grain size analysis provides the grain size distribution, and it is required in classifying the soil.

Specifications:

This test is specified in IS: 2720 (Part 4) – 1985 – Method of test for soil (Part 4-Grain size analysis).

Equipments Required:

a) Sieves of sizes: 4.75 mm, 2.0 mm, 1.0 mm, $600\square$, $300\square$, $150\square$ and $75\square$. That is,

I.S 460-1962 is used. The sieves for soil tests: 4.75 mm to 75 microns.

- b) Thermostatically controlled oven.
- c) Trays, sieve brushes, mortar with a rubber covered pestle, etc.
- d) Mechanical sieve shaker etc.

Theory:

The grain size analysis is widely used in classification of soils. The data obtained from grain size distribution curves is used in the design of filters for earth dams and to determine suitability of soil for road construction, air field etc. Information obtained from grain size analysis can be used to predict soil water movement although permeability tests are generally used. The method is applicable to dry soil passing through 4.75 mm size sieve less than 10 % passing through 75-micron sieve.

Percentage retained on any sieve = (weight of soil retained / total weight) $\square 100$

Cumulative percentage retained = sum of percentages retained on any sieve

on all coarser sieves

Percentage finer than any sieve = 100 percent minus cumulative Size, N Percentage retained.

Procedures:

- (a) Take a representative sample of soil received from the field and dry it in the oven.
- (b) Use a known mass of dried soil with all the grains properly separated out. The maximum mass of soil taken for analysis may not exceed 500 g.
- (c) Prepare a stack of sieves. Set the sieves one over the other with an ascending order (sieves having larger opening sizes i.e., lower numbers are placed above the one with smaller opening sizes i.e., smaller numbers). The very last sieve is #200 (75 \square sieve). A pan is attached to the lowest 75 \square sieve to collect the portions passing #200 sieves and fit the nest to a mechanical shaker.
- (d) Make sure sieves are clean. If many soil particles are stuck in the openings try to poke them out using brush.
- (e) The whole nest of sieves is given a horizontal shaking for 10 min in sieve shaker till the soil retained on each reaches a constant value.
- (f) Determine mass of soil retained on each sieve including that collected in the pan below.

Table:

The test results obtained from a sample of soil are given below.

Mass of soil taken for analysis $W = \underline{\hspace{1cm}} gm$

S.No.	IS Sieves (mm)	Particle size (mm)	Mass Retained (gm)	% Mas Retained	Cumulative % Mass Retained	% Finer
	4.75	4.75				
	2.00	2.00				
	1.00	1.00				
	0.600	0.600				

0.300	0.300		
0.150	0.150		
0.075	0.075		
Pan			

Cumulative		
mass retained	W'=	gm

Graph (Grain Size in mm vs. Percentage Finer in %):

Draw graph of log sieve size vs. % finer. The graph is known as grading curve. Corresponding to 10%, 30% and 60% finer, obtain diameters from graph these are D10, D30, D60, using these obtain Cc and Cu which further represent how well the soil is graded i.e. whether the soil is well-graded, gap-graded or poorly graded.

Result:

Uniformity coefficient, Cu =

Coefficient of curvature, Cc =

Percentage gravel =

Percentage sand =

Percentage silt =

Verification/Validations:

For the soil to be well graded the value of coefficient of uniformity Cu has to be greater than 4 and Cc should be in the range of 1 to 3. So higher the value of Cu the larger the range of the particle sizes in the soil. So if the Cu value is high it indicates that the soil mass consists of different ranges of particle sizes.



DETERMINATION OF pH OF SOIL

Aim:

To determine the pH value of soil sample.

Apparatus Required:

- pH meter
- 100ml glass beaker
- stirring rods
- wash bottle containing distilled water

Procedure:

- The given soil sample is sieved through 425 micron IS sieve.
- Take 30g of the passed soil in a 100ml beaker.
- 75ml of distilled water shall be added to the beaker. The suspension shall be stirred for a minute.
- Then the beaker shall be covered with a glass cover and allowed to stand for one hour with a glass cover and allowed to stand for one hour with occasional stirring.
- It shall be again stirred well immediately before testing.
- The pH meter shall be calibrated by means of standard buffer solution.
- The electrodes shall be washed with distilled water dried with the help of an ordinary filter paper and then immersed in the soil suspension.
- Three readings of the pH for the soil suspension shall be recorded with brief stirring in between each reading.
- The electrode shall be removed from the soil suspension immediately and washed with distilled water.

Result:

pH value of the given soil sample is _____

Observation:

pH value of distilled water =
pH value of soil + distilled water, trial 1 =
trial 2 =
trial 3 =

Average pH value of soil + distilled water =

DETERMINATION OF ELECTRICAL CONDUCTIVITY OF SOIL

Aim:

To determine the electrical conductivity of given soil to obtain the amount of dissolved salts presents in the soil.

Apparatus Required:

- Glass beaker, stirrer
- Weighing balance
- Measuring pan
- Conductivity meter

Procedure:

- 30g of soil passing through 425micron IS sieve is taken in a glass beaker.
- 75ml of distilled water is added to it and mixed thoroughly with stirrer until uniform mix is obtained.
- The conductivity meter is calibrated using standard solution.
- The conductivity rod is immersed in the prepared solution and readings are noted.
- The same procedure is repeated twice for two other identical soil specimen.

Result:

The electrical conductivity of the given soil sample =

Observation:

SAMPLE	ELECTRICAL CONDUCTIVITY VALUE		
SAMIFLE			
1			
2			
3			
Average Value			

1 millisimon (ms) = 1 milli mho