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SUBSURFACE CHARACTERIZATION AND MONITORING TECHNIQUES:

A DESK REFERENCE GUIDE

Volume II: The Vadose Zone, Field Screening and Analytical Methods Appendices C and D

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This document is not intended to be a guidance or support document for a specific regulatory program. Guidance documents are available from EPA and must be consulted to address specific regulatory issues.

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- American Society for Testing and Materials (ASTM), Philadelphia, PA: Figures 1.3.5, 3.4.6, 5.5.2a and b, 5.6.2b, and A.1.
- American Society of Agronomy/Soil Science Society of America, Madison, WI: Figures 1.6.3, 4.2.1b, 4.2.2, 4.2.3, 6.1.4b, 6.1.7, 6.2.4, 6.2.6, 6.2.7, 6.3.3b, 7.1.1, 7.2.1, 7.2.4, 7.2.2, 7.2.3, 7.2.6, 7.3.2, 7.3.5, 7.3.6, 7.3.7, 7.3.8, 7.5.4, 8.3.1d, 8.3.5a, 9.1.2a, 9.1.3, 9.2.5, 9.2.7, 9.4.4, 9.4.5a, 9.5.2, 10.3.5, 10.6.3, and 10.6.4.

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Williams and Wilkins, Baltimore, MD (Soil Science): Figure 9.3.3.

INTRODUCTION

Many EPA programs, including those under the Resource Conservation and Recovery Act (RCRA) and the Comprehensive Response, Compensation, and Liability Act (CERCLA), require subsurface characterization and monitoring to detect ground-water contamination and provide data to develop plans to prevent new contamination and remediate existing contamination. Hundreds of specific methods and techniques exist for characterizing, sampling, and monitoring the saturated and unsaturated zones at contaminated sites. Existing field methods are often refined and new methods are continually being developed. This guide is designed to serve as a single, comprehensive source of information on existing and developing field methods as of early 1993. Appendix C provides some suggestions on the best places to obtain information on new developments that occur after this guide is completed.

USE OF THIS GUIDE

As the title "Desk Reference Guide" implies, this is not a how-to handbook for the field. Instead, the guide provides, in a single document, enough information about specific techniques to make some judgements concerning their potential suitability for a specific site and also gives information on where to go to find more detailed guidance on how to use the technique. This guide can be used in two major ways:

- 1. Development of Site Characterization and Monitoring Plans. Each subsection listed in the table of contents represents a one-to-two page summary of a specific technique or several related techniques. A table at the beginning of each of the 10 major sections (summarized below), provides general comparative information on all methods covered in the sections, and cross-references relevant methods covered in other parts of the guide. In the summary tables, **boldfacing** is used to identify those techniques that are most commonly used. These tables might also be helpful in identifying new, or less common methods that might be of value for specific objectives or site conditions. Within a grouping of method summary sheets, techniques are listed in approximate order of frequency of use.
- 2. Overview of Specific Methods. Individuals who are unfamiliar with specific methods that are being used or proposed to be used at a hazardous waste site can find a concise description of the method, its applications, major advantages and disadvantages in its use, and major reference sources where more detailed information can be found about the method. To locate information on a specific method, the table of contents should be used to identify the section in which the method is located. If the term used to describe the method is not included in the table of contents, go to the summary table at the beginning of the appropriate section of the guide. If the summary table does not use the term, peruse the listing of alternative names for techniques in the individual summary sheets. For example, the hydraulic percussion drilling method is not listed in neither the table of contents or the summary, and requires looking through the individual summary sheets in Section 2.1 (Drilling Methods), until Section 2.1.6 is reached, which identifies the hollow-rod method as an alternative term for hydraulic percussion.

GUIDE ORGANIZATION AND FORMAT

Site characterization, monitoring, and field screening are related activities for which there might not be a clear dividing line. Generally, site characterization methods involve one-time field point measurements and sampling (or continuous measurements in the case of some geophysical methods) of physical and chemical properties of the subsurface, or multiple measurements to characterize seasonal variations at the site. Monitoring methods, on the other hand, involve sampling or measurements at a single point or the same area over time. Many methods can be used for both site characterization and monitoring, and site characterization activities can continue after monitoring begins to further refine subsurface interpretations. Field screening is a form of site characterization that involves the use of rapid, relatively low-cost field methods (typically chemical) in the field during site characterization to assist in the selection of locations for permanent monitoring well installations or for guiding remediation activities. Field analytical methods are distinguished from field screening methods by having a higher degree of precision and accuracy than field screening methods. This distinction in discussed further in the introduction to Section 10. This guide includes two volumes. The first volume covers solids and ground water and the second volume covers the vadose zone. The site characterization, monitoring, and field screening methods covered in the guide are divided into 10 major sections, which are described below. Because site characterization generally precedes monitoring, earlier sections of the guide tend to cover site characterization methods, while later sections cover monitoring. Finally, field screening and analytical methods are covered in Section 10.

Section 1 (Remote Sensing and Surface Geophysical Methods) covers more than 30 airborne and surface geophysical methods that are often valuable during the initial phases of site characterization. These methods can provide preliminary information on the subsurface to provide guidance on placement of boreholes for direct observation of the subsurface and installation of permanent monitoring wells. A number of these methods can also be useful for monitoring the movement of contaminant plumes.

Section 2 (Drilling and Solids Sampling Methods) covers 20 drilling methods, and a variety of powerdriven and hand-held devices for sampling soils and geologic materials. The section also briefly identifies important soil physical properties that are described in the field.

Section 3 (Geophysical Logging of Boreholes) covers more than 40 borehole logging and sensing techniques for the physical and chemical characterization of the subsurface.

Section 4 (Aquifer Test Methods) covers 10 methods for measuring ground-water well levels or pressure, pumping and slug tests, six categories of ground-water tracers, and several other techniques for measurement of aquifer properties that might be needed for modeling ground-water flow and contaminant transport.

Section 5 (Ground-Water Sampling Devices and Installations) covers more than 20 types of portable ground-water sampling devices and different types of permanent well installations for portable sampling devices. Appendix A (Design and Installation of Monitoring Wells) provides more detailed information on such installations. Section 5 also includes various types of portable and fixed in situ sampling devices and installations. General ground-water sampling methods are covered in Appendix B.

Section 6 (Vadose Zone Hydrologic Properties (I): Water State) covers over 20 methods for measuring vadose zone soil water potential, moisture content, and other soil hydrologic characteristics.

Section 7 (Vadose Zone Hydrologic Properties (II): Infiltration, Conductivity, and Flux) covers four approaches to measuring or estimating infiltration and approximately 30 methods for measuring unsaturated and saturated hydraulic conductivity and water flux in the vadose zone.

Section 8 (Vadose Zone Water Budget Characterization Methods) covers a large number of methods for obtaining data that might be required for water budget calculations to assess contaminant transport in the vadose zone. This includes 37 methods for obtaining various types of hydrometeorologic data, and 16 methods for measuring or estimating transpiration or evapotranspiration.

Section 9 (Vadose Zone Soil-Solute/Gas Sampling and Monitoring Methods) covers six indirect methods for monitoring soil solute movement, more than 20 methods for direct sampling of soil solutions, and a variety of methods for soil gas sampling and gaseous phase characterization in the vadose zone. The section also summarizes a number of methods to measure or estimate soil solute and gas flux in the vadose zone.

Section 10 (Field Screening and Analytical Methods) covers a large number of techniques and groups of techniques for field screening and analysis: Chemical field measurement (three summary sheets), sample extraction procedures (five summary sheets), gaseous phase analytical techniques (five summary sheets), luminescence/spectroscopic techniques (four summary sheets); wet chemistry methods (four summary sheets), and other techniques (five summary sheets). More than 280 specific field methods are covered in this guide. The large number of methods precludes detailed coverage of any single method, which is often available from other sources. Instead, each method has a single-page summary in a uniform format that includes:

- 1. General method category title.
- 2. Method title.
- 3. Other names used to describe method.
- 4. Uses at contaminated sites.
- 5. Method/procedure/device description.
- 6. Method selection considerations.
- 7. Frequency of use.
- 8. Standard Methods/Guidelines (ASTM or other sources that give detailed instruction for use of the specific method).
- 9. Sources for additional information (which provides comparative information where other methods for similar applications are available).

The frequency of use ratings are very approximate, and actual usage might vary from region to region. Similarly, the summary tables at the beginning of each section should not be relied upon as definitive. Specific instrumentation or variants of techniques covered in this guide might have different characteristics than indicated in the summary tables. A specific method that has been rarely used might be suited for certain site-specific conditions. Conversely, site-specific conditions might make a widely-used technique a poor method of choice. When in doubt, obtaining the opinion of more than one person familiar with a particular technique is advisable.

Wherever possible, one or more figures or tables that illustrate instruments or how a method is used are included with summary sheets. These figures and tables have the same number as the section to which they are related (i.e., Figure 1.1.1 and Table 1.1.1 are located after Section 1.1.1 on visible and near infrared remote sensing). Each major section has a brief introduction that defines major concepts and provides an overview of methods covered in the section. Summary tables and figures at the beginning of each section, and index reference tables near the end of a section are numbered in sequence (i.e., Tables 1-1 to 1-3 provide summary information on remote sensing and geophysical methods, and Tables 1-4 and 1-5 provide an index to references contained at the end of the section).

SOURCES OF ADDITIONAL INFORMATION

As indicated above, two types of references are given for each method. First, if ASTM, EPA, or other standard methods, protocols, or guidelines related to the method have been promulgated, or are being developed, these are identified. Otherwise, references that give detailed instructions on how to use the method are cited, if available.

Secondly, major references that provide information on the use of the method in the context of groundwater and hazardous waste site investigations are listed. All references are in a single section. EPA documents are indicated (with EPA and NTIS numbers). Appendix C (Guide to Major References on Subsurface Characterization and Monitoring) provides annotated descriptions of more than 70 major books and reports and over 80 published conference and symposium proceedings that can serve as information sources for general and specific aspects of soil quality and ground-water field screening, characterization, and monitoring.

The following EPA documents are recommended for use as companions to this guide (all of which are available for no cost from U.S. EPA's Center for Environmental Research Information (see Appendix C for ordering address): Ground-Water Handbook, Volume 1: Ground Water and Contamination; Volume 2: Methodology (U.S. EPA, 1990 and 1991a), Site Characterization for Subsurface Remediation (U.S. EPA, 1991b), Handbook of Suggested Practices for the Design and Installation of Ground-Water Monitoring Wells (Aller et al., 1991), Description and Sampling of Contaminated Soils: A Field Pocket Guide (Boulding, 1991), and Use of Airborne, Surface, and Borehole Geophysical Techniques at Contaminated Sites: A Reference Guide (U.S. EPA, 1993). Other EPA documents that are available from NTIS and commercially published references that can be of potential value are too numerous to be named individually here. Appendix B should provide guidance concerning other publications that might be worth obtaining.

REFERENCES

- Aller, L. et al. 1991. Handbook of Suggested Practices for the Design and Installation of Ground-Water Monitoring Wells. EPA/600/4-89/034, 221 pp. Also published in 1989 by the National Water Well Association, Dublin, OH in its NWWA/EPA series, 398 pp. [Nielsen and Schalla (1991) contain a more updated version of the material in this handbook that is related to design and installation of groundwater monitoring wells.]
- Boulding, J.R. 1991. Description and Sampling of Contaminated Soils: A Field Pocket Guide. EPA/625/12-91/002, 122 pp.
- Nielsen, D.M. and R. Schalla. 1991. Design and Installation of Ground-Water Monitoring Wells. In: Practical Handbook of Ground-Water Monitoring, D.M. Nielsen (ed.), Lewis Publishers, Chelsea, MI, pp. 239-331.
- U.S. Environmental Protection Agency (EPA). 1990. Handbook Ground Water. Volume I: Ground Water and Contamination. EPA/625/6-90/016a, 144 pp.
- U.S. Environmental Protection Agency (EPA). 1991a. Handbook Ground Water. Volume II: Methodology. EPA/625/6-90/016b, 141 pp.
- U.S. Environmental Protection Agency (EPA). 1991b. Site Characterization for Subsurface Remediation. EPA/625/4-91/026, 259 pp.
- U.S. Environmental Protection Agency (EPA). 1993. Use of Airborne, Surface, and Borehole Geophysical Techniques at Contaminated Sites: A Reference Guide. EPA/625/R-92/007.

SECTION 6

VADOSE ZONE HYDROLOGIC PROPERTIES (I): WATER STATE

Water state in the subsurface is measured in terms of hydraulic head in the saturated zone (see Section 4.1) and negative pressure potential or suction in the vadose zone. Water movement in the vadose zone is determined by the interaction of three major types of energy potentials: (1) Matric potential (the attraction of water to solids in the subsurface, (2) osmotic potential (the attraction of solute ions to water molecules), and (3) gravitational potential (the attraction of the force of gravity toward the earth's center). Matric and osmotic potentials are negative and serve to inhibit the movement of water when the vadose zone is unsaturated. Unsaturated flow occurs, however, whenever the force of gravity on a water molecule exceeds matric plus osmotic potential. Water flow in the vadose zone is strongly influenced by the moisture content, with flow decreasing as moisture content decreases.

Table 6-1 provides summary information on six major techniques for measuring soil water potential and a dozen methods for measuring soil moisture content. The measurement of soil water potential and moisture content in the vadose zone is intimately connected, and a specific measurement technique can be classified as measuring potential or moisture content, depending on the perspective of the writer in the literature. Either measurement can be used to obtain the other if a **moisture characteristic curve** has been developed (Section 6.3.1). **Porous cup tensiometers** are the most commonly used method for measuring soil water potential in the vadose zone. The **gravimetric** method is most commonly used to measure moisture content from soil samples, and the neutron probe and gamma gamma methods are most commonly used for in situ measurement of soil moisture. The relatively recent commercial availability of **dielectric** or capacitance sensors (Section 6.2.3) is likely to increase the use of this method, which provides accuracy similar to the neutron probe without some of the disadvantages of nuclear methods (i.e., radioactive sources). Similarly, **time domain reflectometry**, a relatively new method (Section 6.2.4), is becoming more widely used with the advent of commercially available units. All methods for vadose zone measurement of water content or matric potential have limitations with respect to soils contaminated with nonaqueous phase liquids, due to interference effects.

Other field-measurable hydrologic properties of the vadose zone, which might be of use in evaluating contaminant transport include water sorptivity and diffusivity (Section 6.3.2) and available water capacity (Section 6.3.3). Sorptivity and diffusivity are properties that are significant in evaluating infiltration of water into the subsurface (discussed in more detail in Section 7.1). Available water capacity is a measure of the ability of soil to store water.

Table 6-1 Summary Information on Vadose Zone Water State Measurement and Monitoring Methods

Method	Property Measured	Accuracy/ Range	Sections	
Vadose Zone Soil Water Potential	Measurement ^a			
Porous Cup Tensiometers	Capillary pressure	0 to -85 kPa ^b	6.1.1	
Thermocouple Psychrometers	Relative humidity	0 to -80 kPa° -200 to -8,000 kPa ^b -100 to -5,000 kPa°	6.1.2	
Water Activity Meter	Relative humidity	0 to -31.600 kPa	6.1.2	
Resistance Sensors	Resistance	-50 to -1,500 kPa°	6.1.3	
Gypsum Blocks	Resistance	0 to -30 kPa ^b	6.1.3	
Fiberglass/Nylon Cells	Resistance	No limits ^b	6.1.3	
Electrothermal Methods	Heat transfer	0 to -200 kPa	6.1.4	
Osmotic Tensiometers	Osmotic + pressure potential	0 to -1,500 kPa ^b	6.1.5	
Filter-Paper Method	Water content	-10 to -100,000 kPa	6.1.6	
Electro-Optical Sensors	Optical properties	0 to -2,400 kPa	6.2.6	
Vadose Zone Soil Water Content 1	Measurement*			
Gravimetric	Weight	đ	6.2.1	
Gamma-Gamma	Radiation	đ	6.2.2, 3.3.2	
Neutron Moisture Probe	Radiation	đ	6.2.2, 3.3.3	
Dielectric Sensors	Dielectric	đ	6.2.3	
Time Domain Reflectometry	Dielectric	d	6.2.4	
Nuclear Magnetic Resonance	Magnetic field	d	3.2.4, 10.6.3	
Electro-Optical Sensors	Optical properties	d	6.2.6	
CAT Scan	Radiation	d	6.2.7	
Thermal Infrared	Remote sensing	d	1.1.3	
Active Microwave	Remote sensing	d	1.1.4	
Four-Electrode Method	Resistivity	d	9.1.1	
Salinity Sensors	Conductivity	đ	9.1.3	
Electromagnetic Induction	Conductivity	đ	9.1.4	

Boldface = most commonly used methods.

*Moisture content can be determined from measurement of soil water potential and vice versa by the use of a moisture characteristic curve, which relates matric potential to water content (Section 6.3.1). The pascal is the Standard International unit for measuring pressure used by the Soil Science Society of America. The bar is commonly used as a pressure unit in vadose zone investigations: 1 kPa = 1 centibar.

Indicated by Rehm et al. (1985).

Indicated by Bruce and Luxmoore (1986).

"Most methods for measuring moisture content are accurate to around 1%. Gravimetric methods and nuclear methods can be accurate to 0.1% or less.

6.1 VADOSE ZONE SOIL WATER POTENTIAL

6.1.1 Porous Cup Tensiometers

<u>Other Names Used to Describe Method</u>: Capillary potentiometer, soil hygrometer, soil moisture meter, transiometer. Tensiometers often are described according to the type of device that is used to measure pressure: Vacuum gauge, water manometer, mercury-water manometer, or electrical pressure transducer.

<u>Uses at Contaminated Sites</u>: Measuring water (matric) potential and gradients in the unsaturated zone; irrigation scheduling; performing root zone delineation; developing moisture characteristic curves (see Section 6.3.1); can be used to measure and monitor changes in moisture content if matric potential-water content relationship is known.

<u>Method Description</u>: Many designs for tensiometers have been developed. Most have the following basic elements: (1) A porous tip or cup attached to a barrel or connective tube, (2) a removable air tight cap for filling the tensiometer with water, and (3) a device to measure pressure in the water in the porous cup. The ceramic cup (or other material, such as fritted glass) is placed in the soil, filled with water, and the unit is sealed. Pores in the cup form a continuum with the pores in the soil, and water moves into or out of the tensiometer until equilibrium is reached. The measured pressure corresponds to the water pressure in the soil. Figure 6.1.1 shows three types of porous cup tensiometers. A transiometer is a type of porous cup tensiometer in which a pressure transducer is placed inside the porous cup, rather than at the surface.

<u>Method Selection Considerations</u>: The useful range of tensiometers is 0 to 0.85 bars capillary pressure when the ambient atmospheric pressure is around 76 centimeters of mercury. Advantages: (1) Provide continuous in-place measurements of the same soil material over time; (2) are relatively inexpensive and simple; (3) transducer unit responds fairly rapidly to water content changes and can be used for automatic data collection; and (4) transiometers can be used to measure soil water potential in both saturated and unsaturated conditions. Disadvantages: (1) Units fail at the air entry value of the ceramic cup, generally about -0.8 atmospheres; (2) unit will not operate properly unless good contact is made between cup and soil; (3) are sensitive to temperature changes; (4) water content estimates prone to error resulting from uncertainty of moisture-matric potential relationship (hysteresis causes different curves depending on soil is wetting or drying); (5) difficult to install at great depth in the vadose zone; (6) air in the system causes errors in measurement, and special efforts, such as using deaired water, are required to minimize such problems; (7) lower air pressures at higher elevations reduce the operating range; (8) operation will be affected if the surface tension characteristics of chemical liquid wastes in the vadose zone differ from that of water; and (9) multiple calibration curves are required for soil moisture monitoring in stratified media.

Frequency of Use: Widely used for pressure measurement; usually not recommended for water content measurement.

Standard Methods/Guidelines: ASTM (1991), Cassel and Klute (1986).

Sources for Additional Information: Brakensiek et al. (1979), Gairon and Hadas (1973), Holmes et al. (1967), Morrison (1983), Rehm et al. (1985), Stannard (1986), Troolen et al. (1986-transiometer), Wilson (1980, 1981). See also, Table 6-2.



550 12

Figure 6.1.1 Three types of porous-cup tensiometers (Morrison, 1983, by permission).

6.1 VADOSE ZONE SOIL WATER POTENTIAL

6.1.2 Thermocouple Psychrometers

Other Names Used to Describe Method: Spanner/Peltier psychrometer, Richards-Ogata/wet-loop psychrometer, thermocouple hygrometer, in situ hygrometer.

<u>Uses at Contaminated Sites</u>: Measuring water potential (sum of osmotic and matric potential) and gradients in the unsaturated zone; estimating water content (if moisture characteristic curve is developed, see Section 6.3.1); measuring soil water flux in the vadose zone (see Section 7.5.2).

<u>Method Description</u>: Soil water potential is calculated based on measurement of relative humidity within the soil voids. A basic psychrometer unit consists of: (1) A porous bulb, with a chamber in which the relative humidity of the soil is sampled, (2) a sensitive thermocouple, (3) a heat sink, and (4) a reference electrode. Two major types are available, wet bulb and dew point; both types rely on cooling of the thermocouple junctions by the Peltier effect, but differ in how temperature is controlled once the dew point of the sample is reached. With in situ measurements of soil water potential, the thermocouple is protected by a cup-shaped device that maintains a void in the soil. Calibration curves relating relative humidity to water potential, osmotic potential, and temperature (if temperature in the subsurface varies) need to be developed in the laboratory. Figure 6.1.2 illustrates: (a) A basic Spanner, and (b) a modified Spanner-type psychrometer.

Method Selection Considerations: The dew point method is more accurate than the wet bulb method. The useful range is 10 to 70 bars capillary pressure. Advantages: (1) In situ pressure measurements are possible for very dry soils in arid regions; (2) continuous recording of pressures is possible; (3) can be interfaced with portable or remote data collection systems; and (4) depth is no limitation (installations have gone as deep as 300 feet). Disadvantages: (1) Water content estimates prone to errors due to hysteresis; (2) even in very dry soils, the relative humidity is high, making accurate calibration difficult; (3) good contact between bulb and surrounding material might be difficult to achieve; (4) provide only point measurements; (5) accurate calibration curves for deep regions of the vadose zone might be difficult to obtain; (6) instruments are expensive, fragile, and require great care in installation; (7) contamination of the chamber interior or thermocouple can result in erroneous readings; (8) interference from dissolved solutes is likely in calcium-rich waste and acid media and can cause thermocouple wire corrosion problems; (9) perform very poorly in very wet media (water pressure >1 bar); (10) accuracy of near-surface measurements is adversely affected by diurnal changes in heat flux; (11) unsealed cup units are susceptible to attack by fungi and bacteria; and (12) ceramic cup psychrometers respond slowly to rapid changes in moisture content.

Frequency of Use: Widely used in agricultural research; sometime used at hazardous waste sites in the arid west.

Standard Methods/Guidelines: Rawlins and Campbell (1986).

Sources for Additional Information: Morrison (1983), Rehm et al. (1985), Thompson et al. (1989), Wilson (1981). See also, Table 6-2.



(b)

Figure 6.1.2 Thermocouple psychrometers: (a) Spanner psychrometer (Morrison, 1983, after Meyn and White, 1972, by permission); (b) Double-loop, temperature-compensating psychrometer (Morrison, 1983, after Meeuwig, 1972, by permission).

6.1 VADOSE ZONE SOIL WATER POTENTIAL

6.1.3 Electrical Resistance Sensors

Other Names Used to Describe Method: Four-electrode soil moisture probe, electrical resistance blocks, porousblock method, soil moisture blocks.

<u>Uses at Contaminated Sites</u>: Measuring/monitoring water potential and water content in the unsaturated zone; monitoring of soil freezing.

<u>Method Description</u>: In the **porous-block method**, two electrodes (Figure 6.1.3a and b) are imbedded in a porous block (nylon cloth, fiberglass, or casting plaster), or multi-electrode probes can be used (Figure 6.1.3c). Calibration curves are first developed by placing the porous block in soil typical of the area to be measured and resistivity is plotted against changes in matric potential. In the field, the porous blocks can be placed in a hole and buried, or horizontally in the side of a trench, and the blocks are allowed to equilibrate with the surrounding soil. Matric potential then can be monitored by taking resistance measurements, using the calibration curves to convert the measurements to pressure. Water content also can be monitored either by using the procedure described above to develop a calibration curve for water content or by using a moisture characteristic curve if a resistance-water potential calibration curve has been developed (Section 6.3.1).

<u>Method Selection Considerations</u>: Advantages: (1) Are inexpensive and relatively easy to install; (2) measurements can be recorded from many units over a large area using an automated recording system; and (3) can be calibrated for either suction or water content. **Disadvantages**: (1) Calibration procedures can be complicated and time consuming if accurate measurement of water potential for evaluation of hydraulic gradient is required; (2) restricted water flow at the interface between the smooth face of a porous black creates some problems for measurements in coarse soil material; (3) small changes in electrolyte concentration of the soil water (which might well occur at contaminated sites) will affect resistivity readings; (4) measurements are made in equilibrium with matric potential, so moisture content is inferred from matric potential rather than actual moisture content; (5) gypsum sensors might dissolve in the subsurface; (6) water content estimates are prone to error resulting from uncertainty of moisture-matric potential relationship (hysteresis causes different curves depending on soil is wetting or drying); and (7) are rather insensitive to moisture changes in the wet range.

<u>Frequency of Use</u>: Commonly used for irrigation timing and other qualitative field-monitoring programs. Less common for accurate measurement of soil hydrologic properties.

Standard Methods/Guidelines: Campbell and Gee (1986), Gardner (1986).

Sources for Additional Information: Bouyoucos (1960), Everett et al. (1983), Gairon and Hadas (1973), Holmes et al. (1967), Morrison (1983), Rehm et al. (1985), Schmugge et al. (1980). See also, Table 6-2.



- (c)
- Figure 6.1.3 Electrical resistance sensors: (a) Rectangular soil moisture block (Morrison, 1983, by permission); (b) Fiberglass and Monel soil moisture sensor with thermistor (Morrison, 1983, after Colman and Hendrix, 1949, by permission); (c) Multielectrode probe (Morrison, 1983, after Perrier and Marsh, 1958, by permission).

6.1 VADOSE ZONE SOIL WATER POTENTIAL

6.1.4 Electrothermal Methods

Other Names Used to Describe Method: Thermal diffusivity, heat diffusion/dissipation sensors.

<u>Uses at Contaminated Sites</u>: Measuring water potential and gradients in the unsaturated zone; estimating water content (if moisture characteristic curve is developed, see Section 6.3.1); measuring soil temperature.

<u>Method Description</u>: Similar to the resistivity method (Section 6.1.3), except that thermal diffusivity of an implanted porous sensor, which is in equilibrium with the surrounding soil, is measured. A known amount of heat is applied in the center of the sensor and the rate of dissipation is measured, which is a function of water content. Major types of sensors include: (1) Porous-block type with embedded electrical elements (Figure 6.1.4a), (2) direct-contact type with electrical elements in direct contact with the soil, and (3) modified direct-contact probe or cell, in which the heating wire is enclosed in a protective sheath with high thermal conductivity (Figure 6.1.4b). Calibration curves of matric potential vs. temperature difference are obtained in the laboratory with soils from the site using a pressure plate apparatus. The matric potential is related to water content by preparing a moisture characteristic curve (Section 6.3.1).

<u>Method Selection Considerations</u>: Useful range is 0 to 2 bars capillary pressure. Advantages: (1) Are simple; (2) can be interfaced with data acquisition systems for remote collection of data; (3) measurements are independent of salt content of soil; (4) calibration appears to remain constant; (5) can be used to measure soil temperature as well as matric potential; and (6) are useful for measurement of water contents in the dry range. **Disadvantages:** (1) Water content estimates subject to hysteresis; (2) calibration is required for each change in texture; and (3) might be difficult to install at depth in the vadose zone and to maintain good contact between the sensor and medium.

Frequency of Use: Uncommon.

Standard Methods/Guidelines: Campbell and Gee (1986).

Sources for Additional Information: Morrison (1983), Thompson et al. (1989), Wilson (1981). See also, Table 6-2.





(b)

Figure 6.1.4 Electrothermal sensors: (a) Porous-block type (Morrison, 1983, after Phene et al., 1971b, by permission); (b) Modified direct-contact probe (Sophocleous, 1979, by permission).

6.1 VADOSE ZONE SOIL WATER POTENTIAL

6.1.5 Osmotic Tensiometers

Other Names Used to Describe Method --

Uses at Contaminated Sites: Measuring of combined osmotic and pressure potential.

<u>Method Description</u>: An osmotic tensiometer uses a confined solution of polyethylene glycol, rather than deaired water, as the reference solution and a semipermeable membrane, which separates the confined solution from the soil water (Figure 6.1.5). Small ions and molecules in the soil water are able pass through the membrane, and once equilibrium is attained between the soil water and the reference solution, a pressure transducer measures subsequent soil moisture-related pressure changes.

<u>Method Selection Considerations</u>: Advantages: Allows differentiation of osmotic and pressure components of soil water potential if used with porous cup tensiometer. **Disadvantages**: (1) Are susceptible to fluid leakage and instrument drift; (2) require long equilibration times (hours to days); and (3) are sensitive to temperature changes.

<u>Frequency of Use</u>: Uncommon for reasons cited above. Thermocouple psychrometers are the preferred method for measuring combined osmotic and pressure potential.

Standard Methods/Guidelines --

Sources for Additional Information: Bocking and Fredland (1979), Morrison (1983), Peck and Rabbidge (1966a,b, 1969), Rehm et al. (1985).



Figure 6.1.5 Osmotic tensiometer (Morrison, 1983, after Peck and Rabbidge, 1966a, by permission).

6.1 VADOSE ZONE SOIL WATER POTENTIAL

6.1.6 Filter-Paper Method

Other Names Used to Describe Method --

Uses at Contaminated Sites: Measuring soil water potential and/or moisture content.

<u>Method Description</u>: The filter-paper method involves the collection of soil cores at different locations and/or depths. Each soil core is placed in a sealed container in contact with filter paper, which has been pretreated with 3% pentachlorophenol dissolved in methanol to prevent microbial degradation. In the laboratory, the samples are maintained at a constant temperature for at least 1 week to allow equilibration of moisture between the soil and the filter paper. Gravimetric water content of the soil and filter paper is determined using the oven-drying method (see Section 6.2.1). The matric potential then is calculated using a calibration equation (Figure 6.1.6).

<u>Method Selection Considerations</u>: Advantages: (1) Reasonably accurate over a wide range of matric potentials; (2) requires minimal and inexpensive equipment; (3) cores can be used to directly measure moisture content and to measure bulk density; and (4) simplicity allows taking a large number of measurement to characterize spatial variability. Disadvantages: (1) Soil core collection is destructive and does not allow repeated measurements at exactly the same location (Figure 6.2.1 in the next section shows patterns for sequential sampling, if this method is used); and (2) different filter papers might require development of separate calibration curves.

Frequency of Use: Has commonly been used in studies of western rangeland hydrology.

Standard Methods/Guidelines: ASTM (1992b).

Sources for Additional Information: McQueen and Miller (1968a, 1968b), Sorenson et al. (1989) cite 18 references on this method.



Figure 6.1.6 Calibration equations (for above and below field capacity) used to determine soil matric potential from filter-paper water content (Sorenson et al., 1989).

6.1 VADOSE ZONE SOIL WATER POTENTIAL

6.1.7 Water Activity Meter

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Measuring soil water potential in very dry soils.

<u>Method Description</u>: A water activity meter (another term for relative humidity) can be used to measure soil water potential in a soil using principles similar to thermocouple psychrometers (Section 6.1.2). A small amount of soil (4 centimeters in diameter and 0.5 centimeters thick) on a slide tray is placed in a measuring chamber (Figure 6.1.7). The sample temperature is monitored using a built-in infrared thermometer and the dew point of the water vapor above the sample is measured using a cooled mirror. The dew point and sample temperature are recorded on a data logger and converted to relative humidity, using an algorithm that accounts for temperature differences between the soil sample and chamber as both are equilibrating at room temperature. Gee et al. (1992) obtained good results with a commercially available water activity meter for measuring soil water potential for soil textures ranging from sand to clay.

<u>Method Selection Considerations</u>: Advantages: (1) Provides rapid (3 to 5 minutes/sample), accurate measurements over a wide range of soil water potentials (-4 to <-2,640 kPa); (2) allows measurement of soil water potential in soils that are too dry from tensiometer measurements (<-85 kPa); (3) instrumentation is relatively simple and less subject to errors than thermocouple psychrometers for measuring low potentials; and (4) commercially available instrumentation can be readily used; Disadvantages: Requires collection of soil samples, so limited to relatively shallow depths if time-series monitoring is desired (see 6.2.1 for possible sampling patterns).

Frequency of Use: Only recently applied to measurement of water potential in soil samples.

Standard Methods/Guidelines: --

Sources for Additional Information: Gee et al. (1992).



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Figure 6.1.7 Schematic of a water activity (relative humidity) meter (Gee et al., 1992, by permission).

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6.2 VADOSE ZONE MOISTURE CONTENT

6.2.1 Gravimetric Methods

Other Names Used to Describe Method: Oven-drying method, carbide/gas pressure method.

Uses at Contaminated Sites: Measurement of soil moisture content.

<u>Method Description</u>: **Oven-dry method**: The mass of a sample collected in the field is weighed before and after oven drying, typically at 105°C, the difference being the water content. Other methods of drying, such as a microwave oven and direct heating using a hotplate, stove, or blowtorch, can be used for more rapid, but less accurate measurements in the field. **Carbide method**: A soil sample of known weight is placed in a container with calcium carbide. The calcium carbide reacts with water, releasing a gas. After completion of the reaction, the gas pressure, registered on a gage, is converted into water content on a dry weight basis. Since all gravimetric moisture measurements require destructive sampling, the careful design of the sample collection sequence is required to measure changes in moisture content over time (Figure 6.2.1). **Other gravimetric methods**: Other techniques of drying and soil moisture extraction include: (1) Centrifugation, (2) pressure plate extraction, and (3) desiccation. Section 9.3.4 further discusses these and other methods of soil water extraction from solids samples.

Method Selection Considerations: Standard Oven-Dry Method Advantages: (1) The most accurate available method and serves as the standard method for the calibration of all other moisture determination techniques; (2) is simple; (3) provides a direct measurement of the mass of water. Standard Oven-Dry Method Disadvantages: (1) Obtaining representative moisture values in a heterogeneous profile is difficult, requiring a large number of replicate samples for each depth increment; (2) is destructive, requiring removal of samples for laboratory analysis and thus preventing additional measurements at the same sites; (3) expensive if large numbers of samples are required; (4) plotted vertical moisture profiles will not be accurate if water is moving rapidly through the vadose zone, because water distribution profile is changing as samples are being taken; (5) samples from contaminated sites might require special handling if hazardous contaminants are present; (6) not suitable for nongranular media (i.e., fractured rock, carbonates); and (7) sample collection might be difficult in indurated layers, such as fragipans, when soil is very dry (soil difficult to penetrate) or very wet (soil will not remain in sampling tool), and when soils are frozen. Other Drying Methods Advantages: Generally faster than standard oven-dry method. Other Drying Methods Disadvantages: (1) Might not be as accurate as standard oven-drying; (2) with microwave oven, sample might explode and be lost if power level is too high; and (3) other disadvantages are the same as for oven drying. Carbide Method Advantages: (1) Can be used in the field and is more rapid than oven-drying; (2) initial capital investment is lower. Carbide Method Disadvantages: (1) Might not be as accurate as standard oven-drying; and (2) other disadvantages are same as for oven drying.

Frequency of Use: Widely used. The ASTM oven-dry method is the standard by which the accuracy of other moisture measurement methods are evaluated.

Standard Methods/Guidelines: Gardner (1986); Standard oven-dry method: ASTM (1990); Microwave oven method: ASTM (1987); Direct heating method: ASTM (1989a); Carbide method: ASTM (1989b).

Sources for Additional Information: Everett et al. (1983), Morrison (1983), Thompson et al. (1989), Wilson (1981). See also, Table 6-2.

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6	1	2	3	4	5
3	4	5	6	1	2
5	6	1	2	3	; 4
2	3	4	5	6	1

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(b)

Figure 6.2.1 Arrangement of boreholes for gravimetric soil-moisture sampling: (a) Rectangular microplots; (b) along perimeters of polygons (Brown et al., 1983)

6.2 VADOSE ZONE MOISTURE CONTENT

6.2.2 Nuclear Methods

Other Names Used to Describe Method: Neutron probe, gamma transmission/double-tube gamma method.

Uses at Contaminated Sites: Measuring and monitoring of changes in soil moisture content.

<u>Method Description</u>: The neutron method, which measures moisture content based on the interaction between neutrons and hydrogen atoms in water molecules, is discussed in Section 3.3.3, and the gamma-gamma method for measuring soil moisture is discussed in Section 3.3.2. Soil moisture using the neutron method can be measured using either a surface neutron probe (Figure 6.2.2a) or a depth probe (see Figure 3.3.3). Near-surface soil moisture measurements usually involve the gamma-transmission method, in which a gamma photon source and detectors are lowered simultaneously down two parallel boreholes (Figure 6.2.2b). In boreholes, the gamma-scattering method is used (see Figure 3.3.2).

<u>Method Selection Considerations</u>: Neutron probe: See Section 3.3.3; Double-tube gamma method: See Section 3.3.2.

<u>Frequency of Use</u>: Neutron probes are commonly used for monitoring of soil moisture in the near surface. The double-tube gamma method is less common.

Standard Methods/Guidelines: Neutron probe: ASTM (1988, 1992a), Gardner (1986).

Sources for Additional Information: Table 3-4 in Section 3 provides an index of over 100 references on the neutron method and around 40 references on gamma-gamma logging methods.



(b)

Figure 6.2.2 Nuclear methods for soil moisture measurement: (a) Cross-section of surface neutron probe (Morrison, 1983, after DeVries and King, 1961, by permission); (b) Double-tube gamma method for soil moisture content determination (Rehm et al., 1985, Copyright © 1985, Electric Power Research Institute, EPRI EA-4301, Field Measurement Methods for Hydrogeologic Investigations: A Critical Review of the Literature, reprinted with permission).

6.2 VADOSE ZONE MOISTURE CONTENT

6.2.3 Dielectric Sensors

Other Names Used to Describe Method: Radio frequency/microwave techniques, capacitance techniques, capacitive sensors, 'fringe' capacitance, resonance capacitance, in situ permittivity meter.

Uses at Contaminated Sites: Measuring soil moisture content.*

<u>Method Description</u>: Basic principles are similar to the induced polarization surface geophysical method (see Section 1.2.3 for discussion of frequency domain and time domain IP methods), except that sensors are placed below the ground surface. A variety of capacitive sensors (Figure 6.2.3a,b) have been developed that measure the dielectric properties of soil, which are primarily related to water content. These sensors depend upon specific electrode configurations and detailed calibration. **Dielectric probes**, which measure vertical soil moisture profiles in a cased holed similar to neutron probes (Section 3.3.3), are a relatively recent development. Dielectric probes have significant advantages over neutron probes and other nuclear methods for measuring soil moisture (see below).

Method Selection Considerations: Sensor Advantages: (1) With accurate calibration, can provide accurate values for soil moisture; (2) can be placed at any depth for obtaining moisture profile data; (3) a wide variety of sensor configurations, from very small to large, are possible, allowing some control over the sensor volume of influence; and (4) capacitive sensors have high precision and the property they measure (dielectric constant) is primarily related to water content. Sensor Disadvantages: (1) The moisture sensor must be implanted properly to minimize disturbance to the soil; (2) long-term reliability and maintenance of the calibration is uncertain, especially if the ionic concentration of the soil water changes; and (3) cost of readout devices and interfaces with remote collection platforms is high. Probe Advantages: (1) Provide better resolution in measuring vertical soil moisture profiles than neutron probes; (2) are less expensive than neutron probes and time domain reflectometry sensors; (3) are as accurate as neutron probes without having to deal with radioactive materials; and (4) can be used to accurately determine position of a wetting front and ground-water level in soil. Probe Disadvantages: (1) Special care is required to make sure that there are no air gaps outside the access tube, because relatively limited radial penetration gives more weight to measurements near the borehole compared to neutron probe; (2) less sensitive at high moisture contents than low moisture contents; and (3) air-soil interface affects accuracy of measurements of in the upper 20 to 50 centimeters of soil.

<u>Frequency of Use</u>: Numerous prototypes have been developed. Relatively recent development of commercially available units means that this method is likely to be used more commonly in the future.

Standard Methods/Guidelines: --

Sources for Additional Information: Morrison (1983), Schmugge et al. (1980). See also, Table 6-2.

*Capacitance sensors are classified here as moisture sensors because they are most commonly calibrated to measure soil moisture. They could just as easily be classified as a matric potential measurement technique because they operate by moisture moving into the sensor in response to the matric potential gradient.



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(a)



(b)

Figure 6.2.3 Capacitance sensors: (a) Capacitance probe (Morrison, 1983, after Thomas, 1966, by permission); (b) Cylindrical sensor (Morrison, 1983, after Wobschall, 1978, by permission).

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6.2 VADOSE ZONE MOISTURE CONTENT

6.2.4 Time Domain Reflectometry

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Measuring soil moisture content; estimating soil bulk electrical conductivity.

<u>Method Description</u>: Use of time domain reflectometry to measure soil moisture content is a relatively recent development that shows great promise for field applications. Volumetric water content can be determined based on measuring the travel time and the attenuation of the amplitude of an electromagnetic pulse launched along one or more transmission lines (coaxial, two-, three-, or four-rod probes) embedded in the soil. Portable probes can be used to make multiple near-surface measurements or in situ probes of varying length can be installed vertically to different depths, or horizontally at different depths in the side of a trench (Figure 6.2.4). The TDR trace can be recorded either on a photograph of the oscilloscope display or on an X-Y recorder. The measured dielectric constant is converted to volumetric water content using an empirically derived equation that can be applied to many soils. Electrical conductivity also can be estimated from the attenuation of the signal (Section 9.1.4).

<u>Method Selection Considerations</u>: Advantages: (1) With accurate calibration, can provide accurate values for soil moisture; (2) can be placed at any depth for obtaining moisture profile data; (3) a wide variety of sensor configurations, from very small to large, are possible, allowing some control over the sensor volume of influence; (4) readily amenable for use with automatic data acquisition systems; and (5) available from several commercial sources. **Disadvantages**: (1) The moisture sensor must be implanted properly to minimize disturbance to the soil; (2) long-term reliability and maintenance of the calibration is uncertain, especially if the ionic concentration of the soil water changes; and (3) cost of readout devices and interfaces with remote collection platforms is high.

Frequency of Use: Relatively new method with good potential for field applications.

Standard Methods/Guidelines: --

Sources for Additional Information: See Table 6-2.



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Figure 6.2.4 Diagram of vertical and horizontal TDR probe installations for soil moisture monitoring at different depths (Topp and Davis, 1985a, by permission).

6.2 VADOSE ZONE MOISTURE CONTENT

6.2.5 Nuclear Magnetic Resonance (NMR)

Other Names Used to Describe Method: Nuclear-magnetic logging.

Uses at Contaminated Sites: Measuring porosity, permeability, moisture content, pore-size distribution, available water.

<u>Method Description</u>: A magnetic field is induced using a pulsed, direct current, polarizing field to align a fraction of the nuclei of hydrogen atoms (protons). When the polarizing field is shut off, the probe records the precession of the protons into the Earth's magnetic field. The proton relaxation time is short for fluids in solids or bound to a surface, but is much longer for fluids free to move in pore spaces. Figure 6.2.5a shows components of a pulsed nuclear magnetic resonance sensor and Figure 6.2.5b shows a prototype unit for in situ measurements of soil moisture. Installation involves digging a test pit to the desired depth, driving a thin-walled plastic tube into the bottom, and excavating around the tube to a depth of about 4 centimeters. The sensor is slipped over the tube and seated firmly and the excavation is backfilled with a coaxial cable running to the surface, which is plugged into the instrumentation for inducing the magnetic field and measuring the response when it shut off.

<u>Method Selection Considerations</u>: Advantages: (1) More precise characterization of free and bound water and porosity than other logging methods; and (2) prototype units for in situ measurement of soil moisture have been developed. Disadvantages: (1) Use limited to large boreholes (generally >7 inches) filled with drilling mud (magnetite powder usually has to be added to the mud to eliminate the borehole contribution to the log); (2) installation procedures are relatively complex for in situ units; and (3) equipment availability might be a problem.

<u>Frequency of Use</u>: Not widely used for petroleum applications and relatively unknown for ground-water applications. Potentially very useful if borehole conditions are appropriate.

Standard Methods/Guidelines: --

Sources for Additional Information: NMR general: Abragam (1961), Schlichter (1963), see also, Section 10.6.3; Borehole applications: See Section 3.2.4; Soil moisture applications: Morrison (1983), and references indexed in Table 6-2.



Figure 6.2.5 Nuclear magnetic resonance: (a) Components of a pulsed nuclear magnetic resonance sensor and associated instrumentation; (b) Prototype in situ nuclear magnetic resonance sensor (Morrison, 1983, after Matzkanin and Gardner, 1974, by permission).

6.2 VADOSE ZONE MOISTURE CONTENT

6.2.6 Electro-Optical Sensors

Other Names Used to Describe Method: Electro-optical switch sensor, CdS photoresistor sensor.

Uses at Contaminated Sites: In situ monitoring of soil water content and/or matric potential.

<u>Method Description</u>: Several types of electro-optical sensors have been developed that can use changes in the optical properties of different materials at different moisture contents to measure soil moisture and/or matric potential. The electro-optical switch sensor involves placement of a nylon filter disk in the gap of an electro-optical switch. (Figure 6.2.6a). An infrared light emitting diode (IR LED) sends a signal that passes through the filter disc and is sensed by a photo diode. The sensor is calibrated by measuring the response in the soil at known moisture content and/or matric potentials. A second type of sensor involves the placement of porous glass or nylon disks of different pore-size grades between a CdS photoresistor cell and light emitting diode (Figure 6.2.6b). The use of different pore-size disks allows continuous measurement of electrical response over a wide range of moisture contents. Calibration procedures are similar to those for the electro-optical switch sensor.

<u>Method Selection Considerations</u>: Advantages: (1) Potential for low cost and high physical stability and reliability; (2) can be calibrated to measure both moisture content and water potential over a wide range of moisture contents and matric potentials (electro-optical switches are better than CdS photoresistors for direct measurement of matric potential); and (3) electronic circuitry allows automatic data acquisition and analysis. Disadvantages: (1) New technique with limited operational and field experience; (2) equipment is not yet readily available (although both types of devices can be readily made using off-the-shelf materials); and (3) separate calibrations might be required for changes in soil texture.

Frequency of Use: New technique with potential for widespread use.

Standard Methods/Guidelines: --

Sources for Additional Information: Alessia and Prunty (1986), Cary et al. (1989, 1991)



Figure 6.2.6 Diagrams of two electro-optical soil-water sensors: (a) Electro-optical switch with nylon disk using infrared-light transmission; (b) CdS cell (photoresistor) with layered fritted glass using visible-light transmission (Cary et al., 1991, by permission).

6.2 VADOSE ZONE MOISTURE CONTENT

6.2.7 Computerized Axial Tomography (CAT)

Other Names Used to Describe Method: Computer assisted tomography, CAT scanning, x-ray computed (computer) tomography, computed tomographic scanning, CT scanning, x-ray CT, gamma-ray attenuation CAT, nuclear tomography.

<u>Uses at Contaminated Sites</u>: Potential for measuring spatial distribution of soil moisture, bulk density, and soil macroporosity; detecting roots, seeds, insects.

<u>Method Description</u>: CAT scanning systems can use single or multiple sources of gamma radiation or x-rays. Detectors can be on the same probe as the source, or placed in adjacent boreholes. The detectors measure the attenuated signal, and counts in the desired energy ranges are discriminated by a single channel analyzer. Signals are processed using tomographic theory (see also, Section 3.4.5) to allow three dimensional analysis of variations of the parameter of interest. Figure 6.2.7 illustrates the operation of a CT scanner used for scanning soil cores in the laboratory.

<u>Method Selection Considerations</u>: Relatively new method, which has shown promising results in laboratory studies, but has not yet been tested for field applications.

Frequency of Use: Usefulness in the field not yet demonstrated.

Standard Methods/Guidelines: ASTM (1991b).

Sources for Additional Information: Anderson et al. (1988), Phogat et al. (1991).



Figure 6.2.7 Schematic of how a CT scanner measures the attenuated x-ray beams passing through a detection aperture containing a soil core. The x-ray source and detector array rotate clockwise around the detection zone (Anderson et al., 1988, by permission).

6.3 OTHER SOIL HYDROLOGIC PROPERTIES

6.3.1 Soil Moisture-Potential-Conductivity Relationships

Other Names Used to Describe Method: Moisture characteristics curves: Water retention function, specific retention, water content-matric potential relationship, capillary pressure-saturation curve, capillary-moisture relationship.

<u>Uses at Contaminated Sites</u>: Estimating water content from soil water potential measurements; estimating soil water potential from soil moisture measurements; estimating hydraulic conductivity from soil water potential or soil moisture measurements; modeling of contaminant flow in the vadose zone.

Method Description: Soil moisture (usually represented by the symbol "b"), soil matric potential (usually represented with the symbol " ϕ "), and soil hydraulic conductivity are all intimately related. Once the relationship between two of these properties have been established for a soil horizon, then measurement of one parameter allows calculation of the other parameters. The soil moisture characteristic curve (see above for other terms used to describe this relationship) is a commonly used relationship to define soil hydrologic properties. An important property of this relationship is that it is subject to hysteresis (i.e., the relationship is different depending on whether the soil is wetting or drying). Figure 6.3.1a shows the moisture characteristic curve for a sandy soil and illustrates the effect of hysteresis. In the field, the moisture characteristic curve is determined by monitoring soil water content (using methods described in Section 6.2) and soil water potential (using methods described in Section 6.1) during the wetting or drying cycle of a soil. Jury et al. (1978) provide an example of developing a moisture characteristic curve in the field using tensiometers and neutron-probe measurements. Shani et al. (1987) describe a reliable and quick method for estimating this relationship in the field using a dripper method (Section 7.2.5). In a similar manner, $K(\theta)$ (hydraulic conductivity as a function of moisture content) and $K(\phi)$ (hydraulic conductivity as a function of soil-water potential [pressure head]) can be measured. These relationships also are subject to hysteresis as shown in Figure 6.3.1b. Estimation from other soil properties: The hydrologic properties of soils are strongly related to physical properties, such as particle-size distribution, porosity, and bulk density. Empirical relationships between physical and hydrologic properties can be used to estimate soil moisture-potential relationships based on measurement of physical properties, provided that the soils are similar to the soils from which the empirical relationships have been derived. Section 7.2.8 provides additional information on estimation of unsaturated hydraulic conductivity using physically and empirically-based equations and relationships, many of which can be related to moisture characteristic curves. Mualem and Friedman (1991) have developed an equation that relates soil electrical conductivity (from saturation extract--see Section 9.3.5) to water content, which can be used to estimate soil water content of samples for a wide range of coarse and stable structured soils when no other data are available. Reference index Table 9-3 (EC-Salinity Calibrations), identifies other references that discuss the relationship between moisture content and electrical conductivity. The moisture characteristic also can be estimated from sorptivity measurements (Section 6.3.2).

Method Selection Considerations: The soil moisture-potential relationship is required input for many vadose hydrologic models. Field measurement of moisture characteristic curves using conventional methods is complicated and time-consuming, although the recently developed dripper method (Section 7.2.5) now provides a simpler and more rapid alternative for field measurement. Laboratory measurements using undisturbed core samples are simpler, but a large number of cores might be required to adequately characterize spatial variability. Empirical relationships based on other soil physical properties are the simplest and least expensive method, but probably are the least accurate method unless soil physical properties are very similar to the soils from which the empirical relationships were derived.

<u>Frequency of Use</u>: Field measurement is uncommon. Usually measured in the laboratory or estimated using empirical relationships.

Standard Methods/Guidelines: Field: Bruce and Luxmore (1986), Shani et al. (1987); Laboratory: ASTM (1968), Klute (1986); Empirical equations/relationships: See Table 6-3.

Sources for Additional Information: See Table 6-3.



(b)

Figure 6.3.1 Relationships between soil moisture, matric potential, and hydraulic conductivity: (a) Moisture characteristic curves for a sandy soil during wetting and drying; (b) K-matric potential curves showing effect of hysteresis during wetting and drying (Everett et al. 1983, after Freeze and Cherry, 1979).

6.3 OTHER SOIL HYDROLOGIC PROPERTIES

6.3.2 Water Sorptivity and Diffusivity

Other Names Used to Describe Method: --

<u>Uses at Contaminated Sites</u>: **Sorptivity**: Estimating diffusivity-moisture relationship, moisture characteristic curve, and hydraulic conductivity as a function of matric potential; estimating soil water distribution. Diffusivity: Characterizing soil transmission/storage properties; calculating infiltration (some equations).

Method Description: Soil diffusivity is a single parameter of unsaturated soil that relates the hydraulic conductivity and water storage properties of a soil and can be calculated as either a function of changes in soil water potential or water content. Hydraulic diffusivity can be an important parameter in infiltration equations (Section 7.1.4). Sorptivity is a measure of the capacity of a porous medium to absorb a wetting liquid. The greater the value, the larger the volume of water that can be absorbed, and the more rapidly it will be absorbed. Sorptivity decreases from a maximum value (dependent on the soil physical properties) to zero as water content/matric potential increase to the point of saturation. Sorptivity is closely related to hydraulic conductivity and soil water diffusivity, and is sometimes used to calculate diffusivity. Field Measurement of Diffusivity: Any field method for measuring unsaturated hydraulic conductivity as a function of matric potential, which measures changes in water content with time and changes in matric potential with time, can be used to determine diffusivity (see Table 7-1). Field measurement of sorptivity: Green et al. (1986) describe two methods for measurement of sorptivity: (1) Ponded infiltration: A single- or double-ring infiltrometer (see Section 7.3.1) is filled with water and cumulative infiltration is measured as the head of ponded water falls with time; (2) Constant-head porous Plate: Similar to ponded infiltration method, except that a constant-head device delivers water to the soil through a porous plate in contact with the soil. This process results in a slight negative pressure at the bottom of the porous plate, preventing water from entering large pores or cracks. The second method is a variant of the tension infiltrometer (see Section 7.2.3). Table 7-1 identifies other methods for measuring saturated and unsaturated hydraulic conductivity, which also can be used to measure sorptivity.

<u>Method Selection Considerations</u>: Sorptivity: The ponded infiltration method is simple and rapid, but works only when there is negligible flow of water through large cracks or channels. The constant-head porous plate method also is simple, rapid and reliable, and should be used any time flow through large pores is a concern. See also, appropriate subsections in Section 7, as identified in Table 7-1.

Frequency of Use: Relatively uncommon in routine field applications.

Standard Methods/Guidelines: Green et al. (1986).

Sources for Additional Information: See Table 6-3 and additional references identified under tension infiltrometers in reference index Table 7-3.

6.3 OTHER SOIL HYDROLOGIC PROPERTIES

6.3.3 Available Water Capacity

Other Names Used to Describe Method: Field capacity, water holding capacity.

<u>Uses at Contaminated Sites</u>: Evaluating water storage in the rooting zone and the movement of contaminants from the rooting to the vadose zone in response to precipitation events.

Method Description: Available water capacity is the difference between field capacity (the amount of water remaining in a soil 2 or 3 days after having been wetted and after free drainage is negligible) and water held in the soil at the permanent wilting point, PWP (the point at which plants generally are unable to extract additional water from the soil--around 15 bars suction = -1,500 kPa). This represents the amount of water that is available to plants for growth. The field procedure involves wetting soil test plots and measuring water content using one of the methods identified in Table 6-1 (gravimetric, neutron, or gamma-gamma are the most commonly used) when the soil is at field capacity at different depth increments. Alternatively, natural changes in soil moisture can be monitored over an extended period of time. Figure 6.3.3a shows that this approach can result in a range rather than an exact percentage for field capacity. Determination of the PWP requires laboratory tests using the Sunflower method, in which water is withheld from a dwarf sunflower growing in a sample of the depth horizon of interest until it wilts, at which time the soil water content is determined. Alternatively, PWP can be approximated using a pressure plate apparatus to withdraw water from a sample of the depth increment of interest until matric potential is -15 bars, at which time water content is measured. Figure 6.3.3b show several ways in which the resulting data can be plotted, and illustrates the difference that rooting depth of plants present in a soil can make in the amount of water that is likely to be removed from the soil by evapotranspiration. Available water capacity also can be estimated from a moisture characteristic curve (Figure 6.3.1), and from soil texture (Figure 6.3.3c). The particle-size ranges of the texture classes shown in Figure 6.3.3c are shown in Figure 2.5.1.

<u>Method Selection Considerations</u>: There is no good alternative to the procedure described above for accurate measurement of in situ field capacity, although approximations using laboratory analysis of undisturbed soil cores or centrifugation of disturbed soil samples can be obtained.

Frequency of Use: Uncommon.

Standard Methods/Guidelines: Cassel and Nielsen (1986).

Sources for Additional Information: Richards (1965).









(c)



Available water capacity: (a) Estimation of field capacity (28 to 30 percent) by repeated measurement of soil moisture in situ (Dunne and Leopold, 1978, From: *Water in Environmental Planning* by Dunne and Leopold, Copyright © 1978 by W.H. Freeman and Company, reprinted with permission), (b) Upper and lower limits of available water: (A) Measured in 0.15-meter increments, (B) limits expressed for this profile for a 0.6-meter rooting depth, (C) and for a 1.5-meter rooting depth; FC = field capacity, PWP = permanent wilting point (Cassel and Nielsen, 1986, by permission); (c) Chart for estimating field capacity and available water capacity based on soil texture (Dunne and Leopold, 1978, From: *Water in Environmental Planning* by Dunne and Leopold, Copyright © 1978 by W.H. Freeman and Company, reprinted with permission).

Торіс	References			
Soil Water (Matric) Potential				
General	Hendrickx et al. (1990-variability), Wilkinson and Klute (1962-temperature effects); <u>Reviews</u> : Bouyoucos (1960), Brakensiek et al. (1979), Mullins (1991), Richards (1949)			
Porous Cup Tensiometers	Colman et al. (1946), Cummings and Chandler (1940), Dennehy and McMahon (1989), Hendrickx and Nieber (1985), Huber and Dirksen (1978), Hunter and Kelley (1946), McKim et al. (1980b), Miller (1951), Oaksford (1978-manometer), Perrier and Evans (1961), Rehm et al. (1987), Richards (1942), Richards and Gardner (1936), Richards and Neal (1937), Richards et al. (1938, 1973), Rogers (1935), Savvides et al. (1977- mercury), Thomas and Phillips (1991), Towner (1980), Wendt et al. (1978); <u>Reviews:</u> Cassel and Klute (1986), Hendrickx (1990), Richards (1949), Schmugge et al. (1980), Stannard (1986, 1990); <u>Transducer type</u> : Anderson and Burt (1977), Bianchi (1962), Burt (1978), Elzeftawy and Mansell (1975), Enfield and Gillaspy (1980), Fitzsimmons and Young (1972), Gillham et al. (1976), Klute and Peters (1962), Leonard and Low (1962), Long (1982), Long and Huck (1980), Marthaler et al. (1983), Rice (1969), Thiel et al. (1963), Thony and Vachaud (1980), Watson (1965, 1967), Watson and Jackson (1967-temperature effects), Williams (1978); <u>Recording Tensiometers/Automatic Data Acquisition</u> : Anderson and Burt (1977), Bianchi and Tovey (1968), Burt (1978), Enfield and Gillaspy (1980), Long and Huck (1980), Lowery et al. (1986), Nyhan and Drennon (1990), Rice (1969), Walkotten (1972), Williams (1978); <u>Moisture Measurement</u> : Cummings and Chandler (1940), McKim et al. (1980b), Troolen et al. (1986); <u>Snowpack</u> : Colbeck (1976), Wankiewicz (1978); <u>Wick <u>Tensiometer</u>: Gee and Campbell (1991)</u>			
Thermocouple Psychrometers	<u>Review</u> : Savage and Cass (1984); <u>Papers</u> : Barrs and Slaytor (1965), Box (1965), Brown (1970), Brown and Collins (1980), Brown and Johnson (1976), Campbell (1972, 1979), Campbell et al. (1968), Chow and deVries (1973), Dalton and Rawlins (1968), Daniel (1979), Daniel et al. (1981), Enfield and Hsieh (1972), Enfield et al. (1973), Hoffman et al. (1969, 1972), Hsieh and Hungate (1970), Hsieh et al. (1972), Ingvalson et al. (1970), Jones et al. (1990), Koorevar and Janse (1972), Korven and Taylor (1959), Lambert and van Schilfgaarde (1965), Lang and Trickett (1965), Lopushinsky (1971), Lopushinsky and Klock (1971), Madsen et al. (1986), Meeuwig (1972), Merril and Rawlins (1972), Merril et al. (1968), Meyn and White (1972), Monteith and Owen (1958), Moore and Caldwell (1972), Peck (1968), Rawlins (1966), Rawlins and Dalton (1967), Richards (1949), Richards and Caldwell (1987), Richards and Ogata (1958), Roundy (1984), Spanner (1951), Van Heveren (1972-humidity in snow), Van Heveren and Brown (1972), Wiebe et al. (1971, 1977), Zanstra (1976), Zollinger et al. (1966); <u>Texts</u> : Brown and Van Heveren (1972), Campbell (1977), Fritschen and Gay (1979); <u>Measurement Interpretation</u> : Campbell and Gardner (1971), Lang (1967); <u>Calibration</u> : Brown and Collins (1980); <u>Water Activity Meter</u> : Gee et al. (1992)			
Resistance Sensors	Anderson and Edlefsen (1942), Atchison and Butler (1951), Becker et al. (1946), Bourget et al. (1958), Bouyoucos (1949, 1953, 1954), Bouyoucos and Mick (1940, 1947, 1948), Colman and Hendrix (1949), Croney et al. (1951), Cummings and Chandler (1940), Daniel et al. (1992), Dennehy and McMahon (1989), El-Samie and Marsh (1955), Haise and Kelley (1946), Hancox and Walker (1966), Kemper and Amemiya (1958), Michelson and Lord (1962), Pereira (1951), Perrier and Marsh (1958), Rehm et al. (1987), Richards and Weaver (1943), Salaruddin and Khasbardar (1967), Schlub and Maine (1979), Slater (1942), Tanner and Hanks (1952), Williams (1980); <u>Automatic Data Acquisition</u> : Armstong et al. (1985); <u>Calibration</u> : Atchison and Butler (1951), Kelley (1944), Shaw and Baver (1939a); <u>Moisture Measurement</u> : Cummings and Chandler (1940)			

Table 6-2 (cont.)

Topic	References				
Soil Water (Matric) Potential (cont.)					
Electrothermal Methods	Aldous and Lawton (1952), Beck et al. (1971), Blackwell (1954, 1956), Bloodworth and Page (1957), Bloomer and Ward (1979), Cummings and Chandler (1940), Daniel et al. (1992), DeVries (1952, 1953), DeVries and Peck (1958a,b), Fritton et al. (1974), Fuchs and Hada (1973), Fuchs and Tanner (1968), Gardner et al. (1991), Hooper (1952), Hooper and Leeper (1950), Jaeger (1958), Kubo (1953), Momin (1945), Phene et al. (1971a, 1971b, 1973), Philip (1961), Shaw and Baver (1939a,b), Slusarchuk and Fougler (1973), Sophocleous (1979), Van Duin and DeVries (1954), Wechsler et al. (1965); <u>Calibration</u> : Kelley (1944), Overgaard (1970), Shaw and Baver (1939a); <u>Moisture Measurement</u> : Cummings and Chandler (1940)				
Soil Moisture Content					
General	Hendrickx et al. (1990-variability), Reinhart (1961-physical factors), Yates and Warrick (1987-estimation with cokriging); <u>Reviews</u> : Bouyoucos (1952), Johnson (1962), McKim et al. (1980a), Postlethwaite and Trickett (1956), Schmugge et al. (1980), Taylor (1955), Wilson (1971)				
Gravimetric	Hawley et al. (1982), Hendrickx (1990), McKim et al. (1980b), Rehm et al. (1987), Reynolds (1970a,b,c)				
Time Domain Reflectometry	Ansoult et al. (1985), Baker and Allmaras (1990), Baker and Lascano (1989), Brisco et al. (1992), Chudobiak et al. (1979), Cole (1977), Dalton (1989), Dalton and van Genucthen (1986), Dalton et al. (1984), Dasberg and Dalton (1985), Dasberg and Hopmans (1992-calibration), Davis and Annan (1977), Davis and Chudobiak (1975), Elrick et al. (1992), Fellner-Feldegg (1969, 1972), Heimovaara et al. (1988), Hokett et al. (1992), Hook et al. (1992), Kachonoski et al. (1990, 1992), Nadler (1991), Nadler et al. (1991), Patterson and Smith (1981), Redman et al. (1991), Reeves and Elgezawi (1992), Smith and Tice (1988), Stein and Kane (1983), Tektronix (1987), Topp and Davis (1981, 1985a,b), Topp et al. (1980a,b, 1982a,b, 1984, 1988), Van Loon et al. (1990-electrical conductivity), Yanuka et al. (1988), Zegelin et al. (1989); <u>NAPL Detection</u> : Brewster et al. (1992); <u>Leak Detection</u> : Davis et al. (1984)				
Dielectric Sensors	Bell et al. (1987), Birchak et al. (1974), Brisco et al. (1992), Dean et al. (1987), DePlater (1955), Hancox and Walker (1966), Kuráž (1981), Kuráž and Matousek (1977), Kuráž et al. (1970), Layman (1979), Mack and Brach (1966), Matthews (1963), Matzkanin et al. (1979), McKim et al. (1979, 1980b), Roth (1966), Selig and Mansukhani (1975), Selig et al. (1975), Thomas (1966), Troxler Electronic Laboratories (1992), Walsh et al. (1979), Wobschall (1978); <u>Soil Dielectric Properties</u> : Cihlar and Ulaby (1974), Hipp (1974), Hoekstra and Delaney (1974), Smith-Rose (1933), Wang and Schmugge (1978), see also, references in Section 1.5.1				
Nuclear Magnetic Resonance	Soil Moisture: Andreyev and Martens (1960), Matzkanin and Gardner (1974), Prebble and Currie (1970), Rollwitz (1965), Smith and Tice (1988), Tice et al. (1981), Wu (1964); <u>Borehole</u> : See Section 3.2.4				

 Table 6-3 Reference Index for Measurement and Estimation Soil Hydrologic Properties Other than Hydraulic

 Conductivity

Topic	References				
General Soil-Water Relationships	Bouwer and Jackson (1974), Childs (1969), Day et al. (1967), Gairon and Hadas (1973), Hendrickx (1990), Holmes et al. (1967), Marshall (1960), Nielsen et al. (1972), Reeve and Carter (1991), Richards (1965), Rode (1965), Wiebe et al. (1971)				
Soil-Water Retention	Measurement: Madsen et al. (1986), Richards (1965), Su and Brooks (1980); Equations: Bumb et al. (1991), Gillham et al. (1979), McKee and Bumb (1984); see also, Section 7.2.8 and Table 7-5; Estimation from Other Soil Properties: Ahuja et al. (1985), Alessi et el. (1992), Arya and Paris (1981), Bruce (1972), Brust et al. (1968), Carsel and Parrish (1988), Clausnitzer et al. (1992), DeJong (1982), Gregson et al. (1987), Gupta and Larson (1979), Haverkamp and Parlange (1986), Hendrickx (1990), Hendrickx et al. (1991), McQueen and Miller (1974), Mishra and Parker (1990), Mishra et al. (1989), Puckett et al. (1985), Rawls and Brakensiek (1985), Rawls et al. (1982), Rogowski (1971, 1972), Ross et al. (1991), Saxton et al. (1986), Schuh et al. (1988), Topp and Zebchuck (1979), Tyler and Wheatcraft (1989), Vereecken et al. (1992), Williams et al. (1992), Yoshida et al. (1985); <u>Temperature Effects</u> : Haridasan and Jensen (1972), Nimmo and Miller (1986); <u>Hysteresis</u> : Nimmo (1992)				
Sorptivity	Bridge and Ross (1985), Brutsaert (1976), Chong (1983), Chong et al. (1982), Clothier and White (1982), Dirksen (1975), Kutilek and Valentova (1986), Parlange (1971, 1975a,b), Philip (1955), Reichardt and Libardi (1974), Reynolds and Elrick (1990), Smiles (1977), Smiles et al. (1981, 1982), Talsma (1969), Topp and Zebchuck (1979), White (1979), White and Perroux (1987, 1989), White et al. (1989)				
Diffusivity	Bruce and Klute (1956), Brutsaert (1976, 1979), Cassel et al. (1968), Dirksen (1975), Gardner (1970), Gardner and Mayhugh (1958), Hillel and Gardner (1970), Jackson (1963), Klute (1965, 1972), Miller and Bresler (1977), Parlange (1975a,b), Perroux et al. (1981), Philip (1955), Reichardt et al. (1972), Roberts (1984), Scotter and Clothier (1983), Smiles (1977), Smiles and Harvey (1973), Weeks and Richards (1967)				

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SECTION 7

VADOSE ZONE HYDROLOGIC PROPERTIES (II): INFILTRATION, CONDUCTIVITY, AND FLUX

Characterization of water movement in the vadose zone is complicated by the fact that hydraulic conductivity varies as a function of pressure potential and moisture content. The introduction to Section 6 discusses the types of energy potentials that affect flow of water in the vadose zone. Various terms are used to describe hydraulic conductivity in the vadose zone:

- 1. Saturated hydraulic conductivity (K_{sat}) is the hydraulic conductivity at saturation with no entrapped air. This state rarely is achieved in the vadose zone, except, perhaps, in the zone of seasonal fluctuation of an unconfined water table.
- 2. Field-saturated hydraulic conductivity (K_{ta}), also called the satiated hydraulic conductivity, is the hydraulic conductivity when entrapped air is present, which can be as much as 50 percent below the true K_{sat} (Reynolds and Elrick, 1986). Methods for measuring saturated hydraulic conductivity above the water table usually measure K_{tb}. Another term, K_(sat), has been proposed by Bouma (1982) for hydraulic conductivity measurements of the soil matrix without macropore flow (see Column-Crust method, Section 7.3.8 and Figure 7.3.8b[c]). K_(sat) will be less than K_{sat} or K_{tb} because water flows more rapidly in macropores than in the soil matrix. The term K_{sat} often is loosely used for reporting measurements that should more accurately be termed K_{tb}.
- 3. Unsaturated hydraulic conductivity (K_{unsat}) is the hydraulic conductivity of soil at negative pressure potentials. $K(\phi)$ is the term usually used to describe the hydraulic conductivity-pressure potential function, and $K(\theta)$ to describe the hydraulic conductivity-moisture content function. Complete characterization of K_{unsat} requires measuring hydraulic conductivity at a range of moisture contents to develop a $K(\theta)$ curve or at a range of pressures to develop a $K(\phi)$ curve (see Section 6.4.1). These functions are subject to hysteresis (i.e., K_{unsat} might differ at the same water content or matric potential, depending on whether the soil is wetting or drying [Section 6.4.1]).

Infiltration

The infiltration capacity of a soil is a critical element of water budget calculations because it affects how much precipitation that reaches the ground surface enters the soil and how much moves off a site as surface runoff. The infiltration rate generally is the same as the unsaturated and saturated hydraulic conductivity, except that some processes, such as the initial moisture content (see Figure 7.1.4), crusting, or sediment clogging, might cause different infiltrations at the ground surface compared to the subsurface with all other soil factors being equal. Table 7-1 summarizes information on eight methods for measuring or estimating infiltration grouped into four categories: (1) Impoundment methods, where infiltration is below a water surface (Section 7.1.1); (2) land surface methods (Section 7.1.2), (3) watershed methods for estimating infiltration over larger areas (Section 7.1.3), and (4) infiltration equations (Section 7.1.4). In most situations infiltration can be estimated using empirical relations or infiltration suing other measured variables, which can be measured with an air-entry permeameter (Section 7.3.4) and unsaturated hydraulic conductivity/pressure head relations (Section 6.3.1).

Measurement of Unsaturated Hydraulic Conductivity

Table 7-1 summarizes information on nine methods for measuring or estimating unsaturated hydraulic conductivity from field measurements. Most of these methods can be used to develop $K(\phi)$ or $K(\theta)$ relationships, which once established, allow subsequent monitoring to focus on either changes in pressure potential or moisture content. The instantaneous profile method (Section 7.2.1) is the most commonly used method for accurate measurement of unsaturated hydraulic conductivity in the field. Various draining profile methods (Section 7.2.2) are simpler and less expensive to use if the simplifying assumptions apply to the site of interest. Another common procedure is to collect undisturbed core samples and measure K_{unsat} in the laboratory (Klute and

Technique	K ₆ or K	K Direction ^b	Other Parameters Measured	Section	Tables
Infiltration (see also, Sections 7.2.3,	, 7.2.5, 7.2.6, 7.3.1, 7.3	3.4)			
Seepage Meters	Saturated	Undefined	I	7.1.1	
Instantaneous Rate	Saturated	Undefined	I	7.1.1	
Impoundment Water Budget	Saturated	Undefined	I	7.1.1	
Sprinkler Infiltrometer	Saturated	Vertical	I	7.1.2	
Infiltration Test Basins	Saturated	Undefined	I	7.1.2	
Watershed Average	Undefined	Undefined	I	7.1.3	
Watershed Empirical Relations	Undefined	Undefined	I	7.1.3	
Infiltration Equations	Both	Vertical	I	7.1.4	7-5
Unsaturated Hydraulic Conductivity	,				· •
Instantaneous Profile	Unsaturated	Vertical	D, F, K(ø), R	7.2.1	7-3
Draining Profile Methods	Unsaturated	Vertical	D, F, K(ϕ), R, S	7.2.2	7-3
Tension Infiltrometers	Both	Vertical	I, D, F, K(¢), R, S	7.2.3	7-3
Crust-Imposed Steady Flux	Unsaturated	Vertical	I, F, K(φ)	7.2.4	7-3
Sprinkler/Dripper Methods	Unsaturated	Vertical	I, F, K(φ), R, S	7.2.5	7-3
Entrapped Air Method	Unsaturated	Vertical	I, F	7.2.6	7-3
Parameter Identification	Both	Undefined	R	7.2.7	7-3
Empirical Equations	Both	Undefined	Varies	7.2.8	7-5
Column-Crust	Both	Vertical	F, K(φ)	7.3.8	7-4
Saturated Hydraulic Conductivity A	bove Shallow Water	Table			
Cylinder Infiltrometers	Saturated	Vertical	I. S	7.3.1	7-4
Constant Head Borehole			-, -		
Infiltration	Saturated	Horizontal	S	7.3.2	7-4
Guelph Permeameter	Both	Vert./Hor.	K(φ), S	7.3.3	7-4
Air-Entry Permeameter	Both	Vertical	I, K(ø), S	7.3.4	7-2, 7-4
Double Tube	Saturated	Vertical	-	7.3.5	7-2, 7-4
Cylinder Permeameter	Saturated	Vertical		7.3.6	7-2, 7-4
Infiltration Gradient	Saturated	Vertical ^d		7.3.7	7-4
Cube	Saturated	Vert./Hor.		7.3.8	7-2, 7-4
Column/Monoliths	Saturated	Vertical	-	7.3.8	7-2, 7-4
Boutwell Method	Saturated	Vert./Hor.		7.3.9	
Velocity Permeameter	Saturated	Vertical		7.3.10	
Percolation Test	*	•		7.3.11	7-4
CP Porous Probe	Saturated	Horizontal		2.2.2	
Collection Lysimeter	Saturated	Vertical	F	9.3.1	
Saturated Hydraulic Conductivity Al	bove Deep Water Ta	blc°			
USBR Single Well	Saturated	Undefined		7.4.1	
USBR Multiple-Well	Saturated	Horizontal		7.4.2	
Stephens-Neuman Single Well	Saturated	Undefined		7.4.3	
Air Permeability	Saturated	Undefined		7.4.4	
Packer Tests	Saturated	Vert/Hor.		4.3.3	

Table 7-1 Summary Information on Vadose Hydraulic Conductivity Techniques*

ŧ

D = diffusivity; F = Flux; I = Infiltration; $K(\phi)$ = hydraulic conductivity-pressure head relationship; R = Retention (pressure-moisture relationship); S = Sorptivity.

*Most methods for measuring or estimating unsaturated hydraulic conductivity also can be used to measure water flux in the vadose zone. Section 7.5 discusses the application of these and other methods for measuring soil water flux.

^bDirectional ratings are qualitative in nature. Different references might give different ratings depending on site conditions and criteria used to define directionality.

These methods measure <u>field-saturated</u> or <u>satiated</u> hydraulic conductivity (K_{L}) , which is lower than <u>saturated</u> hydraulic conductivity, due to the presence of entrapped air.

Differentiation of vertical and horizontal is possible when used with double tube method.

The percolation test does not provide an accurate measure of saturated hydraulic conductivity. See Table 7-4 for sources on information on the relationship between percolation test results and K_{tas} .
Dirksen, 1986). ASTM (1990a) provides guidance on selecting field methods for measuring unsaturated hydraulic conductivity in the vadose zone.

Measurement of Saturated Hydraulic Conductivity

Table 7-1 summarizes information on 10 methods for measuring K_{b} above a shallow water table (Section 7.2), and 5 methods for measuring K_{b} above a deep water table (Section 7.3). The cylinder or ring infiltrometer (Section 7.3.1) is a widely used method that measures both infiltration and K_{b} at the soil surface. Most other shallow methods require a borehole and devices at the surface to control the flow of water into the hole to achieve steady state infiltration before measurements are taken. The constant-head borehole infiltration or shallow-well pump-in method (Section 7.3.2) and the Guelph permeameter (Section 7.3.3) probably are the most commonly used methods for measuring K_{b} . Most of these methods are restricted to depth of 2 meters or less, but recently developed compact constant-head permeameter (Section 7.3.2) can be used to depths of 10 meters. Most methods for measuring K_{b} above a deep water table require drilling or relatively large diameter boreholes (at least 6 inches) and a large supply of water, which can be pumped into the borehole. ASTM (1990a) provides guidance on selecting field methods, for measuring saturated hydraulic conductivity above and below a water table.

Measurement of Water Flux in the Vadose Zone

Various methods are available to measure or estimate the amount of water that passes through the vadose zone and enters the ground-water system. A water budget (Section 7.5.1) uses a mass balance by measuring inflows, outflows, and storage changes in the area of interest. More often, a simplified water budget approach can be used, in which only changes in soil moisture or matric potential are measured (Section 7.5.2). A variety of tracers, such as chloride and tritium, can be used to estimate the rate of recharge and water flux (Section 7.5.3). Localized water flux can be measured using a soil-water flux meter (Section 7.5.4). A variety of methods for measuring the velocity of water flow in the vadose zone are described in Section 7.5.5. Finally, a variety of physical and empirical equations can be used in combination with the methods above, or using site-specific data on hydraulic conductivity or soil physical characteristics, such as texture and bulk density. Tile drains or collection lysimeters (Section 9.3.1) also can be used to measure water flux in the vadose zone, provided the area of vertical infiltration is known and lateral ground-water flow can be excluded or quantified.

K _{sat} Method	hoitonsa and and a soundware have ion carbon and a sound a sou
Column method	• (•) •
Cube method	(.) .
Drain-cube method	• •
Air entry perm.	
Cylinder perm.	
Double-tube	
Augerhole method	
Piezometer method	
Four-holes method	

Source: Amoozegar and Warrick (1986), after Bouma (1983)

Dirksen, 1986). ASTM (1990a) provides guidance on selecting field methods for measuring unsaturated hydraulic conductivity in the vadose zone.

Measurement of Saturated Hydraulic Conductivity

Table 7-1 summarizes information on 10 methods for measuring K_5 above a shallow water table (Section 7.2), and 5 methods for measuring K_5 above a deep water table (Section 7.3). The cylinder or ring infiltrometer (Section 7.3.1) is a widely used method that measures both infiltration and K_5 at the soil surface. Most other shallow methods require a borehole and devices at the surface to control the flow of water into the hole to achieve steady state infiltration before measurements are taken. The constant-head borehole infiltration or shallow-well pump-in method (Section 7.3.2) and the Guelph permeameter (Section 7.3.3) probably are the most commonly used methods for measuring K_5 . Most of these methods are restricted to depth of 2 meters or less, but recently developed compact constant-head permeameter (Section 7.3.2) can be used to depths of 10 meters. Most methods for measuring K_5 above a deep water table require drilling or relatively large diameter boreholes (at least 6 inches) and a large supply of water, which can be pumped into the borehole. ASTM (1990a) provides guidance on selecting field methods, for measuring saturated hydraulic conductivity in the vadose zone, and Table 7-2 provides comparative information on nine methods for measuring saturated hydraulic conductivity above and below a water table.

Measurement of Water Flux in the Vadose Zone

Various methods are available to measure or estimate the amount of water that passes through the vadose zone and enters the ground-water system. A water budget (Section 7.5.1) uses a mass balance by measuring inflows, outflows, and storage changes in the area of interest. More often, a simplified water budget approach can be used, in which only changes in soil moisture or matric potential are measured (Section 7.5.2). A variety of tracers, such as chloride and tritium, can be used to estimate the rate of recharge and water flux (Section 7.5.3). Localized water flux can be measured using a soil-water flux meter (Section 7.5.4). A variety of methods for measuring the velocity of water flow in the vadose zone are described in Section 7.5.5. Finally, a variety of physical and empirical equations can be used in combination with the methods above, or using site-specific data on hydraulic conductivity or soil physical characteristics, such as texture and bulk density. Tile drains or collection lysimeters (Section 9.3.1) also can be used to measure water flux in the vadose zone, provided the area of vertical infiltration is known and lateral ground-water flow can be excluded or quantified.

7.1 INFILTRATION

7.1.1 Impoundment Methods

Other Names Used to Describe Method: Seepage meters (SCS, USBR, Bouwer-Rice), instantaneous rate method, water budget method.

Uses at Contaminated Sites: Measuring infiltration of surface water impoundments into the ground.

<u>Method Description</u>: Seepage meters are sealed infiltrometers placed in the bottom of a channel or pond that is connected by a tube to a small reservoir of water in a container, which can be raised or lowered in relation to the water surface of the impoundment (Figure 7.1.1). When the small reservoir is raised above the level of the natural water surface, the rate of fall is measured, or relative changes in pressure head inside the seepage meter and the water outside are measured. Infiltration rate can be calculated from these measurements. Types include the SCS, U.S. Bureau of Reclamation, and Bouwer-Rice seepage meters. The instantaneous rate method involves shutting down all inflows and discharges from a pond and observing the drop in water level. Assuming evaporation is negligible and there is no ground-water recharge to the pond, the rate of decline in water level is the infiltration (Wilson, 1982). The water budget method requires measurement of inflow to the pond from all sources, precipitation, discharge, and evaporation, and the relationship between head vs. surface area. When these parameters are known, infiltration can be solved from the water budget equation (Bouwer, 1978). Section 7.5.1 discusses the water budget method further, and Figure 7.5.1 illustrates an annual cumulative water balance used to determine seepage from a wastewater lagoon. Other methods for characterizing interactions between surface impoundments and ground-water systems include shoreline monitoring wells, mini-piezometers, well points, and core samples (Wolf et al., 1991).

Method Selection Considerations: Table 7.1.1 provides some general guidelines for selecting techniques for evaluating surface-ground water interactions. Seepage Meter Advantages: Relatively inexpensive and simple to operate. Seepage Meter Disadvantages: (1) A large number of measurements are required to obtain average infiltration rates, especially in unlined ponds with variable texture; (2) some underwater work is required to install the unit, which might be hazardous to personnel in waste ponds with toxic chemical; (3) measurements must be obtained on sides and bottoms of ponds and installation is difficult in ponds with steep-sided slopes; and (4) cannot be used in frozen ponds. Instantaneous Rate Advantages: Simple and inexpensive way to measure average infiltration rate. Instantaneous Rate Disadvantages: Results might be inaccurate if there is ground-water recharge or rates of decline are slow enough for evaporation to become significant. Water Budget Advantages: Can be used in most hydrogeologic settings. Water Budget Disadvantages: (1) Time consuming and expensive; (2) will not work where the water table is able to rise above the level of water in the pond; (3) requires accurate estimation of evaporation, which is not easy, especially if impoundment contains chemicals that change evaporation properties; (4) installing inflow and outflow measuring devices might be difficult at some sites; (5) errors in measurements of any auxiliary parameter affect the accuracy of estimated infiltration; and (6) calculation of changes in storage is difficult where water levels change slowly (can be overcome with special techniques for very accurate measurement such as laser equipment).

Frequency of Use: Seepage meters are most commonly used. Water budget is rarely used due to complexity and cost.

Standard Methods/Guidelines: Seepage meter: Bouwer and Rice (1963); Water budget: Bouwer (1978).

Sources for Additional Information: Seepage meter: Bouwer (1978, 1986), Kraatz (1977), Wilson (1982); Other: Everett et al. (1983), Wilson (1982), Wolf et al. (1991).



Figure 7.1.1 Schematic of seepage meter in open channel with a falling-level reservoir and U-tube manometer (Bouwer, 1986, by permission).

Use	Shoreline Monitoring Well	Seepage Meter	Mini- Piezometer	Well Point	Core Sample	
Determination of hydraulic gradient	в, с	NA	A	A	NA	
Determination of hydraulic conductivity of sediments	в, с	в, с	с	A	с	
Flux between ground water/surface water systems	D	с	в, с	В, С	NA	
Determination of long term interaction of ground and surface water	В	D	С	В	NA	
Collection of flux samples for field screening analysis	с	с	А	A	NA	
Collection of flux samples for lab analysis	с	D	с	Α	NA	
Estimation of sediment transport properties	NA	NA	C (porosity only)	C (porosity only)	A	
NA - not applicable A - good performance in most conditions B - acceptable when used in conjunction w/ another technique C - acceptable under certain conditions D - poor choice						

Source: Wolf et al. (1991), by permission

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7.1 INFILTRATION

7.1.2 Land Surface Methods*

Other Names Used to Describe Method: Infiltration test basins, sprinkler infiltrometer, cylinder infiltrometer.

<u>Uses at Contaminated Sites</u>: Measuring of ground-water recharge; determining soil infiltration capacity for land treatment of wastewater; calculating sorptivity (sprinkler infiltrometer [see also, Section 6.4.2]).

<u>Method Description</u>: Infiltration test basins: Large cylinder infiltrometers (Figure 7.1.2a and b) or basins (20 feet by 20 feet) are constructed at several locations in a field and flooded with water. Measurements are similar to cylinder infiltrometer for measuring infiltration rates. Sprinkler infiltrometer: Nozzles or drop-formers are used to simulate the size and fall velocity of natural raindrops over a plot, which is set up so that surface runoff can be accurately measured (see also, Section 7.2.5 and Figure 7.2.5). The difference between the amount of water applied and the surface runoff is the infiltration rate.

<u>Method Selection Considerations</u>: Infiltration can be estimated using empirical relations or infiltration equations (Section 7.1.4), using other measured variables that can be measured with an air-entry permeameter (Section 7.3.4), and from soil moisture content, if the $K(\phi)$ relationship is known (see Section 6.4.1). Infiltration test basins are relatively simple and provide more representative measurements than cylinder infiltrometers, but are relatively expensive, time consuming, and water availability can be a problem. Sprinkler infiltrometers are relatively complex, expensive to operate, and are not well adapted to routine field applications. Relatively recent developments of more portable equipment might make this a more attractive method (see Section 7.2.5).

<u>Frequency of Use</u>: Test basins are used primarily for the design of full-scale projects for the land treatment of municipal wastewater. Sprinkler infiltrometers have been widely used in agricultural research, but have not been commonly used for contaminated site characterization.

Standard Methods/Guidelines: Sprinkler infiltrometer: Peterson and Bubenzer (1986), Test basin: U.S. EPA (1981).

Sources for Additional Information: Dunne and Leopold (1978), Thompson et al. (1989), Wilson (1982). Sprinkler infiltrometers: Bertrand (1965), Clothier et al. (1981b), Dunne and Leopold (1978), Grierson and Oades (1977), Hamon (1979), Parr and Bertrand (1960), Peterson and Bubenzer (1986), Sidle (1979), Tovey and Pair (1963), U.S. EPA (1981), Zegelin and White (1982). Peterson and Bubenzer (1986) summarize information on over 30 rainfall simulator and sprinkler-infiltrometer studies and cite 66 references, which are not listed here on this topic. See also, Section 7.2.5. Cylinder infiltrometers: Bureau of Reclamation (1978), Haise et al. (1956), Hills (1971), Parr and Bertrand (1960); See also, Section 7.3.1 and Table 7-4. Test basins: Abele et al. (1980), Nielsen et al. (1973), Parr and Bertrand (1960), U.S. Army Corps of Engineers (1980).

*See also, cylinder infiltrometers (Section 7.3.1).



(a)



(b)

Figure 7.1.2 Infiltration test basin: (a) Groove preparation for flashing (berm); (b) Schematic of finished installation (U.S. EPA, 1981, after U.S. Army Corps of Engineers, 1980).

7.1 INFILTRATION

7.1.3 Watershed Methods

Other Names Used to Describe Method: Average infiltration method, point infiltration method, empirical relations.

Uses at Contaminated Sites: Estimating infiltration over large areas for water budget studies.

<u>Method Description</u>: Average infiltration method: Infiltration is estimated by measuring the rainfall duration and intensity from individual precipitation events, and subtracting the measured runoff. The difference between the two values is assumed to be the infiltration. Figure 7.1.3 illustrates how infiltration capacity curves are developed for a small watershed. Empirical relationships: Musgrave and Holtan (1964) have grouped soils into four basic classes and summarized infiltration rates for a large number of different soil types within these classes. The U.S. Soil Conservation Service has used this classification system to develop some empirical relationships for estimating infiltration based on soil-vegetation types to approximate infiltration over large watershed areas (SCS, 1975). Huggins and Monke (1966) developed an empirical infiltration equation in which infiltration is a function of soil moisture. Rankl (1990) has developed a point infiltration watershed model for estimating runoff using infiltration estimates based on soil types and several empirical infiltration parameters.

<u>Method Selection Considerations</u>: Methods for estimating infiltration in watersheds generally do not have enough accuracy for site specific applications. Empirical relationships might be useful when combined with limited infiltrometer measurements to obtain a gross approximation of infiltration.

<u>Frequency of Use</u>: Uncommon, mainly because site investigations tend to cover areas that are smaller than entire watersheds.

Standard Methods/Guidelines: --

Sources for Additional Information: Average infiltration method: Dunne and Leopold (1978); Empirical relations: Bras (1990-Huggins-Monke and SCS methods), Huggins and Monke (1966), Musgrave and Holtan (1964), Rankl (1982, 1990), SCS (1975); Other: Parr and Bertrand (1960).



Figure 7.1.3 Average infiltration method of computing an infiltration capacity curve for a small drainage basin (Dunne and Lcopold, 1978). Bursts of rainfall plotted in the upper diagram (a) cause separate hydrograph rises (b). Each burst provides one point on the infiltration capacity curve (c). (From: *Water in Environmental Planning* by Dunne and Leopold, Copyright © 1978 by W.H. Freeman and Company, reprinted with permission).

7.1 INFILTRATION

7.1.4 Infiltration Equations

Other Names Used to Describe Method: Green-Ampt, Richards, Philip equations (numerous other solutions and refinements have been derived from these equations); parametric infiltration equations; Horton equation; Huggins-Monke equation (see Section 7.1.3).

Uses at Contaminated Sites: Obtaining indirect estimations of soil infiltration rates.

Method Description: An alternative to direct measurement of infiltration is to measure variables required for analytical equations, such as the Richards', Green-Ampt, and Philip's equations. Variables typically required for these equations include the hydraulic conductivity of the wetted zone, pressure head at the wetting front, and sorptivity of the soil. Most of these variables can be measured in situ with an air entry permeameter (Section 7.3.4), from unsaturated hydraulic-conductivity/pressure head relations (Section 6.3.1), and from infiltrometer measurements (Section 7.3.1). When estimating infiltration, it also is important to take the initial water content of the soil into account. Infiltration rates in a dry soil will be initially higher, and take a longer time to reach saturated hydraulic conductivity than infiltration into a soil that already is relatively wet (Figure 7.1.4). The Horton empirical equation for infiltration has been commonly used by hydrologists, but has a basic problem in that it does not satisfy the theoretical requirement that the initial infiltration be of infinite value. It might be suitable for describing infiltration when water is applied by rain or sprinkling for short time periods. Most infiltration equations have been derived from the study of soil physics. The Green-Ampt equation is satisfactory for describing infiltration into initially dry coarse-textured soils, and requires data on the hydraulic conductivity of the wetted zone and an estimate of the critical pressure head of soil for wetting. The Philip's equation is a two-parameter algebraic equation derived from the Richards' basic partial differential equation for unsaturated flow, and requires measurement or estimation of sorptivity and an infiltration curve. Numerous solutions and refinements of the basic Green-Ampt and Richards' equations have been developed in recent years, as well other approaches, such as parametric infiltration equations. Each equation or model has its own assumptions and soil moisture conditions that must be satisfied. For example, the Broadbridge-White model (Broadbridge and White, 1988) spans a wide range of known soil hydraulic properties. Table 7-5 identifies over 50 references dealing with equations and models for infiltration and unsaturated flow in the vadose zone and this literature should be reviewed to identify the most appropriate equation or model.

<u>Method Selection Considerations</u>: All infiltration equations require field measurement and characterization of the spatial variability of the required parameters for accurate estimation of infiltration. Advantages: Might be the best method for evaluating vertical infiltration rates of soils that contain restricting layers at some depth. Disadvantages: If infiltrating water contains sediment or suspended solids, the reduction infiltration rate due to the accumulation of solids on the soil surface must be estimated.

<u>Frequency of Use</u>: The Green-Ampt and Philip's equations are probably the most commonly used. As noted above, numerous refinements and alternatives to these equations, which might merit consideration, have been developed in recent years.

Standard Methods/Guidelines: --

Sources for Additional Information: Bouwer (1986), Green and Ampt (1911), Philip (1957a, 1969), Thompson, et al. (1989). See also, references in Table 7-5 and Section 7.2.8 (Unsaturated Hydraulic Conductivity, Physical/Empirical Equations and Relationships).



Figure 7.1.4 The effect of the initial water content of soil on infiltration rates (Everett et al., 1983).

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7.2 UNSATURATED HYDRAULIC CONDUCTIVITY

7.2.1 Instantaneous Profile Method

Other Names Used to Describe Method: Unsteady drainage flux, plane of zero flux, instantaneous rate method, hot-air method (Arya et al., 1975, as cited by Bouma, 1982).

<u>Uses at Contaminated Sites</u>: Measuring unsaturated hydraulic conductivity and diffusivity for vadose zone contaminant transport evaluation. Also can be used for monitoring water flux (Section 7.5.2) and developing moisture characteristic curves (Section 6.3.1) or K-matric potential relationships.

<u>Method Description</u>: A field plot (Figure 7.2.1a) or a double-ring infiltrometer is placed on a soil plot (Figure 7.2.1b) and instrumented with a battery of tensiometers at different depths for measuring water pressures (see Section 6.2.1) and an access tube for neutron moisture logging (see Section 3.3.3 and 6.3.2). The soil is wetted to saturation throughout the study depth. Wetting is stopped and the surface covered to prevent evaporation. Water pressure and water content are measured at intervals as the soil drains. Any combination of methods for measuring soil water potential (see Section 6.2) and soil moisture content (see Section 6.3 and Table 6-1) can be used for this method. Tensiometer/soil core method: A variant of the instantaneous profile method in which only changes in soil water pressure are monitored in the field after the soil is wetted. Soil cores are collected from the depth increments that tensiometers have been placed and moisture characteristic curves are measured in the laboratory. Hydraulic conductivity at different matric potentials is calculated from the field-measured tensiometer data and the moisture characteristic curve. The entrapped air method (Section 7.2.6) also can be considered a variant of this method.

Method/Device Selection Considerations: Instantaneous Profile Advantages: (1) Simple and reasonably accurate at each measuring site; and (2) suitable for stratified soils. Instantaneous Profile Disadvantages: (1) Provides hydraulic conductivity values only for draining profiles and values will be different during wetting cycles; (2) time consuming and relatively expensive, especially if site variability requires a large number of sites to obtain mean values; (3) does not provide reliable data near saturation (0 to -15 centimeters) because of rapidly changing and poorly defined pressure head gradients; (4) primarily measures vertical conductivity and will underestimate flux if horizontal conductivity exceeds vertical conductivity; (5) interactions between wastewater and solids might affect results (such as dispersion of clays or clogging); and (6) not suitable for percolating water with sufficient concentration of chemical wastes (such as nonaqueous phase liquids) to change its physical properties that affect infiltration rates. Tensiometer/Soil Core Advantages: Similar to instantaneous profile method except that field data collection is less time consuming and expensive because only soil-water pressure is monitored. Tensiometer/Soil Core Disadvantages: Similar to instantaneous profile method, except that use of laboratory measurements on soil cores might not accurately reflect in situ conditions.

Frequency of Use: Probably the most commonly used field method for accurate measurement of unsaturated hydraulic conductivity.

Standard Methods/Guidelines: ASTM (1990a), Bouma et al. (1974), Green et al. (1986).

Sources for Additional Information: Bouwer and Jackson (1974), Everett et al. (1982, 1983), Hendrickx (1990), Thompson et al. (1989), Wilson (1980). See also, Table 7-3.



(a)



Figure 7.2.1 Instantaneous profile method: (a) Planar view of field plot; (b) Double-ring infiltrometer with multipledepth tensiometers (Green et al., 1986, by permission).

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7.2 UNSATURATED HYDRAULIC CONDUCTIVITY

7.2.2 Draining Profile Methods

Other Names Used to Describe Method: Simplified unsteady drainage flux method, unit gradient method, Theta (θ) method, flux method, CGA-method, water content measurement method (flux), tensiometric simplified functions method.

Uses at Contaminated Sites: Estimating unsaturated hydraulic conductivity and flux of water in the vadose zone.

Method Description: Unsaturated hydraulic conductivity: A number of approaches have been developed to simplify the instantaneous profile method so that only soil moisture content or soil water potential needs to be monitored in the field. Procedures are similar to the instantaneous profile method in that the soil is wetted until steady-state infiltration (field saturated) conditions are reached at the test plot or double-ring infiltrometer, at which time wetting is stopped and the surface covered to prevent evaporation. Changes in the draining profile are observed as a function of time either by monitoring soil water content at different depths, or by monitoring soil matric potential at different depths. Different equations are used to calculate hydraulic conductivity as a function of soil water content. In the theta (θ) method, changes in soil water content with time at different depths are used in the calculations. Figure 7.2.2a illustrates use of this method for a single soil horizon. In the flux- and CGA-methods, different formulas involving changes in average water content over the depth of interest are used to calculate hydraulic conductivity as a function of soil-water suction (Figure 7.2.2b). Flux in the vadose zone: Monitoring of changes in water content over time (neutron logging, tensiometers, resistance blocks, and psychrometers) allows calculation of the water flux for a given depth (Wilson 1980, 1982). See Section 7.5.2 for further discussion of flux measurement using these methods.

Method Selection Considerations: Moisture Profile Advantages: (1) Simpler instrumentation allows measurements to be made at more points than with more complex methods, allowing statistical analysis to characterize soil variability; and (2) works well on coarse- and fine-textured homogenous materials. Moisture Profile Disadvantages: (1) Point measurements are less accurate than instantaneous profile and crust methods; (2) most methods assume a unit hydraulic gradient and will not work if the assumption does not apply; and (3) separate measurements of matric potential-water content relationships are required. Pressure Profile Advantages: (1) Simpler instrumentation allows measurements to be made at more points than with more complex methods, allowing statistical analysis to characterize soil variability; (2) the assumption of unit hydraulic gradient is not required; (3) measurement of matric potential-water content relationships are not required; (4) a one-time measurement of the soil water content profile allows estimates of drainage fluxes and soil water storage in the profile with time and as a function of average matric potential; and (5) works well in coarse- and fine-textured soils and soil profiles with stratification. Pressure Profile Disadvantages: (1) Point measurements are less accurate than instantaneous profile and crust methods; (2) reliable, frequent tensiometric data at small time and depth intervals, especially at low suctions, are required; (3) accurate determination of the representative field-saturated hydraulic conductivity is required; and (4) curves are somewhat less accurate for depths greater than around 100 centimeters. General Disadvantages: (1) Generally requires uniform drainage over shallow water tables (in deeper soils the upper profile can be draining while the lower profile is wetting, so flux will not equal drainage); (2) chemical conditions affecting methods to measure water content changes might introduce errors (i.e., chlorine in solution affecting neutron logging); (3) drainage in well-structured soils might occur more rapidly than in soil blocks where water content changes are measured, resulting in underestimation of water flux; and (4) a large number of measurements is required to characterize spatial variability.

Frequency of Use: Relatively new methods with good potential for more extensive field application due to their relative simplicity.

Standard Methods/Guidelines: θ and flux methods: Libardi et al. (1980); CGA-method: Chong et al. (1981); Pressure profile: Ahuja et al. (1988). See also, Section 7.2.7 (Parameter Identification).







(b)

Figure 7.2.2 Draining profile methods: (a) Theta method involves plotting the change in water content over time to determine empirical constant beta--hydraulic conductivity at any water content can then be calculated if steady state water content (θ_o) and steady state hydraulic conductivity (k_o) are known (Libardi et al., 1980, by permission); (b) Hydraulic conductivity as a function of soil-water suction: Calculations using the pressure profile method compared to detailed Darcian analysis (Ahuja et al., 1988, by permission).

Sources for Additional Information: Everett et al. (1983), Green et al. (1986), Hendrickx (1990), Wilson (1980, 1982). See also, Table 7-3.

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7.2 UNSATURATED HYDRAULIC CONDUCTIVITY

7.2.3 Tension Infiltrometers

Other Names Used to Describe Method: Tension disc permeameter, suction permeameter, porous plate infiltrometer, Guelph infiltrometer, sorptivity method.

Uses at Contaminated Sites: Measuring infiltration, sorptivity, hydraulic conductivity; characterizing macropore soil water flow and mean pore size.

Method Description: The tension infiltrometer originally was developed to measure soil sorptivity and diffusivity (see Section 6.3.2), but relatively recent improvements in instrument design (Figures 7.2.3a and b) have made this a versatile device for measuring and estimating a variety of soil hydrologic properties. The tension disc permeameter has three main components: (1) A nylon membrane that rests on the ground surface, (2) a calibrated reservoir, and (3) a bubble tower, which is used to control the starting tension in the calibrated reservoir (Figure 7.2.3a). At the beginning of the test, the water reservoir is full of water, and the water level in the bubble tower is set at a height to achieve the desired starting tension. The stopcock in the bubbler tower is opened to start the test, allowing air to enter the reservoir as water moves through the membrane into the soil. Multiple tests can be run in several ways by varying: (1) The starting tension, (2) the pore size of the nylon membrane, and (3) the size of the disc-membrane. Sorptivity is calculated from the rate at which the water level in the calibrated reservoir falls during the first 3 minutes, and hydraulic conductivity is determined when the infiltration rate reaches a steady flux. Measurements of sorptivity and hydraulic conductivity at different starting tensions allows development of a hydraulic conductivity-matric potential curve. The positive pressure permeameter (Figure 7.2.3b) looks similar to the tension permeameter, but operates quite differently. The unit is attached to a stainless steel cylinder, which is driven far enough into the ground to prevent water from leaking around the side. The supply pressure is the distance between the air bubble exit point and the soil surface, and can be adjusted by screws. The air entry side tube is filled with enough water to fill the space between the central water reservoir and the soil. This water is rapidly deposited on the soil surface by opening the side tube stopcock to start infiltration, and the rate of fall of water in the central calibirated water reservoir is measured.

<u>Method Selection Considerations</u>: Advantages: (1) Simpler than instantaneous profile, draining, and steady-flux methods because knowledge of initial water potential or content is not required, which eliminates requirements for installation of tensiometer or neutron access probes not required; (2) lower cost than more complex methods allows more extensive characterization of spatial variability of soil hydraulic characteristics; (3) rings do not need to be driven into the soil surface, avoiding possible disturbance of soil structure and allowing use of the method on rocky soils; and (4) control of tension at the surface allows characterization of flow in different pore sizes. Disadvantages: (1) Accurate measurements might be difficult in very wet and highly permeable soils; (2) methods requiring solution of simultaneous equations might be susceptible to errors resulting from soil heterogeneity; (3) measurements with instruments using different radii surface disks might be affected by spatial variability associated with different soil surfaces; and (4) measurements sample a relatively shallow depth of the soil surface (different depths can be tested by excavation, but the process become more cumbersome and time-consuming).

<u>Frequency of Use</u>: Tension infiltrometers have gained rapid acceptance in the last few years and are likely to become a standard tool for in situ determination of saturated and near-saturated soil hydraulic properties near the soil surface.

Standard Methods/Guidelines: Perroux and White (1988).

Sources for Additional Information: See Table 7-3.



Figure 7.2.3 Disc permeameters: (a) For supplying water at pressures less than or equal to zero; (b) For supplying water at positive pressures (Perroux and White, 1988, by permission).

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7.2 UNSATURATED HYDRAULIC CONDUCTIVITY

7.2.4 Crust-Imposed Steady Flux

Other Names Used to Describe Method: Unit-hydraulic gradient method, crust test.

<u>Uses at Contaminated Sites</u>: Measuring vertical unsaturated hydraulic conductivity during wetting portion of moisture characteristic curve.

<u>Method Description</u>: A pedestal soil about 25 cetimeters in diameter and 30 centimeters high is exposed by excavation, the exterior of which is covered with aluminum foil, and a tensiometer is inserted into the pedestal (Figure 7.2.4). Crusts with varying hydraulic conductivity by varying percentages of gypsum and sand or sand and quick-setting hydraulic cement. Each test run uses a crust placed on the soil surface, which is then covered by an infiltration ring with an air-tight cover, which fits tightly over the pedestal. A water source supplies water to the infiltration ring assembly at a constant head, with the crust controlling the flow of water to the soil pedestal to a rate below the maximum possible infiltration rates. Pressure head is monitored near the surface and at depth to determine when steady-state unsaturated flow has been reached. Successive steady-state flow systems, with increasing levels of saturation, are achieved by using crusts with increasingly higher permeabilities. Multiple tests allow plotting of hydraulic conductivity as a function of pressure head.

<u>Method Selection Considerations</u>: Advantages: (1) Measurements and calculations are reasonably simple; (2) a high degree of accuracy can be achieved if the crusts are carefully prepared; (3) can be used on sloping land surfaces; and (4) measurements can be made on large undisturbed soil columns to include effects of soil structure and other macroporosity, which might be missed by laboratory measurements of soil cores. Disadvantages: (1) Measurements are time and labor intensive; (2) a unit hydraulic gradient must exist in a vertical direction for measurements to be accurate (a reasonable assumption if steady-state flow is reached and the soil material is homogeneous); (3) only records the wetting portion of the soil water retention curve (see Section 6.4.1), so the effects of hysteresis are not determined; and (4) measurements apply to a relatively small area of soil.

Frequency of Use: Uncommon.

Standard Methods/Guidelines: ASTM (1990a), Green et al. (1986).

Sources for Additional Information: Bouma et al. (1974), Hendrickx (1990), Thompson et al. (1989), Wilson (1982). See also, Table 7-3.



Figure 7.2.4 Schematic diagram of field installation of the measurement apparatus for the crust-imposed steady flux method: M = constant-head device, Sc = wing nut, PC = plastic cover, W = water inlet, A = air outlet, RG = rubber gasket, C = gypsum-sand crust, Ca = tensiometer cap, Cy = metal cylinder with sharpened edge, H = height of mercury column above mercury pool, and G = height of mercury pool above tensiometer porous cup, P (Green et al., 1986, after Baker, 1977, by permission).

7.2 UNSATURATED HYDRAULIC CONDUCTIVITY

7.2.5 Sprinkler/Dripper Methods

Other Names Used to Describe Method: Sprinkler-imposed steady flux, dripper method.

<u>Uses at Contaminated Sites</u>: Sprinkler-impose steady flux: Measuring vertical unsaturated hydraulic conductivity during wetting portion of moisture characteristic curve; Dripper method: Measuring of saturated hydraulic conductivity and sorptivity and estimating hydraulic conductivity-matric potential function, $K(\phi)$, and matric-potential (ϕ)-moisture (θ) function.

Method Description: The sprinkler-imposed steady flux method is similar in principle to the crust-imposed steady flux method (Section 7.2.4). A sprinkler (Figure 7.2.5) is used to apply a steady rate of water to the soil surface, which is below the rate sufficient to saturate the soil. Soil moisture content is monitored using a neutron access tube, and matric potential is measured using tensiometers placed at different depths. Moisture content and pressure head is measured when steady-state flow conditions are achieved. K is calculated by dividing flux per unit area by the hydraulic gradient. Successively higher sprinkler flux rates are used to create the next steady-state flow system. Typically, the vertical gradient is unity. The dripper method is a relatively new and different method for measuring and estimating a variety of soil hydrologic properties. A water storage bottle with Marriott type burette is connected to button drippers (used commercially for drip irrigation) in a cluster-like arrangement, which allows different rates of constant discharge by plugging different numbers of drippers. The drippers are located in the center of a level and relatively smooth plot (about 0.8 meters square). When water flow begins, the diameters of the horizontal wetted and ponded zones are measured until a constant value is reached (i.e., the water dripping onto the soil moves downward rather than outward on the soil surface). When a steady state is reached, the rate of dripping is increased and the diameter measured until it stabilizes again. Sorptivity is determined by measuring the horizontal wetting front advance from the ponded zone borders as a function of time. The hydraulic conductivity-matric-potential-water content functions are estimated from measurements of the saturated area on the soil surface and the distance from the ponded radius to the wetting front as a function of dripper discharge rate for several rates using equations such as Brooks-Corey (Section 7.2.8).

Method Selection Considerations: Sprinkler-Imposed Steady Flux Advantages: Measurements apply to a larger sample area than the crust method. Sprinkler-Imposed Steady Flux Disadvantages: (1) As with the crust method, K is determined only during wetting; (2) unlike the crust method, it works only at relatively high moisture contents; and (3) sprinklers are relatively expensive and cumbersome to use. Dripper Method Advantages: (1) Equipment is much simpler and more portable than conventional sprinkler devices; (2) rock fragments in the soil do no pose a limitation (rock at the soil surface might create problems); and (3) several hydrologic parameters are measured (infiltration, sorptivity, K_{ts} , $K[\phi]$, and ϕ - θ functions). Dripper Method Disadvantages: Requires a flat, relatively dry soil.

Frequency of Use: Uncommon.

Standard Methods/Guidelines: Sprinkler-imposed steady flux: Green et al. (1986); Dripper method: Shani et al. (1987).

Sources for Additional Information: Hendrickx (1990), Thompson et al. (1989). See also, Table 7-3 and references in Section 7.1.2.



Figure 7.2.5 Layout of sprinkler infiltrometer (U.S. EPA, 1981, after Tovey and Pair, 1963).

7.2 UNSATURATED HYDRAULIC CONDUCTIVITY

7.2.6 Entrapped Air Method

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Measuring unsaturated hydraulic conductivity and flux in the vadose zone.

<u>Method Description</u>: This method is a variant of the instantaneous profile method (Section 7.2.1). An initially saturated column of porous material, in the process of draining to a water table at its base, is rewet at its upper surface at an appropriate time, causing an increase in the pore air pressure in the zone of entrapped air in the profile between the wetting and draining fronts. When steady state is reached, soil-water pressure is measured at different depth increments in the column to calculate pressure-head gradients along the bell-shaped water content profile caused by the zone of entrapped air. The water content profile is measured directly, or inferred from a separately measured moisture characteristic curve. Since the flow is steady and the flow rate is known, hydraulic conductivity over a range of water contents can be readily calculated. Figure 7.2.6 illustrates the types of data plots that are used in this method.

<u>Method Selection Considerations</u>: Advantages: (1) Instrumentation for the instantaneous profile method also can be used for this method; and (2) total time for data collection might be somewhat shorter than for instantaneous profile method. Disadvantages: (1) Generally does not work well in fine-grained soils; and (2) requires more closely spaced instrumentation for moisture and matric potential measurement than a conventional instantaneous profile method.

Frequency of Use: Uncommon. More widespread use in coarse-grained soils might be merited.

Standard Methods/Guidelines: Watson (1967).

Sources for Additional Information: Bouwer and Jackson (1974). See also, Table 7-3.



Figure 7.2.6 Entapped air method (Watson, 1967, by permission): Hydraulic head (a) and water content profiles (b) are measured at the same time after draining soil has been rewetted and the entrapped air moved downward into the soil. The hydraulic gradient (c) is determined from the hydraulic head profile and conductivity values are determined by dividing the steady-state flux by the gradient, and these values are plotted against the corresponding water content (c) to develop the $K(\theta)$ relationship.

7.2 UNSATURATED HYDRAULIC CONDUCTIVITY

7.2.7 Parameter Identification

Other Names Used to Describe Method: Parameter estimation/optimization.

Uses at Contaminated Sites: Estimating hydraulic conductivity at different water contents and other hydraulic properties from limited data.

<u>Method Description</u>: Results of one field or laboratory test are used to estimate hydraulic conductivity. Transient cumulative discharge of water from an initially saturated core (or in situ soil) are measured as a function of time. Numerical models coupled to statistical optimization routines analyze the result of the test by adjusting parameter values in the model until the measured response fits the model. Dane and Hruska (1983) used parameter estimation methods to estimate unsaturated hydraulic conductivity with varying hydraulic head using the draining profile method (see Section 7.2.2).

<u>Method Selection Considerations</u>: Advantages: (1) Relatively fast and inexpensive; and (2) measuring moisture content and hydraulic head as a function of time is not mandatory (but doing so will reduce the degree of uncertainty). Disadvantages: Incorrect solutions can result if incorrect models for soil hydraulic properties are used.

Frequency of Use: Relatively new method, which is being used with increasing frequency.

Standard Methods/Guidelines: --

Sources for Additional Information: Hendrickx (1990), Thompson et al. (1989). See also, Table 7-5.

7.2 UNSATURATED HYDRAULIC CONDUCTIVITY

7.2.8 Physical/Empirical Equations and Relationships

Other Names Used to Describe Method: --

<u>Uses at Contaminated Sites</u>: Estimating saturated and unsaturated hydraulic conductivity from other known or estimated soil parameters.

<u>Method Description</u>: Numerous empirical equations have been developed for estimating unsaturated hydraulic conductivity from other soil properties, such as: Pore-size distribution and moisture characteristic curves. Mualem (1986) classifies formulas into three major categories: (1) <u>Empirical</u> forms of $K(\phi)$ and $K(\theta)$ relationships; (2) <u>macroscopic</u> models, which derive an analytical formula for the $K(\theta)$ relationship; and (3) <u>statistical</u> models, which primarily rely on the soil moisture retention curve (see Section 6.4.1) as an analogy to the pore radii distribution function. The Childs-Collis George, Marshall, Millington-Quirk, Brooks-Corey, Mualem, and van Genuchten equations are well-known equations based on statistical models. Table 7-5 identifies over 30 references (Empirical Equations/Models), which cover theoretical aspects of these equations, and also 25 references, which focus on the estimation of soil hydraulic properties from soil physical properties.

<u>Method Selection Considerations</u>: Relatively fast; each empirical equation has its own application and limitations based upon the assumptions of the equations. Mualem (1986) provides guidance on which methods to use based on the type of soil data that are known or can be estimated.

Frequency of Use: Fairly Common. The Brooks-Corey (1964), Mualem (1976a), and Van Genuchten (1980) are among the more commonly used formulas in current use.

Standard Methods/Guidelines: Mualem (1986).

Sources for Additional Information: Bouwer and Jackson (1974), Hendrickx (1990), Thompson et al. (1989). See also, Table 7-5.

7.3 SATURATED HYDRAULIC CONDUCTIVITY (SHALLOW)

7.3.1 Cylinder Infiltrometers

Other Names Used to Describe Method: Open and sealed single-ring and double-ring infiltrometers.

<u>Uses at Contaminated Sites</u>: Measuring infiltration rates/potential, saturated hydraulic conductivity, and sorptivity; estimating ground-water recharge.

<u>Method Description</u>: An open ended cylinder (10 to 30 centimeters in diameter) is driven into the ground to a depth ranging from 5 to 50 centimeters. A shallow ponded depth (1 to 2 centimeters) is maintained in the cylinder for a long enough time to allow steady-state (saturated-flow) infiltration to develop. The rate at which water is added to maintain the ponded depth, or a constant head in the cylinder, is a direct measure of the maximum infiltration rate for the soil. One or two rings (with water maintained in both the inner and outer rings) can be used and the rings can be open or sealed (Figure 7.3.1a). Where infiltration rates are very slow, as in clay soils or testing of clay liners, sealed double-rings (Figure 7.3.1b) are recommended for measuring infiltration rates (Sai and Anderson, 1991). Sorptivity can be determined from infiltrometer measurements by plotting the rate of infiltration versus time during the first few minutes when flow is unsaturated (see Section 6.4.2).

<u>Method Selection Considerations</u>: Ring infiltrometers are the recommended method for testing the hydraulic conductivity of compacted soils (Sai and Anderson, 1991). Advantages: (1) Are simple, inexpensive, and portable; and (2) sealed ring infiltrometers can be used to evaluate macropore flow, but the process is more cumbersome than using a tension infiltrometer (see Section 7.2.3). Disadvantages: (1) Tend to overestimate natural infiltration due primarily to lateral divergence of flow with depth (especially single rings); (2) provide point measurements only, so numerous tests are required to characterize spatial variability; (3) results might be misleading if water used during the test is not similar to that which normally infiltrates (i.e., wastewater might reduce infiltration by clogging compared to rainwater); and (4) shallow impeding layers might promote lateral movement of water in preference to truly vertical flow, resulting in overestimation of intake rates over larger areas.

<u>Frequency of Use</u>: Standard method for measuring compacted soils. Less commonly used to measure infiltration potential of natural soils (see Section 7.1.1).

Standard Methods/Guidelines: Cylinder infiltrometer: Bouwer (1986); Double-ring: ASTM (1988, 1990a), Johnson (1963); Sealed double-ring infiltrometer: ASTM (1990a,b), U.S. EPA (1989).

Sources for Additional Information: Thompson et al. (1989), Wilson (1982). See also, Table 7-4.







Open, Double Ring



Sealed, Single Ring



Sealed, Double Ring

(a)



(b)

Figure 7.3.1 Cylinder infiltrometers: (a) Open and sealed single- and double-ring infiltrometers; (b) Details of sealed double-ring infiltrometer (U.S. EPA, 1989).

7.3 SATURATED HYDRAULIC CONDUCTIVITY (SHALLOW)

7.3.2 Constant-Head Borehole Infiltration

Other Names Used to Describe Method: Shallow-well pump-in, constant-head infiltrometer, borehole permeameter, dry/inverted auger hole method, compact constant-head (CCH) permeameter.

<u>Uses at Contaminated Sites</u>: Mainly measuring the horizontal component of saturated hydraulic conductivity in unsaturated soil.

<u>Method Description</u>: A hole is bored to the desired depth and a constant head of water is maintained in the hole (Figure 7.3.2). The test also can be used with a screened well point. When water flow into the soil reaches steady state conditions (i.e., water flow is constant to maintain constant head), the flow is measured. Hydraulic conductivity is calculated from equations using the following measurements: (1) Steady-state injection rate, (2) radius of the borehole, (3) height of water in the borehole, and (4) depth from the bottom of the borehole to the top of the impermeable layer. The calculated rate is the average hydraulic conductivity for the portion of the hole that was tested and, in a uniform soil, the measured rate of flow is dominated by the horizontal conductivity.

<u>Method Selection Considerations</u>: Advantages: (1) Recently developed compact constant-head permeameter can be used to depths up to 10 meters; (2) can be used in rocky or gravelly soil; and (3) tests a larger volume of soil compared to the Guelph permeameter. **Disadvantages**: (1) Test requires presence of an impermeable layer below the bottom of the borehole; (2) large quantities of water might be required; (3) a single test can take several days to complete; (4) requires soil that can maintain an open borehole; (5) smearing of the auger hole walls will result in underestimation of conductivity; and (6) measurements using water might not be applicable for evaluating potential for moving sewage wastewater or chemical waste liquids through the soil (can be overcome by using fluids in the test that are similar to the fluids of concern).

Frequency of Use: Commonly used method.

Standard Methods/Guidelines: Amoozegar and Warrick (1986, Section 29-3.2), ASTM (1990a).

Sources for Additional Information: Bureau of Reclamation (1978), Hendrickx (1990), Thompson et al. (1989). See also, Table 7-4.



Figure 7.3.2 Diagram of constant head device and geometry of the shallow well pump-in set up (Amoozegar and Warrick, 1986, by permission).

7.3 SATURATED HYDRAULIC CONDUCTIVITY (SHALLOW)

7.3.3 Guelph Permeameter

Other Names Used to Describe Method: Constant head well permeameter.

Uses at Contaminated Sites: Measuring saturated and unsaturated hydraulic conductivity and sorptivity in unsaturated soil.

<u>Method Description</u>: The Guelph Permeameter is a constant-head apparatus designed for small-diameter boreholes (2 to 5 centimeters). A device, which controls hydraulic head and measures the injection rate, is inserted into an uncased borehole. Constant head is maintained until steady-state flow is achieved. The design differs for models used in high conductivity and low conductivity porous media (Figure 7.3.3a and b). A vertical profile of K can be developed by repeating the test at various depths. Measurements typically represent an average of horizontal and vertical hydraulic conductivity.

<u>Method Selection Considerations</u>: Advantages: (1) Only requires one operator and is fast (usually ranges from 5 to 60 minutes); (2) relatively small volumes of water are required; (3) other parameters, such as unsaturated hydraulic conductivity and sorptivity, can be estimated; and (4) is commercially available. Disadvantage: (1) A limited volume of soil is tested, so replication and multiple tests are required to characterize spatial variability; (2) requires materials that can maintain an open borehole; (3) smearing of clay on borehole walls will result in measurements lower than the actual K; (4) rests on bottom of hole, which might impede vertical water flow, especially in small diameter holes; (5) unsaturated hydraulic conductivity and sorptivity measurements are based on assumptions that will have varying degrees of validity for different porous media; (6) depth limited to about 2 meters; and (7) measurements using water might not be applicable for evaluating potential for movement of sewage wastewater or chemical waste liquids through the soil (can be overcome by using fluids in the test that are similar to the fluids of concern).

Frequency of Use: This is a relatively new technique, which has gained rapid acceptance.

Standard Methods/Guidelines: ASTM (1990a), Reynolds and Elrick (1986).

Sources for Additional Information: Hendrickx (1990), Thompson et al. (1989). See also, Table 7-4.



Figure 7.3.3 Schematic of Guelph permeameter: (a) Model 1 for high conductivity porous media; (b) Model 2 for low conductivity porous media (Reynolds and Elrick, 1986, by permission).

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7.3 SATURATED HYDRAULIC CONDUCTIVITY (SHALLOW)

7.3.4 Air-Entry Permeameter

Other Names Used to Describe Method: --

<u>Uses at Contaminated Sites</u>: Measuring saturated hydraulic conductivity in unsaturated soil; estimating of $K(\phi)$ and $K(\theta)$ relationships.

<u>Method Description</u>: A cylinder 20 to 30 centimeters in diameter and over 10 centimeters long is driven about 10 centimeters into the soil. A layer of sand is placed inside the cylinder, and the cylinder is sealed with a topplate assembly and water is supplied to the cylinder from a reservoir (Figure 7.3.4). An air valve allows air to escape from the cylinder until the cylinder is completely filled, at which time it is closed. When the wetting front reaches the bottom of the cylinder below the soil surface, the supply of water is shut off and a valve attached to a vacuum gage is opened. The time required for the wetting front can be estimated by a few trials before the procedure is started, or alternatively, it can be detected using a fine tensiometer probe. The pressure inside the cylinder decreases to a minimum (the **air-entry value**), at which time air begins to bubble up through the soil. At this point, the equipment is removed and the depth of wetting front is determined by digging. The air-entry pressure can be calculated from pressure measurements and the depth of the wetting front, which can in turn be used to calculate saturated hydraulic conductivity. Bresler et al. (1978) describe a method for estimating unsaturated hydraulic conductivity as a function of water content and matric potential using the air-entry value obtained using an air-entry permeameter.

<u>Method Selection Considerations</u>: More sensitive to vertical than horizontal K. Advantages: (1) Is fast (around 1 hour), requires a small volume of water (around 10 liters), and is relatively simple to use; and (2) tests larger volume of soil than the Guelph permeameter. Disadvantages: (1) Multiple tests are required to characterize spatial variability; (2) the presence of macropores and cracks might cause problems; (3) measurements using water might not be applicable for evaluating potential for movement of sewage wastewater or chemical waste liquids through the soil (can be overcome by using fluids in the test that are similar to the fluids of concern); (4) gravel within 10 to 20 centimeters of the ground surface can cause problems in placement of the cylinder. See Table 7-2 for additional information.

Frequency of Use: Fairly widely used.

Standard Methods/Guidelines: ASTM (1990a), Bouwer (1966).

Sources for Additional Information: Amoozegar and Warrick (1986), Bouma (1983), Hendrickx (1990), Thompson et al. (1989). See also, Table 7-4.



Figure 7.3.4 Air entry permeameter (Thompson et al., 1989, after Bouwer, 1966, Copyright © 1989, Electric Power Research Institute, EPRI EN-6637, *Techniques to Develop Data for Hydrogeochemical Models*, reprinted with permission).

7.3 SATURATED HYDRAULIC CONDUCTIVITY (SHALLOW)

7.3.5 Double Tube Method

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Measuring saturated hydraulic conductivity in unsaturated soil.

<u>Method Description</u>: An auger hole is dug to the desired depth and cleaned with special tools. An outer tube is pushed into the bottom of the hole about 5 centimeters and an inner tube and a top-plate assembly are installed in the outer tube, with the inner tube pushed about 2 centimeters into the bottom of the hole (Figure 7.3.5). Each tube has a standpipe above the tube for observing the water levels in each tube. Both the inner and outer tubes are filled with water and equal head pressure is maintained in both by adjusting the water level in the inner tube, if necessary. After saturation of the bottom of the hole is achieved (usually after 1 hour for fine-textured soils), two sets of measurements are taken: (1) Water flow is shut off to the inner tube, and (2) the water level in both tubes is brought back to the starting level, and the water level in the outer tube is controlled so that it falls at the same rate as the inner tube. Saturated hydraulic conductivity is calculated using the two head versus time graphs plotted from the measurements.

<u>Method Selection Considerations</u>: Advantages: (1) Is commercially available; and (2) characterization of anisotropic soils is possible when the method is combined with the infiltration gradient method (Section 7.3.7) Disadvantages: (1) Is relatively complex and time-consuming (depending on the permeability of the soil the test procedures takes from 2 to 6 hours to complete, and requires over 200 liters of water for each test); (2) is not suitable for rocky soils; (3) multiple measurements are required to characterize spatial variability; (4) is less accurate than other available methods (see Table 7-2); and (5) measurements using water might not be applicable for evaluating potential for movement of sewage wastewater or chemical waste liquids through the soil (can be overcome by using fluids in the test that are similar to the fluids of concern).

Frequency of Use: Fairly uncommon.

Standard Methods/Guidelines: ASTM (1990a), Amoozegar and Warrick (1986).

Sources for Additional Information: Bouma (1983), Hendrickx (1990). See also, Table 7-4.


Figure 7.3.5 Diagram of equipment used for double-tube method (Amoozegar and Warrick, 1986, by permission).

7.3 SATURATED HYDRAULIC CONDUCTIVITY (SHALLOW)

7.3.6 Cylinder Permeameter

Other Names Used to Describe Method: Permeameter, ring permeameter.

Uses at Contaminated Sites: Measuring saturated hydraulic conductivity in unsaturated soil.

<u>Method Description</u>: A cylinder 45 to 50 centimeters in diameter and greater than 35 centimeters long is placed in a dug hole, which is wider than the cylinder. The cylinder is driven about 15 centimeters into the soil, and four tensiometers are placed symmetrically around the cylinder 10 centimeters from its sides and about 23 centimeters below the bottom of the hole (Figure 7.3.6). The hole and inside of the cylinder are maintained at a depth of about 15 centimeters The tensiometers are monitored until they read zero (saturation is achieved), at which time the rate of flow of water into the soil from the cylinder is measured. Conductivity is measured using Darcy's equation.

<u>Method Selection Considerations</u>: Advantages: (1) Is relatively simple; and (2) calculations are easy. Disadvantages: (1) Is time-consuming and requires in excess of 100 liters of water; (2) is not suitable for rocky soils; (3) measurements are not very accurate; and (4) measurements using water might not be applicable for evaluating potential for movement of sewage wastewater or chemical waste liquids through the soil (can be overcome by using fluids in the test that are similar to the fluids of concern). See Table 7-2 for additional information.

Frequency of Use: Uncommon.

Standard Methods/Guidelines: Boersma (1965).

Sources for Additional Information: Bouma (1983), Bureau of Reclamation (1978), Hendrickx (1990), Roberts (1984), U.S. EPA (1981), Winger (1960).



Figure 7.3.6 Schematic diagram of equipment for the cylinder permeameter method (Boersma, 1965, by permission).

7.3 SATURATED HYDRAULIC CONDUCTIVITY (SHALLOW)

7.3.7 Infiltration Gradient Method

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Measuring saturated hydraulic conductivity in unsaturated soil.

<u>Method Description</u>: This method combines elements of the cylinder permeameter (Section 7.3.6) and the double tube methods (Section 7.3.5). Two concentric cylinders are placed in an auger hole with small, fast-reacting piczometer tubes placed at different depths inside the inner tube (Figure 7.3.7). Changes in vertical hydraulic gradient are recorded as the hydraulic head in both tubes is kept equal and varied from 20 to over 200 centimeters When combined with the double tube method in the same hole, vertical and horizontal hydraulic conductivity components can be separated out.

<u>Method Selection Considerations</u>: Advantages: (1) Measures primarily vertical hydraulic conductivity; and (2) when used with the double tube method, vertical and horizontal components of hydraulic conductivity can be differentiated. Disadvantages: (1) Requires about 3 hours to complete and about 100 liters of water; (2) is not suitable for stony soils; (3) measurements using water might not be applicable for evaluating potential for movement of sewage wastewater or chemical waste liquids through the soil (can be overcome by using fluids in the test that are similar to the fluids of concern).

Frequency of Use: Uncommon.

Standard Methods/Guidelines: Bouwer (1978).

Sources for Additional Information: Bouwer and Jackson (1974). See also, Table 7-4.





7.3 SATURATED HYDRAULIC CONDUCTIVITY (SHALLOW)

7.3.8 In Situ Monoliths

Other Names Used to Describe Method: Column method, cube method.

<u>Uses at Contaminated Sites</u>: Measuring vertical saturated hydraulic conductivity; measuring K_{mt} in soils with continuous macropores (column-crust method).

Method Description: Column method: A soil column (30 centimeters in diameter and 30 centimeters thick) is carved out in situ and encased in gypsum or resin. Water is applied to the top of the column until steady-state infiltration is reached. Flow through the column is measured volumetrically, either by collecting outflow from a column that has been detached from the soil, or by measuring the flow rate once steady-state infiltration has been reached. Cube method: This is a variant of the column method, which allows measurement of both vertical and horizontal saturated hydraulic conductivity. A cube of soil (30 centimeters by 30 centimeters by 30 centimeters) is excavated in situ and encased in gypsum. The cube is removed, and vertical hydraulic conductivity is measured using procedures similar to the column method. Next, the open ends of the cube are sealed with gypsum, the cube is turned sideways, and the gypsum removed from the top and bottom for a second measurement of hydraulic conductivity (Figure 7.3.8a). Column-crust method: This combines elements of the crust test (Section 7.2.4) with the column method in order to differentiate between the macropore and soil matrix components of saturated flow. A column of soil is excavated in situ and tensiometers are placed in the column before it is encased in gypsum. Macropore flow is measured by adding water until steady-state infiltration is reached with the column detached (Figure 7.3.8b-A). A light crust then is placed on the column and water applied until steady infiltration is reached at zero pressure head (Figure 7.3.8b-C). The latter measurement represents K_{ut} without macropore flow. Macropore flow is the difference between the first and second measurements. By using crusts of different thicknesses, unsaturated hydraulic conductivity can be measured with this method as well.

Method Selection Considerations: Advantages: (1) Are relatively simple; (2) calculations are simple and accurate; (3) cube method allows accurate measurement of vertical and horizontal saturated-hydraulic conductivity; and (4) column-crust method allows differentiation of macropore and soil matrix saturated flow. Disadvantages: (1) Preparation and execution are relatively time consuming; and (2) measurements using water might not be applicable for evaluating potential for movement of sewage wastewater or chemical waste liquids through the soil (can be overcome by using fluids in the test that are similar to the fluids of concern). See Table 7-2 for additional information.

Frequency of Use: Uncommon.

Standard Methods/Guidelines: --

Sources for Additional Information: Bouma (1983). See also, Table 7-4.



(a)



(b)

Figure 7.3.8 Monolith methods: (a) Gypsum covered cube of soil used to measure vertical and horizontal saturatedhydraulic conductivity (Bouma and Dekker, 1981, by permission); (b) Schematic representation of three types of flux measurements using the column-crust method (Bouma, 1982, by permission).

7.3 SATURATED HYDRAULIC CONDUCTIVITY (SHALLOW)

7.3.9 Boutwell Method

Other Names Used to Describe Method: --

<u>Uses at Contaminated Sites</u>: Measuring vertical and horizontal components of saturated hydraulic conductivity at the ground surface, especially clay liners.

Method Description: The Boutwell method is a two-stage falling-head borehole test used to calculate vertical and horizontal hydraulic conductivity. In Stage I, a borehole is cased, grouted, and filled with water (Figure 7.3.9a). The casing and standpipe are filled with water and flow out of the bottom of the borehole is monitored until steady-state conditions are reached. In Stage II, the hole is extended beyond the bottom of the borehole, with the ratio of the length to diameter of the uncased zone between 1 and 1.5 (Figure 7.3.9a). The casing and standpipe are reassembled and the rate of fall of water in the stand pipe is monitored until steady-state conditions are reached again. Sai and Anderson (1991) provide the equations for calculating vertical and horizontal hydraulic conductivity.

<u>Method Selection Considerations</u>: Advantages: (1) Is relatively fast, inexpensive, simple, and convenient to use; (2) can measure very low hydraulic conductivities $(1 \times 10^9 \text{ meters/second})$; (3) allows determination of vertical and horizontal components of hydraulic conductivity. **Disadvantages**: (1) Measures small volume, so might miss soil macropores and other flaws in soil liner construction; (2) short test periods do not allow entrapped air to dissolve; (3) method does not account for the effects of soil suction; and (4) effects of incomplete and variable suction are not known.

Frequency of Use: Relatively new method, which has not been widely tested.

Standard Methods/Guidelines: Sai and Anderson (1991).

Sources for Additional Information: Boutwell and Derric (1986).



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Figure 7.3.9 Schmatic diagram of Boutwell borehole permeameter: (a) Stage I; (b) Stage II (Sai and Anderson, 1991).

7.3 SATURATED HYDRAULIC CONDUCTIVITY (SHALLOW)

7.3.10 Velocity Permeameter

Other Names Used to Describe Method: Velocity head permeameter, falling head permeameter.

Uses at Contaminated Sites: Measuring vertical saturated hydraulic conductivity.

<u>Method Description</u>: The velocity permeameter estimates hydraulic conductivity based on the rate of fall of water in a head tube above a soil core enclosed within a coring tube (Figure 7.3.10a). This is a falling-head test in which data on change of water level in the head tube is entered into small programmable calculator equipped with a timing module. The data on varying rates of fall are used to calculate a series of hydraulic conductivity values, which are plotted against time since the test began (Figure 7.3.10b). The field saturated hydraulic conductivity is the lowest value on the graph.

<u>Method Selection Considerations</u>: Advantages: Is a relatively simple and rapid method (about an hour), provided the velocity of the fall of water in the head tube can be measured accurately (accuracy increases as the ratio of the soil-core diameter to the head-tube diameter increases). Disadvantages: (1) Maintaining a seal around the edges of the coring device might be difficult under high liquid heads; and (2) field measurements and data reduction require a skilled operator.

Frequency of Use: Relatively new method, which has not been widely tested.

Standard Methods/Guidelines: --

Sources for Additional Information: Kanwar et al. (1987), Sai and Anderson (1991).



Figure 7.3.10 Velocity permeameter: (a) Front view and operational schematic; (b) Measured hydraulic conductivity versus time (Sai and Anderson, 1991).

7.3 SATURATED HYDRAULIC CONDUCTIVITY (SHALLOW)

7.3.11 Percolation Test

Other Names Used to Describe Method: Perc test, falling head test.

Uses at Contaminated Sites: --

<u>Method Description</u>: This test is similar to the constant head shallow-well pump-in method, except that a constant head is not maintained for the test. A 6-inch diameter hole is augered or dug to the depth of interest and 2 inches of gravel are placed in the bottom to prevent scouring by water poured into the hole (Figure 7.3.11). Water is maintained at a depth of 12 inches in the hole until the soil around the hole is saturated (generally 4 to 12 hours). The water level is adjusted to 6 inches above the gravel and the amount of fall over a 30 minute period is measured. The water level is adjusted to 6 inches above the gravel after each measurement, and measurements are repeated until two successive water drops do not vary by more than 1/16 inches. Results are reported in minutes/inch or inches/hour.

<u>Method Selection Considerations</u>: Advantages: Is simple and easy to calculate. Disadvantages: (1) Results can be highly variable due to soil moisture conditions at the time of the test and the individual performing the test; and (2) when properly done, still only provides an approximate measure of saturated hydraulic conductivity.

<u>Frequency of Use</u>: Widely used for assessing soil suitability and design of septic tank soil absorption systems for sewage treatment. Not recommended for measuring saturated hydraulic conductivity.

Standard Methods/Guidelines: U.S. EPA (1980).

Sources for Additional Information: See Table 7-4.



Figure 7.3.11 Floating indicator for percolation test (U.S. EPA, 1980).

7.4 SATURATED HYDRAULIC CONDUCTIVITY (DEEP)

7.4.1 USBR Single-Well Methods

Other Names Used to Describe Method: Gravity permeability tests.

Uses at Contaminated Sites: Estimating saturated hydraulic conductivity in deep boreholes in the vadose zone.

<u>Method Description</u>: Water is pumped into a borehole at a rate that maintains a uniform water level in a basal test section. Saturated hydraulic conductivity is estimated from appropriate curves and equations based on: (1) Dimensions of the hole and inlet pipes, (2) length in contact with formation, (3) height of water above the base of the borehole, (4) depth to water table, and (5) intake rate at steady state. Method 1 (Figure 7.4.1a) uses an open borehole of 6 inches or more in diameter. The bottom of a feed pipe and observation pipe are set near the bottom of the borehole, and the open portion of the borehole is filled with gravel pack if required to maintain stability. Where gravel pack is required for stability, 40 feet is about the maximum depth that this test can be economically carried out. Method 2 (Figure 7.4.1b) uses a perforated casing for the depth of interest into which water is pumped and an observation pipe set near the top of the perforations. The casing is sunk by drilling, jetting or driving, whichever give the tightest fit. This method generally is less accurate than Method 1 for unconsolidated materials, but might be the only practical method for determining permeabilities in streambeds or lakebeds below water. Method 2 will not work because it is the least accurate of the three methods.

<u>Method Selection Considerations</u>: All methods require some form of a casing advancement drilling method. Advantages: (1) Allows estimating saturated hydraulic conductivity at great depths in the vadose zone; (2) a series of tests as the borehole is deepened allows developing of profile of K values; and (3) can be conducted in unconsolidated formations where packer testing (Section 4.2.3) might not be feasible. Disadvantages: (1) K_{sat} tends to be underestimated because solution method assumes the flow region is entirely saturated, which is not true; (2) expensive and time-consuming (especially in dry, coarse-grained material), so multiple tests to adequately characterize spatial variability might be prohibitive; and (3) requires skilled personnel to conduct tests. Method descriptions above indicate specific conditions under which the different methods are used. Packer testing (Section 4.2.3) is probably the preferred method where boreholes are in consolidated rock.

Frequency of Use: Most likely to be used in the western United States where the saturated zone is far below the ground surface.

Standard Methods/Guidelines: Bureau of Reclamation (1981).

Sources for Additional Information: Everett et al. (1982), Schmid (1967), Stephens and Neuman (1982a,b), Wilson (1982), Zanger (1953).



K = coefficient of permeability, feet per second under a unit gradient

- Q=uniform flow into well, ft3/s
- r = radius of test section, ft
- H = height of column of water in well, ft
- A =length of test section, ft (for this method, A = H)

- C_u and C_s = conductivity coefficients $X = \frac{H}{T_u}$ (100) = percent of unsaturated stratum $T_u = U^u D + H =$ distance from water surface in well to water table, ft
- U = thickness of unsaturated permeable bed, ft
- D = distance from around surface to bottom of test section, ft
- I = feed pipe for pouring water into well (a 2-inch standard pipe is usually satisfactory)
- O = observation pipe ($1\frac{1}{4}$ -inch o.d. pipe is satisfactory)

a = surface area of test section (area of wall plus area of battam), ft* Limitations:

 $A \ge 10r$ and $\frac{Q}{q} \le 0.10$



(a)

- K = coefficient of permeability, feet per second under a unit gradient
- Q = steady flow into well, ft³/s
- H=height of water in well, ft
- A = length of perforated section, ft
- r, =outside rodius of casing (radius of hole in consolidated material), ft
- re =effective radius of well = r. (area of perforations)/ (outside area of perforated section of casing) ; $\mathbf{r_i} = \mathbf{r_e}$ in consolidated material that will stand open and is not cased
- C_u and $C_s =$ conductivity coefficients
- T_u=distance from water level in casing to water table, ft
- a = surface area of test section (area of perforations plus area of bottom), ft²; where clay seal is used at bottom, a= area of perforations
- S = thickness of saturated permeable material above an underlying relatively impermeable stratum, ft
- $X = \frac{H}{T_u} (100) = percent of unsaturated stratum U = thickness of unsaturated material above water table, ft$
- D = distance from ground surface to bottom of test section, ft
- O = observation pipe (1 to $1\frac{1}{4}$ inch pipe)

Limitations: S \geq 5A, A \geq 10r, and $\frac{Q}{a} \leq 0.10$ Notes:

In zone 3, H is the difference in elevation between the normal water table and the water level in the well. In zones 2 and 3, if a clay seal is placed at the bottom of the casing, the factor $4 \frac{\Gamma_i}{\Gamma_e}$ is omitted from the equations. Where the test is run with "A" as an open hole, $\frac{\Gamma_i}{\Gamma_e} = 1$ and $(C_s + 4 \frac{\Gamma_i}{\Gamma_e}) = (C_s + 4)$.

(b)

Figure 7.4.1 USBR single-well hydraulic conductivity tests: (a) Method 1; (b) Method 2 (Bureau of Reclamation, 1981).

7.4 SATURATED HYDRAULIC CONDUCTIVITY (DEEP)

7.4.2 USBR Multiple-Well Method

Other Names Used to Describe Method: Gravity permeability test.

<u>Uses at Contaminated Sites</u>: Estimating saturated hydraulic conductivity where lenses of slowly permeable material are widespread.

<u>Method Description</u>: A 6-inch intake well and at least three observation wells are installed to the top of an impermeable layer (Figure 7.4.2). Water is pumped into the central well at a steady rate and changes in water levels in the piezometers are measured. K_{ast} is calculated using the appropriate curves and equations.

<u>Method Selection Considerations</u>: Advantages: Results can be used to estimate lateral flow rates in perched ground-water regions. Disadvantages: (1) Is expensive and time-consuming; and (2) requires trained personnel.

Frequency of Use: Relatively uncommon.

Standard Methods/Guidelines: Bureau of Reclamation (1981-Method 4).

Sources for Additional Information: Wilson (1982).

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K = coefficient of permeability, feet per second under a unit gradient

Q = uniform flow into intake well, ft³/s

 $r_1, r_2, and r_3 = distance from intake well to observation holes, ft$

h₁,h₂, and h₃ = height of water in observation holes r₁, r₂, and r₃ respectively, above elevation of top of impermeable layer, ft

H= height of column of water in intake pipe above top of impermeable stratum, ft

U= distance from ground surface to impermeable bed, ft

Figure 7.4.2 USBR multiple-well method (Bureau of Reclamation, 1981).

7.4 SATURATED HYDRAULIC CONDUCTIVITY (DEEP)

7.4.3 Stephens-Neuman Single-Well Method

Other Names Used to Describe Method: Unsteady flow permeability test.

Uses at Contaminated Sites: Estimating saturated hydraulic conductivity in the deep vadose zone.

<u>Method Description</u>: Water is pumped into a well drilled to the depth of interest, and changes in water level with time are measured and used to estimate steady-state infiltration, rather than pumping until steady-state infiltration is achieved, as in the USBR single-well tests (Section 7.4.1). Empirical formulas based on numerical simulations using the unsaturated characteristics of four soils allows correction for unsaturated flow conditions during the test.

<u>Method Selection Considerations</u>: Advantages: (1) Provides more accurate estimation of the saturated hydraulic conductivity of unsaturated soil than the USBR single-well methods; (2) less time is required for the test because steady-state flow conditions are not required; (3) allows estimating saturated hydraulic conductivity at great depths in the vadose zone; (4) a series of tests as the borehole is deepened allows developing of a profile of K values; and (5) can be conducted in unconsolidated formations where packer testing (Section 4.2.3) might not be feasible. Disadvantages: The cost of drilling deep boreholes makes it difficult to characterize spatial variability of hydraulic conductivity with this method. Packer testing (Section 4.2.3) probably is the preferred method where boreholes are in consolidated rock.

Frequency of Use: Uncommon.

Standard Methods/Guidelines: Stephens and Neuman (1980, 1982c).

Sources for Additional Information: Everett et al. (1982), Wilson (1982).

7.4 SATURATED HYDRAULIC CONDUCTIVITY (DEEP)

7.4.4 Air Permeability Method

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Estimating saturated hydraulic conductivity in the deep vadose zone.

<u>Method Description</u>: Air pressure changes in the subsurface in response to change in the barometric pressure at the land surface are measured in specially constructed piezometers (Figure 7.4.4). Pressure response data combined with information on the air-filled porosity allow calculation of air permeability. If the Klinkenberg effect is small, hydraulic conductivity can be calculated from air permeability. Section 9.4.4 provides further information of methods for measuring air permeability in shallow zones.

<u>Method Selection Considerations</u>: Advantages: Can be used to estimate hydraulic conductivity in layered materials in the vadose zone. Disadvantages: (1) Is indirect; (2) soils must be dry, since too much soil water inhibits air flow; (3) is expensive and time consuming; (4) is complex, requiring trained personnel; (5) in finegrained materials, the permeability to air is greater than the hydraulic permeability because of the Klinkenberg effect; and (6) presence of clays with high shrink-swell make it difficult to accurately calculate hydraulic conductivity from air permeability.

Frequency of Use: Uncommon.

Standard Methods/Guidelines: Weeks (1978).

Sources for Additional Information: Everett et al. (1982), Wilson (1982).



Figure 7.4.4 Schematic diagram of manifold and connections to a piezometer nest to determine air pressure at selected depths in the vadose zone (adapted from Weeks, 1978).

7.5 WATER FLUX (UNSATURATED ZONE)

7.5.1 Water Budget Methods

Other Names Used to Describe Method: Water balance method, water content method.

<u>Uses at Contaminated Sites</u>: Estimating leachate generation by percolating water in the subsurface; estimating solute velocity.

Method Description: Water flux: The method itself involves calculations of water flux in the subsurface based on inflow (precipitation), outflow (runoff, evapotranspiration), and changes in storage (water content). Parameters, which must be estimated or measured in the field, include: (1) Precipitation (Sections 8.1.1 and 8.1.2), (2) evapotranspiration (Sections 8.3 and 8.4), and (3) available water capacity (Section 6.3.3) or changes in water content or soil matric potential with time (using methods described in Sections 6.2 and 6.3). Figure 7.5.1 shows a cumulative water balance used to determine seepage from a wastewater lagoon to ground water. Simplified versions of this approach include: (1) The instantaneous profile method (Section 7.2.1); (2) Wilson (1980) describes a variant of this method, which provides a profile-specific water budget by measuring changes in water content at different depths by assuming all terms of the water budget calculation are zero except for flux and soil-water storage changes (similar to draining profile method, Section 7.2.2); and (3) the Thornthwaite method, which can be used with climatic data (monthly precipitation and temperature) and soil water holding characteristics. Vadose-zone solute-transport models involving the soil rooting zone are based primarily on water budget principles and allow estimation of water and contaminant flux. Velocity: Everett et al. (1983) describe a simplified method for estimating vertical travel time to a water table where the vadose zone is very thick. Depth of penetration (dv2) equals depth of percolating water during a specified time period (dw), divided by the volumetric water content at field capacity (θ): $dv^2 = dw/\theta$. Dividing this value into the thickness of the vadose zone provides an estimate of how long it will take water to percolate below the rooting zone to reach the water table if no preferential flow paths occur.

<u>Method Selection Considerations</u>: Advantages: (1) Provides estimates of flux for an entire area, rather than point estimates; and (2) relatively simple if most parameters in the water budget equation can be estimated with acceptable accuracy or set equal to zero. **Disadvantages**: (1) Accurate field measurement of some parameters, such as evapotranspiration, is difficult and field measurement of all required parameters is expensive and time consuming; (2) errors in measurement or estimation of components (inflow and outflow, evapotranspiration, rainfall, and ambient temperature) might accumulate in flux estimates; (3) difficult to use where water table is high and changes in water storage are minimal; (4) contaminant chemical reactions in soil solution, which change water transmission and water holding properties, reduce accuracy of estimates; (5) flux calculations based on soilwater storage changes will be underestimated in highly structure soils where water flow occurs primarily in cracks and macropores; and (6) in poorly leveled fields, water might pond in low spots, and run off rapidly in other areas, resulting in actual local fluxes, which can vary considerably from average fluxes calculated assuming uniform water application.

Frequency of Use: Relatively common (most vadose zone computer models use some form of water budget)

Standard Methods/Guidelines: Wagenet (1986).

Sources for Additional Information: Everett et al. (1983), Wilson (1980, 1982). See also, Table 7-6.



Figure 7.5.1 Schematic annual cumulative water balance to determine seepage from a wastewater lagoon to ground water (Wells, 1988, by permission).

7.5 WATER FLUX (UNSATURATED ZONE)

7.5.2 Soil Moisture/Matric Potential Methods

Other Names Used to Describe Method: Hydraulic gradient/unit hydraulic gradient methods, instantaneous profile method (Section 7.2.1), draining profile methods (7.2.2).

Uses at Contaminated Sites: Estimating water flux in the vadose zone.

Method Description: A variety of methods are available to estimate flux in the vadose, based on measurements of changes in soil moisture and/or matric potential with depth and over time. Depending on the specific method, various types of calibration curves, such as matric potential versus water content and hydraulic conductivity as a function of matric potential and/or water content, can be used. The instantaneous profile method (Section 7.2.1) and various draining profile methods (Section 7.2.2) can be used to calculate water flux. The hydraulic gradient method uses the basic approach of the instantaneous profile method (Section 7.2.1), except that evapotranspiration and infiltration of natural precipitation can be allowed. Hydraulic gradients in the unsaturated zone are measured in the subsurface by installing tensiometers or psychrometers (Sections 6.2.1 and 6.2.2). For each textural change, calibration curves are required to relate negative pressure measurement to water content (moisture retention curves, see Section 6.4.1) and water content to unsaturated hydraulic conductivity (see methods allowing measurement of the $K(\theta)$ function in Table 7-1). The unit hydraulic gradient method is similar to the hydraulic gradient method, except that a hydraulic gradient of 1 is assumed, requiring only one pressure measuring unit at each depth of interest. Curves relating water content to matric potential (Section 6.3.1), and water content to hydraulic conductivity, allow calculation of the amount of water flowing at the time of each measurement, and measurements taken over time allow calculation of water flux. Alternatively, curves directly relating hydraulic conductivity to matric potential can be used (see Table 7-1).

Method Selection Considerations: Instantaneous Profile: See Section 7.2.1. Draining Profile: See Section 7.2.2. Hydraulic Gradient Advantages: Allows accurate measurement over a relatively large area. Hydraulic Gradient Disadvantages: (1) Is relatively expensive to install enough units to characterize spatial variability for statistical analysis; (2) generally is restricted to shallow depths in the vadose zone and might not be suitable for ponds or landfills; (3) results are subject to hysteresis in the calibration curves (i.e., water content-pressure relations differ depending on whether the soil is wetting or drying.); (4) requires obtaining calibration curves (water content versus matric potential and hydraulic conductivity as a function of water content/matric potential) for each change in texture; (5) requires measurement units in depthwise increments throughout the vadose zone, and gradients across layers might suggest vertical flow when horizontal flow is actually predominant; and (6) might not be suitable at sites underlain by fractured media. Unit Hydraulic Gradient Advantages: Simpler and less expensive than the unit hydraulic gradient method because fewer calibration relationships are required. Unit Hydraulle Gradient Disadvantages: (1) A large number of units are still required to characterize spatial variability; (2) the assumption of unit hydraulic gradients might not apply, particularly in layered media; (3) might not be suitable for ponds or landfills; and (4) as with the hydraulic gradient method, calibration measurements are required for each change in texture and results are subject to hysteresis in calibration curves.

Frequency of Use: Commonly used in research applications, less commonly used for monitoring flux at contaminated sites.

Standard Methods/Guidelines: Everett et al. (1983) describe steps, equations, and sample calculations for several draining profile and hydraulic gradient methods.

Sources for Additional Information: Bouwer and Jackson (1974), Everett et al. (1983), Wilson (1980, 1982). See also, Table 7-6.

7.5 WATER FLUX (UNSATURATED ZONE)

7.5.3 Tracers

Other Names Used to Describe Method: Chloride mass balance, bomb-pulse radionuclides (tritium, chlorine-36), stable isotopes (deuterium and oxygen-18), other tracers.

Uses at Contaminated Sites: Estimating flux and velocity.

Method Description: A wide variety of tracers can be used to estimate flux and velocity in the vadose zone. The chloride mass-balance flux method is a geochemical technique in which vertical profiles of chloride concentration are developed by analysis of soil samples. Flux is calculated based on assumed flux contributed to the soil from precipitation. The tritium and chloride-36 flux methods are used to identify water that has infiltrated in the last 30 to 40 years (see Section 4.3.5). The technique involves extracting soil water from core samples and analyzing for tritium concentration (liquid scintillation counting technique) or extracting chloride as AgCl for analysis of chloride-36 on using a tandem accelerator mass spectrometer. The stable isotope flux method is a relatively new method based on the movement of deuterium and oxygen-18 in water molecules through the vadose zone (the same isotopes have long been used to date ground water [see Section 4.3.4]). Water from soil cores is extracted using a vacuum distillation procedure and the soil water is processed using CO2/H2O equilibration or hydrogen reduction for analysis of stable isotope ratios on a ratio mass spectrometer. Velocity methods: A conservative tracer (iodide, bromide) is introduced into the liquid source. Samples obtained from suction samplers and/or free drainage samplers at successive depths are used to plot tracer breakthrough. Artificial tracers are used by applying a known amount of a conservative tracer, such as chloride or bromide, to the ground surface and collecting samples (from vertically spaced suction and/or free drainage samplers) at intervals to trace the speed of flow. Analysis of changes in concentration with time also allows estimation of flux by mass balance analysis.

Method Selection Considerations: Chloride Mass-Balance Advantages: Is relatively inexpensive and easy to use. Chloride Mass Balance Disadvantages: Is inaccurate if the following key assumptions do not apply: (1) Average rate of chloride deposition from precipitation to the soil is constant; and (2) chloride does not move below the root zone by preferential flow paths. Bomb-Pulse Advantages: Good method for determining whether water has infiltrated in the last 30 to 40 years. Bomb-Pulse Disadvantages: Extraction and analytical techniques are relatively complex and expensive and required equipment might not be readily available. Stable Isotope Advantages: In addition to estimating recharge rate, other soil-water movement processes, such as evaporation and liquid/vapor flux, can be estimated. Stable Isotope Disadvantages: (1) Extraction and analytical techniques are relatively complex and expensive and required equipment might not be readily available; (2) requires sampling to be done during a lengthy period of little or no precipitation, or infiltration into the soil from other sources will occur (i.e., the method is restricted to arid and semi-arid areas); and (3) requires vertical movement of soil water because significant lateral soil-water movement would invalidate assumptions used to calculate flux. Artificial Tracer Velocity Advantages: (1) Direct and simple method; (2) reflect flow in actual pores if freedrainage samplers are used; and (3) more accurate than methods requiring measurements of parameters in Darcy's equation. Artificial Tracer Velocity Disadvantages: (1) Use of nonconservative tracers (i.e., tracers that move slower than the velocity of water) will underestimate flux/velocity; (2) use of suction samplers might alter flow field and suction samplers cannot be used to sample soil water in very dry soil; (3) in structured media, actual velocity might be higher than measured because of flow in cracks (can be dealt with by also using zerotension samplers [Section 9.3.1]); (4) if velocities are slow, long time periods might be required for tests; and (5) average velocity of water-borne tracers might not be the same as average velocity of chemical liquids.

Frequency of Use: Tracer velocity: Relatively common.

Standard Methods/Guidelines: ASTM Draft Guide for Comparison of Techniques to Quantify the Soil-Moisture Flux (in preparation).

Sources for Additional Information: Everett et al. (1983). See also, Table 7-6 and Sections 4.3.6 and 4.3.7.

7.5 WATER FLUX (UNSATURATED ZONE)

7.5.4 Soil-Water Flux Meters

Other Names Used to Describe Method: Soil water flowmeter, direct flow/intercepting/hydraulic-resistance type, thermal/heat probe type.

<u>Uses at Contaminated Sites</u>: Measuring soil-water flux (amount of water moving through a unit cross-sectional area of soil in a unit time period).

<u>Method Description</u>: Two major types of instruments have been developed for direct measurement of in situ unsaturated soil-water flux: (1) Units that measure flow directly (intercepting meters), and (2) thermal meters, which measure the movement of a thermal pulse in a porous cup. The intercepting-type hydraulic-resistance meter, first developed by Cary (1968, 1970) and refined by Dirksen (1972, 1974), involves intercepting part or all of the soil-water flux and determining its magnitude by measuring the hydraulic-head loss across the inflow and outflow portions of the meter. Tensiometers are installed nearby to monitor head loss in the undisturbed soil and hydraulic resistance of the valve in the meter adjusted to match conditions in the soil. A recent refinement combines features of the methods by Dirksen (1974) and by Duke and Haise (1973-see Section 9.2.6). In this instrument, soil-water flow is intercepted by a porous plate in which the suction is automatically adjusted to maintain the same matric potentials above the plate and in the surrounding undisturbed soil (Figure 7.5.4). Both the hydraulic-resistance type and suction type meters require excavation of a pit and installation of the meter in the side of the pit at the desired depth.

Method Selection Considerations: Advantages: (1) Information on hydraulic conductivity or hydraulic gradient is not required; (2) can provide reasonable direct water flux measurements if properly used; (3) most useful for localized and specialized studies; (4) suction-type meter overcomes most of the major disadvantages of other types by eliminating the need for extensive laboratory or in situ calibrations, is not adversely affected by air bubbles, and can sample flux over a larger area. Disadvantages: (1) Is relatively expensive and complex method; (2) localized nature of measurement does not allow estimating flux over large areas unless many flux meters are installed; (3) soil is disturbed during installation of most types and might interrupt normal soil-water flow patterns; (4) calibration procedures are tedious, especially for multilayered media (Dirksen hydraulic-resistance type and suction type do not require much calibration); (5) requirement for trench installation limits use to relatively shallow depths; (6) hydraulic-resistance type meters require fairly wet soils to perform effectively and the presence of air bubbles in soil water or in filter cloth and tubing will reduce flow into the meter; (7) most types involve the measurement of flow in disturbed soil, and meters are especially difficult to install in layered media without affecting flow lines; (8) most units require contact with relatively fine-grained porous media and will not work well in coarse-grained or fractured media (not a problem with suction type); and (9) thermal meter will give erroneous readings if chemical waste fluids have different heat conducting properties than water and have not been thoroughly tested in the field.

Frequency of Use: Uncommon.

Standard Methods/Guidelines: Hydraulic-resistance type: Wagenet (1986); Suction type: van Grinsven et al. (1988).

Sources for Additional Information: Bouwer and Jackson (1974), Everett et al. (1983), Wilson (1980). See also, Table 7-6.



Figure 7.5.4 Schematic diagram of suction soil-water flux meter components as used in field; P₁, P₂, and P₃ are tensiometers (van Grinsven et al., 1988, by permission).

7.5 WATER FLUX (UNSATURATED ZONE)

7.5.5 Velocity Estimation

Other Names Used to Describe Method: Velocity-flux/velocity-long-term infiltration calculation, velocity from suction cups.

Uses at Contaminated Sites: Measuring or estimating velocity with which water travels in the vadose zone.

<u>Method Description</u>: Flux or long-term infiltration calculation: Velocity can be calculated by dividing flux values obtained by methods described above (Sections 7.5.1 to 7.5.4), or dividing the long-term infiltration rate (as determined using methods in Sections 7.1.1 or 7.1.4) by average water content. Both methods assume that: (1) Hydraulic gradients are unity, (2) an average water content can be determined, (3) flow is vertical, and (4) a homogenous media exists. Indirect estimates of velocity can be obtained using suction samplers (Section 9.2). Apparent vertical velocity is estimated by observing the time it takes a wetting front from a surface source to reach vertically placed suction samplers, as indicated by a change from little or no soil-water retrieval during sampling to ready collection of soil water during suction. Section 7.5.1 (flux water budget methods) describes a simplified method for estimating velocity using water budget data, and 7.5.3 (tracers) describes use of tracers to estimating velocity.

<u>Method Selection Considerations</u>: Flux/Infiltration Calculation Advantages: (1) Is simple and inexpensive when coupled with other methods; and (2) is suitable for making a preliminary estimate of travel time of pollutants in the vadose zone. Flux/Infiltration Calculation Disadvantages: (1) Underestimates velocity in structured media; (2) is not valid if perching layers cause lateral flow; (3) for multi-layered media, an average moisture content value might be difficult to obtain; (4) might be difficult to obtain equivalent water content values where liquid wastes have different properties than water.

Frequency of Use: Flux/infiltration calculations: Relatively common.

Standard Methods/Guidelines: Flux calculation: Bouwer (1980), Wilson (1980); Suction cup: Everett et al. (1983).

Sources for Additional Information: Wilson (1982), Everett et al. (1983); Case studies: Biggar and Nielsen (1976), Jury and Sposito (1985).

7.5 WATER FLUX (UNSATURATED ZONE)

7.5.6 Physical and Empirical Equations

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Estimating water flux in the vadose zone.

<u>Method Description</u>: A soil-physics based approach to quantifying soil-water flux requires measurement or estimation of hydraulic characteristic data and the use of physically or empirically-based equations to calculate flux. Any technique that measures hydraulic conductivity as a function of water content or matric potential (see Section 6.3.1 and Section 7.2 generally) allows calculation of water flux if the appropriate parameter (water content or matric potential) is monitored. Most of the methods in Section 7.5.2 (Soil Moisture/Matric Potential Methods) use this approach in one way or another. Numerous physically- and empirically-based equations have been developed to model infiltration and flow in the unsaturated zone. Sections 7.1.4 (Infiltration Equations) and Section 7.2.8 (Physical/Empirical Equations and Relationships) provide an overview of these approaches. The catalog-of-hydraulic-properties approach involves the use of "typical" hydraulic properties associated with physical soil properties, such as texture, porosity, and bulk density, to estimate both saturated and unsaturated hydraulic conductivity, provided that physical characteristics of the soils of interest are similar to soils for which data are available.

<u>Method Selection Considerations</u>: Physical and empirical equations: See Sections 7.1.4 and 7.2.8. Catalog-ofhydraulic-properties advantages: (1) Simple, quick, and can be used to estimate relative variations in hydraulic conductivity caused by stratification; and (2) is good for sensitivity analysis. Catalog-of-hydraulic-properties disadvantages: Might be prone to large errors because of lack of comparability between soil properties and because of spatial variability in soil properties.

<u>Frequency of Use</u>: Most methods for measuring soil hydraulic properties are based on, or require the use of, one or more physical and/or empirical models. Estimation of hydraulic properties from other soil physical properties is commonly used to obtain "ballpark" estimates of flux.

Standard Methods/Guidelines: --

Sources for Additional Information: Warrick et al. (1977). See generally, references for Sections 7.1.4, 7.2.8, and Table 7-5.

Table 7-3 Reference Index for Measurement of Unsaturated Hydraulic Conductivity and Flux in the Vadose Zone

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Topic	References
Reviews	Bouma (1977), Bouma et al. (1974), Bouwer and Jackson (1974), Dirksen (1991), Green et al. (1986), Hendrickx (1990), Hillel and Benyamini (1974), Stephens and Neuman (1982a), U.S. EPA (1986)
Instantaneous Profile	Ahuja et al. (1976), Arya et al. (1975), Baker et al. (1974), Cassel (1974), Dane (1980), Davidson et al. (1969), Flühler et al. (1976), Hillel and Benyamini (1974), Hillel et al. (1972), Hsieh and Enfield (1974), Klute (1972), Nagnal and DeVries
	(1976), Nielsen and Biggar (1973), Nielsen et al. (1964, 1973), Ogata and Richards (1957), Richards et al. (1956), Roberts (1984), Rose and Krishnan
	(1967), Rose et al. (1965), Schuh and Cline (1990), Shouse et al. (1992), Simmons et al. (1979), Stone et al. (1973), Stoner (1985), Unlu et al. (1989, 1990), van Bavel et al. (1968), Warrick and Amoozegar-Fard (1980), Watson (1966); In Situ
	Soil Block: Cheng et al. (1975), Luxmoore et al. (1981); <u>Tensiometers/Soil Cores</u> : Cassel (1971), Carvallo et al. (1976), Miller et al. (1965)
Draining Profile	<u>Moisture Profile</u> : Chong et al. (1981), Dane (1980), Dane and Hruska (1983), Libardi et al. (1980), Luxmoore et al. (1981), Sisson et al. (1980); <u>Pressure</u> <u>Profile</u> : Ahuja et al. (1980, 1982, 1988), Schuh et al. (1984), Wall and John (1982)
Tension Infiltrometers	Designs: Ankeny et al. (1988), Perroux and White (1988); <u>Hydraulic Conductivity</u> : Ankeny et al. (1991), Baumgartner et al. (1987), Clothier and Smettem (1990), Cook (1991), Elrick et al. (1987), Havlena and Stephens (1992), Reynolds and Elrick (1991), Sai and Anderson (1991), Smettem and Clothier (1989), Warrick (1992), White and Perroux (1987, 1989); <u>Sorptivity/Diffusivity</u> : Chong and Green (1983), Clothier and Smettem (1990), Clothier and White (1981), Dirksen (1975),
	Russo and Bresler (1980), Smettem and Clothier (1989), Smiles and Harvey (1973), Walker and Chong (1986), White and Perroux (1987, 1989); <u>Infiltration/Macroporosity</u> : Ankeny et al. (1990), Clothier et al. (1981a), Jarvis et al. (1984), Watson and Luxmoore (1986), Wilson and Luxmoore (1988).
Crust-Imposed Steady Flux	Anderson and Bouma (1977-laboratory), Baker (1977), Baker and Bouma (1976), Booltinck et al. (1991), Bouma (1975), Bouma and Denning (1972), Bouma et al. (1971), Hillel and Gardner (1969, 1970), Reinds (1988-laboratory), Roberts (1984), Spaans et al. (1990), Stoner (1985)
Sprinkler/Dripper Methods	<u>Sprinkler-Imposed Flux</u> : Chong (1983-sorptivity), Hillel and Benyamini (1974), Hills et al. (1989), McQueen (1963), Morin et al. (1967), Rawitz et al. (1972), Reinds (1988-laboratory), van de Pol et al. (1977), Youngs (1964-laboratory); <u>Dripper Infiltrometers</u> : Bridge and Ross (1985), Shani et al (1987); <u>Infiltration Rates</u> : See Section 7.1.2.
Entrapped Air Method	Dixon and Linden (1972), Peck (1965), Starr et al. (1978), Takagi (1960); see also, references on effects of entrapped air on hydraulic conductivity in Table 7-4
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Table 7-4 Reference Index for Measurement of Saturated Hydraulic Conductivity in the Vadose Zone

Торіс	References
Saturated Hydraulic Conductivity (Above Shallow Water Table)	
Reviews	Amoozegar and Warrick (1986), Boersma (1965), Bouwer and Jackson (1974), Hamilton et al. (1981), Hendrickx (1990), Kessler and Oosterbaan (1974), Lambe (1955), Sai and Anderson (1991), Stephens et al. (1988), Winger (1960), Youngs (1991); <u>Method Comparisons</u> : Havlena and Stephens (1992), Lee et al. (1985), Reynolds et al. (1983), Roberts (1984), Sai and Anderson (1991), U.S. EPA (1986); <u>Chemical Effects on Clays</u> : Brown (1988), Roberts (1984)
Effect of Entrapped Air	Bouwer (1966, 1978), Bouwer and Jackson (1974), Chahal (1964), Corey (1957), Jarrett and Fritton (1978), McWhorter et al. (1973), Peck (1969), Stephens et al. (1984)
Temperature Effects	Chahal (1964), Constantz (1982), Giakoumakis and Tsakiris (1991), Haridasan and Jensen (1972), Hopmans and Dane (1986)
Cylinder Infiltrometers	Aronovici (1955), Bouwer (1963), Burgy and Luthin (1956), Dixon (1975-sealed infiltrometer), Havlena and Stephens (1992), Johnson (1963), Priksat et al. (1992), Reynolds and Elrick (1990), Roberts (1984), Sai and Anderson (1991), Scotter et al. (1982), Swartzendruber and Olsen (1961a,b); <u>Compacted Liner Tests</u> : Daniel (1984, 1989), Daniel and Trautwein (1986), Elsbury et al. (1988), Panno et al. (1991), Pederson et al. (1988), Rogowski (1990), Sai and Anderson (1991), U.S. EPA (1989), Youngs (1991)
Constant-Head Borehole Infiltration	Amoozegar (1989a,b), Banton (1993), Boersma (1965), Bouwer (1978), Elrick and Reynolds (1992), Fritton et al. (1986), Havlena and Stephens (1992), Heinen and Raats (1990), Kanwar et al. (1987), Philip (1985a), Picornell and Guerra (1992), Reynolds et al. (1983, 1985), Stephens et al. (1987, 1988), Talsma (1987), Zanger (1953)
Guelph Permeameter	Elrick and Reynolds (1992), Elrick et al. (1987, 1988), Havlena and Stephens (1992), Heinen and Raats (1990), Jabro and Fritton (1990), Lee et al. (1985), Logsdon et al. (1990), Reynolds and Elrick (1985a, 1985b, 1986, 1987), Reynolds et al. (1983), Sai and Anderson (1991), Stephens et al. (1988), Talsma (1987), Talsma and Hallam (1980), Wilson et al. (1989)
Air-Entry Permeameter	Aldabagh and Beer (1971), Bouma (1983), Bouwer (1966, 1978), Bresler et al. (1978-K ^{misst}), Havlena and Stephens (1992), Lee et al. (1985), Reynolds et al. (1983), Roberts (1984), Russo and Bresler (1980), Sai and Anderson (1991), Shani et al. (1987), Stephens et al. (1988), Topp and Binns (1976), U.S. EPA (1981), Youngs (1991)
Double Tube Method	Boersma (1965), Bouma (1971, 1983), Bouma and Hole (1971), Bouwer (1961, 1962, 1964a, 1978), Bouwer and Rice (1964, 1967), Brust et al. (1968), Kessler and Oosterbaan (1974), U.S. EPA (1981)
Infiltration Gradient	Bouwer (1964a, 1978), Bouwer and Jackson (1974), Bouwer and Rice (1967), Rice (1967)

Table 7-4 (cont.)

Торіс	References
In Situ Monoliths	<u>Cube Method</u> : Bouma and Dekker (1981), Roberts (1984); <u>Column Method</u> : Baker and Bouma (1975), Bouma (1980), Bouma et al. (1976, 1979, 1981), Vroon et al. (1988); <u>Column-Crust Method</u> : Bouma (1982); <u>Monoliths</u> : Jager and van der Voort (1966), Mielke (1973), Sai and Anderson (1991), Stibbe et al. (1970), Tzimas (1979)
Percolation Test	Barbarick et al. (1976), Chan (1976), Elrick and Reynolds (1986), Hill (1966), Jabro and Fritton (1990), U.S. EPA (1980), U.S. PHS (1969); <u>Percolation Test</u> <u>Relationship to Ksat</u> : Bicki et al. (1988), Bouma (1971), Fritton et al. (1986), Healy and Laak (1973), Jabro and Fritton (1990), Mellon (1973), Paige and Veneman (1993), Winneberger (1974)
Saturated Hydraulic Conductivity (Deep Water Table)	See Section 7.4

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Table 7-5 Reference Index for Physical and Empirical Equations and Models of Hydraulic Properties in the Vadose Zone

Topic	References
Infiltration/Saturated Hydraulic Conductivity	
Infiltration Theory/Equations	Bouwer (1964b), Brandt et al. (1971), Broadbridge and White (1988), Childs (1967), Clothier et al. (1981b), Hanks and Bowers (1962), Hanks et al. (1969), Hogarth et al. (1989), Horton (1935, 1939, 1940), Knight (1983), Knight and Philip (1974), Kutilek (1980), Panikar and Nanjappa (1977), Parlange (1972), Parlange and Smith (1976), Parlange et al. (1982, 1985), Parr and Bertrand (1960), Philip (1954, 1957a,b, 1958a,b, 1969, 1973, 1975, 1985b, 1989a,b), Philip and Knight (1974), Pullan (1990), Raats (1973), Reichardt et al. (1972), Richards (1931, 1965), Rijtema and Wassink (1969), Rubin and Steinhardt (1963), Rubin et al. (1964), Sharma et al. (1980), Stallman (1967), Swartzendruber (1987a,b), Swartzendruber and Clague (1989), Swartzendruber and Hogarth (1991), Talsma and Parlange (1972), Warrick (1985), Warrick and Hussen (1993), White and Broadbridge (1988), White and Sully (1987), White et al. (1989), Wilson and Luthin (1963), Wooding (1968)
Unsaturated Hydraulic Conductivit	Y
Parameter Identification	Dane and Hruska (1983), Hornung (1983), Kool and Parker (1988), Kool et al. (1985, 1987), Parker et al. (1985), Ravi and Jennings (1990-laboratory measurements), Sisson et al. (1980), van Dam et al. (1992), Van Genuchten et al. (1989), Zachmann (1981, 1982)
Empirical Equations/Models (See also, Table 6-3)	Empirical Equations: Bresler et al. (1978), Brooks and Corey (1964, 1966), Gardner (1958), Laliberte et al. (1966-values for use with BC equation), Messing (1989), Raats and Gardner (1971), Ritjema (1965), Wind (1955); <u>Macroscopic Models</u> : Irmay (1954), Mualem (1978); <u>Statistical Models</u> : Burdine (1953), Childs and Collis-George (1950), Marshall (1958), Millington and Quirk (1959, 1961, 1964), Mualem (1976a), Mualem and Dagan (1978), Purcell (1949), Rieu and Sposito (1991a,b), Ross and Smettem (1993), Vachaud (1967), Van Genuchten (1979, 1980), Weeks and Richards (1967); <u>Use/Comparisons</u> : Bruce (1972), Brust et al. (1968), Brutsaert (1967), Green and Corey (1971), Jackson (1972), Jackson et al. (1965), Kunze et al. (1968), Nielsen et al. (1960), Rogers and Klute (1971), Roulier et al. (1972), Stockton and Warrick (1971),
Hydraulic Properties from Soil Physical Properties	
(See also, Table 6-3)	Ahuja et al. (1984), Alexander and Skaggs (1987), Anderson and Bouma (1973), Basak (1972), Bloeman (1980), Burdine (1953), Campbell (1974), Clapp and Hornberger (1978), Clausnitzer et al. (1992), de Jong (1982), Hanks et al. (1969), Laliberte and Corey (1967), Marshall (1958), Mason et al. (1957), McCuen et al. (1981), Mehuys et al. (1975), Mishra et al. (1989), Mualem (1976b), Puckett et al. (1985), Rawls and Brakensiek (1985), Rawls et al. (1982), Reichardt et al. (1975), Rogowski (1972), Saxton et al. (1986), Schuh et al. (1988), Tyler and Wheatcraft (1989), Van Genuchten and Nielsen (1985), White and Perroux (1989-sorptivity), Williams et al. (1992), Wösten and Van Genuchten (1988)

Table 7-6 Reference Index for Water Flux Methods

Торіс	References
General Reviews	Allison (1987), Gee and Hillel (1988), Roth et al. (1990), Simmons et al. (1979), U.S. EPA (1986-vadose zone travel time), Wagenet (1986)
Water Budget	Fenn et al. (1975), Gee and Hillel (1988), Jensen (1974), Kmet (1982), Simmers (1987), Sokolow and Chapman (1974), Warrick and Amoozegar-Fard (1981), Zepp and Belz (1992); <u>Thornthwaite Method</u> : Dunne and Leopold (1978), Thornethwaite and Mather (1957), Wilmott (1977); <u>Case Studies</u> : Aguilar and Aldon (1991), Dreiss and Anderson (1985), Fenn et al. (1975), Forslund and Daily (1990), Mather and Rodriguez (1978), Orr et al. (1990), Panno et al. (1991), Young and Clapp (1989)
Soil Moisture/Potential	Case Studies: Aguilar and Aldon (1991), Enfield et al. (1973), LaRue et al. (1968), Simmons et al. (1979), van Bavel et al. (1968)
Tracers	<u>Chloride</u> : Allison (1987), Allison and Huges (1978, 1983), Allison et al. (1985), Johnston (1987), Knowlton et al. (1992), Scanlon (1991), Sharma and Hughes (1985), Sukhija et al. (1988), Walker et al (1991); <u>Tritium</u> : Allison and Huges (1978), Evans et al. (1976), Frissel et al. (1974), Knowlton et al. (1992), Phillips et al. (1988); <u>Other</u> : Allison et al. (1985), Frissel et al. (1974), Knowlton et al. (1992), Sharma and Hughes (1985)
Soil-Water Flux Meters	<u>Thermal</u> : Byrne (1971), Byrne et al. (1967, 1968); <u>Hydraulic Resistance</u> : Cary (1968, 1970, 1971, 1973), Dirksen (1972, 1974); <u>Suction-Hydraulic Resistance</u> : van Grinsven et al. (1988).

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SECTION 8

VADOSE ZONE WATER BUDGET CHARACTERIZATION METHODS

Water movement and transport of contaminants in the vadose zone is determined by the amount of **precipitation** that enters the ground by **infiltration**, and the amount of water that is removed from the soil by **evaporation** from bare soil or by **evapotranspiration** where vegetation covers the soil. This section contains information on more than 50 techniques for measuring or estimating: (1) Hydrometeorological parameters, and (2) evaporation and evapotranspiration for water budget calculations in the vadose zone and shallow ground-water systems. Methods for measuring and estimating infiltration are covered in Section 7.1.

Hydrometeorological Data

Table 8-1 provides some general summary information on 38 techniques for measuring six major hydrometeorological parameters and identifies sections of this guide were more detailed information can be found. Precipitation is a primary input into water budget calculations, and devices for measuring precipitation fall into two main categories: (1) Manual gages (Section 8.1.1), and (2) recording gages (Section 8.1.2). Measurement of humidity (Sections 8.1.3 and 8.1.4) might be required during field work for protection of health and safety and are required with most micrometeorological methods for measuring evapotranspiration (Section 8.4). Other hydrometeorological measurements might be required for monitoring weather conditions, such as temperature (Sections 8.2.1 and 8.2.2), windspeed (Section 8.2.3), and wind direction (Section 8.2.4). Measurement or estimation of these same parameters, as well as atmospheric pressure (Section 8.2.5) and insolation or radiation measurement (Sections 8.2.6 and 8.2.7), might be required in order to quantify the evapotranspiration component of water budget studies (discussed further below). Although numerous techniques and devices have been developed for hydrometeorological measurements, most of the parameters of interest usually can be estimated for purposes of vadose zone water budget studies by using data from nearby weather stations or interpolations using hydrometeorological tables or maps. Consequently, only those methods relevant to health and safety (temperature, humidity, windspeed, and direction) are likely to be used routinely during site investigations. Table 8-1 identifies the specific hydrometeorological techniques or devices that are most commonly used for site investigations. ASTM (1986) provides guidance on determining the operational comparability of meteorological measurements.

Evaporation and Evapotranspiration

Water that reaches the earth's surface can return to the atmosphere either by evaporation from free water surfaces or bare soil, or by transpiration by plants. The term evapotranspiration (ET) specifically refers to the combined effects of evaporation and transpiration from the land surface, but also might be used loosely to refer to the combined effects of evaporation from water and soil surfaces and transpiration. ET is a critical component of vadose zone water budget calculations, and is one of the most difficult of these components to measure accurately. The numerous methods that have been developed for measuring or estimating ET can be broadly classified as water budget or balance methods and micrometeorological methods. Table 8-1 summarizes information on 10 water balance methods and 6 micrometeorological methods, and identifies specific applications for each method (water evaporation, bare soil evaporation, evapotranspiration, and transpiration). Most of these methods are too complex and time-consuming for routine site investigations.

Lysimeters (Section 8.3.1) and soil moisture monitoring (Section 8.3.2) probably are the most commonly used methods for measuring evapotranspiration where site-specific data are required. Most vadose zone hydrologic models use empirical equations (Section 8.4.1) and use data from nearby weather stations data and published maps. The physically-based Penman equation (and various methods developed as refinements and adaptations of the Penman equation) probably is the most commonly used method for estimation of evaporation and/or evapotranspiration, where some measurements of meteorological data are feasible but the more complex measurements and instrumentation of other micrometeorological methods are not feasible.

Table 8-1 Summary Information on Vadose Zone Water Budget Characterization Methods

Technique	Parameters Measured	Manual/ Automatic	S/A/R	Section	Tables
Water-Related Hydrometeorological	Measurements		Α		۰.
Sacramento Gage	Rain	Manual	± 1 mm*	8.1.1	8-2
Storage Gage	Rain	Manual		8.1.1	8-2
Automatic Wet/Dry Collectors	Rain/Snow	Either		8.1.1	8-2
Weighing Gage	Rain/Snow	Automatic	-	8.1.2	8-2
Tipping Bucket Gage	Rain	Automatic		8.1.2	8-2
Float Gage	Rain	Automatic	π	8.1.2	8-2
			S/A/R		
Sling Psychrometer	Humidity	Manual	0.1/0.5/	8.1.3	8-2
Aspirated Psychrometer	Humidity	Either	0.02/0.1/	8.1.3	8-2
Thermocouple Psychrometer	Humidity	Either	?	6.1.2	6-1, 6-3
Mechanical Hygrometers	Humidity	Either	1.0/5.0/20 to 100%	8.1.4	8-2
Dew-/Frost-Point Hygrometer	Humidity	Either	.05/0.25/	8.1.4	8-2
Dew Cell/Probes	Humidity	Automatic	0.5/2.0/10 to 100%	8.1.4	8-2
Electric Hygrometers	Humidity	Either	0.5/2.0/5.0 to 98%	8.1.4	8-2
Diffusion Hygrometers	Humidity	?	?	8.1.4	8-2
Absorption Spectra Hygrometers	Humidity	?	?	8.1.4	8-2
Other Hydrometeorological Measurements			A/R		
Liquid-in-Glass Thermometer	Temperature	Manual	+0.5°C/-40 to +60 ^b	8.2.1	
Bi-Metal Thermometer	Temperature	Either	<u> </u>	8.2.1	
Bourdon Tube Thermometer	Temperature	Either	Ħ	8.2.1	
Thermocouple	Temperature	Either	-	8.2.2	
Metallic Resistance Bulb	Temperature	Either	*	8.2.2	• <i>*</i>
Thermistor	Temperature	Either	•	8.2.2	
Cup Anemonieters	H windspeed	Fither	1.0 to 50/+0.5m/s*	8.2.3	8-2
Windmill Anemometers	V-H windspeed	Either	*	8.2.3	8-2
Pressure Anemometers	H windspeed	Manual	•	8.2.3	8-2
Hot-Wire Anemometer	V-H windspeed	Automatic		8.2.3	8-2
Acoustic Anemometer	V-H windspeed	Automatic		8.2.3	8-2
Wind Vanes	Direction	Either	$0.5 \text{ to } 50/\pm 5^{\circ}$	8.2.4	8-2
Wind Cones	Direction	Manual		8.2.4	8-2
			S/A		
Mercury Barometer	Air pressure	Manual	?	8.2.5	
Altimeter	Air pressure	Manual	$\frac{1}{2} hPa/+0.2\%$	8.2.5	
Precision Aneroid	Air pressure	Either	0.5 hPa/?	8.2.5	
	F				
Thermonile Domensor	Global rad	Automatia	$A + 0.1 to 0.5 m W/m^2$	826	8-7
Emetallo Doranometer	Global rad.	Either	$\pm 1.0 \text{ mW/cm}^2$	826	8-2
Photopolitaio Pyranometer	Global rad.	Fither	21.0 mw/cm	826	8-2
Net Dediometers	Net flow	Either	2	827	8-2
Purheliometers	Direct red	Fither	2	827	8-2
r yn reuollieuera	Direct ran.	LAUICI	÷	0.2.1	0-2

Boldface = Most commonly used instruments/methods.

S = Sensitivity = The smallest fraction of a division on a scale on which a reading can be made directly or by estimation; A = Accuracy = The closeness with which an observation approaches the true value; R = Range of relative humidity that can be measured.

*Recommended accuracy by World Meteorological Organization. Less precise measurements might be acceptable, depending on the purpose of measurements.

⁶Range and accuracy of specific thermometers can range considerably, value shown is the recommended specification in U.S. EPA (1987b).

Table 8-1 (cont.)

Technique	Parameters Measured	Manual/ Automatic	Accuracy	Section	Tables
Evapotranspiration (Water Balance	Methods)	<u> </u>			
Lysimeters	WE,SE,ET,T	Either	Moderate to high*	8.3.1	8-3
Soil Moisture Monitoring	SE,ET,T	Manual	Moderate to high	8.3.2	8-3
Water Budget Methods	WESE ET,T	Manual	Low to high	8.3.3	8-3
Evaporation Pans	WE	Manual	Moderate	8.3.4	8-3
Evaporimeter	SE	Manual	High [•]	8.3.5	
Atmometers	SE.T	Manual	Moderate	8.3.5	8-3
Chloride Tracer	SE.ET.T	Manual	Moderate	8.3.6	
Ground-Water Fluctuation	SELET	Manual	Moderate	8.3.7	8-3
Other Transpiration Methods	T	Manual	Moderate to high [*]	8.3.8	8-3
Thermal Infrared	WE,SE,ET	Either	Low to moderate	1.1.3	1-3
Evapotranspiration (Micrometeoro)	logical)				
Empirical Equations	WE,SE,ET,T	Manual	Moderate to high	8.4.1	8-3
Physically-Based Equations	WE SE ET	Either	Moderate to high	8.4.2	8-3
Mass Transfer Methods	WE,ET,T	Either	Moderate to high	8.4.3	8-3
Energy Budget Methods	WE,SE,ET,T	Either	Moderate to high	8.4.4	8-3
Profile/Gradient Method	WE,SE,ET	Either	Low to moderate	8.4.5	8-3
Eddy Correlation	WEET	Either	High	8.4.6	8-3

Boldface = Most commonly used methods.

WE = Water evaporation; SE = Bare soil evaporation; ET = Evapotranspiration; T = transpiration.

*For high accuracy, numerous measurements at different locations might be required to adequately characterize the variability of evapotranspiration.

8.1 WATER-RELATED HYDROMETEOROLOGICAL DATA

8.1.1 Precipitation (Nonrecording Gages)

Other Names Used to Describe Method: Standard gage, sacramento gage, conduit gage.

Uses at Contaminated Sites: Monitoring of site conditions during field work; measuring precipitation for water budget analysis.

Device Description: Nonrecording gages require visual observation or manual measurement to record the amount of precipitation, even though some types might involve automated handling of collected precipitation. Sacramento gage: An 8-inch diameter receiving funnel, routes precipitation into a measuring tube with a crosssectional area one-tenth that of the gage. The funnel attached to the collector both directs the precipitation into the tube and minimizes evaporation loss (Figure 8.1.1). Accumulated precipitation is measured periodically. Snow and other forms of frozen water are melted before measurement in order to give the equivalent amount of rainfall. The receiving cylinder can be clear with graduated markings for direct readings, or depth is measured using a measuring stick. Storage gages are similar to funnel gages, except that the storage container is large enough to store the seasonal catch and oil or other evaporation suppressing material is added to reduce evaporation between measurement. Automatic wet/dry precipitation collectors are specialized nonrecording instruments, where chemical and/or radioactive analysis of precipitation is required. The collector is built with a sensor, which detects the onset and cessation of precipitation, and automatically releases a lid to open and cover the collector, which prevents evaporation of the samples collected between precipitation events. Other manual gages: A wide variety of inexpensive gages, with various shapes for openings and graduated scales for measuring precipitation, are available.

Device Selection Considerations: Sacramento Gage Advantages: (1) Is inexpensive and easy to use; and (2) has no moving parts or electronic equipment to malfunction. Sacramento Gage Disadvantages: (1) Accurate characterization of precipitation events requires measurement after each precipitation event, which is difficult unless personnel are readily available to take readings at the required intervals; and (2) tend to underestimate precipitation that falls as snow. Storage Gages: Used at inaccessible sites where seasonal measurements are adequate for data needs. Automatic Precipitation Collectors: Used to collect bulk samples of precipitation for chemical analysis. Standard collectors require manual recording of the precipitation that is collected, but recent advances allow both automated recording of precipitation amounts and collection of snow and rain samples for chemical analysis (Purcell and Brown, 1991).

Frequency of Use: The Sacramento gage is the standard nonrecording gage used in the United States.

Standard Methods/Guidelines: NWS Specification No. 450.2301.

Sources for Additional Information: Brakensiek et al. (1979), Brock and Nicolaidis (1984), Lockhart (1989a), Malone (1951), National Weather Service (1972), U.S. EPA (1985), U.S. Geological Survey (1980), WMO (1971). See also, Table 8-2. Most of the general hydrology texts listed in Table 8-3 also discuss methods for measuring precipitation.



Figure 8.1.1 Typical non-recording rain gage (Kazmann, 1988, by permission).

8.1 WATER-RELATED HYDROMETEOROLOGICAL DATA

8.1.2 Precipitation (Recording Gages)

Other Names Used to Describe Devices: Weighing, Fergusson, or universal gage; tipping bucket gage; float gage.

<u>Uses at Contaminated Sites</u>: Measuring precipitation at remote sites or where accurate characterization of the amount and intensity of precipitation is required for water budget analysis.

<u>Device Description</u>: Weighing gage: A mechanical recording device is attached to a scale, which provides continuous weight measurements of precipitation that enters a cylinder gage (Figure 8.1.2a). Changes in weight are recorded on a chart recorder. Tipping bucket gage: A pair of small containers designed so that when a certain amount of rainfall (typically 0.01 inches) falls in one of the containers, it tips, and moves the other container into position to receive the next rainfall (Figure 8.1.2b). When the collection container empties into a storage container, an electrical contact is closed and the event is recorded on an electronic data logger. Float gages are cylinder gages equipped with a float and a recording device to automate measurement (used in Great Britain). Special features, which can be used with any gage, include: (1) Shields to improve collection efficiency of snow, (2) heaters to melt frozen precipitation so it will not clog the collectors or funnels of the gage, and (3) suppressants to reduce evaporation losses.

Device Selection Considerations: Weighing Gage Advantages: (1) Are very reliable; (2) equipment is readily available; and (3) measures both rain and frozen precipitation. Weighing Gage Disadvantages: (1) Manual reading of the chart recorder is required; (2) collection container usually must be emptied manually; and (3) measurements of snow might not be accurate (accuracy can be increased by shielding [Simmons and Bigelow (1990)]. Tipping Bucket Gage Advantages: (1) Data are generated electronically, which facilitates data analysis, and (2) is reliable and equipment is readily available. Tipping Bucket Gage Disadvantages: (1) Requires more maintenance than weighing gages; (2) is not accurate for measuring snowfall; and (3) requires power source for recording.

<u>Frequency of Use</u>: The weighing gage is the official precipitation measurement device of the National Weather Service. Tipping bucket gages are both readily available and widely used.

Standard Methods/Guidelines: Weighing gage: NWS Specification No. 450.2201.

Sources for Additional Information: Brakensiek et al. (1979), Brock and Nicolaidis (1984), Lockhart (1989a), Malone (1951), National Weather Service (1972), U.S. EPA (1985), U.S. Geological Survey (1980), WMO (1971). See also, Table 8-2. Most of the general hydrology texts listed in Table 8-3 also discuss methods for measuring precipitation.





Figure 8.1.2 Recording rain gages: (a) Typical weighing rain gage (Kazmann, 1988, by permission); (b) Typical tipping bucket rain gage (Dunne and Leopold, 1978, from: *Water in Environmental Planning* by Dunne and Leopold, Copyright © 1978 by W.H. Freeman and Company, reprinted with permission).

8.1 WATER-RELATED HYDROMETEOROLOGICAL DATA

8.1.3 Humidity Measurement (Psychrometers)

Other Names Used to Describe Device: Sling psychrometer, dry-bulb/wet-bulb thermometer, aspirated psychrometer, thermocouple psychrometers.

<u>Uses at Contaminated Sites</u>: Estimating effective air temperature when it is very hot; psychrometers are required for several micrometeorological evapotranspiration methods (profile, eddy correlation, mass transfer).

<u>Device Description</u>: Psychrometers operate on the principle of reduction of temperature by evaporation.* Sling pychrometer: A dry-bulb/wet-bulb thermometer (two matched mercury-in-glass thermometers mounted on a metal frame with the bulb of one covered by a moistened wick) is attached to a handle with a chain so that the thermometer can swing around to equilibrate (Figure 8.1.3). Charts are used to determine relative humidity based on the difference in temperature between the two thermometers. Readings from a static dry-bulb/wet-bulb thermometer also can be used, but are not quite as accurate. Aspirated psychrometers are dry-bulb/wet-bulb thermometers in which a motor-driven fan or blower draws air over the thermometers at a constant rate. As with the sling psychrometer, humidity is determined using charts. Thermocouple psychrometers are discussed in Section 6.1.2.

<u>Device Selection Considerations</u>: Sling psychrometers are accurate, readily available, and easy to use. Aspirated psychrometers provide greater accuracy (Table 8-1), but require a power source and involve more complex installation procedures, such as use of a radiation shield. Humidity should be monitored whenever use of protective clothing in hot temperatures creates a possibility of heat stress.

Frequency of Use: Commonly used.

Standard Methods/Guidelines: ASTM (1982, 1984a).

Sources for Additional Information: Berry et al. (1945), Lockhart (1989a), Spilhaus and Middleton (1973), U.S. EPA (1987a,b), U.S. Geological Survey (1980), Wexler (1965), WMO (1971). See also, Table 8-2.

*Note that the terms "psychrometer" and "hygrometer" might be used interchangeably in the published literature. In this guide, the term psychrometer is applied to methods involving evaporation and hygrometer to any other method of measuring humidity.





8.1 WATER-RELATED HYDROMETEOROLOGICAL DATA

8.1.4 Humidity Measurement (Hygrometers)

Other Names Used to Describe Device: Mechanical hygrometer, dew-point or frost-point hygrometer, dew cell or dew probe, electric hygrometers (resistance, capacitance, or Dunmore Cell), diffusion hygrometer, absorption spectra hygrometers (infrared, ultraviolet, alpha radiation).

<u>Uses at Contaminated Sites</u>: Estimating effective air temperature when it is very hot; hygrometers required for several micrometeorological evapotranspiration methods (profile, eddy correlation, mass transfer).

Device Description: Hygrometers* include a wide variety of instruments that measure humidity by methods other than evaporative effects on temperature (psychrometry, previous section). Mechanical hygrometer: Operates on a similar principle to a bi-metal thermometer (Section 8.1.1), except that materials with differing response to air moisture (hair, wood, and natural and synthetic fibers) are used. Mechanical hygrometers usually are read manually, but can be attached to chart recorders. Dew-point and frost-point hygrometers measure the temperature at which dew or frost condenses from the air on a cooled surface, usually a polished mirror. The temperature can be converted into vapor pressure from vapor-pressure formulations or tables. For relative humidity, the dry-bulb temperature also must be measured, and measurement of atmospheric pressure is required for calculating the mixing ratio. Dew cells operate on the principle that the equilibrium vapor pressure of a saturated solution is a function of the temperature of the solution. The dew cell consists of a temperature sensor surrounded by a wick impregnated with a saturated solution of a salt, such as lithium chloride. A control circuit maintains the solution at the temperature at which the equilibrium vapor pressure of the solution is equal to the vapor pressure of the ambient air. The output from the sensor is indicated on a dial or is recorded on a chart, which is calibrated in terms of the dew-point temperature of the ambient air. Figure 8.1.4 illustrates a typical dew cell sensor housing and transmitter. Electric hygrometers measure changes in resistance or capacitance of a thin film of hygroscopic material. Most instruments consist of a sensor and a measuring circuit with the output indicated on a meter or recorded. The response of the sensor is an empirical function of relative humidity and temperature. Diffusion hygrometers involve the diffusion of moisture through porous membranes. Absorptionspectra hygrometers use the absorption spectra of water vapor, in response to infrared, ultra-violet, or alpha radiation.

Device Selection Considerations: Mechanical hygrometers are simple and inexpensive, but the least accurate of available methods. Dew-/frost-point hygrometers are the most accurate of available methods. Dew cells are less accurate than sling psychrometers, but can be adapted for automatic data collection. Electric hygrometers are comparable to dew cells in terms of accuracy, allow automatic data collection, and have the added advantage being able to measure a somewhat wider range of relative humidity. Diffusion and absorption spectra hygrometers are very accurate but require frequent attention and are expensive to purchase and maintain.

<u>Frequency of Use</u>: Mechanical hygrometers are widely used when a high degree of accuracy is not required. Diffusion and absorption spectra hygrometers are used primarily for specialized research purposes

Standard Methods/Guidelines: ASTM (1982, 1983, 1985b).

Sources for Additional Information: Berry et al. (1945), Lockhart (1989a), Spilhaus and Middleton (1973), U.S. EPA (1987a,b), U.S. Geological Survey (1980), Wexler (1957, 1965), Wexler and Brombacher (1951), WMO (1971). See also, Table 8-2.

• Note that the terms "psychrometer" and "hygrometer" might be used interchangeably in the published literature. In this guide, the term psychrometer is applied to methods involving evaporation and hygrometer to any other method of measuring humidity.



Figure 8.1.4 A typical dew cell sensor housing and transmitter (Lockhart, 1989a).

8.2 OTHER HYDROMETEOROLOGICAL DATA

8.2.1 Air Thermometry (Manual)

Other Names Used to Describe Device: Liquid-in-glass thermometer; deformation thermometers: Bi-metallic (flat spiral, single helix, multiple helix) and filled systems/Bourdon tubes (liquid-filled, vapor-pressure systems, gas-filled systems, mercury-in-steel).

<u>Uses at Contaminated Sites</u>: Measuring air temperature for calculating evaporation and relative humidity; health and safety monitoring for potential heat or cold stress.

<u>Device Description</u>: Liquid-in-glass thermometer: Liquid in a sealed glass tube expands and contracts in response to changes in temperature, and changes in the level read from a calibrated scale. The most common liquid-in-glass thermometer is the mercury thermometer (Figure 8.2.1a), which measures to -38.9°C or -38.0°F. Other liquids can be used if extremely low temperatures must be measured (spirit thermometers, ethyl alcohol freezes at -117°C, and mercury-thallium thermometers record to -59°C). Deformation Thermometers: Metals with different coefficients of expansion (bi-metallic, Figure 8.2.1b), or filled systems in which liquid, gas, or mercury in a sealed, coiled metal tube (Bourdon tube, Figure 8.2.1c), expand and contract in response to temperature changes, which are recorded by a moving pointer or pen on a calibrated scale. The accuracy of filled-systems depends on the extent to which the differential responses of different components in the system are compensated for. The most accurate types have full compensation, others provides for compensation of the detecting element only. See section 8.1.4 for discussion of radiation shields for air temperature measurements. Bi-metallic and filled-system thermometers can be used for continuous recording of temperature changes by attaching them to a rotating drum recorder.

Device Selection Considerations: Unless required data can be obtained easily and cost-effectively with manual temperature readings, these methods are not recommended. Liquid-in-Glass Advantages: (1) Have a simple design; (2) are easy to use; (3) are inexpensive; and (4) are accurate. Liquid-in-Glass Disadvantages: (1) Are very fragile; (2) have a relatively long time constant (the time required to respond to a temperature change is relatively long). Bimetallic Advantages: Are rugged. Bimetallic Disadvantages: (1) Severe mechanical shock or vibration can cause distortion resulting in large shifts in their calibration; (2) have time constant about the same as liquid-in-glass thermometers; and (3) are less accurate and more expensive the liquid-in-glass thermometers. Filled System Advantages: (1) Fundamental simplicity allows rugged construction; and (2) bulb and detection element can be separated by some distance. Filled System Disadvantages: (1) Are sensitive to severe shock, vibration, or other forms of mechanical abuse; and (2) capillary tube is not highly flexible or convenient to handle.

<u>Frequency of Use:</u> Liquid-in-glass thermometers are commonly used for monitoring temperature conditions under which field personnel are operating.

Standard Methods/Guidelines: ASTM (annual).

Sources for Additional Information: Brock and Nicolaidis (1984), Hardy and Fisher (1972-Chapter 1), Lockhart (1989a), Meteorological Office (1956-Chapter 1), Spilhaus and Middleton (1973), National Weather Service (1975-Chapter A.9), Stevens et al. (1975), Thompson et al. (1989), U.S. Geological Survey (1980), WMO (1971-Chapter 4, 1974-Chapter 4).



Figure 8.2.1 Manual thermometers: (a) Liquid-in-glass; (b) Deformation thermometer with helical-type bimetal elements; (c) Deformation thermometer with Bourdon tube (Stevens et al., 1975).

8.2 OTHER HYDROMETEOROLOGICAL DATA

8.2.2 Air Thermometry (Electric)

Other Names Used to Describe Device: Thermocouple, wire bobbin probe, wire resistance probe, wire bobbin bulb, wire resistance bulb, thermistor.

Uses at Contaminated Sites: Measuring temperature of air, soil, and/or water (see also, Sections 1.6 and 3.5.2).

<u>Device Description</u>: A thermocouple is a circuit made of two dissimilar metals (See Figure 1.6.1). A current is produced in the circuit when the two junctions are at different temperatures. Maintaining one junction at a known temperature and exposing the other allows sensitive and accurate measurement of temperature, provided that the temperature is calibrated. The two major types of electrical-resistance thermometers are: (1) Metallic resistance thermometers, which pass an electrical current through a wire (platinum and nickel-iron being the most commonly used wires), the resistance of which is proportional to temperature; and (2) thermistors, which are glass insulated semiconductors with a negative coefficient of resistance such that electrical resistance varies sharply with changes in temperature. For all types of thermometers, measurement of ambient air temperature requires some form of shielding so that the air temperature measurements are not influenced by radiant heat. Figure 8.2.2 provides examples of 14 types of radiation shields.

Device Selection Considerations: All electrical temperature measuring devices are well suited for electronic data logging. Thermocouples or thermistors are the recommended method for temperature measurement when automatic data recording is desired. Thermocouple Advantages: (1) Can be separated a considerable distance from the measuring instrument; (2) have very rapid response time (slower in water because they have to be cased); and (3) are relatively inexpensive. Thermocouple Disadvantages: (1) Measuring instruments used with thermocouples are relatively expensive; and (2) insertion of electric leads of different metals between the thermocouple and the measuring can cause errors as a result of extraneous voltages. Resistance Thermometer Advantages: (1) Both types can be separated a considerable distance from the measuring instrument; (2) metallic resistance thermometers are more sensitive to small temperature changes than thermocouples; and (3) thermistors are less expensive that metallic resistance bulbs and even more sensitive. Resistance Thermometer Disadvantages: (1) Metallic thermometers have slightly longer response time than thermocouples to changes in temperature; and (2) thermistor's response to temperature might change with time, requiring recalibration.

Frequency of Use: Thermistors and thermocouples are most commonly used.

Standard Methods/Guidelines: ASTM (annual).

Sources for Additional Information: Brock and Nicolaidis (1984), Hardy and Fisher (1972-Chapter 1), Lockhart (1989a), Meteorological Office (1956-Chapter 1), Spilhaus and Middleton (1973), National Weather Service (1975-Chapter A.9), Stevens et al. (1975), Thompson et al. (1989), U.S. Geological Survey (1980), WMO (1971-Chapter 4, 1974-Chapter 4).



Figure 8.2.2 Examples of various types of radiation shields for air temperature measurements (Lockhart, 1989a, after McKay and McTaggart-Cowan, 1977).

8.2 OTHER HYDROMETEOROLOGICAL DATA

8.2.3 Wind Speed

Other Names Used to Describe Device: Cup/bridled cup, windmill (air meter, propeller/aerovane), pressure anemometers (hand-held/pith-ball wind meter, Dines), hot-wire anemometer; acoustic/sonic anemometer, contact anemometer, condenser-discharge anemometer.

<u>Uses at Contaminated Sites</u>: Evaluating transport of atmospheric pollutants or dust from disposal sites; evaluating evaporation rates; evaluating wind chill for field work in the winter.

Device Description: Numerous specific types of anemometers have been developed to measure wind speed. Six major types of anemometers are described here. Cup anemometers consist of three or four cups mounted around a vertical axis on radial arms at equal angles, which allow the anemometer to be equally responsive to wind in any direction (Figure 8.2.3). The vertical shaft transfers the motion of the cups either to a counter or to a generator for electronic recording. Windmill anemometers include: (1) Propeller anemometers with helicoidal vanes, which rotate about an axis and drive a miniature generator with an electrical output that is proportional to the wind speed, and (2) air meters with flat vanes that records the number of linear feet (or meters) of air that has passed the instrument during its exposure. Propeller anemometers are usually combined with a wind vane to maintain an orientation directly into the wind, but sometimes are built with three propellers oriented at right angles to each other to measure horizontal and vertical components separately. Manually operated pressure anemometers consist of a thin tube open at one end. The pressure change produced by air moving across the opening is proportional to the wind speed. In a variant of this, a pith ball rises in a graduate tube. Hot-wire and acoustic (or sonic) anemometers are very precise instruments, which measure velocity by measuring the change in resistance of a heated tungsten wire and accurately measuring sound velocity, respectively. A contact anemometer actuates an electrical contact at a rate that depends on windspeed. The number of contacts during a given time is indicated by the number of flashes of a lamp or sounds of a buzzer. A condenserdischarge anemometer is a type of contact device with an electrical circuit that indicates average windspeed.

<u>Device Selection Considerations</u>: Propeller and cup anemometers are the most common types because they are rugged, reliable, and accurate to within a few percent or less. Both are well suited for electronic data logging. Propeller-type anemometers can measure wind speeds up to 200 miles per hour; cup-type anemometers measure up to 100 miles per hour and can be constructed to be extremely sensitive to slight changes in speed. Pressure anemometers are not recommended unless manual measurement is acceptable. Hot-wire and acoustic anemometers are for specialized applications where accurate measurement of turbulence is required.

Frequency of Use: Both propeller and cup-type anemometers are widely used.

Standard Methods/Guidelines: ASTM (1985a, 1990).

Sources for Additional Information: Hardy and Fisher (1972-Chapter 3), Lockhart (1989a), Meteorological Office (1956-Chapter 5), National Weather Service (1975-Chapter A10), Spilhaus and Middleton (1973-Chapter 6), Thompson et al. (1989), U.S. EPA (1987a,b), U.S. Geological Survey (1980). See also, Table 8-2.



Figure 8.2.3 Portable hand cup anemometer for measuring windspeed (Cameron et al., 1966).

8.2 OTHER HYDROMETEOROLOGICAL DATA

8.2.4 Wind Direction

Other Names Used to Describe Method: Wind cone/sleeve/sock, vanes (flat-plate, aerodynamic-shaped, splayed, bivanes).

Uses at Contaminated Sites: Assessing possible directions of air-borne contaminant transport and deposition.

<u>Method Description</u>: Wind direction can be determined visually by observing the direction of movement of any freely moving substance or object, such as smoke or ribbons attached to poles. Wind cones are made of a tapered fabric sleeve, which is shaped like a truncated cone and pivoted to a standard at its larger end. Various types of wind vanes also can serve as indicators of wind direction. A flat-plate vane is mounted on a horizontal shaft, which is attached to a vertical bearing shaft that is free to rotate (Figure 8.2.4). Aerodynamic-shaped vanes use an airfoil section instead of a flat plate, and are usually heavier than the flat plate type. Splayed vanes have two flat plates joined at a small angle at the end of the horizontal shaft, and react to small changes in the wind somewhat better than flat-plate or aerodynamic vanes. Bivanes consist of two light-weight airfoil sections mounted orthogonally on the end of a counter-balanced rod, which is free to rotate in the horizontal and vertical planes and is used in turbulence studies to record horizontal and vertical components of wind. Wind roses can be developed from manual recording of wind direction at specified intervals or automatic recorders attached to wind vanes. The frequency with which the wind blows in various directions can be useful information in designing soil sampling plans where a point source has released contaminants to the air that have been deposited at downwind locations.

<u>Method Selection Considerations</u>: Some kind of wind direction indicator should be used any time site activities could result in release of contaminants to the air.

<u>Frequency of Use</u>: Commonly used for health and safety purposes; less common for obtaining hydrometeorological applications.

Standard Methods/Guidelines: ASTM (1985a).

Sources for Additional Information: Hardy and Fisher (1972-Chapter 3), Lockhart (1989a), Meteorological Office (1956-Chapter 5), National Weather Service (1975-Chapter A10), Spilhaus and Middleton (1973-Chapter 6), Thompson et al. (1989), U.S. EPA (1987a,b), U.S. Geological Survey (1980). See also, Table 8-2.





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Figure 8.2.4 Sample of dynamic response of some wind vanes (Lockhart, 1989a).

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8.2 OTHER HYDROMETEOROLOGICAL DATA

8.2.5 Atmospheric Pressure

Other Names Used to Describe Method: Mercury barometers (Fortin-type, fixed-cistern type), aneroid barometer, altimeter.

<u>Uses at Contaminated Sites</u>: Interpreting ground-water level measurements; required for several methods of measuring or estimating evapotranspiration using micrometeorological method (Section 8.4); required for calculations involving humidity measurement (Sections 8.1.3 and 8.1.4); estimating altitude in remote locations.

Method Description: There are two major types of instruments for measuring atmospheric pressure. Mercury barometers use changes in the level of mercury in a container to measure changes in atmospheric pressure. The Fortin-type mercury barometer is used by the National Weather Service as the official station pressure instrument. A cistern containing mercury has a pointer made of noncorrodible materials, such as ivory or stainless steel, projected down from the roof. The level of mercury within the cistern is raised or lowered by turning a thumb screw beneath the cistern, until it just touches the tip of the pointer (called the ivory point, index point, or zero point). Pressure is read from mercury in a graduated column connected to the cistern that can be read to a thousandth of an inch or a tenth of a millibar with a vernier on the scale. Aneroid barometers measure pressure by the response of a capsule that is practically evacuated of gas. The response can be measured either by deflection of a spring connected to the cell, by the change in curvature of a Bourdon tube, or by a change in natural resonant frequency. The barometer must be temperature compensated at a given pressure level by adjusting the residual gas in the aneroid or by a bimetallic-link arrangement. Altimeters are aneroid barometers that have a pointer and a dial calibrated for elevation or pressure readings (Figure 8.2.5). Precision aneroids can be of the direct-reading kind, similar to altimeters, but are designed for more accurate measurements. A relatively recent development is the accurate digital-readout precision aneroids, which use electronic indicators rather than mechanical linkages. Sensor types used in these instruments can be a fused quartz Bourdon tube (quartz barometer), an aneroid capsule with which the natural frequency as related to pressure is measured (vibrating diaphragm barometer), or the conventional aneroid capsule in which spring deflection is measured.

<u>Method Selection Considerations</u>: Fortin barometers are very accurate (can be read to a thousandth of an inch), but require permanent installation. Aneroid barometers have the main advantage of being portable. Disadvantages include requirements for periodic calibration against mercury barometers, and the requirement for temperature compensation.

Frequency of Use: Not commonly used at contaminated sites.

Standard Methods/Guidelines: ASTM (1984c).

Sources for Additional Information: Brock and Nicolaidis (1984), Lockhart (1989a), U.S. Geological Survey (1980), U.S. Weather Bureau (1963b), National Weather Service (1975-Chapter 8), WMO (1971).





8.2 OTHER HYDROMETEOROLOGICAL DATA

8.2.6 Solar Radiation (Pyranometers)

Other Names Used to Describe Methods: Thermopile, photovoltaic, bimetallic (Robitzsch-type, actinograph), thermo-electric pyranometers, incident solar radiation meter, solarimeter.

<u>Uses at Contaminated Sites</u>: Global radiation data are needed for some empirical and physically-based equations for estimation of evaporation and evapotranspiration (Sections 8.4.1 and 8.4.2).

<u>Device Description</u>: A pyranometer measures global solar radiation (direct plus diffuse radiation falling on a horizontal surface, see Figure 8.2.6a), and is the most commonly measured type of radiation. It does not measure terrestrial or atmospheric radiation. Most pyranometers incorporate a sensor that responds to the temperature difference caused by differential absorption of radiation of a black surface and a white surface (Figure 8.2.6b). The most commonly used temperature sensor is a thermopile, but bimetallic sensors also can be used. Photovoltaic pyranometers use silicon cells that respond to solar radiation by generating an electric current, which is proportional to the amount energy hitting the cell. WMO (1971) has established criteria for classification of pyranometers according to physical response characteristics, with 1st class being the most sensitive and 3rd class being the least sensitive. A net pyranometer measures the net upward and downward solar radiation flux through a horizontal surface.

Device Selection Considerations: Thermopile Advantages: (1) A variety of instruments of this type have been developed and are commercially available; (2) are the most accurate and responsive of available instruments (most are 1st or 2nd class); (3) the thermopile pyranometer is the standard instrument to use if direct measurements are required; and (4) can be readily configured for output to an electronic recording device. Bimetallic Advantages: (1) Are simple; (2) can be attached to a chart recorder for continuous recording; and (3) are suitable for measurements in which daily or longer interval data are acceptable. Bimetallic Disadvantages: (1) Are relatively inaccurate (3rd class) compared to thermopile pyranometers; (2) have relatively slow response time; and (3) require use of temperature-correction factor or some temperature compensation mechanism. Photovoltaic Advantages: (1) Are simple and inexpensive; (2) have a nearly instantaneous response; (3) have high current output, which can be used for automatic data recording; and (4) use can be acceptable as long as integration periods are 1 day or longer. Photovoltaic Disadvantages: Least accurate of available methods due to variations in sensitivity to different wavelengths.

Frequency of Use: Rare. Often estimated from nearby meteorologic station or from charts or maps.

Standard Methods/Guidelines: --

Sources for Additional Information: Brock and Nicolaidis (1984), Carter et al. (1977), Coulson (1975), Latimer (1972), Lockhart (1989a), Monteith (1972), Norris (1974), Selcuk and Yellott (1962), Thompson et al. (1989), U.S. Army (1975), U.S. Geological Survey (1980), WMO (1971). See also, Table 8-2.







(b)

Figure 8.2.6 Measurement of solar radiation: (a) Kinds of insolation and types of measuring instruments (U.S. Geological Survey, 1980); (b) Features of a typical pyranometer (Carter et al., 1977).

8.2 OTHER HYDROMETEOROLOGICAL DATA

8.2.7 Solar Radiation (Other Radiometers)

Other Names Used to Describe Method: Net radiometer/pyrradiometer, pyrheliometer (Angstrom electrical compensating, silver-disk, absolute, operational)

<u>Uses at Contaminated Sites</u>: Measuring net radiation flux for energy budget measurements of evapotranspiration (Section 8.4.4).

<u>Method Description</u>: Other radiometers measure different types of radiation. **Pyrradiometers** measure total radiation falling on a horizontal surface (combined solar, atmospheric and terrestrial radiation), and are similar in design to pyranometers (Section 8.2.6). **Net pyrradiometers** or **radiometers** are designed to measure the difference between downward and upward total radiation. Most commercially available net radiometers are made with a small disc-shaped thermopile covered by polyethylene hemispheres. **Pyrheliometers** measure the intensity of direct solar radiation and normal incidence, and are mounted in trackers that keep the devices pointed toward the sun as the traverses from east to west (Figure 8.2.6a and 8.2.7).

<u>Method Selection Considerations</u>: Unlikely to be used unless an energy budget method for computing evapotranspiration is used.

Frequency of Use: Uncommon for site characterization. Net radiometers are sometimes used in air-pollution related programs.

Standard Methods/Guidelines: ASTM (1984b).

Sources for Additional Information: Brock and Nicolaidis (1984), Carter et al. (1977), Coulson (1975), Latimer (1972), Lockhart (1989a), Monteith (1972), Norris (1974), Selcuk and Yellott (1962), Thompson et al. (1989), U.S. Army (1975), U.S. Geological Survey (1980), WMO (1971). See also, Table 8-2.



Figure 8.2.7 Features of a typical pyrheliometer and tracking mount (Carter et al., 1977).

8.3 EVAPOTRANSPIRATION (WATER BALANCE METHODS)

8.3.1 Lysimeters

Other Names Used to Describe Method: Pan/filled-in lysimeters (nonweighable, weighing, hydraulic/floating), monolith/soil block lysimeter, monolith/soil block evapotranspirimeter, microlysimeter.

<u>Uses at Contaminated Sites</u>: Measuring evaporation from vegetated soil (pan lysimeter) or unvegetated soil (microlysimeter), in order to separate out the transpiration component of evapotranspiration (ET).

Method/Device Description: A lysimeter consists of a block os soil, usually planted with some vegetation that is enclosed in a container, which isolates the lysimeter hydrologically from its surroundings. Lysimeters used for sampling soil solutions are discussed in Section 9.3.1 (Free-Drainage Samplers). There are three main types of filled-in lysimeters, in which disturbed soil is used for measuring ET: (1) Nonweighing lysimeters (Figure 8.3.1a); (2) hydraulic or floating lysimeters, which rest on rubber bags or other water-filled tubing or bolsters that allow recording of changes in pressure in response to changes in weight (Figure 8.3.1b); and (3) weighing lysimeters, in which changes in moisture contents are measured by changes in the weight of the soil block (Figure 8.3.1c). A typical pan lysimeter is 1 meter in diameter (range from 0.1 to 10 square meters) and range from 0.5 to 3 meters deep. Soil and vegetation representative of the area are placed in the lysimeter with the surface level the same as the surrounding soil. Monolith lysimeters are constructed of undisturbed soil. In nonweighing lysimeters, changes in soil moisture are determined by various soil moisture determination methods, such as neutron-moisture logging, gamma-ray transmission, electrical resistance blocks, or tensiometers (see Section 6.3). Weighing and hydraulic lysimeters measure changes in moisture content by recording changes in the total weight of the lysimeter over time with a sensitive scale or transducer. Most lysimeters record ET over relatively large areas. An exception is the microlysimeter, where a thin-walled cylinder is pushed into the soil, the sample is removed, sealed at the bottom, and weighed. The sample is replaced in the original hole to subject it to the same evaporative conditions as the soil, and is removed periodically for reweighing (Figure 8.3.1d).

<u>Method Device Selection Considerations</u>: Pan Lysimeter Advantages: (1) Probably are the most accurate of the water balance methods; (2) allow measurement of ET from a medium or large area; and (3) cost is moderate to low. Pan Lysimeter Disadvantages: (1) Are relatively complicated to install; and (2) must be surrounded by a considerable area of the same vegetation to avoid horizontal diversion in energy for ET. Microlysimeter Advantages: (1) Measure evaporation under a wide range of soil moisture conditions; and (2) are inexpensive and easy to use. Microlysimeter Disadvantages: Have small areal coverage.

Frequency of Use: Lysimeters are a commonly used method, if field measurement of ET is required.

Standard Methods/Guidelines: Pan lysimeter: Aboukhaled et al. (1982); Microlysimeter: Boast (1986).

Sources for Additional Information: Dunne and Leopold (1978), Sharma (1985), Thompson et al. (1989), U.S. Geological Survey (1982), Veihmeyer (1964). See also, Table 8-3.



(a)



Figure 8.3.1 Lysimetric methods: (a) Nonweighing, drainage type; (b) Weighing float type; (c) Spring-balance weighing type (Dunne and Leopold, 1978, from: *Water in Environmental Planning* by Dunne and Leopold, Copyright © 1978 by W.H. Freeman and Company, reprinted with permission); (d) Procedure for microlysimeter determination of evaporation (Boast, 1986, after Boast and Robertson, 1982, by permission).

8.3 EVAPOTRANSPIRATION (WATER BALANCE METHODS)

8.3.2 Soil Moisture Budget

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Measuring evapotranspiration (ET).

<u>Method Description</u>: Soil moisture content is measured over the entire root zone using one or more methods described in Section 6.3, before and after irrigation events. Assuming that irrigation brings the soil to field capacity, the initial moisture content after irrigation will be the available water capacity of the root zone. The evapotranspiration rate is the difference in moisture content between the two sampling periods divided by the time interval.

<u>Method Selection Considerations</u>: Advantages: Relatively simple method with the added advantage that soil moisture monitoring is often required for other objectives (see Section 6). Disadvantages: (1) Requires uniform soil type and texture and a water table deep enough that it does not influence the soil root zone; (2) precipitation events will disrupt the method; and (3) calculation of ET requires adjustments (modulation) to account for the fact that ET rates might change as soil moisture decreases.⁴

Frequency of Use: Probably the oldest and most commonly used method for determining ET.

Standard Methods/Guidelines: --

Sources for Additional Information: Thompson et al. (1989), U.S. Geological Survey (1982), Veihmeyer (1964). See also, Table 8-3.

*There is not universal agreement on the need for such corrections (see, for example, Veihmeyer and Hendrickson, 1955), although Gray (1973) reviews some of the literature on this question and recommends that "modulated" values of ET be used when doing soil moisture budget calculations. See also, references identified in Table 8-3.

8.3 EVAPOTRANSPIRATION (WATER BALANCE METHODS)

8.3.3 Water Budget Methods

Other Names Used to Describe Method: River basin water balance, inflow-outflow measurement, integration method.

Uses at Contaminated Sites: Estimating evaporation and evapotranspiration (ET).

<u>Method Description</u>: Inflow-outflow method: All inflows (precipitation), outflows (surface runoff, ground water leaving basin), and changes in storage in a watershed, are measured or estimated except for ET. ET is calculated using a water-balance equation. Figure 8.3.3 illustrates the components of the water balance equation. Figure 8.4.1 compares evaporation from a lake in Canada, which was computed using a water budget, to six other methods. Integration method: Evaporation and ET for an area is calculated by the summation of the products of ET for each crop times its area, plus the ET of natural vegetation times its areas, plus water-surface evaporation times its surface areas, plus evaporation from bare land times its areas. This method requires knowledge of unit ET and the areas of various classes of agricultural crops, natural vegetation, bare land, and water surfaces. Often this can be done using sequential remote sensing data (satellite, airphotos) to identify crop/vegetation patterns (see Raymond and Rezin, 1989, for a recent example of this approach).

<u>Method Selection Considerations</u>: Advantages: Water budget methods can be manageable to difficult, with moderate to low cost. Disadvantages: (1) Small errors in measuring or estimating various components of the water-balance equation (such as deep percolation) can cumulatively result in a large error in the calculated ET value; (2) suitable for application to a specific site only if ET at the site can be assumed to be close to the average ET for the watershed or area of interest; and (3) use of water budgets to calculate evaporation from lakes is not recommended for time periods of less than 1 month in duration if the estimate is expected to be within plus or minus 5 percent of the actual amount (Gray, 1973).

Frequency of Use: Commonly used in hydrologic studies; rarely used at the site-specific level.

Standard Methods/Guidelines: --

Sources for Additional Information: Bras (1990), Dunne and Leopold (1978), Rosenberg et al. (1983), Sharma (1985), Thompson et al. (1989), U.S. Geological Survey (1982), Veihmeyer (1964). See also, Table 8-3.



Figure 8.3.3 Water balance equation and schematic diagram for a hillside or a small catchment (Dunne and Leopold, 1978). P = precipitation; I = interception; AET = actual evapotranspiration; OF = overland flow; ΔSM = change in soil moisture; ΔGWS = change in ground-water storage; GWR = ground-water outflow. Solving for ET requires measurement or estimation of other elements in the equation.

8.3 EVAPOTRANSPIRATION (WATER BALANCE METHODS)

8.3.4 Evaporation Pans

Other Names Used to Describe Method: Class A land pan, U.S Bureau of Plant Industry sunken pan, Colorado sunken pan, U.S. Geological Survey floating pan, insulated evaporation pan.

<u>Uses at Contaminated Sites</u>: Estimating of evaporation from water impoundment surfaces; can also be used to indirectly estimate potential evapotranspiration (ET) (Veihmeyer, 1964).

<u>Method Description</u>: The standard U.S. Weather Service Class A pan is built of unpainted galvanized iron. It is 4 feet in diameter, 10 inches deeps, and mounted 12 inches above the ground on a wooden frame (Figure 8.3.4a). The rate of evaporation of water from the pan is measured. Precipitation also must be measured to correct for additions to the pan. A pan coefficient is used (generally from .70 to .75) for large bodies of water, to estimate actual evaporation from the water body of interest (Figure 8.4.1 illustrates the tendency of Class-A pans to overestimate actual evaporation). The insulated evaporation pan is constructed of fiberglass with 8 centimeters of freon-blown polyethylene (Figure 8.3.4b). The insulation reduces effects of climate and season on variability of coefficients used to calculate actual evaporation. Other commonly used types of evaporation pans include the U.S. Bureau of Plant Industry sunken pan, Colorado sunken pan, and U.S. Geological Survey floating pan.

<u>Method Selection Considerations</u>: Advantages: (1) The Class A pan is the standard method for measuring evaporation; (2) data on pan evaporation for the vicinity of a site in question might be published or available; and (3) insulated pans allow use of standard coefficients. Disadvantages: (1) Several years of data are required to characterize seasonal and annual variations in evaporation; (2) use of incorrect pan coefficient can bias results; (3) coefficients measured using noninsulated evaporation pans can vary with location, climate, or season; (4) cannot be used when temperature is below freezing; and (5) sunken pans are difficult to install and maintain, they tend to collect trash, leaks are hard to detect, and it is difficult to evaluate heat loss from the pan to the surround soil. Floating pans probably give the best estimates of lake evaporation (see Figure 8.4.1), but are not widely used due to operational difficulties (inaccessibility and water splashing into or out of the pan).

Frequency of Use: Uncommon for site specific field measurement. Other methods usually are available for estimating evaporation. Pan evaporation data are commonly used to estimate potential ET.

Standard Methods/Guidelines: Class A pan: National Weather Service (1972). Insulated pan: U.S. Geological Survey (1982).

Sources for Additional Information: Dunne and Leopold (1978), Thompson et al. (1989), U.S. Geological Survey (1982), Veihmeyer (1964). See also, Table 8-3.



(a)



(b)

Figure 8.3.4 Evaporation Pans: (a) U.S. Weather Bureau Class A land pan (after Veihmeyer, 1964); (b) Cross section of National Weather Service insulated evaporation pan (U.S. Geological Survey, 1982).

8.3 EVAPOTRANSPIRATION (WATER BALANCE METHODS)

8.3.5 Evaporimeters and Atmometers

Other Names Used to Describe Method: First-stage evaporimeter, Piche/Bellani/Livingston/Wilde atmometer.

<u>Uses at Contaminated Sites</u>: Evaporimeter: Measuring evaporation from unvegetated soil; Atmometers: Measuring latent evaporation (mainly a measure of the drying power of the air).

<u>Method/Device Description</u>: Evaporimeter: A flat, soil-covered tray 0.1 meters in area is connected to a constant suction water supply (Figure 8.3.5a). The rate of water loss from the water supply equals the evaporation rate. Atmometer: A water-filled glass tube that has an open end through which water evaporates from a filter paper (Piche type) or porous plate (Bellani type, Figure 8.3.5b). The tube supplying water is graduated to read evaporation in millimeters. Atmometer measurements require different conversion factors related to evaporation rate and location to estimate evaporation from water bodies.

Method Selection Considerations: Evaporimeter Advantages: Are relatively simple and easy to use. Evaporimeter Disadvantages: (1) Only measure evaporation during the stage when evaporation equals potential evaporation; and (2) have small areal coverage. Atmometer Advantages: (1) Are inexpensive; (2) are portable and easily maintained and installed; (3) are representative of conditions affecting moisture loss from plants; and (4) require a small amount of water to operate. Atmometer Disadvantages (1) Value for estimating evaporation loss from water bodies is questionable because they are more responsive to windspeed than radiant energy; (2) observations are difficult to interpret; (3) Class-A pans are better for estimating evaporation from lakes; and (4) cannot be used when temperature is below freezing.

<u>Frequency of Use</u>: Evaporimeters: Uncommon. Generally measurement or estimates of total evapotranspiration will meet the requirements for most water budget calculations. Atmometers commonly are used in agricultural studies but their use has not been reported at contaminated sites.

Standard Methods/Guidelines: --

Sources for Additional Information: Evaporimeters: Adams et al. (1976), Arkin et al. (1974), Boast (1986). Atmometers: U.S. Geological Survey (1982), Veihmeyer (1964). See also, Table 8-3.



Figure 8.3.5 Evaporation measurement instruments: (a) Top view and cross section of the first-stage evaporimeter tray (Boast, 1986, after Arkin et al., 1974, by permission); (b) Set of black-and-white Livingston atmometers (after Veihmeyer, 1964).

8.3 EVAPOTRANSPIRATION (WATER BALANCE METHODS)

8.3.6 Chloride Tracer

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Indirectly estimating evapotranspiration (ET).

<u>Method Description</u>: The chloride content of precipitation and shallow ground water samples is measured at intervals to obtain average chloride concentrations of the precipitation and ground water. Annual ET is calculated by multiplying the ratio of chloride concentration in precipitation to chloride in ground-water times the long-term average precipitation.

<u>Method Selection Considerations</u>: The following site conditions need to apply if this method is to be used: (1) There is a shallow water table; (2) chloride in the ground water comes only from precipitation; and (3) runoff is negligible. Laboratory analysis of samples is required and collection of precipitation samples results in moderate to high cost.

Frequency of Use: Uncommon.

Standard Methods/Guidelines: --

Sources for Additional Information: Sharma (1985), Thompson et al. (1989).

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8.3 EVAPOTRANSPIRATION (WATER BALANCE METHODS)

8.3.7 Ground-Water Fluctuation

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Indirectly estimating evapotranspiration (ET).

<u>Method Description</u>: Aquifer storage values are measured or estimated, and continuous measurement of waterlevel fluctuations are continuously measured or measurements are taken at sufficiently close intervals to plot diurnal fluctuations in ground-water level (Figure 8.3.7). Estimation of average ET rates requires continuing measurements over months. A variant of this approach in floodplain areas is to analyze diurnal fluctuations in stream hydrographs to estimate daily ET rates (Reigner, 1966), or based flow recession curves for monthly estimates of ET (Langbein, 1942).

<u>Method Selection Considerations</u>: This method requires; (1) A shallow water-table, (2) uniform coarse or medium soil texture that results in measurable, diurnal fluctuations in water table in response to ET, and (3) limited precipitation unless precipitation is accurately measured as well. Where conditions are suitable, the cost is moderate to low.

Frequency of Use: Uncommon.

Standard Methods/Guidelines: Davis and DeWiest (1966).

Sources for Additional Information: Thompson et al. (1989), U.S. Geological Survey (1982), Veihmeyer (1964). See also, Table 8-3.



Figure 8.3.7 Estimation of evapotranspiration by phreatophytes from daily water-level fluctuations in a water well (Davis and DeWiest, 1966, reprinted by permission of John Wiley & Sons, Inc. from *Hydrogeology* by S.N. Davis and R.J.M. DeWiest, Copyright © 1966).

8.3 EVAPOTRANSPIRATION (WATER BALANCE METHODS)

8.3.8 Other Transpiration Methods

Other Names Used to Describe Method: Enclosures, physico-biological methods, heat-pulse method, radioisotopes.

<u>Uses at Contaminated Sites</u>: Directly or indirectly measuring the transpiration component of evapotranspiration is not likely to be required.

<u>Method Description</u>: The transpiration component of evapotranspiration can be measured or estimated by a number of the methods discussed elsewhere in this section: Lysimeters (Section 8.3.1, considered one of the best approach), soil moisture depletion (Section 8.3.2), mass transfer methods (Section 8.4.3), and energy balance methods (Section 8.4.4). Other field methods for indirect estimation of transpiration include: (1) Enclosures, in which changes in air moisture resulting from transpiration are measured; (2) heat-pulse methods, where plants with woody stems are heated quickly and the rate of ascent of the heated sap is timed; and (3) injecting radioisotopes into trees and tracing their movement through the plant (see Section 4.4.5 for additional information on radioisotope tracers). Methods for direct measurement of transpiration (such as the use of phytometers, photometers, porometers, thermocouple psychrometry, and corona analysis) is generally are done in the laboratory.

<u>Method Selection Considerations</u>: If transpiration needs to be estimated, lysimeter or soil moisture depletion methods probably are the best for use with water budget studies.

Frequency of Use: Rare.

Standard Methods/Guidelines: --

Sources for Additional Information: See Table 8-3.

8.4 EVAPOTRANSPIRATION (MICROMETEOROLOGICAL METHODS)

8.4.1 Empirical Equations

Other Names Used to Describe Method: Equations are often identified by the names of individuals who developed the equation.

Uses at Contaminated Sites: Estimating evaporation and evapotranspiration (ET).

Method Description: Evapotranspiration: Numerous empirical equations have been developed that allow estimation of potential evapotranspiration (PET) using climatic data, which can be available from nearby weather stations or charts and maps. Once PET is known, empirical factors based primarily on the type of vegetation are used to estimate actual evapotranspiration (AET). Three of the most commonly used empirical methods are described here. Thornthwaite: This equation requires data on mean monthly air temperature. Latitude, month, and average monthly daylight are required to determine adjustment factors to take into account the total numbers of days and hours available for ET. The main advantage of this equation is that it allows general estimates in areas where climatic records and ET data are limited. Blaney-Criddle: This equation requires mean monthly temperature, monthly percentage of daylight hours per year, and an empirical coefficient for the month, which depends on the crop. A modified equation accounts for changes in the sun's zenith angle to correct for reduced power of the sun's rays during winter, allowing use of a single empirical coefficient for crop/vegetation type. Jensen-Haise: This equation requires mean air temperature, solar radiation, and the saturated vapor pressures at the mean maximum and mean minimum temperature for the warmest month of the year. Numerous other empirical equations have been developed (Table 8.4.1 shows eight of these equations), but the above mentioned ones are the most commonly used equations. Evaporation equations: A number of empirical equations have been developed for estimating lake evaporation. Most are based on simple aerodynamic equations, which require measurement or estimation of: (1) Windspeed, (2) vapor pressure of saturated air at the temperature of the water surface, (3) actual vapor pressure of air at some height above the water surface, and (4) empirical constants appropriate to the type of water body. Table 8-3 identifies a number of references that review and present empirical evaporation equations. Figure 8.4.1 shows calculations of evaporation from a lake using three empirical formulas (Nordenson-Kohler-Fox, Lake Hefner "upwind formula," and Meyer formula) with four other methods (Class-A and floating pan, water budget, energy budget, and Penman formula). It is clear from this figure that empirical formulas can yield good results if the appropriate one is used, but can be very far off if the wrong formula is used.

Method Selection Considerations: Evapotranspiration Equation Advantages: (1) Is best for developing monthly, seasonal, or annual consumptive water use values; and (2) is very inexpensive if input data can be obtained from existing meteorological records. Evapotranspiration Equation Disadvantages: (1) Calculations might not be very accurate if site conditions are not typical of conditions upon which the equation is based (use of several equations and comparing the results can be useful for developing an estimated range); (2) should not be used to estimate short-term (hours to days) variations in ET because no allowance is made for variation in wind and relative humidity; and (3) equations tend to overestimate water use during vegetation emergence and underestimate water use for midseason, unless appropriate crop factors are used (such as Blaney-Criddle method). Thornthwaite: Works best in the central and eastern United States for sod with high moisture content in areas with limited advection; is inaccurate if short-term (less than one-month) data are used. Blaney-Criddle: Is widely used in the western United States; requires empirical coefficient for crop or vegetation type (already available for many crops and vegetation types). Jensen-Haise: Was developed for use with irrigated crops in the western United States. Evaporation Equation Advantages: Is very simple and allow estimates from standard meteorological data. Evaporation Equation Disadvantages: (1) Most equations of this type require measurement of the surface temperature of the body of water, which is difficult to obtain; (2) if mean air temperature is used instead, the failure to account for effects of advected energy to the lake on evaporation might cause considerable error because small errors in temperature induce large errors in the calculations; (3) measurement of wind speed and vapor pressure must be taken at heights specified in the equation; and (4) results will be inaccurate if the characteristics of the water body are not similar to the water body for which the empirical constants were developed.

Frequency of Use: All equations are commonly used. See method selection considerations for geographic limitations.

Name	Date	Period for U	Unit for <i>U</i>	Equation	
Hedke (Harding et al., 1930) Lowry-Johnson (1942)	1930 1942	Annual Annual	Fcet Fect	U = k I I U = 0.000156 I I + 0.8	(11-20) (11-21)
Blaney-Morin (1942)	1942	m months	Inches	$U = k \sum_{1}^{m} pt(114-h)$	(11-22)
Thornthwaite and Wilm (1944)	1944	Monthly	Centimeters	$U = 1.6 \left(\frac{10t}{TE}\right)^{a}$	(11-23)
Penman (1948)	1948	Daily	Millimeters	$U = \frac{AH - 0.27E}{A - 0.27}$	(11-24)
				where $E = 0.35(e_a - e_d)(1 + 0.0098w_2)$ $H = R(1 - r)(0.18 + 0.55S) - B(0.56 - 0.092e_d^{0.5})(0.10 + 0.90S)$	
Blaney-Criddle (1950)	1950	m months	Inches	$U = k \sum_{l=1}^{m} pl = kl^{r} \text{ where } F = \sum_{l=1}^{m} pl$	(11-25)
Halkias-Veihmeyer-Hendrickson (1955)	1955	Monthly	Inches	U = SD	(11-27)
Hargreaves (1956)	1956	m months	Inches	$U = \sum_{1}^{m} kd(0.38 - 0.0038h)(t - 32)$	(11-26)

NOTATION

A = slope of saturated-vapor-pressure curve of air at absolute temperature in °F, or de_a/dt in nm Hg/°F (Fig. 11-8)

B = a coefficient depending on temperature (Table 11-7)

D = difference in evaporation between white and black atmometers in cm³

d = monthly daytime coefficient dependent upon latitude (Table 11-9)

 e_{α} = saturation vapor pressure at mean air temperature in mm Hg (Fig. 11-7) e_{d} = saturation vapor pressure at mean dew point (i.e., actual vapor pressure in the air) in mm Hg, being equal to e_{α} multiplied by relative humidity in per cent E = daily evaporation in mm

- a bar evaluation in min
 b = mean monthly relative humidity at noon, in Eq. (11-26), or annual mean relative humidity in per cent, in Eq. (11-22)
 II = accumulated degree-days above minimum growing temperature for growing season, in Eq. (11-20); or accumulated degree-days of maximum daily temperature above 32°F for growing season, in Eq. (11-21); or daily heat budget at surface in mm of water, in Eq. (11-24)
- k =annual, scasonal, or monthly consumptive use coefficient p =per cent of daytime hours of the year, occurring during the period, divided by 100 (Table 11-4) r =estimated percentage of reflecting surface

r = estimated percentage of reflecting surface R = mean monthly extraterrestrial radiation in mm of water evaporated per day (Table 11-6) $S = \text{estimated ratio of actual duration of bright sunshine to maximum possible duration of bright sunshine; or slope of regression line between D and U in Eq. (11-27)$ $<math>TE = \text{Thornthwaite's temperature-efficiency index, being equal to the sum of 12 monthly values of heat index i = (t/5)^{1.514}$, where t is mean monthly temperature in °C t = mean monthly temperature in °F, in Eqs. (11-22), (11-25), and (11-26), or in °C in Eq. (11-23) U = evapotranspiration or consumptive use for given period

 w_2 = mean wind velocity at 2 m above the ground in miles/day, or equal to w_1 (log 6.6/log h), where w_1 is measured wind velocity in miles/day at height h in ft

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Figure 8.4.1 Comparison of 1961 cumulative measured and computed evaporation for Weyburn reservoir, southern Saskatchewan, using eight methods (McKay and Stichling, 1961).

Standard Methods/Guidelines: --

Sources for Additional Information: See Table 8-3.

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8.4 EVAPOTRANSPIRATION (MICROMETEOROLOGICAL METHODS)

8.4.2 Physically-Based Equations (Penman and Related Methods)

Other Names Used to Describe Method: Combination method, Penman (combination) equation, Penman-Monteith equation.

Uses at Contaminated Sites: Indirect field method for estimating evapotranspiration (ET).

Method Description: Physically-based process equations combine energy balance (Section 8.4.4) and aerodynamic transport of water vapor (Section 8.4.3) to calculate potential evapotranspiration (PET). Specific parameters that must be measured in the field vary slightly, depending on the equation, but can include: Surface temperature, surface resistance, saturation vapor pressure at mean air temperature, actual vapor pressure, mean monthly solar radiation, and wind velocity. Although these equations are physically-based, they require the measurement or estimation of various empirical constants. The Penman equation (Penman, 1948) was the first equation developed using this approach, and used weekly mean climatic data in empirically derived expressions for the energy and aerodynamic components. Figure 8.4.1 illustrates lake evaporation computed using the Penman formula compared to six other methods. Various modifications have been suggested since then, with the Penman-Monteith equation (which eliminated the need for surface temperature measurement) being the most commonly used. The theory of a complementary relationship between actual evapotranspiration (AET) and PET (Bouchet, 1963) has contributed to the further development of physically-based evaporation and ET models. In arid areas PET always exceeds AET, but as the amount of moisture available for removal from the soil increases, AET increases and PET decreases (because moisture in the air reduces the capacity for further additions of water vapor), until they converge on a value that is called wet environment evapotranspiration (WET). The Morton or Complementary Relationship Areal Evapotranspiration (CRAE) model developed for calculating WET, replaces the wind function in the Penman equation with a vapor transfer coefficient. The Brutsaert-Stricker, or Advection-Aridity Evaporation model, has been developed for calculating evaporation and ET in arid areas.

Method Selection Considerations: Penman and Related Equations Advantages: (1) Empirical constants in the equations can be obtained from published tables and graphs, rather than being determined from additional measurements for a specific site; and (2) work well for daily or larger periods in relatively humid areas where horizontal heat divergence in negligible, there is a good vegetative cover, and water is not limiting. Penman and Related Equations Disadvantages: (1) Field measurements for a number of meteorological parameters are required and are relatively expensive (although generally less expensive than other micrometeorological methods); and (2) serious discrepancies can occur in dry areas where advected heat accounts for a significant proportion of ET, unless locally determined empirical correction factors are developed. CRAE Model: Provides comparable results to the Penman equation, with the advantage that wind speed measurement is not required. Advection-Aridity Model: Works well for daily evaporation predictions. The main advantage is that it does not require surface resistance, soil moisture content, or other land surface measures of aridity.

<u>Frequency of Use</u>: Equation has been widely in England and to some extent in the eastern part of the United States. Not recommended for routine field applications and assessments.

Standard Methods/Guidelines: --

Sources for Additional Information: See Table 8-3.

8.4 EVAPOTRANSPIRATION (MICROMETEOROLOGICAL METHODS)

8.4.3 Mass Transfer Methods

Other Names Used to Describe Method: Dalton's law.

Uses at Contaminated Sites: Measuring evaporation.

<u>Method Description</u>: Mass transfer methods use semi-empirical equations for calculating evaporation as a function of: (1) Windspeed, often called the wind function, (2) saturation vapor pressure calculated from the temperature of the water surface, and (3) vapor pressure of the air. The wind function represents the combined effect of many variables and requires the estimation or measurement of one or more empirical constants and a mass-transfer coefficient.

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<u>Method Selection Considerations</u>: Advantages: Once the wind function has been determined, fewer measurements are required than for energy balance methods. **Disadvantages**: (1) Most accurate results require taking measurements in the center of a water body, which is difficult; (2) requires calibration with independently determined evaporation estimates; and (3) mass transfer methods for determining evapotranspiration (ET) generally require very complex instrumentation and well-trained personnel.

Frequency of Use: Widely used for measuring evaporation; rarely used for ET.

Standard Methods/Guidelines: --

Sources for Additional Information: Bras (1990), Dunne and Leopold (1978), U.S. Geological Survey (1982). See also, Table 8-3.

8.4 EVAPOTRANSPIRATION (MICROMETEOROLOGICAL METHODS)

8.4.4 Energy Budget Methods

Other Names Used to Describe Method: Energy balance/Bowen ratio method.

Uses at Contaminated Sites: Indirect field method for estimating evaporation and evapotranspiration (ET).

<u>Method Description</u>: The total energy available for evaporation or ET is measured. The equation for net radiation at the earth's surface is rearranged to solve for ET. Figure 8.4.4 illustrates the various components of the heat budget equation for a vegetated soil surface. Required field measurements include humidity (vapor pressure) and temperature profile above the ground or water surface, net radiation (Section 8.2.7), and soil heat flux (Section 1.6.3). Temperature gradients usually are measured using thermocouples. Humidity gradient is measured either by using two psychrometers or hygrometers (Sections 8.1.3 and 8.1.4) positioned at different elevations above the vegetative cover, or two tubes for collecting of samples for which moisture content is measured.

<u>Method Selection Considerations</u>: ET Advantages: (1) Is accurate in high humidity environments (within 5 to 10 percent of actual); and (2) can be used on hilly as well as flat terrain and for a wide variety of vegetation types, such as croplands and forests. ET Disadvantages: (1) Is expensive because of the large number of parameters that must be measured; (2) is less accurate where humidity is low; (3) heat divergence, sampling techniques, and advection can cause problems; (4) weekly instrumentation maintenance is required; (5) measurements over months or years are required to obtain average ET values; and (6) the energy required for photosynthesis (around 5 to 10 percent) is difficult to measure accurately, so it must be estimated. Lake Evaporation Disadvantages: (1) Does not consider flow of heat through the bottom of the lake, which can be significant in shallow lakes; (2) does not account for effects due to radiative diffusivity, stability of the air, and spray; and (3) is strongly affected by the ability to evaluate the advective energy component.

<u>Frequency of Use</u>: Well accepted for research applications. Not recommended for routine field applications and assessments.

Standard Methods/Guidelines: --

Sources for Additional Information: Bowen (1926), Bras (1990), DeVries and Afgan (1975), Dunne and Leopold (1978), Robins (1965), Rosenberg et al. (1983), Sharma (1985), Thompson et al. (1989), U.S. Geological Survey (1982). See also, Table 8-3.



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Figure 8.4.4 The heat budget equation and diagram of energy balance over a vegetated surface (Gray, 1973, after King, 1961). R_N S, K, and N must be measured or estimated to solve for evapotranspiration.

8.4 EVAPOTRANSPIRATION (MICROMETEOROLOGICAL METHODS)

8.4.5 Profile/Gradient Method

Other Names Used to Describe Method: Aerodynamic/vapor transfer method.

Uses at Contaminated Sites: Indirect field method for estimating evapotranspiration (ET).

<u>Method Description</u>: The profile or gradient method relates the vertical gradients of humidity and horizontal wind velocity to the rate of evaporation or ET from the underlying surface. Field measurements include: (1) The humidity gradient above the vegetative cover, and (2) wind profiles to estimate a momentum transfer coefficient (Km). The turbulent transfer coefficient for water vapor (Ke) might need to be measured to determine an empirical coefficient in the equation used to calculate ET to account for observed differences between the two transfer coefficients. The Thornthwaite-Holtzman equation is the most widely used formula for calculating evaporation using this method.

<u>Method Selection Considerations</u>: Advantages: (1) Once required coefficients have been determined, only windspeed and humidity gradient need be measured; and (2) works best in large, flat areas with uniform plant cover. Disadvantages: (1) Requires relatively complicated humidity and wind profile measurements; (2) the turbulent transfer coefficient for water vapor (Ke) also must be measured, unless it can be assumed to equal the momentum transfer coefficient (Km); (3) is less suitable for areas that are aerodynamically unstable because of rough vegetation cover or topography; and (4) is less accurate than mass transfer and energy budget methods because calculation sensitivity of instruments is more critical for accurate results and errors are more likely from adverse boundary conditions.

<u>Frequency of Use</u>: Sometimes used for short-term intensive studies, but not recommended for routine field applications and assessments.

Standard Methods/Guidelines: --

Sources for Additional Information: Sharma (1985), Thompson et al. (1989), U.S. Geological Survey (1982), Veihmeyer (1964). See also, Table 8-3.

8.4 EVAPOTRANSPIRATION (MICROMETEOROLOGICAL METHODS)

8.4.6 Eddy Correlation Method

Other Names Used to Describe Method: Eddy flux method.

Uses at Contaminated Sites: Indirect field method for estimating evaporation and evapotranspiration (ET).

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<u>Method Description</u>: Accurate, closely spaced instantaneous measurements of vertical wind velocity and humidity are averaged over a period of 1/2 hour or more. Water vapor flux (ET) is calculated from an equation relating deviations of humidity and vertical wind velocity from the mean. Extremely sensitive instrumentation, such as a propeller anemometer or sonic anemometer, is required for vertical wind measurements. Infrared hygrometry or wet-bulb/dry-bulb psychrometers usually are used for humidity measurements. On sloping surfaces, threedimensional wind measurements are required.

<u>Method Selection Considerations</u>: Advantages: Is the most direct means of measuring ET; (2) is independent of atmospheric conditions or types of underlying surfaces; (3) is accurate in low and high humidity environments. **Disadvantages**: Requires expensive and delicate instrumentation.

Frequency of Use: Well accepted for short-term research applications. Not recommended for routine field applications and assessments.

Standard Methods/Guidelines: --

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Sources for Additional Information: Rosenberg et al. (1983), Sharma (1985), Thompson et al. (1989), U.S. Geological Survey (1982). See also, Table 8-3.

Table 8-2 Reference Index for Hydrometeorological Data Collection and Measurement Methods

Торіс	References	
Climatic Data Sources/Uses	Eder et al. (1989), Hatch (1988), Whiting (1975, 1976)	
Meteorological Tables	Letestu (1966), List (1966)	
General References	ASTM (1986), Berry et al. (1945), Brakensiek et al. (1979), Brock and Nicolaidis (1984), Brunt (1944), Fritschen and Gay (1979), Hardy and Fisher (1972), Huschke (1970), Lockhart (1989a), Malone (1951), Meteorological Office (1956), Monteith (1972), National Weather Service (1972, 1975), Spilhaus and Middleton (1973), Tanner (1963), UNESCO (1969), U.S. Army (1975), U.S. Geological Survey (1980), U.S. Weather Bureau (1955), WMO (1971, 1973, 1974, 1975), WMO-IASH (1965)	
EPA Guidance Documents	U.S. EPA (1985, 1987a,b)	
Precipitation Gages/Samplers	Gilman (1964), Neff (1977), Purcell and Brown (1991), Simmons and Bigelow (1990)	
Precipitation Analysis	Butler (1957), DeWiest (1966), Kazmann (1988), Skeat (1969), Wisler and Brater (1959); <u>Frequency/Probability Maps</u> : Thomas and Whiting (1977), U.S. Department of Commerce (1961)	
Wind Speed/Direction	ASTM (1985a, 1990), Finkelstein et al. (1986a,b), Hayashi (1987), Lockhart (1985a,b, 1987, 1989b), Snow et al. (1989), Stearns (1985), Turner (1986)	
Humidity	ASTM (1982, 1983, 1984a, 1985b), U.S. Weather Bureau (1963a), Wexler (1957, 1965), Wexler and Brombacher (1951)	
Solar Radiation	ASTM (1984b), Carter et al. (1977), Coulson (1975), Elsasser and Culbertson (1960-atmospheric radiation table), Gates (1962), Kennedy (1949-pyrheliometers), Latimer (1972), Norris (1974), Selcuk and Yellott (1962), Suomi and Kuhn (1958); Estimation methods: Anderson and Baker (1967), Koberg (1964)	

Topic	References	
Hydrology Texts Covering ET	ASCE (1952), Branson et al. (1981), Bras (1990), DeWiest (1966), Dunne and Leopld (1978), Eagleson (1970), Gray (1973), Kazmann (1988), Linsley et al. (1949, 1982), Skeat (1969), Viessman et al. (1977), Wisler and Brater (1959); <u>Symposia</u> : Sokolow and Chapman (1974)	
General Reviews	Anderson et al. (1950), Barry (1973), Bennett and Linstedt (1978), Black et al. (1969), Brutsaert (1982), Christian et al. (1970), Criddle (1958), Doorenbos and Pruitt (1977), Evans (1962), Gangopadhyaya et al. (1966), Hamon (1961), Hanks and Ashcroft (1980), Hide (1954), Hillel (1982), Jensen (1974), Kittredge (1941), Levine (1959), Lowry and Johnson (1942), Monteith (1965), Robins (1965), Robins and Haise (1961), Rosenberg et al. (1968), Saxton and McGuiness (1982), Sharma (1985), Stephens and Stewart (1964), Tanner (1967, 1968), Thompsen et al. (1989), Thornthwaite (1948), U.S. Geological Survey (1982), Veihmeyer (1964), Webb (1975), WMO (1966)	
Water Balance Methods (See also,	Tables 4-3 and 7-6)	
Lysimetric Methods	Aboukhaled et al. (1982), Harrold (1966), Kohnke et al. (1940), Pelton (1961), Robins (1965), Tanner (1967), van Bavel (1961), Visser (1962); <u>Nonweighable Lysimeters</u> : Colman and Hamilton (1947), Evans (1971), Gilbert and van Bavel (1954), Mather (1954), Patric (1961), Robinson (1970), Stevenson and van Schaik (1967); <u>Weighable Lysimeters (see also, monolith lysimeters, Table 9-4)</u> : Harrold and Dreibelbis (1951, 1958), Katul and Parlange (1992), Mustonen and McGuinness (1968), Pruitt and Angus (1960), Ritchie and Burnett (1968), Rosenberg et al. (1967), van Bavel and Myers (1962), van Bavel and Reginato (1965), van Hylckama (1966, 1968), Williamson (1963), Wind Hzn (1958); <u>Hydraulic Lysimeters</u> : Black et al. (1968), Dagg (1970), DeBoodt et al. (1966), Ekern (1967), Forsgate et al. (1965), Hanks and Shawcroft (1965); <u>Lysimeters</u> (<u>Unspecified</u>): Blad and Rosenberg (1974), Blaney et al. (1930), King et al. (1956), Kittredge (1941), Makkink (1957), Martin and Rich (1948), McGuiness and Bordne (1972), Young and Blaney (1942); <u>Microlysimeters</u> : Abramova (1968), Al-Khafaf et al. (1978), Boast and Robertson (1982), Shawcroft and Gardner (1983), Staple (1974), Walker (1983)	
Soil Moisture Budget	Bowman and King (1965), Bresler and Kemper (1970), DeBoodt et al. (1966), Hillel (1971), Idso et al. (1975), Jenson (1974), Ligon (1969), Lomen and Warrick (1978), McGowan and Williams (1980), Rose (1966), Rose and Krishnan (1967), Slaytor (1967), Tanner (1967, 1968), van Bavel and Stirk (1967); <u>Methods of</u> <u>Modulating Potential Rates to Predict Soil Moisture Withdrawal</u> : Holmes (1961), Robertson and Holmes (1959), Taylor and Haddock (1956)	
Water Budget Methods	Eagleson (1978a,b); Evaporation: Anderson (1954), Hanks et al. (1969), Harbeck and Kennon (1954), Horton (1943b), Langbein et al. (1951), McKay and Stichling (1961), Winter (1981); Inflow-Outflow: Blaney et al. (1938, 1942), Jensen (1967), Lowry and Johnson (1942), Wilcox (1960), Yin and Brook (1992); Integration/Leaf Area Index Methods: Blaney et al. (1938, 1942), Hanks (1974), Jensen et al. (1970), Kristensen (1974), Raymond and Rezin (1989), Ritchie (1974); Watersheds: Hewlett et al. (1969), Lee (1970), Row and Reimann (1961), Williams (1940); Floodplains: Bowie and Kam (1968), Culler (1970), Gatewood et al. (1950), Hanson et al. (1972), Horton (1973-liteature review), Langbein (1942), Reigner (1966), Taylor and Nickle (1933)	

Topic	References
Water Balance Methods (cont.)	L .
Pan Evaporation	ASCE (1934), Bouwer (1959), Doorenbos and Pruitt (1977), Gangopadhyaya et al. (1966), Jensen (1974), Kohler et al. (1955), McKay and Stichling (1961), Mortenson and Hawthorn (1934), Mukammal (1961), Mukammal and Bruce (1960), Nordenson and Baker (1962), Peck and Munro (1976), Pruitt (1960), Rohwer (1931, 1934), U.S. Weather Bureau (1955), Young (1947); <u>Modified Energy Budget with Insulated Pan</u> : Cummings (1940), Kohler and Parmele (1967), U.S.Geological Survey (1982); <u>Pan Coefficients</u> : ASCE (1934), Ficke et al (1977), Hall (1934), Kohler (1954), Rohwer (1931, 1934), Sonmor (1963), State of California (1973), White (1932), Young (1947); <u>Pan Evaporation Maps</u> : Horton (1943a), Kohler et al. (1959); <u>ET Estimates from Pan Evaporation</u> : Mortenson and Hawthorne (1934), Pruitt (1960), Pruitt and Jensen (1955), Robertson and Holmes (1956), Stanhill (1962), Yin and Brook (1992)
Atmometers	Abbe (1935), Halkias et al. (1955), Livingston (1935), Livingston and Haasis (1929), Mukammal (1961), Mukammal and Bruce (1960), O'Connor (1955), Sonmor (1963), State of California (1973)
Ground-Water Fluctuation	Blaney et al. (1933), Davis and DeWiest (1966), Gatewood et al. (1950), Troxell (1936), Weeks and Sorey (1973), White (1932)
Transpiration	Cohen et al. (1981), Jarvis et al. (1981), Koch et al. (1971), Reicosky and Peters (1977), Veihmeyer (1964). Also, U.S. Geological Survey (1982) contains over 50 other references on methods for measuring or estimating transpiration.
Micrometeorological Methods	
General	Cruff and Thompson (1967), DeVries and Afgan (1975), Ficke (1972), Halstead and Covey (1957), Hanks and Ashcroft (1980), Harbeck (1952), Hillel (1980, 1982), Hughes (1967), Lemon et al. (1957), Penman (1963), Penman et al. (1967), Szeicz (1975), Tanner (1967, 1968), Thom (1975), Van Wijk and De Vries (1954); <u>Bare Soils</u> : Black et al. (1969), Fuchs et al. (1969)
Empirical ET Equations	<u>Reviews</u> : Bras (1990), Criddle (1958), Cruff and Thompson (1967), Eagleson (1970), Gray (1973), Jensen (1966a), Pierson and Jackman (1975), Pruitt and Doorenbos (1977), Robins and Haise (1961), Sharma (1985), Tanner (1967), Thompson et al. (1989), U.S. Geological Survey (1982), Veihmeyer (1964); - <u>Blaney-Morin/(Modified) Blaney-Criddle</u> : Blaney (1959), Blaney and Criddle (1950, 1962), Blaney and Morin (1942), Blaney et al. (1952), Criddle (1958), Cruf and Thompson (1967), Dunne and Leopold (1978), Pruitt and Doorenbos (1977), State of California (1973), Stephens and Stewart (1964), U.S. Weather Bureau (1905), Yin and Brook (1992); Jensen-Haise: Jensen (1966b), Jensen and Haise (1963), Jensen et al. (1970); <u>Thornthwaite</u> : Dunne and Leopold (1978), Pelton et al. (1960). Stephens and Stewart (1964). Thornthwaite
	an (1900), Stephens and Stewart (1904), Inornthwatte (1931, 1948), Inornthwatte and Mather (1955, 1957), Thornthwaite and Wilm (1944), Yin and Brook (1992); <u>Others</u> : Behnke and Maxey (1969), Benson et al. (1992), Christiansen (1968), Christiansen and Hargreaves (1969), Gardner (1958), Halkias et al. (1955), Harding et al. (1930), Hargreaves (1956), Hargreaves and Samani (1985), Holdridge (1962), Kincaid et al. (1979), Lowry and Johnson (1942), Makkink (1957), Munson (1962), Priestly and Taylor (1972), Ritchie (1972), Saxton and McGuiness (1982), Tanner and Jury (1976)

Table 8-3 (cont.)

Topic	References
Empirical Evaporation Equations	<u>Reviews</u> : Bras (1990), Helfrich et al. (1982), McKay and Stichling (1961), Weisman (1975); <u>Specific Equations</u> : Harbeck (1962), Kohler (1954), Kohler et al. (1955), Kuzmin (1957), Marciano and Harbeck (1954), Meyer (1915, 1942), Rohwer (1931), Shulyakovsky (1969)
Physically-Based Equations	Benson et al. (1992), Bras (1990), Businger (1956), Chiew and McMahon (1991), Cordova and Bras (1981), Crago and Brutsaert (1992), Duell (1990), Gray (1973), Katul and Parlange (1992), Lemur and Zhang (1990), McKay and Stichling (1961), Monteith (1963), Morton (1978, 1983, 1991), Penman (1948, 1956), Pruitt and Doorenbos (1977), Robins (1965), Rosenberg et al. (1983), Sharma (1985), Staple (1974), Tanner (1968), Tanner and Pelton (1960), Thompson et al. (1989), Turner (1957), U.S. Geological Survey (1982), van Bavel (1966), Veihmeyer (1964); <u>Advection Aridity Evaporation Model</u> : Brutsaert and Stricker (1979), Lemeur and Zhang (1990), Morton (1991), Parlange and Katul (1992); <u>Soil</u> <u>Evaporation Loss Equations</u> : Philip (1957, 1991); <u>Lake Evaporation Equations</u> : Goodling et al. (1976), Weisman and Brutsaert (1973)
Mass-Transfer Methods	Evaporation: Brasklavskii and Vikulina (1954), Ficke (1972), Harbeck (1962), Harbeck et al. (1954, 1958), Hughes (1967), Jobson (1973), Marciano and Harbeck (1954), Munn (1961), Resch and Selva (1979), Richards and Irbe (1969), Sutton (1949), Sverdrup (1946), Thornthwaite and Holzman (1939), Turner (1966), Wunderlich (1972)
Energy Budget Methods	Evaporation: Anderson (1954), Levine (1959), McKay and Stichling (1961), Tanner (1960); Evapotranspiration: Angus and Watts (1984), Aston and van Bavel (1972), Black and McNaughton (1971), Blad and Rosenberg (1974, 1975), Denmead and McIlroy (1970), Dennehy and McMahon (1989), Duell (1990), Fritschen (1965), Jackson et al. (1977), Kohler et al. (1955), Lemon (1960), Munn (1961), Ohmura (1982), Pruitt (1963), Robins (1965), Tanner (1966, 1968, 1988), Webb (1975)
Profile/Gradient Method	Businger et al. (1971), Dyer (1963, 1965, 1974), King (1966), Marciano and Harbeck (1954), Pierson and Jackman (1975), Priestly (1959), Pruitt et al. (1973), Quinn (1979), Szeicz et al. (1969), Thornthwaite and Holzman (1939, 1942); <u>Vapor Transfer Method</u> : Pasquill (1949, 1950), Rider (1954, 1957), Rider and Robinson (1951), Tanner (1960), Veihmeyer (1964)
Eddy Correlation	Christian et al. (1970), Duell (1990), Dyer (1961, 1968), Easterbrook (1969), Gangopadhyaya et al. (1966), Goddard and Pruitt (1966), Goltz et al. (1970), Hicks (1973), Hicks et al. (1973), Jobson (1973), Swinbank (1951), Swinbank and Dyer (1967), Tanner (1966, 1988)

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SECTION 9

VADOSE ZONE SOIL-SOLUTE/GAS SAMPLING AND MONITORING METHODS

Monitoring of soil water in the vadose zone can serve as an early warning system at controlled waste disposal sites that contaminants are entering the subsurface, and can allow actions to be taken before contaminants reach the saturated zone. Methods for sampling and monitoring the vadose zone can be broadly categorized as: (1) Indirect (surface geophysical methods and probes that focus on measuring variations in soil salinity), and (2) direct (in which soil water is collected directly in the field, or extracted from samples of soil solids).

Indirect Soil Salinity Measurements

A variety of methods are available for locating and monitoring areas of high soil salinity. These methods primarily have been developed for agricultural applications to identify saline soils and control irrigation flows where soluble salts can affect crop productivity. Table 9-1 summarizes information on six indirect methods for monitoring soil salinity. The four-probe electrical method is a direct application of the electrical resistivity surface geophysical method (Section 1.2.1), with electrode configurations that measure near-surface resistivity. The electromagnetic induction sensor is an instrument that is specifically designed to measure conductivity in the near surface. The other indirect methods involve placement of probes or sensors in the subsurface. The main advantage of indirect methods is that data can be collected quickly. The main disadvantages are: (1) Instruments must be calibrated for each soil type by collection of samples where salinity is measured directly to obtain quantitative measurement of soil salinity; and (2) actual chemical constituents that are contributing to soil salinity cannot be determined. The four-probe electrical and porous matrix soil salinity sensors are the most commonly used indirect methods.

Direct Soil Solute Sampling Methods

The U.S. Environmental Protection Agency is placing increasing emphasis on vadose zone soil solute sampling as an early warning system to detect movement of contaminants before they reach the saturated zone (Cullen et al., 1992; Durant et al., 1993). Three major types of soil water can be identified in the context of sampling soil water: (1) Macropore or gravitational water, which flows through the soil relatively rapidly in response to gravity (excess of 0.1 to 0.2 bars suction); (2) soil-pore or capillary water, which is held in the soil at negative pressure potentials from around 0.1 to 31 bars of suction; and (3) hygroscopic water that is held at tensions greater than 31 bars suction. Soil-pore water moves through the vadose zone, but at much slower rates than gravitational water (see discussion of potential-conductivity relationships in Section 6.3.1), whereas hygroscopic moves primarily in the vapor form. The term soil solute or solution sampling has been used loosely in the literature to describe most sampling methods, whereas the term soil pore liquid is typically used in a more restricted sense (and is so used here) to apply to sampling of capillary water. The chemistry of soil solute sampling methods can differ significantly, depending the method used. Concentrations of inorganic species generally increase as the matric potential increases. In general, ceramic soil solution samplers (which use suctions up to around 0.8 bars) will collect samples that are most representative of the soil solution for the purpose of evaluating contaminant transport.

There are a large number of specific methods by which soil water can be sampled. Suction samplers draw water from the soil by applying a vacuum. A variety of free-drainage samples collect water percolating through the soil by gravity flow. Other methods include: (1) Use of absorbent materials with retrieval and extraction of water in the laboratory, (2) collection of soil solids with extraction of soil water in the laboratory by a variety of methods, and (3) preparation of a soil saturation extract from a solids sample. Table 9-1 summarizes some information on six types of suction samplers, seven methods of collecting samples by free drainage, and four miscellaneous methods. Table 9-1 also lists collection of soil solids for volatile constituents and soil microorganisms in the vadose zone.

The main advantages of suction samplers is that they are relatively easy to install, and there are essentially no limitations to the depth of sampling when a vacuum-pressure apparatus is used. The main

Method	Sampling Method	Depth Limitation	Chapter Sections
Indirect Salinity Measurement Methods			
Four Probe Electrical	Resistivity	Near surface	9.1.1, 1.2.2
Portable EC Probe	Resistivity	1.5 m	9.1.2
In Situ EC Probe	Resistivity	None	9.1.2
Porous Matrix Salinity Sensors	Resistivity	None	9.1.3
Electromagnetic Induction Sensor	Conductivity	2 m	9.1.4, 1.3.1
Dielectric Sensors	Dielectric	2 m*	9.1.4, 6.2.3
Time Domain Reflectometry Sensor	Dielectric	Up to 20 m	9.1.4, 6.2.4
Neutron Probe	Nuclear	None	3.3.3, 6.2.2
Direct Soil Solute Sampling Methods			
Vacuum-Type Porous Cup	Suction	2 m	9.2.1
Vacuum-Pressure Porous Cup	Suction	45 ft	9.2.2
Vacuum High-Pressure Porous Cup	Suction	300 ft	9.2.2
Vacuum-Plate Sampler	Suction	2 m*	9.2.3
Membrane Filter	Suction	1-4 m ^b	9.2.4
Hollow Fiber	Suction	2 m*	9.2.5
Ceramic Tube Sampler	Suction	2 m*	9.2.6
Capillary Wick Sampler	Capillary	đ	9.2.7
BAT Sampler	Suction	45 ft	5.5.2
Trench Lysimeter	Gravity	đ	9.3.1
Caisson Lysimeter	Gravity	3 m+	9.3.1
Pan Lysimeter	Gravity	đ	9.3.1
Glass Block Lysimeter	Gravity	d	9.3.1
Wicking Type Sampler	Gravity	đ	9.3.1
Tile Drain Outflow	Gravity	50+ ft	9.3.1
Perched Water Table	Gravity	None	9.3.2
Nylon Sponge	Absorbent	Near surface	9.3.3
Ccramic Rod	Absorbent	Near surface	9.3.3
Solid Soil Water Extraction	•	None	9.3.4
Soil Saturation Extract	Slurry	None	9.3.5
SEAMIST	Absorbent	100s ft	9.3.7
Methods for Sampling Sensitive Soil Const	ituents		
Static Soil-Gas Sampling	Absorbent	Near Surface	9.4.1
Soil-Gas Probes	Suction	t	9.4.2
Tank Leak Sensors	Various	Typically <2m	9.4.3
Soil Volatiles/Microorganisms	Core	(9.3.6

Table 9-1 Summary Information on Soil Solute Monitoring and Sampling Methods

Boldface = Most commonly used methods.

*With vacuum sampling apparatus; greater depths would be possible using vacuum-pressure sampling system.

*Upper limit would require modification of system to use vacuum-pressure sampling apparatus.

Sample is collected by free-drainage in all gravity samplers, but suction can be used to bring sample to the surface.

⁴Depth limited by the depth to which a hole or trench can be safely dug for installation of sampler in the sidewall; typically 2 meters or less.

*Various methods can be used to extract soil water from a sample: Squeezing, displacement, displacement/centrifugation, centrifugation, and adsorption.

Depends on density of subsurface material and method of penetration/coring. Soil gas probes used with cone penetration rigs (Sections 2.2.2, 5.5.1, and 5.5.2) can penetrate 100 to 150 feet with favorable soil conditions; greater depths are possible if holes are drilled before insertion of the soil gas probe. Coring depth limits are defined by the type of drilling/coring method used (Sections 2.3 and 2.4).

disadvantage of suction samplers is that they might not collect representative samples. Sampling for organic chemicals, microorganisms, volatile chemicals, and metals is especially problematic due to potential sorption/interferences by the porous cup. Vacuum-type and vacuum-pressure type porous cup samplers are by far the most commonly used types of suction samplers. The main advantage of free-drainage samplers is that relatively large volumes of water, which is representative of water that is actually percolating to deeper zones, is obtained. The main disadvantages are that installation procedures are time consuming and complex and limited to relatively shallow depths. Trench lysimeters with pan collectors are the most commonly used free-drainage samplers. Figure 9-1a illustrates generic vadose zone monitoring installations for an existing hazardous waste landfill and Figure 9-1b illustrates generic vadose zone monitoring installations for a new surface impoundment. Capillary wick samplers (Section 9.2.7) are a relatively new development, which appear to have good potential for collecting more representative samples of soil solutions than either porous cup or freedrainage samplers in the near surface.

Gaseous Phase Characterization

Sampling of soil gases (volatile contaminants or gases such as methane and carbon dioxide, which are indicators of increased microbial activity resulting for organic contaminants) has gained rapid acceptance as a method for preliminary mapping of contaminant plumes in ground water, and monitoring of underground storage tanks. Contaminant plume mapping can be done either by passive sampling, where absorbent collectors are buried for a period of time and retrieved for laboratory analysis (Section 9.4.1), or by using soil-gas sampling probes (Section 9.4.2). Various types of sensors can be used to detect leaks in underground storage tanks (Section 9.4.3). Monitoring of air pressure (Section 9.4.4) and measurement of air permeability (Section 9.4.5) might be required for modeling the transport of contaminants in the vadose zone.

Contaminant Flux

Section 9.5.1 (Solute Flux Methods) briefly describes four methods for estimating the mass transfer of pollutants from the vadose zone to ground water, and Section 9.5.2 (Soil-Gas Flux) describes several methods for estimating soil-gas flux to the atmosphere.



(a)



Figure 9-1 Vadose zone monitoring systems: (a) Generic monitoring design for existing hazardous waste landfill; (b) Water quality monitoring design for a new surface impoundment (Everett et al., 1983).

9.1 SOLUTE MOVEMENT (INDIRECT METHODS)

9.1.1 Four Probe Electrical Resistivity

Other Names Used to Describe Method: Four electrode technique/sensors.

<u>Uses at Contaminated Sites</u>: Measuring in situ soil salinity in the shallow vadose zone; locating brine and chloride plumes; estimating water content.

<u>Method Description</u>: A Wenner four probe electrode array (see Section 1.2.1) is used to detect areas of low electrical resistivity (high conductivity) in the soil. For a given soil type, electrical conductivity of the bulk soil and electrical conductivity of the saturation extract from the soil are directly related. Once a calibration curve has been developed (requiring multiple measurements of both soil conductivity and saturation extract conductivity at different locations in a single soil type), soil conductivity measurements can be related to saturation extract conductivity, which in turn can be related to salinity (see references in Table 9-2).

Method Selection Considerations: Advantages: (1) Is a nondestructive method (once calibration constants have been calculated); (2) readings are obtained rapidly and inexpensively; (3) is useful for detecting the presence of shallow saline ground water; (4) horizontal variations in salinity can be easily measured by lateral transects; (5) vertical changes in salinity can be evaluated by changing the electrode spacing; and (6) a large volume of soil can be measured compared to other methods. Disadvantages: (1) Obtaining calibration relationships can be tedious; (2) accuracy decreases in layered soils; (3) time-series monitoring is difficult due to the requirement of making multiple traverses; (4) is generally limited to shallow depths; (5) does not provide data on specific pollutants; and (6) will not detect pollutants that do not change the electrical conductivity of the subsurface. Water Content Measurement: Moisture content can be estimated from four electrode resistivity measurements if salinity, temperature, and bulk density can be quantified, and calibration curves are developed, however, other simpler and more reliable methods generally are used (see Sections 6.2 and 6.3).

<u>Frequency of Use</u>: Commonly used for identification of saline soils in agricultural studies. DC resistivity methods for detecting conductive contaminant plumes in the deeper subsurface are described in Section 1.2.1.

Standard Methods/Guidelines: Salinity: Rhoades and Oster (1986); Water content: Morrison (1983).

Sources for Additional Information: Everett et al. (1983). See also, Table 9-2.

9.1 SOLUTE MOVEMENT (INDIRECT METHODS)

9.1.2 EC Probes

Other Names Used to Describe Method: Four-electrode salinity probe, electrical conductivity probe*, portable salinity probe, burial type salinity probe, four-electrode conductivity cell.

Uses at Contaminated Sites: Obtaining small volume soil salinity measurements.

<u>Method Description</u>: A cylindrical portable probe containing electrodes at fixed spacing is attached to a rod with a handle (Figure 9.1.2a). A hole the same diameter as the probe is augered, and resistivity is measured at successive depths. Alternatively, a specially dedicated **burial-type** probe is placed permanently in the ground with a cable running to the surface for periodic measurements (Figure 9.1.2b). Calibration of probes is similar to the calibration method for the four probe electrical method (Section 9.1.1). The **four-electrode conductivity cell** is a variant of this approach, in which and undisturbed soil core is collected using a removable lucite columnar insert in a soil-core sampler. The lucite section is removed from the sampler and segmented to form individual cells. Electrodes are inserted into the soil through threaded holes in the lucite cell walls and resistivity is measured.

<u>Method Selection Considerations</u>: Advantages: (1) Salinity changes with depth in stratified soils can be measured; (2) burial probe measurements can be taken at a greater depth than with four electrode method; (3) in-place units allow easy monitoring of changes in salinity with time; (4) are well suited for mapping and diagnosis as well as monitoring; (5) compared to salinity sensor probes, are more versatile, durable, less subject to calibration change, and respond to changes in salinity with less time lag; and (6) can be used to measure different soil volumes. Disadvantages: (1) Developing individual calibration relationships for each strata is time consuming and expensive; (2) use is limited to relatively shallow depths; and (3) provide no data on specific pollutants nor will probes detect pollutants that do not change the electrical conductivity of the subsurface.

Frequency of Use: Primarily used for land treatment areas and irrigated fields.

Standard Methods/Guidelines: Portable probe: Rhoades and van Schilfgaarde (1976), Rhoades et al. (1977); Burial probe: Rhoades (1979).

Sources for Additional Information: Everett et al. (1983). See also, Table 9-2.

*This probe actually measures resistivity, but measurements typically are reported in its reciprocal, conductivity.



(a)



Figure 9.1.2 Electrical conductivity probes: (a) Schematic illustrating the principle of a soil-salinity probe (Rhoades and van Schilfgaarde, 1976, by permission); (b) Installation of in situ soil salinity sensor (Morrison, 1983).

9.1 SOLUTE MOVEMENT (INDIRECT METHODS)

9.1.3 Porous Matrix Salinity Sensors

Other Names Used to Describe Method: Ceramic salinity sensors, in situ salinity sensors.

Uses at Contaminated Sites: Monitoring soil salinity; determining dispersion coefficients from salinity gradients and evapotranspiration; measuring water content measurement.

<u>Method Description</u>: Electrodes and thermistors embedded in porous ceramic are placed in the soil. Many types of sensors have been developed. Figure 9.1.3a illustrates a cylindrical sensor, and Figure 9.1.3b a square salinity sensor. The specific conductance is measured when the soil solution equilibrates with the ceramic. As with the four probe electrical and EC probe, calibration curves that relate signal to salinity and/or water content must be developed to relate conductivity readings to salinity. Temperature also must be measured and used to develop calibration relationships.

<u>Method Selection Considerations</u>: Most suitable for land treatment areas and irrigated fields. Could be installed below ponds before they are filled with water. Advantages: (1) Are simple, easily read and sufficiently accurate for salinity monitoring; (2) readings are taken at same depth and location each time; (3) vertical migration of saline water can be monitored by installing units at different depths; and (4) output can be interfaced with data acquisition systems. Disadvantages: (1) Are more subject to calibration changes than the four-electrode method; (2) are more expensive and less durable than four-electrode method; (3) time lag in response to changing salinity can be several days; (4) cannot be used at soil-water pressures less than about -2 bars; (5) soil disturbance during installation can affect results (salinity readings will be lower compared to undisturbed soil if disturbed soil has greater leaching due to increased permeability); and (6) does not provide data on specific pollutants.

Frequency of Use: Commonly used in agricultural research where continuous monitoring of soil salinity is required.

Standard Methods/Guidelines: Rhoades and Oster (1986), Richards (1966).

Sources for Additional Information: Morrison (1983). See also, Table 9-2.





Figure 9.1.3 Soil salinity sensors: Cylindrial (a); Square (b) (Morrison, 1983, after Enfield and Evans, 1969, by permission).

9.1. SOLUTE MOVEMENT (INDIRECT METHODS)

9.1.4 Electromagnetic Sensors

Other Names Used to Describe Method: Electromagnetic induction sensor, inductive electromagnetic soil conductivity meter, soil conductivity sensor, EM soil salinity sensor, time domain reflectometry (TDR)/dielectric sensors.

Uses at Contaminated Sites: Monitoring soil salinity.

<u>Method Description</u>: The EM soil salinity sensor uses the principles of electromagnetic induction (see Section 1.3.1) to measure electrical conductivity in the soil rooting zone (1 to 2 meters). EM instruments are designed for measurement of conductivity of the near surface. Dielectric sensors (Section 6.2.3) and time domain reflectometry (Section 6.2.4) measure the dielectric properties of the subsurface using probes that transmit and receive electromagnetic signals.

<u>Method Selection Considerations</u>: EM Soil Salinity Sensor Advantages: (1) Equipment is very portable and easy to use; (2) direct contact with the ground is not necessary; and (3) continuous measurements are possible. EM Soil Sensor Disadvantages: Depth of penetration is limited to 1 to 2 meters. Time Domain Reflectometry has the advantage of allowing measurement of both moisture content and electrical conductivity (see Section 6.3.4 for additional discussion of advantages).

<u>Frequency of Use</u>: EM soil salinity sensors are used primarily for agricultural applications for measuring salinity of the soil rooting zone and locating saline seeps. TDR sensors are relatively new but have gained rapid acceptance.

Standard Methods/Guidelines: --

Sources for Additional Information: Salinity sensors: See Table 9-2. TDR: Kachonoski et al. (1992); see also, references listed in Table 6.3.

9.2 DIRECT SOIL-SOLUTE SAMPLING (SUCTION METHODS)

9.2.1 Vacuum-Type Porous Cup

Other Names Used to Describe Method: Suction/soil lysimeter, tension lysimeter, soil-water extractors

<u>Uses at Contaminated Sites</u>: Sampling of soil pore liquids in the vadose zone to characterize contaminated sites or provide early warning of break-through of pollutants at controlled disposal sites.

<u>Method Description</u>: A porous cup or plate (usually ceramic but other materials, such as alundum, fritted glass, and nylon can be used^{*}) is attached to a small diameter tube (usually PVC), which is placed in the soil, making sure that there is good contact with the soil material. A one-hole rubber plug is placed in the other end of the tube and small diameter tubing beginning at the base of the ceramic cup runs through the hole to the surface (Figure 9.2.1a). A vacuum is applied to the small tubing and the soil solution is drawn into a small flask. Tensiometers (Section 6.1.1) can be installed in the vicinity to determine that amount of suction that should be applied during sampling. Figure 9.2.1b illustrates the use of vacuum-type porous cup lysimeters in a barrel lysimeter. A purge-and-trap device at the surface (Figure 9.2.1c) can be used for collection of volatiles from suction samplers.

Method Selection Considerations: Advantages: (1) Allows direct sampling of soil water; (2) successive samples can be obtained from the same depth; (3) is inexpensive and simple; and (4) can be installed below shallow impoundments and landfills prior to construction for monitoring of seepage when the facility is operating. Disadvantages: (1) Generally is limited to depths less than 6 feet; (2) is limited to soil water pressure less than air entry value of the cups (-1 atmosphere or -30 kPa), so will not work in very dry or frozen soils; (3) small volumes sampled might not be representative; (4) only samples pore water, water moving through cracks and macropores might have different chemical composition (can be overcome by also using zero suction samplers [Section 9.3.1]); (5) suction might affect soil-water flow patterns, so installation of tensiometers is required to determine the correct vacuum to apply; (6) samples might not be representative of pore water because method does not account for relationships between pore sequences, water quality and drainage rates; (7) contact between cup and soils difficult to maintain in very coarse textured soils, such as gravels, and exposure to freeze-thaw might break contact with soil; (8) cup might be plugged by solids or bacteria; (9) chemistry of solute might be altered in passage through cup (sorption of metals, ammonia, chlorinated hydrocarbons); (10) PTFE cups have relatively limited operational ranges (up to 7 centibars); (11) dead space, where fluid in the cup is not brought to the surface, might occur if the discharge tube hangs up on the lip of the cup during installation, and some PTFE samplers have a permanent dead space; (12) generally is not suitable for bacterial sampling due to screening and adsorption; and (13) heavy metals might be sorbed on the porous-cup matrix.

Frequency of Use: Very common where near-surface sampling is required.

Standard Methods/Guidelines: ASTM (1992).

Sources for Additional Information: See Table 9-3.

*Teflon also has been used as a material for suction lysimeters, but is not currently recommended because of problems with low bubbling pressure (ASTM, 1992).





Figure 9.2.1 Vacuum-type porous cup lysimeters: (a) Conventional system (Everett et al., 1983); (b) A barrel lysimeter with vacuum porous cup samplers installed within a hazardous waste land treatment facility (Hornby et al., 1986, by permission); (c) System for sampling volatile organics in soil water (Scalf et al., 1981).

9.2 DIRECT SOIL-SOLUTE SAMPLING (SUCTION METHODS)

9.2.2 Vacuum-Pressure Type Porous Cup

Other Names Used to Describe Method: Suction/soil lysimeter, high pressure-vacuum type porous cup sampler, deep pressure vacuum lysimeter, ceramic points.

<u>Uses at Contaminated Sites</u>: Sampling of soil pore liquids in the vadose zone to characterize contaminated sites or provide early warning of break-through of pollutants at controlled disposal sites.

<u>Method Description</u>: Vacuum-pressure type: Similar to vacuum type porous cup, except that a second line is placed in the porous-cup-tipped tube, which ends just below the stopper. The shorter line is connected to a pressure-vacuum source. When the unit is in place, a vacuum is applied to draw soil water into the sampler. Then pressure is applied to push the sample into the flask. Figure 9.2.2a illustrates installation of two vacuum pressure lysimeters in the same hole at two levels. The high pressure-vacuum type sampler is similar to the vacuum-pressure type device, except that the sampler is divided into two chambers connected by the line with a one-way valve (Figure 9.2.2b). When vacuum is applied, the soil solute is pulled into the upper chamber. When pressure is applied to drive the sample into the container at the surface, the one-way valve prevents any of the sample from being pushed out the porous cup. Nightingale et al. (1985) have developed a pressure vacuum-type sampler suitable for both saturated and unsaturated conditions that uses a standpipe rather than a check valve to keep the sample from being forced back into the soil when pressure is applied. With modifications, conventional ceramic porous-cup soil-solution samplers can be used to sample volatile organic compounds in the soil solution (Wood et al., 1981 [see Figure 9.2.1c]).

<u>Method Selection Considerations</u>: In most cases, ceramic vacuum-pressure lysimeters will be the method of choice. Advantages: All the same advantages of the vacuum-type sampler (Section 9.2.1), plus: (1) Can be used at depths below the suction lift of water (down to 50 feet for vacuum-pressure type and down to 300 feet for high pressure-vacuum type); and (2) several units can be installed in the same borehole for sampling soil water at different depths. Disadvantages: Same as for vacuum-type sampler (Section 9.2.1) plus: Some solution is forced back through the walls of the cup when pressure is applied. The high pressure-vacuum type sampler overcomes this problem.

Frequency of Use: Probably the most commonly used method for soil-solute sampling.

Standard Methods/Guidelines: ASTM (1992), Everett (1990), Rhoades and Oster (1986), U.S. EPA (1986).

Sources for Additional Information: See Table 9-3.



Figure 9.2.2 Vacuum-pressure lysimeters: (a) Clustered vacuum-pressure suction cup lysimeters in a single borehole (Everett et al., 1983, after Hounslow et al., 1978); (b) High pressure-vacuum suction cup sampler (Everett et al., 1983).

9.2 DIRECT SOIL-SOLUTE SAMPLING (SUCTION METHODS)

9.2.3 Vacuum-Plate Samplers

Other Names Used to Describe Method: Alundum tension plate.

<u>Uses at Contaminated Sites</u>: Sampling of soil pore liquids in the vadose zone to characterize contaminated sites or provide early warning of break-through of pollutants at controlled disposal sites.

<u>Method Description</u>: Principles are the same as porous cup samplers (Sections 9.2.1 and 9.2.2) except the geometry of the porous material is different. The vacuum plate consists of an Alundum or ceramic disc (range from 4.3 to 25.4 centimeters in diameter) attached to an extraction vacuum extraction tube (Figure 9.2.3). Installations reported in the literature use a vacuum sampling apparatus, but a vacuum-pressure system (Section 9.2.2) could be used as well. Installation is similar to that described in Section 9.3.1 for trench lysimeters.

<u>Method Selection Considerations</u>: Advantages and disadvantages essentially are the same as for porous cup suction samplers (Section 9.2.1 and 9.2.2) with the added advantage that a large sample volume can be obtained without disrupting adjacent flow pattern, and the added disadvantage that trench installation procedures are more complicated

Frequency of Use: Uncommon.

Standard Methods/Guidelines: ASTM (1992).

Sources for Additional Information: See Table 9-3.



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Figure 9.2.3 Vacuum-plate lysimeter (Morrison, 1983, after Cole, 1958, by permission).

9.2 DIRECT SOIL-SOLUTE SAMPLING (SUCTION METHODS)

9.2.4 Membrane Filter

Other Names Used to Describe Method: --

<u>Uses at Contaminated Sites</u>: Sampling of soil pore liquids in the vadose zone to characterize contaminated sites or provide early warning of break-through of pollutants at controlled disposal site.

<u>Method Description</u>: A membrane filter (polycarbonate or cellulose acetate) and a glass fiber prefilter are mounted in a Swinnex-type filter holder (used for filtration of fluids delivered by a syringe [see Figure 9.2.4a]). Installation involves digging a hole to the desired depth (up to 4 meters), and placing glass fiber collectors in the bottom of the hole (Figure 9.2.4b). Glass fiber discs that fit within the filter holder are placed on the fiber collectors and provide a wicking action between the collectors and the filter holder assembly (Figure 9.2.4c). The filter holder is placed in the hole, making sure that the glass fiber prefilter in the holder is in contact with the "wick" discs. The hole is then backfilled. The sample is drawn through a flexible tube attached to the filter holder using suction. The prototype (Stevenson, 1978) has been used at depths to 1 meter using suction apparatus similar to vacuum-type porous cup samplers (Section 9.2.1). Theoretically, installation could be as deep as 4 meters using a vacuum-pressure type apparatus for fluid collection.

<u>Method Selection Considerations</u>: Advantages: (1) Membrane filter is better for determining phosphorus concentrations than porous cup samplers; (2) samples are less susceptible to being drawn back into the soil when soil-moisture tension exceeds the vacuum in the sampler; (3) can be manufactured from inexpensive, readily available materials; (4) the wick-collector system provides contact with a relatively large soil area; and (5) satisfactory sampling rates can be maintained even when parts of the collector sheet become blocked by fine particles. Disadvantages: (1) Installation procedure is more complex than for porous-cup sampler; (2) under very dry soil conditions, the membrane dries out and rapid vacuum loss occurs; (3) depending on the membrane filter composition and manufacturer, a variety of contaminants, such as nitrogen, carbon particulate matter, and sodium, might be contributed to samples (thorough rinsing with distilled water can minimize these contributions); and (4) clogging by biofilm growth is a problem on cellulose acetate membranes (can be controlled, in part, with treatments of silver nitrate and sodium chloride).

Frequency of Use: Uncommon.

Standard Methods/Guidelines: ASTM (1992), Stevenson (1978).

Sources for Additional Information: See Table 9-3.



Preparation of "Swinnex" type filter holder for suction sampler.





Installed sampler, sample receiver, and vacuum indicator.



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Installation of suction sampler showing glass fiber "wick" and "collector" arrangement.

(c)

Figure 9.2.4 Membrane filter sampler: (a) Preparation of Swinnex type filter holder; (b) Installation set-up; (c) Sampler; (Morrison, 1983, after Stevenson, 1978, by permission).

9.2 DIRECT SOIL-SOLUTE SAMPLING (SUCTION METHODS)

9.2.5 Hollow Fiber

Other Names Used to Describe Method: Cellulose-acetate hollow fiber sampler, hollow fiber filters.

<u>Uses at Contaminated Sites</u>: Sampling of soil pore liquids in the vadose zone to characterize contaminated sites or provide early warning of break-through of pollutants at controlled disposal sites.

<u>Method Description</u>: To date, this method has been applied only to soil cores and blocks in the laboratory. Bundles of semipermeable fibers (cellulose-acetate, or hollow fibers produced from a noncellulosic polymer solution) are installed vertically or horizontally (Figure 9.2.5) in a soil core by inserting them down a thin diameter (0.3 centimeter) metal tube. Once the fibers have been pushed into the core, the tube is withdrawn. The fibers also can be placed in a perforated length of PVC tubing that is pushed into place. The hollow fibers are attached to a vacuum pump and suction is applied (based on readings from separately installed tensiometers) to collect the soil solution.

<u>Method Selection Considerations</u>: Not recommended for use in the field at this time due to lack of field testing (Everett, 1990). Advantages: (1) The fibers used have been designed to function as molecular sieves, allowing more precise selection of pore size (macrosolute rejection levels from 500 to 300,000 molecular weight); (2) installing hollow fibers for solute sampling from laboratory core studies requires less disturbance than porous ceramic cup samplers; and (3) encasing a fiber bundle within a perforated plastic tube allows installation of a sampling unit along a long horizontal axis. Disadvantages: (1) Horizontal installation is difficult in the field; (2) cellulose acetate fibers might screen nitrate-nitrogen, phosphorus, and potassium; (3) biological clogging of fibers is a potential problem; (4) suitability for sampling metal and organic contaminants has not been evaluated; and (5) has a narrow operating range (20 to 50 centibars) because large pore diameters result in low bubbling pressure.

Frequency of Use: Uncommon.

Standard Methods/Guidelines: --

Sources for Additional Information: See Table 9-3.



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Figure 9.2.5 Schematic of experimental hollow fiber sampling system (Levin and Jackson, 1977, by permission).

9.2 DIRECT SOIL-SOLUTE SAMPLING (SUCTION METHODS)

9.2.6 Ceramic Tube Sampler

Other Names Used to Describe Method: Ceramic filter candle, vacuum (trough) extractor.

Uses at Contaminated Sites: Sampling soil pore liquids in the rooting zone.

<u>Method Description</u>: The sampling device is similar to the vacuum-type porous cup sampler, except that a porous ceramic candle (around 12 inches long and 1 inch in diameter) is used instead of a porous cup. Installation requires excavation of a vertical trench to below the depth of the rooting zone to provide a work area, as well as further excavation of a horizontal cavity at the desired depth of sampling. The filter candle assembly is placed horizontally in a galvanized sheet metal trough that has the approximate dimensions of the horizontal cavity (Figure 9.2.6). The trough is filled with soil and the assembly is placed in the horizontal hole. Contact with the soil above the trough is ensured by the use of an air pillow or mechanical jack. Sampling is accomplished by using a separately installed tensiometer to measure soil-water tension, and using a vacuum in the system to induce soil-water flow into the trough and candle at the same rate as the surrounding soil. A small diameter tube attached to the other end of the filter candle and extending to the work area in the vertical trench allows rewetting, if necessary.

<u>Method Selection Considerations</u>: Advantages: (1) Allows direct sampling of soil water; (2) successive samples can be obtained from the same depth; and (3) samples both pore water and water flowing through macropores that is intercepted by the trough. Disadvantages: The same as for vacuum-type porous cup samplers, except that disadvantages 4 and 6 do not apply, and the added disadvantage of being more difficult to install.

Frequency of Use: Uncommon, has mainly been used for sampling of irrigation return flow.

Standard Methods/Guidelines: Duke and Haise (1973).

Sources for Additional Information: See Table 9-3.


Figure 9.2.6 Filter candle sampling setup (Everett et al., 1983, after Hoffman et al., 1978).

9.2 DIRECT SOIL-SOLUTE SAMPLING (SUCTION METHODS)

9.2.7 Capillary Wick Sampler

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Soil pore liquid sampling.

<u>Method Description</u>: Capillary wick samplers combine elements of the pan lysimeter (Section 9.3.1) and hollow fiber samplers (Section 9.2.5). A pan with a glass cloth is inserted in the soil below the soil column to be sampled (Figure 9.2.7). Lengths of fiberglass wick are placed in contact with the absorbent material in the pan and suspended vertically over a sampling bottle. When wetted, the wicks apply a continuous tension to the soil pore water equivalent to the wick length (up to -6.0 kPa). The collection chamber can be buried and samples periodically collected through a tube to the surface using a suction sampler (Figure 9.2.7), or the collection bottle can be accessed through a trench installation as with pan lysimeters (Figure 9.3.1a).

Method Selection Considerations: Advantages: (1) Continuous solute samples can be obtained from unsaturated soil without applying suction, minimizing possible affects on volatile contaminants (trench installations); and (2) samples might be more representative of water moving through the soil than samples collected by suction samplers or free-drainage samplers because they can collect soil water from both saturated and unsaturated pores. Disadvantages: (1) Installation is somewhat more complex than for free drainage samplers (Section 9.3.1); (2) solute characteristics might be altered as the solute travels up the wick; (3) is limited to relatively shallow installations (generally 2 meters or less in undisturbed soil); and (4) only custom-built, experimental samplers have been tested to date.

Frequency of Use: Relatively new method that has not been widely tested.

Standard Methods/Guidelines: --

Sources for Additional Information: See Table 9-3.



Figure 9.2.7 Schematic diagram of a capillary wick unsaturated zone soil pore water sampler (Holder et al., 1991, by permission).

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9.3 DIRECT SOIL-SOLUTE SAMPLING (OTHER METHODS)

9.3.1 Free-Drainage Samplers

Other Names Used to Describe Method: Zero-tension samplers, tension-free lysimeter, pan lysimeter, collection lysimeter/manifold collector, trench lysimeter, caisson lysimeter, free-drainage glass block sampler, pan-type collectors, wicking-type sampler.

Uses at Contaminated Sites: Sampling water percolating through the vadose zone; measuring hydraulic conductivity/solute flux.

<u>Method Description</u>: Free-drainage samplers, which intercept and collect water flowing in saturated pores or fractures for delivery to a sample container, are installed in the soil, commonly at depths of interest in the side of a trench or buried culvert. Two major types of installations are possible: (1) **Open trench** or **caisson** (Figures 9.3.1a and b respectively), in which permanent access exists and in which samples are usually collected by simple gravity feed, and (2) **buried trench**, in which the access trench is backfilled after installation and samples are brought to the surface using a suction method (Figure 9.3.1c). Various designs have been developed, including stainless steel troughs, sand-filled funnels, and hollow glass blocks. Geotextile fabric can be used for wicking action. In each case, gravity drainage creates a slightly positive pressure at the soil-sampler interface, allowing the soil water to drip into the sampler. The collection of ground-water outflow from **tile drains** is another way to obtain samples that have recently moved from the vadose zone to the saturated zone. A variant in the tile drain collection method is a collection lysimeter, also called a **manifold collector**, installed at the based of a sanitary landfill to collect leachate (Figure 9.3.1d).

Method Selection Considerations: Advantages: (1) A larger volume of soil can be sampled compared to suction samplers, and the defined surface area might allow quantitative estimates of leachate flux; (2) samples include water moving through both large and small pores and are representative of the soil solute that is actually percolating to greater depths without disturbing natural flow patterns; (3) have less possibility of chemical alteration or loss of volatile compounds from the sample compared to porous-cup samplers; and (4) sampling is continuous without the need for externally applied vacuum. Disadvantages: (1) Installation procedures are time-consuming and complex; (2) samples will not be collected unless gravity flow is occurring; (3) installation under impoundments generally is not feasible; (4) if collection surfaces are not installed perfectly level, a sump or collection area can result in dead space where the soil water cannot be removed; (5) if the collection surface is uneven, the potential exists for cross contamination from residual samples; and (6) safety considerations might limit the depth to which trench lysimeters can be installed. Tile Drain/Collection Lysimeter Advantages: Existing tile drains require no installation, and manifold collectors are relatively easy to install where new landfill pits are excavated. Tile Drain/Collection Lysimeter Disadvantages: (1) NAPLs might not appear in tile drain outflow because they remain above the drain (light NAPLs) or might flow along the bottom of the perched water zone (dense NAPLs); (2) is limited to shallow depth for economic reasons; and (3) the presence of air in the tile lines might alter the chemistry of water flowing into the drain.

Frequency of Use: Relatively uncommon.

Standard Methods/Guidelines: ASTM (1992).

Sources for Additional Information: Wicking-type sampler: Hornby et al. (1986); Trench pan lysimeter: U.S. EPA (1986); Pan lysimeter with tension plate: Shaffer et al. (1979); Free-drainage glass-block sampler: Everett (1990); Collection lysimeters: Sai and Anderson (1991). See also, Table 9-3.



Figure 9.3.1 Free-drainage sampler installations: (a) Trench lysimeter (Parizek and Lane, 1970, by permission); (b) Caisson lysimeter (Morrison, 1983, after Aulenbach and Clesceri, 1980, by permission); (c) Pan lysimeter installation (U.S. EPA, 1986); (d) Leachate collector installed at base of sanitary landfill

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9.3 DIRECT SOIL-SOLUTE SAMPLING (OTHER METHODS)

9.3.2 Perched Water Table

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Characterizing and monitoring vadose zone soil pore liquid.

<u>Method Description</u>: Perched ground water is sampled as representative of water that has percolated through the vadose zone. For shallow perched ground water, samples can be obtained by installing wells (Figure 9.3.2a), piezometer nests (Section 5.4.3), or multilevel samplers (Section 5.6.1), or by installing a tile drainage system and sampling at the outlet. Deeper perched ground water can be obtained by sampling cascading water in existing wells, or by constructing special wells (Figure 9.3.2b). Wells or piezometers screened in perched aquifers are sampled using the appropriate portable ground-water sampling device (see Section 5).

<u>Method Selection Considerations</u>: Advantages: (1) Larger sample volumes are obtained compared to suction and extraction methods (particularly desirable when sampling for organics and viruses); (2) samples are more representative than point samples obtained by suction and extraction methods, because they reflect the integrated quality of water draining from the overlying vadose zone; (3) is cheaper than installing deep wells with batteries of suction samplers; (4) can be located near ponds and landfills without concern about causing leaks; and (5) nested piezometers and multilevel samplers can be used to delineate the vertical and lateral extent of plumes and hydraulic gradients (see Sections 5.4 and 5.6). Disadvantages: (1) Perched zones must be present in the area of concern; (2) detection of perched water deep in the subsurface can be expensive, requiring test wells or geophysical methods; (3) some perched ground water is seasonal and might dry up (backup systems, such as described by Nightingale et al. [1985], are recommended in this situation [see Section 9.2.2]); (4) is most suitable for diffuse sources, such as land spreading areas or irrigated fields; and (5) multilevel sampling is restricted to shallower depths where vacuum pumping is possible.

Frequency of Use: Commonly used, if perched water table is present.

Standard Methods/Guidelines: Sampling from cascading wells: Wilson and Schmidt (1978).

Sources for Additional Information: Everett et al. (1983). See also, Table 9-3.



Figure 9.3.2 Perched water table sampling: (a) Well in perched water table; (b) Cascading water in an idle well (Wilson and Schmidt, 1978, by permission).

9.3 DIRECT SOIL-SOLUTE SAMPLING (OTHER METHODS)

9.3.3 Absorbent Methods

Other Names Used to Describe Method: Cellulose nylon sponge, ceramic rods or points.

Uses at Contaminated Sites: Collecting of soil pore liquids in the vadose zone.

Method Description: Absorbent methods use the ability of a porous material to absorb soil pore water. Cellulose nylon sponge: A sponge is place within a trough, which is positioned against the ceiling of a horizontal tunnel by a series of three-lever hinges. When the sponge has absorbed a certain volume of pore water, the trough is withdrawn and the sponge is placed in a moisture-tight container. In the laboratory, the solution is extracted from the sponge using rollers. Ceramic rods: Tapered ceramic rods or points (90 by 12 millimeters [Figure 9.3.3]) are prepared by boiling in distilled water, drying, and storage in a desiccator. In the field, the rods are taken out, weighed, and driven into the surface soil. After a period of time, the rods are withdrawn and weighed to determine the volume of absorbed water. The points are leached by boiling them in a known volume of distilled water. The solution is analyzed and the original pore water concentration determined from the ratio of water absorbed by the ceramic to the volume of boiling water. Section 9.3.7 describes uses of absorbent collectors with the SEAMIST system.

<u>Method Selection Considerations</u>: Advantages: No clear advantages over alternative methods. Disadvantages: (1) Require near saturated conditions; (2) procedures are relatively complex; (3) representativeness of samples extracted from porous points is questionable.

Frequency of Use: Uncommon.

Standard Methods/Guidelines: --

Sources for Additional Information: Ceramic points: Shimshi (1966); Sponge: Tadros and McGarity (1976).



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Figure 9.3.3 A point made from a discarded ceramic pressure plate used for collecting soil solute samples by absorption; units in millimeters (Shimshi, 1966, by permission).

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9.3 DIRECT SOIL-SOLUTE SAMPLING (OTHER METHODS)

9.3.4 Solids Sampling with Soil-Water Extraction

Other Names Used to Describe Method: Squeezing press, pressure extraction press/filter press, soil press.

<u>Uses at Contaminated Sites</u>: Soil pore water and/or solute sampling during site characterization; soil solute monitoring where suction methods will not work.

<u>Method Description</u>: Soil core samples are collected and soil water is extracted in the field or (more typically) the laboratory by one of a number of methods. Column displacement uses an immiscible fluid that displaces soilpore water in a soil column by gravity. Centrifugation uses a double bottom centrifuge to remove soil water. The displacement/centrifugation method uses a combination an immiscible fluid and a centrifuge. Soil water also can be obtained by squeezing (Figure 9.3.4) or vacuum extraction. The resulting liquid is then analyzed for constituents of interest.

Method Selection Considerations: Advantages: (1) Vertical profiles of concentrations of specific pollutants can be obtained; (2) identification of variations of ionic concentrations in layered sequences is possible; and (3) solids samples can be used for additional analyses, such as grain size, cation exchange capacity, etc. Disadvantages: (1) A large number of samples is required to characterize spatial variability of soil solutes; (2) is expensive if deep sampling is required; (3) changes in soil-water chemistry might occur during preparation and extraction; (4) soil-water samples represent concentrations of moisture content at the time of sampling, ionic concentrations would be different at other moisture contents; (5) is a destructive method, which precludes comparing successive sampling results due to soil variability; (6) core holes might alter infiltration patterns and cause short-circuiting of pollutants to greater depths; and (7) the chemistry of soil water from various extraction methods might differ from the chemistry of soil pore liquid collected using suction and free-drainage samplers (especially if greater pressures are used).

<u>Frequency of Use</u>: Sometimes used during site characterization; rarely used for monitoring because of destructive nature of sampling.

Standard Methods/Guidelines: Squeezing: ASTM (1985).

Sources for Additional Information: See Table 9-3.



Figure 9.3.4 Filter press and chamber assembly for pore water extraction (Luscynski, 1961).

9.3 DIRECT SOIL-SOLUTE SAMPLING (OTHER METHODS)

9.3.5 Solids Sampling with Soil-Saturation Extract

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Measuring water-soluble contaminants and soil minerals.

<u>Method Description</u>: Solids samples are collected using tube samplers or augers, and the electrical conductivity of a saturation extract (prepared in the field or laboratory) is measured. The electrical conductivity measurements are then interpreted in terms of salinity and other properties (Rhoades et al., 1989b,c)

<u>Method Selection Considerations</u>: Advantages: (1) Is a simple procedure; and (2) provides a measure of leaching potential from a soil sample. Disadvantages: (1) A large number of samples is required to characterize spatial variability of soil solutes; (2) is expensive if deep sampling is required; (3) changes in soil-water chemistry might occur during preparation and extraction; (4) sample saturation extract might not be representative of actual soil solution moving through the vadose zone; (5) is a destructive method, which precludes comparing successive sampling results due to soil variability; (6) core holes might alter infiltration patterns and cause short-circuiting of pollutants to greater depths; and (7) the chemistry of soil-saturation extracts will not be comparable to soil pore liquid collected using suction and free-drainage samplers.

Frequency of Use: Commonly used in arid and semi-arid areas where soluble salt concentrations in soils are high.

Standard Methods/Guidelines: Rhoades (1982), SCS (1984, Section 8E).

Sources for Additional Information: See Table 9-3.

9.3 DIRECT SOIL-SOLUTE SAMPLING (OTHER METHODS)

9.3.6 Solids Sampling for Volatile and Microbial Constituents

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Collecting uncontaminated samples for constituents that might be sensitive to exposure to air.

<u>Method Description</u>: Special sampling procedures are required for sampling contaminants that can change in concentration (degassing of volatile compounds) or chemical composition (redox-sensitive chemical species, such as ferrous and ferric iron) when exposed to the air. Similar care is required when sampling for microbiota in the subsurface, especially where oxygen content is low (typically in the zone of saturation). Even where exposure to the air is not a concern for microbiological sampling (typically in the vadose zone), special care is required to make sure that the sample has not been cross contaminated with soil microorganisms from higher soil horizons. The basic procedure involves collection of subsamples of power-driven sample cores (Section 2.4), using smaller diameter corers. Figure 9.3.6a shows suggested locations for microbial and volatile samples from a core. Samples for volatiles should be quickly transferred to the sample container and sealed with no air headspace in the container. Where cores contained anaerobic bacteria and chemical species of concern that are in a reduced state, samples need to be extracted in an oxygen-free environment. Figure 9.3.6b shows a plexiglass field glove box for collecting such samples. Sample containers are sterilized and filled with an inert gas such as nitrogen. In the field, the sealed containers are placed in the field glove box, before the box is filled with nitrogen. The core sample is pushed into the box through an iris port, and a core paring tool is used to collect subsamples in the oxygen-free environment for placement in sample containers.

<u>Method Selection Considerations</u>: Required whenever accurate measurement of volatiles and microorganisms in soil samples is necessary. The more complex glove box procedure should be used when accurate identification of reduced metal species and/or anaerobic microorganisms is required.

Frequency of Use: Relatively uncommon. Should probably be used more commonly.

Standard Methods/Guidelines: Dunlap et al. (1977), Leach et al. (1988).

Sources for Additional Information: Beeman and Suflita (1989), Board and Lovelock (1973), Bordner et al. (1978), Gilmore (1959), Phelps et al. (1989), Russell et al. (1992)



Figure 9.3.6 Solids sampling for microbiological and volatile contaminants: (a) Core subsample (Dunlap et al. 1977); (b) Field sampling glove box (Leach et al., 1988, by permission).

9.3 DIRECT SOIL-SOLUTE SAMPLING (OTHER METHODS)

9.3.7 SEAMIST.

Other Names Used to Describe Method: --

<u>Uses at Contaminated Sites</u>: Sampling of soil-pore liquids and soil gases; measuring air permeability; might eventually be adapted for ground-water sampling.

<u>Method Description</u>: SEAMIST (Science and Engineering Associates Membrane Instrumentation and Sampling Technique) is a recently developed system that involves the placement of a membrane packer in an open borehole (Figure 9.3.7a-c). Soil-gas sampling ports attached to flexible tubing are attached to the membrane to create an in situ multilevel sampling system (Figure 9.3.7d). Multilevel soil pore liquid sampling is accomplished by the placement of absorbent collectors on the outside of the membrane, with leads for measuring electrical resistance running up the inside of the membrane (Figure 9.3.7d). Stabilization of the resistance readings serves as an indicator that the absorbent pad has equilibrated with the moisture content of the borehole wall. The flexible membrane is then retrieved by a reversal of the process shown in Figure 9.3.7a-c, and the absorbent pads are removed for fluid extraction in the laboratory (see Section 9.3.3).

<u>Method Selection Considerations</u>: Advantages: (1) A unit supports the hole wall against the sloughing, eliminating the need for casing and backfilling, provided the borehole is basically stable; (2) multi-level soil-pore liquid and gas sampling from the same borehole is possible, and the method potentially can be used with any type of instrumentation that can be fastened to the membrane fabric; and (3) materials are relatively inexpensive, allowing permanent installation, if desired. Disadvantages: (1) Cannot be used in unstable boreholes (i.e., heaving sands); and (2) is a new technique for which there has been relatively little experience or independent testing.

<u>Frequency of Use</u>: New method for which there is relatively little experience. The U.S Department of Energy is providing research and development funding for this technique.

Standard Methods/Guidelines: --

Sources for Additional Information: Keller (1991, 1992), Keller and Lowry (1990, 1991), Lowry and Narbutobskih (1991), Mallon et al. (1992).



Figure 9.3.7 The basic operation of the SEAMIST system: (a) Insertion of packer; (b) Emplacement of membrane; (c) Enlarged view of bottom emplaced membrane (Lowry and Narbutovskih, 1991, by permission); (d) Collection of pore liquid with an absorbent pad and pore gas via an evacuated tube (Keller and Lowry, 1990).

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9.4 GASEOUS PHASE CHARACTERIZATION

9.4.1 Soil-Gas Sampling (Static)

Other Names Used to Describe Method: Passive sampling.

Uses at Contaminated Sites: Detecting volatile contaminants in the unsaturated zone.

<u>Method Description</u>: Static sampling can be done in two ways. First, an in situ adsorbent (usually an activated charcoal rod) is buried in the soil for a few days to weeks (Figure 9.4.1a and b). The adsorbent is retrieved and analyzed for volatile organic compounds in a laboratory by mass spectrometry or gas chromatography. Second, static grab samples are collected from containers placed in the soil surface, which collect quiescent soil-gas samples. These samples usually are analyzed in the field using portable analytical instruments (see Sections 10.3.1 and 10.3.2).

<u>Method Selection Considerations</u>: Advantages: (1) Is inexpensive and easy to install; (2) laboratory analysis usually provides more precise measurement than field measurement of dynamic samples; (3) if sorption capacity of the sampler is not exceeded, average flux of contaminants to the surface can be calculated; and (4) field operations require minimal training. Disadvantages: (1) Is sensitive to exposure time and insufficient exposure might result in a false negative and overexposure (saturation of sorbent) might mask relative difference in soil-gas contamination at different sampling locations; (2) vertical profiles of soil-gas concentrations are more difficult to obtain than with soil probes; (3) results using in situ absorbent samplers are not available for days to weeks because desorption and laboratory analysis are time consuming; and (4) might not be appropriate for VOCs that have low boiling points (<5 degrees C) or compounds that are prone to thermal decomposition during pyrolysis.

Frequency of Use: Common.

Standard Methods/Guidelines: ASTM (1991a).

Sources for Additional Information: Vroblesky et al. (1992); see also, references for Section 9.4.2.



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9.4 GASEOUS PHASE CHARACTERIZATION

9.4.2 Soil-Gas Probes

Other Names Used to Describe Method: Dynamic grab samples, headspace sampling.

Uses at Contaminated Sites: Detecting and monitoring of volatile organic contaminants in the unsaturated zone.

<u>Method Description</u>: Dynamic soil-gas grab samples are collected from a moving stream of soil gas, which is pumped through a hollow probe that is driven into the soil (Figure 9.4.2a), or from permanently installed tubes at one or more levels in the soil (see Figure 5.5.2c). The probes can be manually or pneumatically driven, or installed in boreholes. Relatively nonvolatile NAPLs can be detected using steam injection (Figure 9.4.2b). The samples usually are analyzed in the field using portable analytical instruments. Grab samples usually are taken at the same depth at a number of surface locations for areal characterization of soil-gas concentrations. Where the vadose zone is thick, or discontinuous impermeable layers exist at a site, samples can be taken at different depths at the same location in order to define vertical changes in soil-gas concentration. An adaptation of the method has been used to detect zone of contaminated discharge to streams using bottom sediment gas bubbles (Figure 9.4.2c).

Method Selection Considerations: Advantages: (1) Is a nondestructive method; (2) hollow-probe samplers allow collection of multiple samples in a relatively short period of time; (3) when combined with on-site gas chromatography, results are available in a matter of minutes; and (4) problems associated with handling and transporting gas samples are minimized. Disadvantages: (1) Grab-sampling results are highly depth dependent and sampling results might be misleading if the correct depth is not sampled (based on site-specific factors, such as moisture conditions, air-filled porosity, and depth to ground water, and compound-specific factors, such as solubility, volatility, and degradability); (2) dynamic sampling perturbs local VOC concentrations as a result of pumping to retrieve sample; and (3) nonvolatile contaminants, if present, will not be detected. Table 9.4.2 provides specifications for a variety of commercially available soil-gas sampling probes.

Frequency of Use: Widely used for preliminary site characterization where volatile contaminants are known or suspected.

Standard Methods/Guidelines: ASTM (1991a). Probe and well sampling: Ford et al. (1984).

Sources for Additional Information: API (1985, 1991), Devitt et al. (1987), Kerfoot (1991), Kerfoot and Barrows (1987), Pitchford et al. (1988), Rector (1991-radon detection), Robbins (1990), Vroblesky and Lorah (1991).



Figure 9.4.2 Dynamic soil-gas sampling systems: (a) Soil vacuum withdrawal (Pitchford et al., 1988); (b) Response of PID detector to soil-gas samples with and without steam injection (Kerfoot, 1991, by permission); (c) Water collection system (Vroblesky and Lorah, 1991, by permission).

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Source: Pollution Equipment News (April 1993)

9.4 GASEOUS PHASE CHARACTERIZATION

9.4.3 Tank/Pipeline Leak Sensors

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Detecting leaks in underground storage tanks and pipelines.

Method Description: Numerous techniques have been developed to detect leaks in underground storage tanks and pipelines (Figure 9.4.3). Inventory monitoring involves identification of discrepancies in tank storage between additions and withdrawals and can be accomplished by manual tank gauging and reconciliation, statistical reconciliation, or using automatic gauging systems. Various methods can be used for soil or ground-water release detection: (1) Sampling of detection wells, (2) soil sampling, (3) using dyes and tracers, and (4) surface geophysics. Vadose zone vapor detection methods include: (1) Grab sampling or soil coring, (2) surface flux chambers (see Section 9.5.2), (3) downhole flux chambers; (4) accumulator systems, and (5) soil-gas probe testing (Section 9.4.2). Secondary containment with interstitial monitoring provides one of the safest leak detection methods because any releases are prevented from entering soil or ground water. Physical test methods include visual inspection using remote cameras (see Section 3.5.7) and tightness testing of tanks and piping. Over 90 leak detection systems are available that involve detection of organic vapors as an indication that underground storage tanks are leaking. Vapor wells and U-tubes are commonly used. More than 200 liquid hydrocarbon and hydrocarbon vapor detectors or sensors are available. Sensor systems can range from systems with alarms that go off when vapors are detected to systems that monitor product flow into and out of the tank and identify discrepancies that might be related to leakage. Table 9.4.2 provides information on a number of commercially available tank leak monitor systems.

<u>Method Selection Considerations</u>: The appropriate state and/or federal regulations should be consulted to determine the types of leak detection systems that should be used.

<u>Frequency of Use</u>: Standard requirement for any new installation of underground storage tanks containing hydrocarbons and other potentially hazardous liquids.

Standard Methods/Guidelines: ASTM (1991b, 1993).

Sources for Additional Information: Boone et al. (1991), Cochran (1987), Durgin and Young (1993), Eckert and Maresca (1992), Eklund and Crow (1987), Fromme et al. (1991), Lyman and Noonan (1990), Maresca and Hillger (1991), Maresca et al. (1991), Morrison and Mioduszewski (1986), Niaki and Broscius (1986), Portnoff et al. (1991), Scheinfeld and Schwendeman (1985), Scheinfeld et al. (1986), Schwendeman and Wilcox (1987), Starr et al. (1991b).



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Figure 9.4.3 Examples of leak detection methods for tanks and piping (Floyd, 1993).

9.4 GASEOUS PHASE CHARACTERIZATION

9.4.4 Air Pressure

Other Names Used to Describe Method: --

<u>Uses at Contaminated Sites</u>: Can be used as an indirect expression of soil structural properties because an air pressure buildup usually affects the relative magnitude of the air permeability and hydraulic conductivity, which are both indices of soil structure.

<u>Method Description</u>: Various types of manometers can be used, including: (1) Pressure transducers, (2) fluid manometers, and (3) aneroid barometers. The manometer is attached to an air-filled access tube that is placed in the soil (Figure 9.4.4). Changes in air pressure with time are measured in response to events like wetting of a dry soil. Care must be taken to ensure that the access tube fits tightly in the surrounding soil.

<u>Method Selection Considerations</u>: Flühler et al. (1986) consider soil-air pressure to be one of the most frequently neglected variables in soil physics research. Air pressure measurements might have value for site characterization, but their utility have not be evaluated in this context.

Frequency of Use: Uncommon.

Standard Methods/Guidelines: Flühler et al. (1986).

Sources for Additional Information: --

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Figure 9.4.4 Diagram of a soil air pressure gauge (Flühler et al., 1986, by permission).

9.4 GASEOUS PHASE CHARACTERIZATION

9.4.5 Gas Permeability and Diffusivity

Other Names Used to Describe Method: Air permeability.

<u>Uses at Contaminated Sites</u>: Measuring gas permeability and diffusivity for modeling vapor transport in the vadose zone and designing vapor extraction remediation systems.

<u>Method Description</u>: The process of testing for air permeability is analogous to a multiple-well ground-water pumping test (Section 4.3.2). A vacuum is applied to a "pumping" well, with a screened interval in the soil zone of interest while changes in air pressure with time are monitored in pressure probes placed in the subsurface (Figure 9.4.5a). Johnson et al. (1990) provide formulas for calculating air permeability from the measured data. Baehr and Hult (1991) describe a more complex installation using multi-level pressure probes and two pumping wells separated by a confining bed. Section 7.4.4 describes use of air permeability in the deep vadose zone to estimated hydraulic conductivity. Gas diffusion is the principle mechanism for exchange of gases between the soil and the atmosphere, and hence is of interest for evaluating the potential for movement of volatile contaminants from the soil to the air. Gas diffusivity is measured in the field by injecting a known concentration of the gas of interest into a sealed cylinder (Figure 9.4.5b). The air in the confined space above the soil is kept stirred with a fan run by a hand drill, and the chamber is sampled over time to determine the change in gas concentration. Diffusivity is then calculated based on the decrease in concentration of the gas with time. A prototype probe for measuring gas diffusivity in a borehole has also been developed (Figure 9.4.5c).

<u>Method Selection Considerations</u>: If measurement of either air permeability or gas diffusivity is required, use of either method is relatively straightforward.

Frequency of Use: Will become increasingly common, especially where vapor extraction remediation activities are involved.

<u>Standard Methods/Guidelines</u>: Air permeability: ASTM Draft Standard Practice for Determining Air Permeability in Soils (Nielsen, 1991); Corey (1986-laboratory measurement using cores); Gas diffusivity: Rolston (1986a).

Sources for Additional Information: Baehr and Hult (1991), Bakker and Hidding (1970), Corey (1957, 1986), Evans and Kirkham (1949), Groenewoud (1968), Grover (1955), Havlena and Stephens (1992), Izadi and Stephenson (1992), Johnson et al. (1990), Kearl et al. (1990), Kirkham (1946), Lowry and Narbutobskih (1991), Marrin et al. (1991), Pirkle et al. (1992), Reeve (1953), Rogers and Nielsen (1991), Springer et al. (1991), Stonetrom and Rubin (1989), Weeks (1978), Weinig (1992); Diffusion: Jellick and Schnabel (1985), Kearl et al. (1988), Rolston et al. (1991).



Figure 9.4.5 Air permeability and gas diffusivity: (a) Schematic of air permeability test system (Johnson et al, 1990, by permission); (b) Schematic diagram of a field diffusion apparatus (Rolston et al., 1991); (c) Illustration of equipment configuration for in situ borehole gas diffusion measurements (Kearl et al., 1988).

9.5 CONTAMINANT FLUX

9.5.1 Solute Flux Methods

<u>Other Names Used to Describe Method</u>: Various solute flux methods are available: (1) Average concentration method, (2) approximate analytic solution method, (3) long-term flux estimation method, and (4) short-term water and solute movement estimation method.

Uses at Contaminated Sites: Estimating the mass transfer of pollutants from the vadose zone to ground water.

Method Description: Several methods are available to calculate solute flux individual constituents or parameters of interest. Average concentration method: Soil-water concentrations from direct samples obtained using one of the methods described in this section for two different times are averaged. Average flux over the time period can be calculated using estimates of water flux during the same time period (using methods described in Section 7.5). Approximate analytical solution method: This is similar to the average concentration method, except an approximate analytic solution that considers diffusion-dispersion is used to estimate solute concentrations rather than measuring them directly. This method requires monitoring of soil-moisture changes with time (Section 6.3), adding a solution to the soil with known concentration, and estimating average water flux (Section 7.5.1) requiring sampling of soil water at selected depth intervals at two times using one of the direct soil-water sampling methods discussed in this section. Short-term water and solute flux estimation method: This is a relatively complex method estimating peak solute concentration in the rooting zone after infiltration and redistribution of the soil water to field capacity.

<u>Method Selection Considerations</u>: These methods primarily have been developed for agricultural applications to calculate flux of nutrients and soluble salts in soil. The average concentration method is the simplest method, with potential use for mass balance analysis of pollutants at contaminated sites, provided that water flux can be estimated. The other methods require field water-budget measurements of varying complexity and are of relatively limited applicability to contaminated sites.

Frequency of Use: Commonly used in agricultural applications; use at contaminated sites increasing.

Standard Methods/Guidelines: Brown et al. (1983), Wagenet (1986).

Sources for Additional Information: Parker and Van Genuchten (1984), Philip (1973), Roth et al. (1990), U.S. EPA (1975).

9.5 CONTAMINANT FLUX

9.5.2 Soil-Gas Flux

Other Names Used to Describe Method: --

<u>Uses at Contaminated Sites</u>: Quantifying sources and sinks of gases within the soil; evaluating soil microbial activity; measuring the rate of flow of gases from volatile subsurface contaminants to the surface.

<u>Method Description</u>: Various methods have been developed: (1) Gas samples taken at different depths and over a period of time are collected and flux is calculated using Fick's law; (2) a closed chamber is placed over the soil surface and the increase in concentration of gas within the chamber is measured as a function of time (Figure 9.5.2); and (3) a capped cylinder is driven into the ground surface (flow-through chamber method). Ambient air is drawn through the chamber and concentration of the gas of interest is measured in both the input and output streams.

<u>Method Selection Considerations</u>: These methods primarily have been used in research related to gases of interest for agriculture (nitrogen and carbon dioxide).

Frequency of Use: Relatively common.

Standard Methods/Guidelines: Ball and Smith (1991), Rolston (1986b).

Sources for Additional Information: Aulach et al. (1991), Gholson et al. (1989), Loftfield et al. (1992), Matthias et al. (1980).



Figure 9.5.2 Diagram of closed chamber for directly measuring gas flux at the soil surface (Rolston, 1986b, after Matthias et al., 1980, by permission).

Topic References Everett et al. (1982, 1983), Kaufman et al. (1981), Rhoades (1978, 1984), Rhoades Overviews and Oster (1986), Wilson (1983), Yadav et al. (1979) Bottraud and Rhoades (1985), Gupta and Hanks (1972), Halvorson et al. (1977), **EC-Salinity** Calibration Klute and Letey (1958), Mualem and Friedman (1991), Rhoades (1980, 1981), Rhoades et al. (1976, 1977, 1989a-c, 1990), Shainberg et al. (1980), van Hoorn (1980) Austin and Rhoades (1979), Bohn et al. (1982), Cameron et al. (1981), Halvorson Four Electrode and Reule (1976), Halvorson and Rhoades (1974, 1976), Nadler (1981, 1991), Nadler and Frenkel (1980), Nadler et al. (1984, 1990), Rhoades and Halvorson (1977), Rhoades and Ingvalson (1971), Roux (1978), van Hoorn (1980); Soil Moisture: Bunnenberg and Kuhn (1980), Edlefson and Anderson (1941), Kirkham and Taylor (1950) EC Probes Nadler et al. (1982), Rhoades (1979), Rhoades and Halvorson (1977), Rhoades and van Schilfgaarde (1976), Shea and Luthin (1961) Austin and Oster (1973), Enfield and Evans (1969), Ingvalson et al. (1970, 1976), Salinity Sensors Kemper (1959), Oster and Ingvalson (1967), Oster and Willardson (1971), Oster et al. (1973, 1976), Reicosky et al. (1970), Rhoades (1972), Rhoades and Oster (1986), Richards (1966), Todd and Kemper (1972), U.S. Soil Salinity Staff (1981), Wesseling and Oster (1973), Wierenga and Patterson (1974), Wood (1978a), Yadav et al. (1979); Temperature Correction Coefficients: Campbell et al. (1948), Richards (1954), Richards and Campbell (1948), Whitney and Means (1897) Cameron et al. (1981), Cook and Walker (1992), Corwin and Rhoades (1982, EMI Sensors 1984), de Jong et al. (1979), Hendrickx et al. (1992), Kachonoski et al. (1988-soil-

water content), McBride et al. (1990), Rhoades and Corwin (1981), Rhoades and

Oster (1986), Williams and Baker (1982), Wollenhaupt et al. (1986)

Table 9-2 Reference Index for Indirect Methods for Monitoring Solute Movement

Topic	References
Overviews	Dorrance et al. (1991), Everett (1990), Everett et al. (1982, 1990), Hornby et al. (1986), Kohnke et al. (1940), Litaor (1988), Morrison (1983), Nagpal (1982), Rhoades and Oster (1986), Robbins and Gemmell (1985), Silkworth and Grigal (1981), U.S. EPA (1986), Wilson (1983, 1990)
Suction Samplers	•
Chemical Effects	Alberts et al. (1977), Anderson (1986), Bottcher et al. (1984), Creasy and Dreiss (1985), Dazzo and Rothwell (1974), England (1974), Grover and Lamborn (1970), Haines et al. (1982), Hansen and Harris (1975), Hornby et al. (1986), Levin and Jackson (1977), McGuire et al. (1992), Peters and Healy (1988), Rhoades and Bernstein (1971), Severson and Grigal (1976), Silkworth and Grigal (1981), Stearns et al. (1980), Suarez (1987), Tasi et al. (1980), Wagner (1962), Wolff (1967), Zabowski and Ugolini (1990)
Physical Effects	Cochran et al. (1970), Kung and Donohue (1991), Morrison and Lowery (1990- sampling radius), Narasimham and Dreiss (1986), Severson and Grilgal (1976), Talsma et al. (1979), van der Ploeg and Beese (1977), Warrick and Amoozegar- Fard (1977)
Porous Cup Cleaning Procedures	Aulenbach and Clesceri (1980), Creasey and Dreiss (1988), Grover and Lamborn (1970), Neary and Tomassini (1985)
Porous Cup (Vacuum)	Ahlert et al. (1976), Alberts et al. (1977), Angle et al. (1991), Ballestero et al. (1990), Barbarick et al. (1979), Barbee and Brown (1986), Bell (1974), Bourgeois and Lavkulich (1972a,b), Brooks et al. (1958), Brown (1986), Burns (1992), Chow (1977a, 1977b-fritted glass), Debyle et al. (1988), de Jong (1976), Dugan et al. (1975), Eleuterius (1980), Grier et al. (1977), Haines et al. (1982), Hansen and Harris (1975), Joslin et al. (1987), Knighton and Streblow (1981), Krone et al. (1951), Miller (1992), Nielsen and Phillips (1958-fritted glass), Quin and Forsyth (1976), Reeves and Doering (1965), Riekerk and Morris (1983), Shuford et al. (1977), Silkworth and Grigal (1981), Smith and Carsel (1986), Starr (1985), Starr et al. (1991a), Suarez (1986), Tyler and Thomas (1977), Wagner (1962, 1965), Wengel and Griffin (1971); <u>Cup Material Comparisons</u> : McQuire and Lowery (1992)
Porous Cup (Vacuum- Pressure)	Apgar and Langmuir (1971), Ball and Coley (1986), Banton et al. (1992), Biggar and Nielsen (1976, 1978), Biggar et al. (1975), Brose et al. (1986), Everett and McMillion (1985), Everett et al. (1984, 1988), Fenn et al. (1977), Gerhardt (1977), Hounslow et al. (1978), Johnson and Cartwright (1980), Johnson et al. (1981), Long (1978), Merry and Palmer (1985), Morrison (1982), Morrison and Szecsody (1985, 1987), Morrison and Tsai (1981), Parizek and Lane (1970), Peters and Healy (1988), Quin and Forsyth (1976-nylon), Rehm et al. (1987), Starr et al. (1978), Tsai et al. (1980), U.S. EPA (1986), Wood (1978b), Wood et al. (1981), Young (1985), Yu et al. (1978), Zimmermann et al. (1978); <u>High-Pressure</u> <u>Vacuum</u> : Bond and Rouse (1985), Wood (1973), Wood and Signor (1975)

Table 9-3 Reference Index for Direct Soil-Water Sampling Methods

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Table 9-3 (cont.)

Торіс	References
Suction Samplers (cont.)	
Vacuum Plates	Chow (1977b), Cochran et al. (1970), Cole (1958), Cole et al. (1961), Duke et al. (1970), Haines et al. (1982), Iskandar and Nakano (1978), Neary and Tomassini (1985), Tanner et al. (1954), van der Ploeg and Beese (1977)
Membrane Filter	Everett (1990), Everett et al. (1982), Stevenson (1978), U.S. EPA (1986), Wagemann and Graham (1974), Wilson (1990)
Hollow Fiber	Everett (1990), Jackson et al. (1976), Levin and Jackson (1977), Morrison (1982), Silkworth and Grigal (1981), U.S. EPA (1986), Wagemann and Graham (1974)
Ceramic Filter Candle	Duke and Haise (1973), Everett (1990), Hergert and Watts (1977), Hoffman et al. (1978), Montgomery et al. (1987), Smith and McWhorter (1977), van Shilfgaarde (1977), U.S. Soil Salinity Laboratory Staff (1981)
Monolith Lysimeters	Belford (1979), Brown (1986), Brown et al. (1974, 1985), Cameron et al. (1992), Merek et al. (1988), Persson and Bergström (1991)
Filled-Type Lysimeters	Tyler (1981), Upchurch et al. (1973)
Capillary Wick Samplers	Boll et al. (1991, 1992), Brown et al. (1988), Holder et al. (1991), Politeka et al. (1992)
Other Direct Soil Water Sampling	Methods
Free-Drainage Samplers	Aulenbach and Clesceri (1980), Barbee and Brown (1986), Boll et al. (1991), Fenn et al. (1977), Haines et al. (1982), Hornby et al. (1986), Jordon (1968), Kmet and Lindorf (1983), Parizek and Lane (1970), Radulovich and Sollins (1987), Rehm et al. (1987), Russell and Ewel (1985), Shaffer et al. (1979), Starr et al. (1991a), Tyler and Thomas (1977), U.S. EPA (1986), Wilson and Small (1973); <u>Caisson Lysimeter</u> : McMichael and McKee (1966), Schmidt and Clements (1978), Schnieder and Oaksford (1986), Schneider et al. (1983); <u>Buried Cup</u> : Miller (1992)
Tile-Drainage Sampling	Richard and Steenhuis (1988), Starr et al. (1991a), Thomas and Barfield (1974), Willardson et al. (1973)
Perched Water Table	Miller (1992), Starr et al. (1991a), Wilson and Schmidt (1978); <u>Multi-Level</u> <u>Samplers</u> : Cherry and Johnson (1982), Hansen and Harris (1974, 1980), Pickens et al. (1981), Smith at al. (1982); see also, Section 5.5.3

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Table 9-3 (cont.)

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Торіс	References
Other Direct Soil Water Sa	ampling Methods (cont.)
Solids Sampling (Soil-	
Water Extraction)	 <u>Solvent/Fluid Column Displacement</u>: Adams (1974), Adams et al. (1980), Barrow (1982), Batley and Giles (1979), Kittrick (1980, 1983); <u>Double-Bottom Centrifuge</u>: Adams et al. (1980), Dao and Lavy (1978), Davies and Davies (1963), Edmunds and Bath (1976), Elkhatib et al. (1986, 1987), Fenn et al. (1977), Gillman (1976), Zabowski (1989), Zabowski and Ugolini (1990); <u>Immiscible Fluid</u> <u>Displacement/Centrifugation</u>: Mubarak and Olsen (1976, 1977), Phillips and Bond (1989), Whelan and Barrow (1980); <u>Squeezing Displacement</u>: Fenn et al. (1977), Lusczynski (1961), Manheim (1966), Patterson et al. (1978); <u>Vacuum</u> <u>Displacement</u>: Fenn et al. (1977), Richards (1954-pressure membrane apparatus), Wolt and Graveel (1986); <u>Unclassified</u>^a: Behel et al. (1983), Brown (1986), Kinniburgh and Miles (1983); Lucas and Reeves (1980), Pratt et al. (1976), Rible et al. (1976), Wellings and Bell (1980), Yamasaki and Kishita (1972)
Solids Sampling (Soil- Saturation Extract)	Barbarick et al. (1979), Campbell et al. (1948), Moran et al. (1978), Rhoades (1981, 1982), Rhoades and Bernstein (1971), Rhoades et al. (1989a-c), Richter and Jury (1986), SCS (1984), Wilson (1983); <u>Chemical Effects</u> : Reitmeier (1946)

*It was not possible to review these references to determine what extraction method was used.

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SECTION 9 REFERENCES

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SECTION 10

FIELD SCREENING AND ANALYTICAL METHODS

The term "field screening" has gained widespread use in recent years to describe a wide variety of methods for chemical characterization of contaminated sites. In this guide, a distinction is made between field screening and field analytical methods. Field screening methods provide an indication of the presence or absence of a particular chemical or chemical class of concern, or provide an indication of whether the chemical or chemical class of concern is above or below a predetermined threshold. Screening methods provide relative concentrations for chemical classes, but rarely provide chemical-specific information. This definition is more restrictive than those usually found in the literature. Field analytical methods include all chemical analysis methods capable of providing chemical-specific quantitative data in the field or non-laboratory setting. Field analytical techniques generally are more rapid and less expensive than similar chemical analyses performed in laboratories with fixed facilities. Field screening and analytical techniques can be classified as portable (require no external power source, are compact, and are rugged enough to be carried by hand into the field), fieldable (require limited external power, are compact, and are rugged enough to be transported in a small van, pick-up, or four-wheel drive), or mobile (are small enough to carry in a mobile laboratory, which is feasible for most analytical instruments although power considerations can be a limitation). The standard by which the sensitivity, precision, and accuracy of field screening techniques are measured are those obtained in fixed-base laboratories in EPA's contract laboratory program (CLP). An intermediate option for analysis of samples is the use of a dedicated laboratory using CLP procedures but involving more rapid turnaround time (as short as overnight) for sample results.

Field Analysis versus CLP Analytical Laboratory

Key advantages of field analytical techniques include: (1) Results can be obtained within hours, compared to the 20 to 40 days required for CLP laboratories, which allows for more rapid definition of the scope of contamination and allows for optimal selection of permanent monitoring wells/locations; (2) lower cost per sample (commonly one-tenth CLP cost) allows for more detailed characterization of contaminant distribution and/or reduced overall costs; and (3) the techniques are best suited for preliminary site characterization, emergency remedial actions, and monitoring of remediation activities. Some general disadvantages of field analytical techniques include: (1) Application of analytical QA/QC procedures is more difficult in the field; (2) generally, less sophisticated instrumentation and disadvantage #1 results in generally higher detection limits and lower precision and accuracy compared to CLP laboratories; and (3) disadvantages #1 and #2 mean that data are more liable to challenge by litigation.

Cost differences between field analysis and laboratory analysis are strongly dependent on the number of samples from a site that must be analyzed, with the cost advantage tending to shift to field analysis as the number of samples increases. For example, if less than 30 to 50 samples are required, laboratory gas chromatograph analyses are likely to be less expensive than using portable or mobile GCs. Similarly, around 50 to 80 samples for field X-ray fluorescence analysis of metals are required to save money over conventional laboratory XRF analyses.

Overview of Specific Techniques

Developments in miniaturization and computer processing of analytical signals and development of innovative analytical techniques mean that almost any instrumental or analytical technique has the potential for being used for field screening. Any attempt to publish a comprehensive compilation of techniques that have been proposed or tested is doomed to be out-of-date before it reaches print. This section, therefore, provides a reasonably comprehensive overview of the state-of-the-art as of early 1993. Table 10-1 provides summary information on over 80 techniques. Techniques are grouped into the following major categories: (1) Routine chemical field measurement techniques (Section 10.1 [chemical sensors covered in Chapter 5 also are indexed under this section in the table]); (2) major sample extraction procedures (Section 10.2); (3) analytical techniques that detect gases or require creation of a gaseous phase during the analytical process if the gaseous phase is not already present (Section 10.3); (4) luminescence, spectrophotometric, and other spectroscopic techniques (Section

Table 10-1 Summary Information on Sample Processing/Analytical Techniques

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Technique/Instrumentation	Technology Status [•]	Sample Matrix	Contaminant Type ^e	Detection Limit ⁴	Section/Table
Chemical Field Measurement Techniqu	ies/Sensors				
ph/Alkalinity/Acidity	I/CP	W,S			10.1.1, 5.5.4
Eh	I/CP	W,S			10.1.2, 5.5.4
Dissolved Oxygen	I/CP	W		ppm	10.1.2, 5.5.4
Temperature	I/CP	W			10.1.3
Electrical Conductance	I/CP	W,S			10.1.3
Filterable Residue	I/CM	W			10.1.3
Other Specific Ion Electrodes	II/CP	W	EA	ppm	5.5.5
Solid/Porous Fiber Optic		W,S,A	VOC	ppm	5.5.0 5.5.6 10.5.2
Immunochemical Fiber Optic		W,S,A	SVO VOCTC	ppo-ppm	5.5.0, 10.5.2 10.6.5
Electrochemical Sensors		w,A	VOCTO	poo-ppm	10.6 5
SAW PIOOCS Bieroelectric Sensor		A A	VOCIO	ppin	10.6.5
Semiconductor Sensors	IV IV	A W	VOC	nnm-%	10.6.5
Seniconductor Sensors			100	ppm-70	10.0.5
Sample Extraction Procedures					
Headspace Analysis	I	Α	VOC		10.2.1
Vacuum Extraction	I/CP	A	VOC		10.2.1
Purge and Trap	I/CP	W	VOC		10.2.2
Solvent Extraction	I/CP	S	SVO,VOC		10.2.3
Thermal Digestion	II/CP	W,S	EA,HM		10.2.4
Thermal Extraction	II/CP	W,S	SVO		10.2.4
Thermal Desorption	III/CP	W,S	VOC,SVO		10.2.4
Supercritical Fluid Extract.	III/CP	W,S	VOC,SVO		10.2.5
Membrane Extraction	IV	W	VOC		10.2.5
Sorbent Extraction	I/CP	A,W	VOC,SVO		10.2.5
Gaseous Phase Analytical Techniques					
Photo-Ionization Detector	I/CP	Α	voc	ppb-ppm	10.3.1
Flame-Ionization Detector	I/CP	Α	VOC	ppb-ppm	10.3.1
Argon-Ionization Detector	III/CP	Α	VOC	100s ppb-ppm	10.3.1
Explosimeter	I/CP	Α	VOC	%	10.3.2
Catalytic Surface Oxidation	I/CP	Α	VOC,TG	ppm-%	10.3.2
Detector Tubes	I/CP	Α	VOC,TG	high ppm	10.3.2
Gas Chromatography (GC)	II/CP,CM	A,W	VOC,SVO,TR	ppb-ppm	10.3.3/Table 10-3
Mass Spectrometry (MS)	II/CF,CM	Α	VOC,SVO,TR	ppm	10.3.4/Table 10-3
GC/MS	II/CM	A	VOC,SVO,TR	ppb	10.3.4/Table 10-3
Ion Trap MS	IV	A	VOC,SVO	ppb-ppm	10.3.4
AA Spectrometry	II/CM	A,W	EA,HM	ppb-ppm	10.3.5/Table 10-3
ICP-AES		A,W	EA,HM	ppo-ppm	10.3.0/Table 10-3
Ion Mobility Spectrometer	II/CP	A	v0C,Sv0,1G	ррі-ррт	10.3.7
Luminescence/Spectroscopic Technique	<u>25</u>				
X-Ray Fluorescence	II/CP,CM	S,W	HM	10s-100s ppm	10.4.1
UV Fluorescence	II	S,W	VOC	sub ppm	10.4.2/Tables 10-3, 10.4.2
Room-Temp. Phosphorimetry	111	S,W	VOC,SVO	ppb-ppm	10.4.2/Table 10.4.2
Synchronous Luminescence	111	W	VOC,SVO	ppm	10.4.2
Synchronous Fluorescence	111	W	VOCSVO	ppm	10.4.2/Table 10.4.2
UV-Visible Spectrophotometry		A,W	VOC	ppb-ppm	10.4.3/Tables 10-3, 10.4.3
Inirared Spectroscopy	11	A,W,S	M,VUC,SVU	ppm-1000s ppm	10.4.3/1 adies 10-3, 10.4.3

Technique/Instrumentation	Technology Status [®]	Sample Matrix	Contaminant Type ^e	Detection Limit ⁴	Section/Table
Luminescence/Spectroscopic Technique	s (cont.)				
FTIR Spectroscopy Scattering/Absorption Lidar Raman Spectroscopy/SERS Near IR Reflectance/Trans. Spect.	II/CP,CM IV II IV	A A W,S S	voc voc voc,svo voc	ррь-% ррт ррь-ррт 100s-1000s ррт	10.4.4/Table 10.4.4 10.4.4 10.4.4/Table 10.4.4 10.4.4/Table 10.4.4
Wet Chemistry					
Chemical Colorimetric Kits Other Colorimetric Methods Titrimetry Immunoassay Colorimetric Kits Ion Chromatography High-Pressure Liquid Chromatography Thin-Layer Chromatography Coulometry Polarography Stripping Voltammetry Radiological	II/CP I/CP I/CP II/CP II II/CM II II II II	W W W W W W W W W	EA,HM,SVO TR EA,HM,TR SVO EA SVO,TR SVO EA,TR EA EA	ppb-100s ppm ppb-100s ppm ppb-100s ppm ppm-100s ppm ppb-ppm ppm ppb-ppm sub-100s ppm sub-100s ppm ppt-ppm	10.5.1/Table 10-3 10.5.1/Table 10-3 10.5.1 10.5.2 10.5.3/Table 10-3 10.5.3 10.5.4 10.5.4 10.5.4
Neutron Activation/INNA	II	S,W	EA,TR	10s ppm	10.6.1, 3.3.5, 3.3.6
Radiation Detectors X-Ray Diffraction Gamma Spectrometry	I/CP II I/CP	s,w A,S,W S S	EA,IIM R,TR M M	varies 	10.6.1, 3.3.1 10.6.1 3.3.2
Other					
Gravimetric Volumetric Nuclear Magnetic Resonance Magnetic Susceptibility Electron Spin Resonance Optical Microscope Scanning Electron Microscope Electron Microprobe Field Bioassessment Toxicity Tests	I/CP I/CP I/CP II II II/CP II II II	W,S S,W,A S,W S,W,A S S S W,S,A	P,TDS P M,P M M M,P M M VOC,HM VOC,HM	% high ppm 	10.6.2 10.6.2 10.6.3, 3.2.4 10.6.3 10.6.3 10.6.4 10.6.4 10.6.4 10.6.6 10.6.6
Biomarkers	III	W,S,A	VOC,SVO,HM	**	10.6.6

Boldface = Most commonly used/proven field techniques.

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"I = Well established and routinely used field technology; II = Well established laboratory technology for which experience in field applications is moderate to limited; III = Relatively well established technology for which there is limited field experience; IV = Developing technology with potentially useful field applications. CP = Commercially available portable instruments; CF = Commercially available fieldable instruments; CM = Commercial/custom mobile laboratories available.

^bA = Air/gaseous matrix; S = Soil/solid matrix; W = Water/aqueous/liquid matrix. Volatile and semivolatiles in water and solid samples can be extracted for analysis by gaseous phase analytical techniques. Similarly, analytes can be extracted from solids samples for analysis using wet chemistry techniques.

EA = Elemental/ionic analysis; HM = Metals; M = Mineralogy; P = Physical characterization; R = Radioisotopes; SVO = Semivolatile organics; TG = Toxic gases; TDS = Total dissolved solids; TR = Tracer studies; VOC = Volatile organic compounds.Ranges for specific instruments and analytes might differ from range shown by orders of magnitude. In general, detection limits for soils will be higher than for ground water. 10.4); (5) wet chemistry techniques (Section 10.5), and (6) radiological and other miscellaneous techniques (Section 10.6).

Some Basic Analytical Concepts

For the nonchemist, terminology used to describe analytical techniques can be bewildering. A further source of potential confusion is that techniques can be used for different purposes in numerous combinations and configurations. For example, a flame ionization detector (FID) can be used by itself as a total vapor detector, or it can be used to detect specific compounds after they have been separated by a gas chromatograph (GC/FID). A gas chromatograph, on the other hand, can be used alone with a FID or other type of detector, or in combination with a mass spectrometer (GC/MS). An understanding of the basic principles of operation of major individual techniques makes it possible to have some idea of how an unfamiliar combination of techniques functions.

A further source of possible confusion is that the different terms can be applied to the same technique. For example, the terms fluorometry, fluorimetry, and spectrofluorometry can be used interchangeably. Furthermore, some terms can be applied to the same technique, but are not necessarily interchangeable. For example the term luminescence can be applied to any technique involving fluorescence, but the term fluorescence is not applicable to all luminescence techniques (which include phosphorescence). The following discussion might be helpful in developing an understanding of some of the basic principles involved in chemical analysis and in sorting out the relationship between similar techniques. It might also be helpful to think of techniques in terms of the major types of analytical signals as summarized in Table 10-2.

Chromatography refers to processes in which individual components of a mixture migrate through a stationary medium at different rates. In analytical chemistry, chromatography refers to a diverse group of separation methods such as gas chromatography (Section 10.3.3) and liquid chromatography (Section 10.5.3) used to separate, isolate, and identify components of mixtures that might otherwise be resolved with great difficulty.

A spectrum is the distribution of the phases of a radiated wave cycle or of the intensity of radiation when some property (frequency, mass, or energy) is allowed to vary. Spectroscopy encompasses a wide range of techniques involving optical instruments used to form and analyze spectra. Spectrometry is a spectroscopic technique in which the instrument measures: (1) The deviation of the refracted rays, and (2) wave lengths and angles between two faces of a prism. Spectrophotometry involves making comparisons of color intensity between corresponding parts of different spectra, or between parts of the same spectrum. Photometry involves the measurement of the intensity of light or the relative intensity of different lights. Luminescence involves the emission of light at temperatures below that of incandescent bodies and includes fluorescence (emission of radiation as a result of absorption of other radiation) and phosphorescence (light given off from slow oxidation of phosphorus).

Table 10-3 provides information on commercial sources for four major classes of analytical instruments: (1) Spectrophotometric instrumentation (atomic absorption, UV/visible, fluorescence, and infrared); (2) chromatographs (gas, ion, and liquid); (3) spectrometers (GC/MS, MS, optical emission, plasma emission, and x-ray); and (4) colorimeters.

Sources of Additional Information

SW-846 (U.S. EPA, 1986b) is the standard reference for solid waste test methods. A field screening methods catalog (U.S. EPA, 1988a) provides information on 26 field screening methods for which protocols have been developed and is available as an expert system for use on a microcomputer. This catalog is in the process of being updated and expanded into a format comparable to SW-846. Table 10-4, at the end of this section, provides an index of references providing overviews of field-screening techniques and more detailed references on sample extraction procedures. Table 10-5, also at the end of this section, provides a fairly detailed index of more than 300 references on specific field screening and other analytical techniques contained in this section.

Table 10-2 Major Analytical Signals and Methods

Signal	Analytical Methods Based on Measurement of Signal
Emission of radiation	Emission spectroscopy (X-ray, UV, visible, electron Auger); fluorescence and phosphorescence spectroscopy (X-ray, UV, visible); radiochemistry
Absorption of radiation	Colorimetry (visible), UV-visible/X-ray/IR spectrophotometry; photoacoustic spectroscopy; nuclear magnetic resonance and electron spin resonance spectroscopy
Scattering of radiation	Turbidimetry; nephelometry; Raman spectroscopy
Refraction of radiation	Refractometry; interferometry; X-ray diffraction
Rotation of radiation	Polarimetry; optical rotatory dispersion; circular dichroism
Electrical potential	Potentiometry; chronopotentiometry
Electrical current	Polarography; amperometry; coulometry; voltammetry
Mass-to-charge ratio	Mass spectrometry
Rate of reaction	Kinetic methods
Thermal properties	Thermal conductivity and enthalpy methods
Mass	Gravimetric analysis
Volume	Volumetric analysis

Boldface = Most commonly used in field screening applications.

Source: Modified from Skoog (1985)

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MANUFACTURER	SPECTROPHOTOMETRIC INSTRUMENTATION	CHROMATOGRAPHS	SPECTROMETERS	COLORIMETERS
ACE GLASS 800/223-4524		LI		
AIR INSTRUMENTS & MEASUREMENTS 818/813-1466, 800/969-4246	IR,UV		OE	
ALLTECH ASSOCIATES 708/948-8600, 800/255-8324	UV	IO,LI	GC	
AMERICAN GAUGE 404/932-0550	IR			
AMERICAN ULTRAVIOLET 908/665-2211	UV			
AMETEK, PROCESS & ANALYTICAL DIV. 302/456-4400, 800/222-6789	AA,UV,IR		MS	co
ANADATA 312/465-2688		GA		
ANALYTE 503/779-0334	AA			
ANARAD 805/963-6583	FL,IR,UV		XR	CO
ASOMA INSTRUMENTS 512/258-6608			XR	
ASTRO INTERNATIONAL 713/332-2484	IR			
BAIRD 617/276-6000			SM	
BALZERS HIGH VACUUM PRODS 603/889-6888	1		MS	T
BASELINE INDUSTRIES 303/823-6661, 800/321-4665		GA		
BOMEM INTERNATIONAL 708/350-0550, 800/888-3847	SI		SM	
BRAINARD KILMAN DRILL CO. 404/469-2720. 800/241-9468		GA		<u> </u>
BUCK SCIENTIFIC 203/853-9444, 800/562-5566	IR.UV.AA	GA		<u> </u>
C F A INSTRUMENTS 201/967-5660			SM	100
CHEMPLEX INDUSTRIES 914/337-4200	+		XB	100
CHBOMATOCHEM 406/728-5897 800/426-7227		11		<u> </u>
COLLOID ENVIRONMENTAL TECHNOLOGIES 708/392-5800			XB	t
COLUMBIA SCIENTIFIC 800/531-5003	FLUV			<u> </u>
DASIBI ENVIRONMENTAL 818/247-7601				<u> </u>
DIONEX 408/737-0700 800/346-6390	11,07,12	1110		
DYNAMATION 313/769-0573				00
E M SCIENCE 609/354-9200_800/222-0342		CH		100-
E S INDUSTRIES 609/082-3616 800/356-6140	·	GA		<u> </u>
EISONS INSTRUMENTS 508/524-1000	<u> </u>	GALL	GC OF PE YR MS	
FOXBORO 800/521-0451	IR I	GA	00,0L,FL,AN,W3	<u> </u>
FOXBORO E M O 508/378-5556		GALL		
GENERAL ANALYSIS 203/852-8000 800/327-2460		04,51		
GOW MAC INSTRUMENTS 008/560-0600		GALL		
H E SCIENTIEIC 813/337-2116	·	UN,LI		0
HN11 SYSTEMS 617/964-6690 800/724-5600		GA	YR	00
HACH 303/569-3050_800/227-4224		94		00
HAMII TON 702/786-7077 800/648-5950		GA	,,,,,	00
HEATH CONSULTANTS 713/047-0202 800/432-9497		GA		
HORIBA INSTRUMENTS 714/250-4811 800/446-7422	IB (IV	<u> </u>	YP	
HOUSTON ATLAS 713/348-1700		GA		
LC M 503/648-2014 800/262-2668		<u> </u>		0
ID ANALYTICAL 415/505-9200 900/427-0701		GALLIO	60	00
INSTRUMENTS SA 008/404-8660 800/438-7730		GA,LI,IO		
INTERNATIONAL LIGHT 508/465-5022	IR IN		06	
IONICS INSTRUMENT DIV 617/026-2500	<u>ei</u>			
ISCO INSTRUMENT DIV 402/464-0221 200/029-4250				
T INDUSTRIES 201//69-6777	10.01			
	111			
LAMOTTE 301/779-2100 900/244-2100				00
1 FAD TECHNOLOGIES 010/020.8814 000/020 0014		64		00
LENANI LADO 500/454.4442		GA		
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11 1 AMALTHOAL INGT RUMENTS 310/490-0900		GA		

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MANUFACTURER	SPECTROPHOTOMETRIC INSTRUMENTATION	CHROMATOGRAPHS	SPECTROMETERS	COLORIMETERS
MCNEILL INTERNATIONAL 800/626-3455			SM	i
MCPHERSON INSTRUMENTS 508/263-7733, 800/255-1055	UV.FL			
METROSONICS 716/334-7300	IR			
MIDAC 714/645-4096	IR			
MILTON ROY, ANALYTICAL PRODUCTS 716/248-4000, 800/654-9955	SLUV			co
MILTON ROY, PROCESS & ENVIRONMENTAL INSTRUMENTS	IR			
714/974-5560				
MONITEK TECHNOLOGIES 510/471-8300	UV			co
NATIONAL DRAEGER 412/787-2207	IR			
O ANALYTICAL 409/690-1711		GA		
PERKIN ELMER 203/762-1000, 800/762-4000	IR.UV.AA.FL	GALL	GC.PE.MS	
PHOTOVAC INTERNATIONAL 518/254-4199		GA		
PROCESS ANALYZEBS 215/736-2596		CH		
QUANTUM ANALYTICS 415/570-5656, 800/992-4199	AA.FL.IR.UV	GA.LLIO	GC.NM.OE.PE.MS	
BESPONSE BENTALS 716/266-3910 800/242-3910	IB	GA	00,111,02,12,110	<u> </u>
S B LINSTRUMENTS 213/214-5090		GA		
SENSIDYNE 813/530-3602 800/451-9444		GA	SM	co
SENTEX SYSTEMS 201/945-3694		GA		
SERVOMEX 617/769-7710, 800/862-0200	IB			
SIEMENS ENERGY AND ALITOMATION 404/740-3931	IR	GA		
SIEBBA MONITOR 408/262-9042	IB		SM	
SHIMADZU SCIENTIFIC 410/381-1227 800/477-1227	IR UV AA FI	GALLIO	GC OF	co
SIEVERS RESEARCH 303/444-2009		GA		
SPECTRA HABDWARE 412/863-7527			PF	
SPECTRA PHYSICS ANALYTICAL 408/432-3333 800/424-7666		11		
SPECTRACE INSTRUMENTS 415/967-6316			XB	
SPECTREX 415/365-6567			OF OF	
SPECTRO ANALYTICAL INSTRUMENTS 508/342-3400_800/548-5809	AA			
SUPREX 412/826-5200		СН		
T N TECHNOLOGIES/MANNING PRODUCTS 512/388-9100, 800/736-0801			XB	
TEXMAR 513/247-7000, 800/543-4461		GA		
TELEDYNE ANALYTICAL 818/961-9221	IR.UV			co
THERMO JARRELL ASH 508/520-1880	AA		OE.PE	
TIMBERLINE INSTRUMENTS 303/494-4104		CH		
TREMETRICS 512/251-1555, 800/876-6711		GA	GC	
TURNER DESIGNS 408/749-0994	FL			
TYTRONICS 617/894-0550	Si			
U V P 818/285-3123, 800/452-6788	UV			
UNOCAL UNIPURE 714/525-9225, 800/323-8647		СН		
VALCO INSTRUMENTS 713/688-9345, 800/367-8424		CH		
VARIAN ANALYTICAL INSTRUMENTS 415/945-2173. 800/926-3000	AA,FL,UV	GA.LI	GC	
VESTEC 713/796-9677		СН		
VIKING INSTRUMENT CORP. 703/758-9339			GC	
WHATMAN 201/773-5800, 800/922-0361		LI		
WYATT TECHNOLOGY 805/963-5904		LI		

KEY

ATOMIC ABSORPTION SPECTROPHOTOMETERS CHROMATOGRAPHS PRODUCED COLORIMETERS PRODUCED FLUORESCENCE SPECTROPHOTOMETERS GAS GC/MS ION INFRARED SPECTROPHOTOMETERS

A HOLACOR

Source: Pollution Equipment News (February, 1993)

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LIQUID MASS SPECTROMETERS NMR SPECTROMETERS OPTICAL EMISSION SPECTROMETERS PLASMA EMISSION SPECTROMETERS SPECTROPHOTOMETRIC INSTRUMENTATION PRODUCED SPECTROMETERS PRODUCED UV/VIS SPECTROPHOTOMETERS X-RAY SPECTROMETERS MS NE SI SI VR

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10.1 FIELD MEASURED GENERAL CHEMICAL PARAMETERS

10.1.1 pH/Alkalinity/Acidity

Other Names Used to Describe Method: --

<u>Uses at Contaminated Sites</u>: pH is used as an indicator during purging before ground-water sampling (see Section C.1) and is a fundamental parameter for chemical characterization of ground water and soils. In addition pH is used to classify corrosivity of wastes (a pH of less than or equal to 2 and greater than or equal to 12.5 is considered hazardous). Alkalinity and acidity are indicators of the buffer capacity of a solution (the resistance to change in pH with the addition of a strong acid or base). Alkalinity is required for chemical equilibrium calculations related to carbonate minerals.

<u>Method Description</u>: The pH is the negative logarithm of the hydrogen ion activity in aqueous solutions and is a significant water quality parameter because it affects solute concentrations perhaps more than any other single variable. Electrometric measurement of pH involves comparison of a glass hydrogen ion electrode in the solution of interest against a reference electrode of known potential by means of a pH meter or other potential measuring device. Measurement of pH in soil and solids by this technique requires preparation of a saturation extract. Colorimetric measurement involves use of reagents or litmus paper and estimation of pH by comparison of the resulting color with color charts. Flow-through cells (see Figure 10.1.2) provide the most accurate measurement of pH because it can be altered when samples are exposed to the atmosphere. The pH electrode and buffer solutions must be about the same temperature as the sample. This can be accomplished by allowing sample water to run over them or by using a portable water bath. Alkalinity and acidity are measured titrimetrically from the initial condition by the addition of a strong base or acid to an inflection point on the titration curve or to a fixed endpoint (titrimetry is discussed further in Section 10.5.1). In ground water, alkalinity is measured as carbonate and bicarbonate.

<u>Method Selection Considerations</u>: Electrometric measurement of pH using pH electrodes and a pH meter is the recommended technique for accurate measurement of both ground water and soil. Colorimetric techniques, which are less precise but somewhat easier to use in the field, are satisfactory for general characterization of soils. Field measurement of alkalinity (as carbonate and bicarbonate) is required for chemical equilibrium calculations related to carbonate minerals because this parameter is subject to change during sample handling. The acidity obtained from titration analysis gives a measure to total ionizable hydrogen that can be used as input to some geochemical computer programs.

<u>Frequency of Use</u>: Field measurement of the pH of ground water should be a standard procedure during sampling. Field measurement of the pH of soil samples often is required for accurate classification of soils and is a useful characterization technique, but is not necessarily required for soil samples collected for laboratory analysis unless redox sensitive species are of special concern.

Standard Methods/Guidelines: See Table 10.1.1.

Sources for Additional Information: Barnes (1964), Garvis and Stuermer (1980), Hem (1985-interpretation), Korte and Ealey (1983), Ritchey (1986), Thompson et al. (1989-Chapter 15).



Figure 10.1.2 Oxidation-reduction status: (a) Eh measuring cell; (b) Flow chamber for determination of dissolved oxygen from a pumped well (Wood, 1976).

Property/Section	Filtration	Sample Collection	Method Description	Reference
Wellhead Ground-Water	Measurements			
Temperature (10.1.3)	No	Dewar flask or flow through	Thermometer	USGS (1980); EPA Method 170.1 (Kopp and McKee, 1983)
рН (10.1.1)	No	Flow through	pH electrodes and meter	USGS (1980); Wood (1976)
Carbonate/Bicarbonate (Alkalinity)(10.1.1)	Membrane	Closed titration vessel	Potentiometric (pH electrode) titration with strong acid	USGS (1980); Wood (1976); D1067-82 (ASTM, 1982)
Acidity (10.1.1)	Membrane	Closed titration vessel	Potentiometric (pH electrode) titration with strong base	D1067-82 (ASTM, 1982)
Eh (Redox Potential) (10.1.2)	No	Flow through	Potentiometric (Pt electrode)	D1498-76 (ASTM, 1976); Wood (1976)
Dissolved Oxygen (10.1.2)	No	Flow through	Potentiometric with oxygen probe or titrimetric	EPA Method 360.1/.2 (Kopp and McKee, 1983); USGS(1980); Wood(1976)
Specific Conductance (10.1.3)	Membrane	Flow through	Wheatstone Bridge conductivity meter	USGS (1980); Wood (1976)
Field Laboratory Ground-	Water Measures	nents		
Filterable Residue (10.1.3)	No	Collect in bottle	Gravimetric	EPA Method 160.1 (Kopp and McKee, 1983)
Nitrate-Nitrite (Separately)(10.1.3)	Membrane	Collect in bottle	Spectrophotometric	EPA Method 353.3 (Kopp and McKee, 1983)
Sulfite (10.1.3)	Membrane	Collect in buffered iodine	Iodine-thiosulfate titration	Method M2 (Radian, 1988)
Soil/Solids				
Soil pH (10.1.1)	-	Grab or core	pH electrodes and meter in saturated paste	Method 8C (SCS, 1984) D4972-89 (ASTM, 1989)
Solid waste pH (10.1.1)	-	Grab or core		EPA Method 9045A [•] (U.S. EPA, 1986b)
Soil Conductivity/ Resistivity (10.1.3)	-	Grab or core	Wheatstone Bridge conductivity meter	Method 8E (SCS, 1984)

Table 10.1.1 Summary of Ground-Water and Soil Measurements to Be Made in the Field

*Revision 1 of this method is dated November, 1990.

Source: Compiled from Boulding (1991) and Thompson et al. (1989)

10.1 FIELD MEASURED GENERAL CHEMICAL PARAMETERS

10.1.2 Redox Potential (Eh)/Dissolved Oxygen

Other Names Used to Describe Method: --

<u>Uses at Contaminated Sites</u>: Characterizing oxidation-reduction conditions in the subsurface for evaluation of potential for mobility of heavy metals and biodegradation of organic contaminants.

Method Description: Redox potential (Eh) is measured electrometrically using a platinum electrode and a reference electrode to provide a reference potential and to provide an electrical connection to the solutions. Measurement of pH and temperature also are required. Eh readings can be strongly affected by exposure to atmospheric oxygen, consequently, flow-through cells must be used (Figure 10.1.2a). It is sometimes difficult to obtain precise readings of Eh because redox couples might not be in mutual equilibrium. More accurate characterization of the redox status of a sample requires analysis of the valence state of redox sensitive species (ferrous/ferric iron, nitrate/nitrite, and sulfate/hydrogen sulfide being most important in natural systems), which involves more complex chemical analytical procedures. Redox status of ground water and soil strongly affect the mobility and toxicity of arsenic, chromium, and selenium. Accurate chemical analysis of valence state is required to confirm Eh measurements. Arsenic and selenjum forms usually are measured using hydride AAS (see Section 10.3.5), and chromium species can be determined colorimetrically (Hach kits for total and hexavalent chromium recently have been developed in cooperation with EPA). Dissolved oxygen (DO) is another indicator of the oxidation-reduction state of an aqueous solution, with low concentrations indicating reducing conditions. In the field, DO is measured electrometrically using a membrane electrode, a reference electrode, and a meter to measure electrode response. As with Eh, flow-through cells are used to prevent alteration of the sample by contact with the atmosphere (Figure 10.1.2b). Dissolved oxygen also can be measured titrimetrically using the Modified Winkler method.

<u>Method Selection Considerations</u>: Along with pH, redox potential and dissolved oxygen are the most significant parameters affecting the chemistry of ground water.

<u>Frequency of Use</u>: Eh and dissolved oxygen in ground-water samples are not measured as routinely as pH, but probably should be.

Standard Methods/Guidelines: See Table 10.1.1.

Sources for Additional Information: Garvis and Stuermer (1980), Hem (1985-interpretation), Holm et al. (1986), Korte and Ealey (1983), Langmuir (1971), Newman and Kimball (1991-DO), Ritchey (1986), Rose and Long (1988-DO), Thompson et al. (1989-Chapter 17).



Figure 10.2.1 Field gas extraction techniques: (a) Vacuum extraction (Hadka and Dickinson, 1988); (b) Diagram of multiple headspace extraction (Ho et al., 1988).

10.1 FIELD MEASURED GENERAL CHEMICAL PARAMETERS

10.1.3 Other Parameters

Other Names Used to Describe Method: Specific conductance/electrical conductivity, temperature, suspended solids (filterable residue), sensitive chemical species (nitrate/nitrite and sulfite).

<u>Uses at Contaminated Sites</u>: Specific conductance: Monitoring of well purge water; performing qualitative assessment of water quality; estimating total dissolved solids; detecting conductance contaminant plumes; performing ionic tracer tests. Temperature: Monitoring of well purge water; correcting for pH and Eh measurements; performing temperature tracer tests; monitoring air temperature. Sensitive chemical species: Performing field measurement for evaluation of water quality. Filterable residue: Characterizing subsurface transport of heavy metals on particles.

Method Description: Temperature measurement techniques are discussed in some detail in other sections of the guide (Sections 1.6.1, 1.6.2, 3.5.2, 8.2.1, and 8.2.3). Temperature of ground water usually is measured with a mercury-filled thermometer, which is placed in a sample that is continuously pumped into a dewar flask. Pumping continues until three identical consecutive readings of temperature are obtained. Specific conductance is measured using a Wheatstone bridge conductivity meter. During well purging, measurement of specific conductance at intervals until there is no significant change between measurements serves as an indication that stagnant water has been completely removed from the well and water quality samples can be collected. Specific conductance typically shows a linear correlation with total dissolved solids, and consequently can be used instead of separate measurement of TDS, provided a correlation curve for the specific area of interest has been developed. Filterable residues are measured gravimetrically after filtering. Sensitive chemical species: The U.S. Geological Survey (1980) recommends that certain sensitive chemical species be analyzed in the field because of potential for alteration with holding times required for laboratory analysis. Sulfite can be analyzed using an iodine-thiosulfate titration, and nitrate/nitrite forms of nitrogen can be analyzed colorimetrically.

<u>Method Selection Considerations</u>: Temperature measurement of ground-water samples is simple, inexpensive, and a necessary complement to pH and Eh measurements. Specific conductance of ground-water samples is simple and inexpensive and is useful for monitoring purge water and estimating total dissolved solids (TDS). A well-designed and constructed monitoring well should produce samples with a minimum of filterable residue, but if significant amounts are present, this measurements should probably be taken, especially if subsurface particle transport of heavy metals is a possibility. Analysis of sensitive species should be performed when they are considered of geochemical significance at the site.

<u>Frequency of Use</u>: Temperature and specific conductance of ground-water samples are standard measurements. Use of specific conductance to estimate total dissolved solids is more commonly used for surface waters, but can be useful for monitoring of contaminant plumes. Filterable residue and analysis of sensitive species are performed less commonly.

Standard Methods/Guidelines: See Table 10.1.1.

Sources for Additional Information: Davis et al. (1985), Hem (1985-interpretation), Korte and Ealey (1983), Thompson et al. (1989).

10.2 CONTAMINANT SAMPLE EXTRACTION PROCEDURES

10.2.1 Gas Headspace/Vacuum Extraction

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Collecting volatile organic compounds in soil gas, soils, and ground water for chemical analysis.

<u>Method Description</u>: Vacuum extraction of pore gases involves the use of a vacuum pump to pull samples of air or soil gas directly into an analytical instrument. Figure 10.2.1a illustrates a vacuum sample probe used to obtain soil-gas samples for analysis by a mass spectrometer (see also, Figure 10.3.3b). Alternatively, a syringe can be used to sample the stream of gas that is created by the vacuum pump (see Figure 9.4.2a). Gas headspace extraction involves the use of a dead space to collect gases that are moving through water or soil, or from a solid or liquid phase to a gaseous phase. This can involve placement of a water or soil sample in a container that is partly filled with air (headspace), and collecting a sample of the headspace gas (usually with a syringe) once the vapors in the sample have equilibrated with the headspace gas. Since not all vapors are likely to degas the first time, multiple headspace extraction is sometimes used (Figure 10.2.1b). Figure 9.4.2c illustrates a field headspace collection device for sampling gases moving through a surface water body.

Method Selection Considerations: Advantages: Is an extremely simple procedure. Disadvantages: Might not result in full extraction of volatiles in ground-water and soil.

<u>Frequency of Use</u>: Vacuum extraction of pore gases and headspace techniques are commonly used for extraction of volatile organic compounds for analysis.

Standard Methods/Guidelines: Headspace techniques: Ford et al. (1984), U.S. EPA (1988b-FM-4 to FM-9, FM-11); Vacuum extraction: U.S. EPA (1988b-FM-12 to FM-14, FM-16, FM-17).

Sources for Additional Information: See Table 10-4.



Figure 10.2.1 Field gas extraction techniques: (a) Vacuum extraction (Hadka and Dickinson, 1988); (b) Diagram of multiple headspace extraction (Ho et al., 1988).

10.2 CONTAMINANT SAMPLE EXTRACTION PROCEDURES

10.2.2 Purge and Trap Methods

Other Names Used to Describe Method: P/T, purge with whole column cryotrapping (P/WCC).

Uses at Contaminated Sites: Extracting volatile organics from soil and water samples.

<u>Method Description</u>: Purge and trap techniques involve the forcing of a gas (usually helium) through a sample of water or soil slurry, which entrains the volatile compounds. The entrained volatiles can be fed directly into the analytical instrument (Figure 10.2.2a) or can be used in combination with a sorbent trap (see Section 10.2.5) to concentrate the samples for later thermal extraction (see Section 10.2.4). Figure 10.2.2b shows a schematic of concentrator/purge and trap device.

<u>Method Selection Considerations</u>: Advantages: Provides better recovery than vacuum/headspace extraction from water and soil samples. Disadvantages: (1) Requires somewhat more complex equipment than vacuum/headspace extraction and also requires a purge gas; and (2) more specialized training is required compared to gas headspace/vacuum extraction.

Frequency of Use: Commonly used for mobile laboratory analysis of volatiles in soil and ground water.

Standard Methods/Guidelines: --

Sources for Additional Information: See Table 10-4.



(a)



(b)

Figure 10.2.2 Purge and trap techniques: (a) Conventional (Wise et al., 1991a); (b) With concentrator (Sherman et al., 1988a).

10.2 CONTAMINANT SAMPLE EXTRACTION PROCEDURES

10.2.3 Solvent/Chemical Extraction/ Microextraction

Other Names Used to Describe Method: Soil-solvent extraction, liquid-liquid extraction, ultrasonication.

Uses at Contaminated Sites: Extracting volatile, semi-volatile, and non-volatile organic compounds from ground water and soils.

<u>Method Description</u>: Solvent extraction procedures involve the use of one or more organic solvents, acids, or other chemical substances and measures, such as filtration and centrifugation, to remove and concentrate the analyte of interest from a soil or ground-water sample (Figure 10.2.3a). Commonly used solvents include acetone, hexane, and methanol. Microextraction procedures require only a very small sample for extraction. Each analyte of concern requires its own specific extraction procedure. Simplified extraction procedures can sometimes be used for field screening purposes. Figure 10.2.3b compares a field screening and standard EPA laboratory extraction procedures for PCBs. Ultrasonication uses ultrasonic sound waves to accelerate the extraction of chemical species into a solvent.

<u>Method Selection Considerations</u>: Advantages: (1) Extraction procedures are compound specific; and (2) simplified extraction procedures have been developed for field screening of PCBs, PAHs, phenols, and pesticides. **Disadvantages**: Depending on the compounds, procedures can be complex and time-consuming.

<u>Frequency of Use</u>: Required for many EPA standard laboratory methods. Standard or simplified field screening extraction procedures are being increasingly used with a variety of field screening and analytical techniques.

Standard Methods/Guidelines: ASTM (in preparation-microextraction), U.S. EPA (1988b-Soil: PCBs, pesticides; Liquid/Liquid: Phenol; Soil or water: Total PNA).

Sources for Additional Information: See Table 10-4.



Figure 10.2.3 Soil extraction: (a) General schematic of organic solvent extraction from soil samples (Overton et al., 1988b); (b) Detailed field screening and EPA 3550/8080 extraction procedures for PCBs (Moy, 1989, by permission).

10.2 CONTAMINANT SAMPLE EXTRACTION PROCEDURES

10.2.4 Thermal Treatment Methods

Other Names Used to Describe Method: Thermal/microwave-assisted digestion, thermal extraction (Ruskan/Pyran thermal chromatograph/Pyrocell), thermal desorption.

<u>Uses at Contaminated Sites</u>: Preparing soil and water samples for instruments requiring a gaseous phase for analysis (Section 10.3); preparing soil and water samples for wet chemistry/colorimetric analysis.

<u>Method Description</u>: Thermal extraction techniques have in common the use of heat to prepare samples for subsequent stages of analysis. This can be as simple as using an electric or microwave oven to dry samples (required for soil moisture content determinations and XRF analysis in the laboratory), to highly sophisticated instruments for vaporizing samples (such as ICP torches for atomic emission spectrometry [see Section 10.3.6]). The term digestion is commonly used when heating is involved in wet chemistry analytical procedures. Figure 10.2.4a shows a thermal extraction device used with a flame ionization detector, and Figure 10.2.4b shows a schematic of a column thermal extractor used with a mass spectrometer.

<u>Method Selection Considerations</u>: Most thermal treatment techniques and devices are small enough that they can be used in mobile laboratories. Thermal digestion is required for many wet chemistry analytical procedures. Thermal extraction/desorption can sometimes be used as an alternative to solvent extraction for analysis of non-gaseous phase samples in analytical instruments, such as gas chromatographs and mass spectrometers, which require a gaseous phase (see Section 10.3).

Frequency of Use: Colorimetric wet chemistry field test kits for liquids, oils, and solids (see Section 10.5.1) often involve an initial digestion step. Use of thermal extraction procedures in conjunction with gas chromatographs and mass spectrometers in mobile laboratories is a relatively new approach that is becoming more common.

Standard Methods/Guidelines: U.S. EPA (1988b-PAHs using GC with heated column).

Sources for Additional Information: See Table 10-4.




(b)

Figure 10.2.4 Thermal extraction devices: (a) Pyrocell for FID; (b) Column thermal extractor for mass spectrometer (Overton et al., 1988b).

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10.2 CONTAMINANT SAMPLE EXTRACTION PROCEDURES

10.2.5 Other Extraction Methods

Other Names Used to Describe Method: Membrane extraction, extraction disks, sorbent/solid phase extraction cartridges, Tenex tubes, cyclohexyl-bonded phase extraction column, supercritical fluid/gas extraction (SFE).

Uses at Contaminated Sites: Extracting contaminants in ground water (sorbent and membrane) and air/soil gases (sorbent).

Method Description: Sorbent extraction involves the contact of air or water through a material, such as granular activated carbon (GAC), polyurethane, or resins, which trap organic compounds by sorption or filtration. Figure 10.2.5a illustrates the use of a polyurethane sorbent for air quality sampling. Bonded sorbents have been used for pesticides, PAHs, and phenols. Resin cartridges can be used for concentration of VOCs obtained from purge and trap (see Figure 10.2.2b). GAC is commonly used for passive soil sampling (see Figures 9.4.1a and b) and sometime to extract volatiles from ground-water samples. Once the sample is collected, a thermal extraction technique (Section 10.2.4) typically is used to extract the concentrated sample for instrumental analysis. Membrane extraction uses extractant fluids containing organic solvents, such as hexane, flowing through a tubular silicone rubber membrane to selectively extract and concentrate organic compounds of interest from a sample flowing outside the tube. In the simplest application, extractant fluid flows directly to the analytical instrument for analysis (Figure 10.2.5b). For more complex samples, additional separation steps might be required. Supercritical fluid extraction (SFE) allows extraction of components from different matrices by means of a supercritical fluid, such as carbon dioxide.

<u>Method Selection Considerations</u>: Sorbent Advantages: (1) Is a simple and inexpensive extraction technique for gaseous and water samples; and (2) is most applicable where preconcentration or precise measurements are required. Sorbent Disadvantages: (1) Concentrations will be underestimated if sorption is not complete or the sorbent becomes saturated; and (2) typically requires a second extraction step for instrumental analysis. Membrane Extraction Advantages: (1) Is a relatively simple technique; and (2) has the potential for automation to eliminate sample handling before it goes into the instrument for analysis. Membrane Extraction Disadvantages: (1) Is limited to aqueous samples; and (2) satisfactory extraction might be difficult with complex samples.

<u>Frequency of Use</u>: Sorbents are being widely used where preconcentration of samples is required. Membrane extraction and SFE are relatively new methods that have not been used extensively.

Standard Methods/Guidelines: Sorbent: U.S. EPA (1988b-FM-10, FM-16).

Sources for Additional Information: U.S. EPA (1988b-FM-D2, FM-D3). See also, Table 10-4.







(b)



10.3 GASEOUS PHASE ANALYTICAL TECHNIQUES

10.3.1 Total Organic Vapor Survey Instruments

Other Names Used to Describe Method: OV detectors, photoionization detector (PID/HNU meter), flame ionization detector (FID/Organic vapor analyzer/OVA), argon ionization detector (AID), combustible-gas indicator (explosimeter [ED]/catalytic surface oxidation device).

<u>Uses at Contaminated Sites</u>: **PID**: Surveying aliphatics and aromatics; **AID**: Surveying aliphatics, aromatics, halomethanes, and halethanes. PIDs, FIDs and AIDs also can be used in combination with a gas chromatograph for detecting specific compounds (Section 10.3.3). Explosimeters are used to test manhole/sewers, pipeline leaks, confined areas in sewage plants, and inside tanks for combustible gases.

Method Description: Photoionization detector (PID): Uses an ultraviolet lamp as an ionizing source and responds to volatile organic compounds that have an ionization potential less than or equal to the lamp. A PID reports concentrations as total ionizable compounds. Flame ionization detector (FID): (Uses a hydrogen flame to ionize organic vapors entering the detector and reports concentrations of total organics as the ppm equivalent to a calibration compound (usually methane).) Argon ionization detector (AID): Similar to a PID, except that an argon lamp is used. Explosimeters use a sensor (hotwire, catalytic, solid state, etc.) to produce a signal, which is processed and displayed as the percentage of the combustible gas present to the total required to reach the lower explosive limit (LEL) and/or the percent combustible gas by volume. Various calibration gases can be used (butane, pentane, natural gas, and petroleum vapors), but methane is the most common.

Method Selection Considerations: Figure 10.3.1 shows sensitivity ranges for organic vapor monitoring instruments. Total Detector Advantages: (1) Are highly portable (FID somewhat less portable than PID) and easy to use; (2) are relatively inexpensive (around \$5,000); (3) FID is sensitive to a larger number of volatile organic compounds than PID (including low molecular weight compounds, such as methane, ethane, and certain toxic gases with high ionization potential, such as carbon tetrachloride and HCN); (4) have very rapid response time (seconds); and (5) AID is the most durable detector. Total Detector Disadvantages: (1) Are non-specific (they indicate if something is present but do not identify); (2) FID is more complicated than PID and requires hydrogen gas; and (3) AID is somewhat less sensitive than FID and PID. Explosimeters are inexpensive and portable.

Frequency of Use: PIDs and FIDs are widely used as survey instruments whenever volatile organics are suspected, and also are commonly used in conjunction with gas chromatographs (see Section 10.3.3). Explosimeters are commonly used where explosive gases are suspected.

Standard Methods/Guidelines: Ford et al. (1984-PID, FID, ED), U.S. EPA (1988b-Section 15).

Sources for Additional Information: Aller (1984-combustible gas indicator). See also, Table 10-5.



SENSITIVITY RANGES FOR ORGANIC VAPOR MONITORING INSTRUMENTS

Figure 10.3.1 Sensitivity ranges for organic vapor monitoring instruments (Moore, 1991).

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10.3 GASEOUS PHASE ANALYTICAL TECHNIQUES

10.3.3 Gas Chromatography (GC)

Other Names Used to Describe Method: Portable, fieldable, or mobile gas chromatograph with a: Flame ionization detector (FID), photoionization detector (PID), argon ionization detector (AID), electron capture detector (ECD), thermal conductivity detector (TCD), flame-photometric detector (FPD), Hall-electrolytic conductivity detector (ELCD), ion trap detector (ITD), microwave [induced] plasma detector (M[I]PD); GC/atomic emission spectroscopy (AES); GC/Fourier transform infrared spectroscopy (FIIR) (see section 10.4.4); GC/mass spectrometry (MS) (see Section 10.3.4).

<u>Uses at Contaminated Sites</u>: **Portable GC**: Assessing volatile organics and other gases (using headspace or purge and trap), soil PAHs, PCP, and PCBs (using extraction techniques). **GC/AES**: Assessing Cl, Br, O, N, P, and S levels.

Method Description: Gas chromatography involves the separation of gaseous constituents on a stationary phase in a column, which is either a solid or liquid held on a solid support. Thermal desorption gas chromatographs (TD-GC), with a unit for vaporizing samples before entering the column, are used when samples are in liquid phases or soil. Once the analytes have been separated in the column, they are eluted one after another, and then enter a detector attached to the column exit. Numerous types of detectors can be used with a gas chromatograph as listed above under other names. An FID or PID (see Section 10.3.1) can be used to detect specific compounds after they have been separated in the GC and FIDs and PIDs commonly are used in portable GCs. The electron capture detector (ECD) is another commonly used detector. Figure 10.3.3 shows several portable GC units. GC commonly is used as a sample preparation step for other types of instrumentation, such as the mass spectrometer (see Section 10.3.4). Relatively new combinations that show promise for use at contaminated sites include: (1) GC/MPD-AES, an experimental technique using GC in combination with a microwave plasma detector (MPD) and atomic emission spectrometry (see Section 10.3.6); and (2) GC/Fourier Transform Infrared (see Section 10.4.4).

Method Selection Considerations: GC Advantages: (1) Are fairly portable; (2) have very good specificity, depending on detector used, with excellent ability to resolve most components in very complex mixtures; (3) have fair sensitivity (ppb to ppm); and (4) inexpensive compared to mass spectrometer (\$10,000 to \$20,000 vs. \$50,000 to \$200,000). GC Disadvantages: (1) Are less sensitive than mass spectrometers; (2) have slower response time than mass spectrometers (tens of minutes vs. seconds) and their calibration can be time-consuming; (3) require a library of retention times to identify compounds and non-target compounds might be difficult to identify if detected analytes are not in the library or the quality of the library match is too low to make positive identification; and (4) require bottled gas. GC/FID: Universal capability in screening samples. GC/MS: Allow better resolution of components in complex mixtures than MS alone and are most commonly used for unequivocal identification of hazardous compounds. GC/FTIR: Allow elucidation of chemical structure and are able to identify additional hazardous compounds not detected by GC/MS. GC/AES Advantages: (1) Allow detection of elements that have been impossible or difficult to monitor with other GC detectors; (2) elementspecific detection can save time in sample preparation; (3) multiple element detection reduces need for GCs with multiple detectors; (4) element ratios can reduce time for interpretation of GC/MS and GC/FTIR data for nontarget compounds; (5) size and weight and other requirements similar to GC/MS field laboratory instrumentation; and (6) detection limits comparable to GC/FID and GC/FPD. GC/AES Disadvantages: Instrumentation still in developmental stages. Other Detectors: ECDs are highly sensitive to halogenated organic molecules and can be used to analyze for PCBs in the presence of unhalogenated hydrocarbons, such as oil. Detectors, such as NPD, ECD and ELCD (Hall detector), have lower detection limits for specific elements.

<u>Frequency of Use:</u> GC is the most well developed and accurate field analytical technique for organic compounds when used with an appropriate detector. The most commonly used detectors include PID, FID, AID, ECD, Hall detector (ELCD), and TCD.



(a)



(b)



Standard Methods/Guidelines: Ford et al. (1984), U.S. EPA (1987-Exhibit 7A-1, mobile lab protocol for organics), U.S. EPA (1988b-TD/GC, GC/ECD, Mobile GC).

Sources for Additional Information: Davis et al. (1985), Nielsen et al. (1992), Szelewski and Wilson (1988), U.S. EPA (1991b, 1992), Weslowski and Alwan (1991). See also, Table 10-5.

10.3 GASEOUS PHASE ANALYTICAL TECHNIQUES

10.3.4 Mass Spectrometry (MS) and GC/MS

Other Names Used to Describe Method: Fieldable/mobile mass spectrometer (MS), mobile tandem MS (MS/MS, MINITMASS), GC/MS, GC/ITMS or ITD (ion trap mass spectrometer or ion trap detector), thermal desorption (TD) or thermal extraction (TE), GC/MS.

<u>Uses at Contaminated Sites</u>: **TD-GC/MS**: Assessing VOCs (water, soil/sediment, soil gas, air), PCBs, PAHs, and pesticides (soil/sediment); **GC/ITMS**: Assessing VOCs (air, water, soil) (Wise et al. [1991a] list detection limits for 30 VOCS in air, and 21 VOCs in water).

<u>Method Description</u>: Mass spectrometry techniques involve conversion of compounds in a sample into charged ionic particles that consist of the parent ion and ionic fragments of the original molecule. Distinctive mass/charge ratios allows for identification of compounds, while the magnitude of ion currents at various mass settings is related to concentration. Major components of a mass spectrometer include: (1) The inlet system, (2) the ion source, (3) the electrostatic accelerating system, and (4) the detector and readout system that gives a mass spectrum recording the numbers of different kinds of ions (Figure 10.3.4a). Mass spectrometers often are used in conjunction with gas chromatography (see Section 10.3.3). Figure 10.3.4 illustrates portable, fieldable, and mobile laboratory mass spectrometers.

<u>Method Selection Considerations</u>: MS Advantages: (1) Have very good specificity in noncomplex matrix; (2) are very sensitive (ppb); (3) have rapid response time (seconds); and (4) very small sample sizes (milligram to microgram) can be used. MS Disadvantages: (1) Have poor resolution in complex mixtures (can be overcome by using GC/MS); (2) are expensive (\$50,000 to \$200,000); (3) are large, heavy, and not very rugged; (4) require high vacuum pumps and a large amount of power; (5) are complex instruments requiring long set up time; (6) require a library of spectra; and (7) calibration procedures are more time-consuming than for GC.

Frequency of Use: Some use in mobile laboratories. Field instruments are in developmental stages.

Standard Methods/Guidelines: U.S. EPA (1988b-GC/ITMS, MS/MS).

Sources for Additional Information: Davis et al. (1985). See also, Table 10-5.





(b)

(a)



Ruska Thermal



Figure 10.3.4 Mass spectrometers: (a) Schematic of man-portable GC/MS system: (A) Vapor inlet/transfer GC column, (B) MSD analyzer, (C) control electronics, (D) portable 386 computer, (E) molecular drag pump, (F) vacuum hose, (G) vacuum reservoir, (H) carrier gas, and (I) 24v DC battery (Meuzelaar et al., 1991), (b) Fieldable mass spectrometer mounted in a 4-wheel drive vehicle (Hadka and Dickinson, 1988), (c) Mobile thermal chromatograph/mass spectrometer (Greenlaw et al., 1989, by permission).

10.3 GASEOUS PHASE ANALYTICAL TECHNIQUES

10.3.5 Atomic Absorption Spectrometry (AAS)

Other Names Used to Describe Method: --

Uses at Contaminated Sites: Analyzing heavy metals, organometallic compounds, and other elements in water and soil/solids.

<u>Method Description</u>: AAS involves the measurement of radiation absorbed by electrons in a vaporized liquid sample. All AAS instruments have the following basic features (Figure 10.3.5a): (1) A light/radiant energy source that emits resonance line radiation; (2) a sample chamber in which the sample is fed as an aerosol and vaporized; (3) a device for selecting only one of the characteristic wavelengths (visible or ultraviolet) of the element being determined; (4) a detector, usually a photomultiplier tube, which measures the amount of absorption; and 5) a readout system (strip chart recorder, digital display, meter, or printer). Techniques for vaporizing the sample include flame (aerosol mixed with fuel and oxidant gas), furnace or electrothermal (sample is deposited at room temperature in a graphite tube and vaporized by heating), hydride generation or derivitization (elements such as As, Se, Sb, and Sn are converted to gaseous hydrides before being vaporized in small quartz tube furnaces), and cold vapor (for mercury only). AAS instruments can have one or two beams (Figure 10.3.5b and c), and more sophisticated (and more expensive) instruments have more than one channel for simultaneous determination of more than one element. Multi-element sequential AAS instruments can be programmed to automatically determine chosen elements sequentially.

<u>Method Selection Considerations</u>: Advantages: (1) Simpler instruments, such as single-beam flame AAS, are relatively inexpensive (around \$6,000 in 1986); (2) operation is very simple and can be partially automated; (3) in many determinations, standardization is easy and straightforward; and (4) have low detection limits (ppb) and high accuracy (furnace AAS has the lowest detection limits; flame AAS is generally 10 to 100 times higher). Disadvantages: (1) Flame AAS can only measure one element at a time and is not well suited for refractory elements, such as boron and vanadium; (2) time required for heating cycle of furnace AAS makes it slow compared to flame AAS; (3) sample preparation requires great care and can be time consuming; and (4) matrix interferences might affect results for specific elements (for example AI, phosphate, and sulfate interfere with Ca determinations).

<u>Frequency of Use</u>: AAS and inductively coupled plasma atomic emission spectrometry (Section 10.3.6) are probably the two most widely used laboratory techniques for elemental analysis of aqueous and solid samples. Many mobile laboratories have AAS for analysis of heavy metals and hydride derivitization. AA has been used less commonly in mobile laboratories for analysis of organometallics.

Standard Methods/Guidelines: U.S. EPA (1988b-FM-1), U.S. EPA (1987-protocol 7A-3).

Sources for Additional Information: Baker and Suhr (1982), Fishman and Friedman (1989), Thompson et al. (1989).



(a)







Figure 10.3.5 Atomic absorption spectrometers: (a) Essential components; (b) Single-beam instrument; (c) Doublebeam instrument with background correction using a deuterium lamp (Baker and Suhr, 1982, by permission).

10.3 GASEOUS PHASE ANALYTICAL TECHNIQUES

10.3.6 Atomic Emission Spectrometry (AES)

Other Names Used to Describe Method: Optical emission spectrometry (OES), flame emission spectrometry (FES)/flame photometry.

Uses at Contaminated Sites: Analyzing heavy metals and other elements in water and soil/solids.

<u>Method Description</u>: AES involves the excitation of electrons in liquid samples and measurement of the radiation emitted when they relax to an unexcited state. Each element emits radiation of a characteristic wavelength and concentrations are proportional to the intensity. AES using a flame as an excitation source, called flame emission spectrometry (FES) or flame photometry, has been in use since the 1860s. A variety of other excitation sources can be used (such as direct current arc, alternating current spark, and direct current discharge plasmas), but the most commonly used source today is the inductively coupled radiofrequency plasma (ICP) torch (Figure 10.3.6).

<u>Method Selection Considerations</u>: Advantages: (1) A large number of elements can be measured simultaneously (10 to 20 for FES and 20 to 35 to ICP-AES), making analysis for any one element very rapid; (2) ICP-AES linear range for detection is greater than AAS, reducing the amount of sample handling and dilution for analysis; (3) FESs are simple and inexpensive to operate; and (4) ICP provides a highly stable, sensitive and relatively interference-free excitation source for solution samples, and is able to handle refractory elements that AAS and FES cannot. Disadvantages: (1) Furnace AAS provides greater sensitivity for arsenic, lead, and selenium; and (2) solids samples requires careful preparation of solutions for analysis.

Frequency of Use: ICP-AES and AAS (Section 10.3.5) are probably the two most widely used laboratory techniques for elemental analysis of aqueous and solid samples.

Standard Methods/Guidelines: --

Sources for Additional Information: Baker and Suhr (1982-FES), Fishman and Friedman (1989), Soltanpour et al. (1982-ICP-OES), Thompson et al. (1989-ICP-AES).



Figure 10.3.6 Plasma torch configuration for ICP-AES (Fishman and Friedman, 1989).

10.3 GASEOUS PHASE ANALYTICAL TECHNIQUES

10.3.7 Ion Mobility Spectrometry (IMS)

Other Names Used to Describe Method: Plasma chromatography.

<u>Uses at Contaminated Sites</u>: IMS: Detecting microorganisms, anilines, nitrosoamines, organophosphorus esters, organic sulfides and arsenicals, selected explosives, and many other organic compounds (Reategui et al. [1988] identify more than 40 organic and inorganic compounds or groups of compounds that can be detected by IMS); GC/IMS: Detecting alcohols, ketones, BTEX, aldehydes, halocarbons, and chlorinated aromatics.

<u>Method Description</u>: IMS resembles a cross between a flame ionization detector and a mass spectrometer. Figure 10.3.7 shows the operation of an IMS cell. A sampling pump draws air though a semipermeable membrane, which is selected to exclude or attenuate possible interferents. The sample is ionized in a reaction region through interaction with a weak plasma of positive and negative ions produced by a radioactive source. A shutter grid allows periodic introduction of the ions into a drift tube, where they separate based on charge, mass, and shape, with the arrival time recorded by a detector. The identity of the molecules is determined using a computer to match the signals to IMS signatures held in memory. If the IMS signature is known it also is possible to program the instrument to detect specific compounds of interest.

Method Selection Considerations: IMS Advantages: (1) Combines the simplicity and sensitivity of ionization detectors (Section 10.3.1) with the ability to distinguish specific compounds in complex matrix; (2) has very good sensitivity (sub ppb to ppm); (3) has very fast response time (seconds); (4) is portable and rugged; and (5) is inexpensive compared to MS and comparable in price to GC (\$5,000 to \$25,000). IMS Disadvantages: (1) Provides specific identification of fewer compounds than GC or MS; (2) is better than MS at identifying certain target compounds in a complex mixture, but GC provides better resolution in this situation; and (3) requires a library of ion mobilities.

Frequency of Use: IMS: Recent development of portable IMS detectors might make the technique an alternative to FID and GC.

Standard Methods/Guidelines: --

Sources for Additional Information: U.S. EPA (1991b, 1992). See also, Table 10-5.



Figure 10.3.7 Theory of operation of ion mobility spectrometer (Reategui et al., 1988).

10.4 LUMINESCENCE/SPECTROSCOPIC ANALYTICAL TECHNIQUES

10.4.1 X-Ray Fluorescence (XRF)

<u>Names Used to Describe Methods</u>: Portable XRF, X-ray fluorescence spectroscopy/spectrometry, x-ray emission spectrography, x-ray spectrochemical analysis.

Uses at Contaminated Sites: Detecting heavy metals and other elements in soil/solids samples. Reported elements As, Cd, Cr, Cu, Fe, Hg, Mn, Ni, Pb, Sn, and Zn.

Method Description: XRF uses primary x-rays to irradiate a solid sample, which causes elements in the sample to emit secondary radiation of a characteristic wavelength (Figure 10.4.1a). Concentration of an element is proportional to the intensity of the secondary radiation emission. Two basic types of detectors are used to detect and analyze the secondary radiation. Wavelength-dispersive XRF spectrometry uses a crystal to diffract the xrays, as the range of angular positions are scanned using a proportional or scintillation detector (see Section 1.5.4 for description of these detectors). Energy-dispersive XRF spectrometry uses a solid-state, Si(Li) detector from which peaks representing pulse-height distributions of the x-ray spectra can be analyzed. It is the latter type of detector that has allowed development of field-portable instruments (Figure 10.4.1b). Figure 10.4.1c shows the effective depth of penetration of various materials. Various terms have been used to describe this technique, but XRF is the most commonly used term in the literature on investigation of contaminated sites.

<u>Method Selection Considerations</u>: Advantages: (1) Is about ofte-tenth the cost of conventional laboratory analyses; (2) sample preparation is minimal compared to conventional analytical techniques; (3) allows simultaneous determination of several elements; and (4) very portable energy-dispersive XRF instruments are now available (Figure 10.4.1b) and more accurate wavelength-dispersive XRF instruments can be used in mobile laboratories. **Disadvantages**: (1) Detection limits for portable instruments (10s to 100s ppm) typically are an order of magnitude higher than ICP-AES; (2) laboratory use with liquid samples requires preconcentration or precipitation, which is time consuming; and (3) the relatively shallow depth of penetration of soil materials (mm) means that collection and processing by grinding samples is generally required to obtain reproducible readings using a portable probe.

<u>Frequency of Use</u>: Along with total/specific organic vapor detectors and gas chromatographs, XRF is the most mature of the portable field screening techniques that have developed in recent years.

Standard Methods/Guidelines: Laboratory XRF: Jones (1982); Field screening: U.S. EPA (1987-Protocol 7A-4), U.S. EPA (1988b).

Sources for Additional Information: Raab et al. (1991), Thompson et al. (1989). See also, Table 10-5.

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Figure 10.4.1 X-ray fluorescence: (a) Schematic indicating the field-portable XRF analytical process (Glanzman, 1988); (b) Process for real time, on-site XRF measurements, data transfer, processing and plotting (Raab et al., 1991, by permission).

10.4.2 LUMINESCENCE/SPECTROSCOPIC ANALYTICAL TECHNIQUES

10.4.2 Other Luminescence Techniques

<u>Names</u> <u>Used to Describe Methods</u>: Fluorometry/fluorimetry/spectrofluorometry: UV fluorescence spectrophotometer; synchronous fluorescence/luminescence (SF/SL); laser fluorometry/laser induced fluorescence (LIF); solid state fluorescence, x-ray fluorescence (Section 10.4.1); room-temperature phosphorimetry (RTP).

<u>Uses at Contaminated Sites</u>: UV and synchronous fluorescence: Performing semiquantitative analysis of semivolatile polynuclear aromatic hydrocarbons (PAHs/PNAs); field screening of BTEX. RTP: Analyzing PCBs.

Method Description: Fluorometry is a photoluminescent technique in which the electronic state of a molecule is elevated by absorption of electromagnetic radiation. When the molecule returns to its ground state, radiation is emitted (typically ultraviolet or visible radiation for most fluorometric techniques) to produce a distinctive excitation and emission spectrum. Instruments used for fluorometric analysis range from simple filter fluorometers to very sophisticated spectrophotofluorometers. These instruments contain four principal components: (1) A source of excitation energy (UV, laser, x-rays, etc.), (2) a sample cuvette, (3) a detector to measure the photoluminescence, and (4) a pair of filters or monochromators for selecting the excitation and emission wavelengths. UV fluorescence has been used in a number of applications for field screening: (1) For semiquantitative analysis of solvent extracted PAHs, (2) for analysis of samples using high performance liquid chromatography (Section 10.5.3), (3) in conjunction with fiber optic sensors (Section 5.5.6), and (4) as a surface contamination detector, in which a non-fluorescing substance sprayed on the ground surface reacts chemically with the contaminant of interest to form a substances that fluoresces with UV excitation. Fiber optic sensors commonly use UV fluorescence (see Section 5.5.6). Synchronous fluorescence, or luminescence, involves the use of both emission and excitation monochromators to record the luminescence signal, which allows greater selectivity in the analysis of environmental samples. RTP is based on detecting the phosphorescence emitted from organic compounds adsorbed on solid substrates at ambient temperatures (conventional phosphorimetry requires cryogenic equipment). A recently developed test for PCBs using RTP involves a rapid extraction procedure (1 to 3 minutes), followed by placement of a few microliters of the sample solution on a filter paper. The sample is dried for about three minutes with a heating lamp and transferred to a spectrofluorimeter equipped with a phosphoroscope. The presence and concentration of PCBs can be determined by the spectral signature and intensity. Table 10.4.2 provides additional information on UV-visible luminescence, synchronous fluorescence, room-temperature phosphorescence, and low-temperature luminescence methods.

<u>Method Selection Considerations</u>: Advantages: (1) Instrumentation is relatively simple and portable; and (2) UV fluorescence can be used for rapid semiquantitative analysis of total PAHs in soil. Disadvantages: Analysis of complex samples can be difficult due to spectral overlap of different luminescing compounds (SF can partly overcome this).

<u>Frequency of Use</u>: Fluorometry in combination with fluorescent dyes probably is the most common technique used in karst limestone tracer studies. PAH-extract/UV fluorescence has been demonstrated as a good field screening technique for semiquantitative analysis of polynuclear aromatic compounds in soil. Synchronous fluorescence and RTP only recently have been tested for field screening of contaminants and are still in the developmental stages.

Standard Methods/Guidelines: U.S. EPA (1988b-total PNA with UV fluorescence).

Sources for Additional Information: Davis et al. (1985-fluorometry), Eastwood and Vo-Dinh (1991). See also, Table 10-5.

Table 10.4.2 General Characteristics of UV-Visible Luminescence, Synchronous Fluorescence, Room Temperature Phosphorescence, and Low Temperature Luminescence Techniques of Field Analysis (See end of Table 10.4.3 for definitions and abbreviations)

Applicability	Advantages	Limitations	Sansitivity	Current Field Applicability	Related Lab Techniques & Sensors
·····	UV-v/s	Luminescence (Fluores	cence and Phosphores	cance)	+ =
Polyaromatic Compounds Fluorescent Dyes	Most Sensitive Method for Trace and Ultratrace Analysis when Applicable	Limited to Compounds with Fairly High Luminescence Yields (Usually PACs, unless	Excellent Sensitivity ppb (pptrillion or Less with Laser	Portable Instruments Available Field Deployable	Luminescence Techniques - Fluorescence - Phosphorescence
Fluorometric Reaction Products	Instrumentation Readily Available	Derivalized) Relatively Unspecific for Structurel	Excitation) Dependent on Quantum Yiekds	Instruments Available Flow-through Oil-Water Monitors and HPLC	- Synchronous - Time and Phase Resolution - Polarization
PCBs	No Interference by Water	Information (Compared to IR)		with Multichannel Detectors	- HT and LT - 3D - Microscopy
Pasticides .	Few Interferences by Nonaromatics	Quantitation Complicated by		Front Surface - RTP	Fiber Optic
Semivolatiles	Some Structural Specificity	Differences in Quantum Yields, Quenching, Microenvironments			Sensors
Nonvolatiles Petroleum Oils	- Enhanced by Special Techniques	Limited Reference Spectra Available			Multichannel Detectors - Diode Arrays - CCDs
	Very Selective - Enhanced by Time and Wavelength Variability Can Distinguish Geometrical Legence				Fluorescence Quenching or Energy Transfer - Indirect Ways to Measure Non- luminescent Molecules
	Geometrical isometris	Synchronoi	Is Fluorescence		L
Increased Specificity for Individual PACs	Increased Specificity	Decrease in Sensitivity with	Good Sensitivity	Portable Instruments	LT Measurements
or PAC Classes in Complex Mixture	Less Spectral Overlap	Narrower Bandpasses and Wavalength Offse	Slightly Lower than t Fluorescence Emission	n Field Deployable	Time and Phase Resolution
Petroleum Oils	Classification of PAHs by Number of Rings	Loss of Vibrational Structure in Spectrum	Dependent on Instrumental	Available	Derivative Remote Monitor
G18050185	Useful for Screening	Need Dual Scanning Monochromators	Dependent on Stokes		under Development
	Combine with Other Luminescence Techniques	Need Polychromatic Source	Shift of Compound		Synchronous Phosphorescence
		Room Temperature	Phosphorescence (RTP))	J~
Most Luminescent PACs, PCBs,	Easy Sample Prep	Oxygen may Quench in Solution	Good Sensitivity	Portable Instruments Under Development	Can Compare with LT Techniques for
PAHs Directly or with	Eliminates Scatter and Fluorescence Background	Less Structure than LTP	Dependent on Quantur	Field Deployable Instruments Available	Time Resolution
Perturber	Longer Lifetimes than Fluorescence	Substrate/Technique Dependent	Dependent on	Front Surface	π.ο
	No Need for Cryo- genic Instrumentation	Quantitation may be Complicated	Perturber	- Filter Paper - TLC Plate	- Micelle Solution - Cyclodextrin
	Useful for Screening Additional Selectivity	Limited Corrected Spectra Available		Dosimetry Easy Sample Prep	
	Due to Perturber		Eluontenan and El-	nhomecenos)	
Inc.	Low Temp	Contraction Contraction	Fuellest Contraction	(Limited Servi Field	1 T Technicus
Luminescent PACs	Higher Sensitivity, Specificity than RT	Gryogenic Apparatus More Complicated	Excellent Sensitivity	Deployability	- Shpolskii Spectra - Laser-line
PCBs	Vibrational Structure	Need Skilled Operator	pptrillion in Optimal Cases		- Site Selection - Matrix Isolation
	Similar to Raman	Lass Reference	Imornived with Laser		I ow Temperatures
	Quantitation Over 6 Orders of Magnitude	Spectral Data than RT	Laser		77 K to 4 K
	Distinguish Isomers	Dependent			
	Very Selective - Enhanced by Time and Wavelength Variability				

Source: Eastwood and Vo-Dinh (1991)

10.4 LUMINESCENCE/SPECTROSCOPIC ANALYTICAL TECHNIQUES

10.4.3 Other Spectrometric/Spectro-Photometric Techniques

Other Names Used to Describe Method: Ultraviolet (UV) spectroscopy, UV spectrophotometry, visible absorption spectroscopy, spectrophotometry, IR infrared (laser diode) spectrometry, photoacoustic spectrometry.

<u>Uses at Contaminated Sites</u>: **IR spectrometry**: Identifying and characterizing amorphous and crystalline inorganic or mineral phases; performing functional-group and qualitative analysis of organic compounds.

<u>Method Description</u>: Spectrophotometry encompasses a number of techniques involving measurement of the absorption spectra of narrow band-widths of radiation (visible and ultraviolet). Colorimetric techniques discussed in Section 10.5.1 require spectrophotometric measurements, as do luminescence techniques discussed in Section 10.4.2. Infrared (IR) spectrometry involves the measurement of infrared radiation absorption bands from low-level transitions between molecular energy levels. Different inorganic and organic functional groups have distinctive absorption spectra that help identify mineral or chemical phases in a sample. Table 10.4.3 provides additional information on UV-visible absorption, dispersive, and near-infrared methods.

<u>Method Selection Considerations</u>: Spectrophotometry: Integral to other techniques covered elsewhere. IR Spectrometry Advantages: Most useful when used in conjunction with x-ray diffraction (XRD) because it is capable of characterizing amorphous inorganic and mineral phases, which cannot be detected by XRD (Section 10.6.1). IR Spectrometry Disadvantages: (1) Results are primarily qualitative and require use of other techniques for definitive identification (quantitative analysis of multicomponent systems is possible, but very difficult); and (2) for solids samples, particle size must be less than the wavelength of the infrared radiation.

<u>Frequency of Use:</u> IR Spectrometry: Relatively common laboratory method for mineralogical study. Use for characterization of soils and waste has been limited.

Standard Methods/Guidelines: White and Roth (1986-IR spectrometry).

Sources for Additional Information: Eastwood and Vo-Dinh (1991).

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Table 10.4.3 General Characteristics of UV-Visible Absorption, Dispersive Infrared, and Near Infrared Techniques for Field Analysis

Applicability	Advantages	Limitations	Sensitivity	Current Field Applicability	Related Lab Techniques & Sensors
<u></u>		UV-vis Al	bsorption		
Polyaromatic Compounds (PACs) Dyes Colorimetric Reaction Products	Matura Technique Instrumentation Readily Available Good Quantitative Accuracy for Single Compounds and Simple Mixtures Few Interferences by Nonaromatics Spectral Data Available	Unspecific (Compared to IR and Luminescence) Extensive Sample Preparation Quantitation may be Affected by Solvent, Polanty, or Medium, Chemical Complexation	Moderate Sensitivity ppm - ppb in Favorable Cases	Portable - Hand-held Colorimeter - Colorimetric Kits Field Deployable Instrumentation with Multichannel Detectors HPLC Detectors	UV-VIS Technique: - FT - Derivative LT Matrix Isolation Reflectance Photoacoustic Spectroscopy Fiber Optic Colonimetric Sensors Multichannel Detectors - Diode Arrays - CCDs
	·	Infrared (L	Dispersive)		
Organic and Inorganic Determination of Specific Functional Groups	Higbly Specific Structural Data on Group Frequencies Mature Technique Instrumentation Widely Available Spectral Libraries Available	Mid/low Sensitivity Water is Interferent Requires Special Optics/Solvents Quantitation Difficultios Week Optical Sources and Detectors	Less Sensitive than UV-vis Absorbance Much Less Sensitive than Fluorescence ppthousand to ppm in Favorable Cases	Portable and Field Instruments Available Portable Unit with Gas Cell Ouantitation of Grease and Oil ATR Attachments for Solids, Oils	FTIR GC/LC-FTIR
		Near l	nfrared	. i	·····
Single Compounds Simple Matrices Organics Overtones	Sources and Optical Materials Better than Mid-IR Optically Good Sensor Materials Can Distinguish Major Components of Simple Matrix Fewer Interferences than Mid-IR	Less Spectral Structure than Mid-IR - Overtone Overlap - Less Specificity - Interpretation Complicated Not Useful for Complex Matrices Signal Processing and Pattern Recognition Required	Low Sensitivity 10-1 ppthousand	Portable Near-IR Instrument with Fiber Optic Probe Characterization ol Oil Bulk Chemical Analysis	Surface/Pollutant Interaction Studies Near IR Sensors Process Control

Portable:

 Battery powered
 Generator powered
 Can fit.in mobile lab

 One person can carry
 Compact, two people can lift (several instruments in mobile lab)
 Complex or fragile instrument

 Little sample prep. (< 10 min.)</td>
 Relatively simple sample prep. (< 1 hr.)</td>
 Often considerable sample prep. (> 1 hr.)

 Instrument cost < \$30,000</td>
 Instrument cost \$30,000
 Instrument cost \$30,000

 Analysis cost < \$30</td>
 Analysis cost \$30 • \$200
 Analysis cost \$ \$200

Field Deployable:

Definitions of abbreviations as used in this table are:

 ATR
 Attenuated Total Relicctance
 PAC
 Polyaromatic Compounds

 CARS
 Coherent Anti-Stokes Raman Spectroscopy
 PAH
 Polyaromatic Hydrocarbons

 CCD
 Charge-Coupled Davice
 PCB
 Polychlorinated Biphenyls

 FTIR
 Fourier Transform-Infrared Spectroscopy
 pb/ppm
 part per billior/part per million (mg/mL, µg/mL)

 GC
 Gas Chromatography
 RTP
 Room Temporature Phosphorescence

 HPLC
 High Performance Liquid Chromatography
 SERS
 Surface-Enhanced Raman Spectroscopy

 IR
 Infrared Spectroscopy
 SFC
 Supercritical Fluid Chromatography

 LC
 Liquid Chromatography
 TLC
 Thin-Layer Chromatography

 LT
 Low Temperature
 UV-vis
 Ultraviolet-Visible Spectroscopy

 NRS
 Normal Raman Spectroscopy
 UV-vis
 Ultraviolet-Visible Spectroscopy

Semi-field Deployable:

Source: Eastwood and Vo-Dinh (1991)

10.4 LUMINESCENCE/SPECTROSCOPIC ANALYTICAL TECHNIQUES

10.4.4 Other Spectroscopic Techniques

Other Names Used to Describe Method: Infrared (IR) spectroscopy, high resolution/long range Fourier transform infrared (FTIR) spectroscopy; Light detection and ranging spectroscopy (LIDAR), including differential scattering LIDAR (DISC) and differential absorption LIDAR (DIAL); IR reflectance/transmission spectroscopy; Raman spectroscopy (RS), (surface enhanced raman scattering (SERS).

Uses at Contaminated Sites: FTIR: Remote monitoring of air contaminants.

Method Description: IR spectroscopy: A field-deployable long-path Fourier transform infrared (FTIR) spectrometer currently is being tested by EPA. The instrument measures the absorption caused by infrared-active molecules. Pollutant inorganic and organic gas concentrations are determined by setting up a retroflector up to 1 kilometer from the spectrometer and transmitting an infrared beam that is returned to the detector (Figure 10.4.4a). Analysis is performed by using a reference spectrum of known concentration and least square fitting routines. The instrument measures various airborne vapors, including both organic and inorganic compounds. Figure 10.4.4b illustrates four applications of IR spectroscopy using differential scattering and absorption LIDAR, techniques that are being tested by the U.S. Army. RS encompasses a variety of techniques that involve detection and analysis of the scattering of radiation. Raman techniques differs from IR spectroscopy by using visible light to obtain structurally unique vibrational and rotational spectra. In the laboratory, RS can be used to identify functional groups to determine mineral phases. SERS is a relatively new analytical technique in which a sorptive surface provides a signal enhancement of up to a million times compared conventional IR spectroscopy, thus greatly reducing the detection limit. Reflectance/transmission spectroscopy in the near and far infrared portion of the electromagnetic spectrum has been proposed for airborne remote sensing identification of surface spills of benzene, toluene, TCE, and gasoline, but has not been field tested. Table 10.4.4 provides additional information about FTIR, normal RS, surface enhanced spectroscopy, and resonance raman methods.

Method Selection Considerations: Long-Path FTIR Advantages: (1) Measurements are rapid (a few minutes), allowing temporal profiles of pollutant gas concentrations; and (2) a range of volumes can be sampled by changing the distance between the instrument and the retroflector. Long-Path FTIR Disadvantages: Instrumentation is still in developmental stages. RS Advantages: (1) Is the best complement to IR spectrometry (Section 10.4.3) because it is able to discern vibrations from functional groups that are not discernible in the IR spectra; (2) resolution allows observation of particles as small as 1 micron, allowing characterization of individual particles; and (3) when combined with high pressure liquid chromotography (Section 10.5.3), can be used with solid and liquid samples to test for nearly all substances on EPA's priority pollutant list, including semivolatiles. RS Disadvantages: (1) Is nondefinitive and qualitative (only identifies functional groups); (2) data interpretation is complex; and (3) instrument availability is limited.

Frequency of Use: Uncommon.

Standard Methods/Guidelines: --

Sources for Additional Information: Eastwood and Vo-Dinh (1991), Thompson et al. (1989-IR and Raman Spectroscopy), U.S. EPA (1991b, 1992-portable FTIR). See also, Table 10-5.







Figure 10.4.4 Several infrared spectroscopic techniques: (a) Schematic of an infrared radiation source, mirror, and FTIR spectrometer equipped with telescopes to allow long-path analysis (Moore et al., 1991); (b) Four applications of differential scattering and absorption Lidar (Mackay, 1991).

Table 10.4.4 General Characteristics of Fourier Transform Infrared, and Raman Spectroscopic Techniques for Field Analysis (See end of Table 10.4.3 for definitions and abbreviations)

Applicability	Adventages	Limitations	Sansitivity	Current Field Applicability	Rolated Lab Techniques & Sensors
e		Infrared (Four	ler Transform)		
Organic and Inorganic Determination of Specific Functional Groups Routinely Used for Real-Time GC and Vapor Analysis	Highly Specific Structural Data on Group Fraquencies Instrumentation Widely Available Real-Time Flow through Vapor Applications - GC-FTIR Spectral Libraries	Less Sensitive than Luminescence Requires Special Optics/Solvents Can Tolerate Some Water (Background Subtraction) Organics Detection 1-10 ppthousand in Water	More Sensitive than Dispersive IR - Signal Averaging ppm to subppm in Favorable Cases	Field and Semi-field Deployable - With or Without GC - Volatiles/Semivolatiles Adaptable to Use with SFC	GC/LC-FTIR Matrix Isolation - LT for Sansitivity Microscopy
					1
·		Normal Reman	Spectroscopy (NRS)		
Organic and Inorganic Aqueous Solutions Biological Matrices Polymers	Specific as IR for Structural Information Different Selection Rules - Complements IR	Fluorescence Interfer- ence in UV-vis Requires Laser Source Relatively Complex Instamentation	Moderate Sensitivity 1000 - 20 ppm	Semi-field Deployable Instruments under Development	Research in: - Aqueous Solutions - Biological Matrices - Polymers Special Raman Techniques
roymus	Fewer Interfarences than IR in vis or near-IR Regions Water and Glass not Interferences Good Optics and Solvents Available Can Handle Unusual Sample Shaney/Firze	Requires Skilled Operator Not as Mature as IR Relatively Poor Limits of Detection			- SERS - Resonance - CARS - Microprobes - Microscopy LT Applications
		Cudese Esheroed Ba	 		ļ
Mary Dation	Constitute Official and	Surface Enhanced Kar	nan Spectroscopy (SEF	(S)	1. <u>.</u>
Many Politicity Demonstrated for: - Pyricine - Pyricine - PAHs - Pesticidas	Array Sensitive than Norre Sensitive than Normal Raman As Sensitive as Luminescence in	Relatively New Tech. Surface/Substrate Material Dependent Reproducibility Requires Laser and	Selected Analytes ppm - ppb in Favorable Cases	Instrumentation under Development	Microprobes Surface Studies
	Favorable Cases No Interference by Water (See Also NRS)	Spocial Substrate Not all Analytes Enhanced Equally Few Spectral Libraries (See Also NRS)			Fiber-Optic Sensors HPLC (under Development) Multichannel Datectors
		Resona	ince Raman		
PACs Absorbing in UV Phenols	/ Specific in Structure May Eliminate Fluoroscance Background (See Also NRS)	Only Chromophore Vibrations Enhanced Limited to UV Ab- sorbing Compounds - Mainly PACs Quantitation Difficult Not Comparable to Other Raman Techniques UV Laser Source	Fair Sensitivity in Favorable Cases with Chromophore Vibrations	Many Practical Difficulties	Chromophore Characterization Biological Application
		Complex Instrumentation (See Also NRS)			

Source: Eastwood and Vo-Dinh (1991)

10.5 WET CHEMISTRY ANALYTICAL TECHNIQUES

10.5.1 Colorimetric Techniques/Kits

Other Names Used to Describe Method: Colorimetry (various field kits using colorimeters/filter photometers/spectrophotometers [see Section 10.4.3]), titrimetry.

<u>Uses at Contaminated Sites</u>: Hach kits: Analyzing Al, Cd, Cr, Co, Cu, Fe, Pb, Mn, Ni, N, P, Ag, and Zn; Hanby kits: Analyzing petroleum hydrocarbons, PAHs; Other kits: Analyzing explosive (TNT/RDX), PCBs, chlorinated organics. Many ground-water tracers can be analyzed using colorimetric techniques.

<u>Method Description</u>: Titrimetry is a wet chemistry procedure by which a solution of known concentration (a standard solution) is added to a water sample or soil-solute extract with an unknown concentration of the analyte of interest until the chemical reaction between the two solutions is complete (the equivalence point of titration). Titrimetry requires an abrupt change in some property of the solution at the equivalence point, which is typically indicated by a change in color produced by an added dye, or by monitoring changes in pH with a meter (electrometric titrations). Colorimetry also involves mixing of reagents of known concentrations with a test solution, but in specified amounts that result in chemical reactions in which the absorption of radiant energy (color of the solution) is a function of the concentration of the analyte of interest. At the simplest level, concentrations can be estimated with visual comparators. Filter photometers can be used for many routine methods that do not involve complex spectra, and precise work is done with spectrophotometers (see Section 10.4.3). Titrimetric and colorimetric techniques are well suited for development of wet chemistry field test kits, and such kits are available commercially for many inorganic and some organic compounds. Figure 10.5.1 shows sample instructions for chromium using a Hach test kit. HNU/Hanby test kits use reagents that can be used in the field without use of a digester. Spectrochem has developed a kit that detects major classes of chemicals in water (chlorinated hydrocarbons, carbamates, and organic phosphated insecticides).

<u>Method Selection Considerations</u>: Titrimetry and Colorimetry Advantages: (1) Procedures are relatively simple and amenable to the development of field test kits for many analytes; (2) best suited for preliminary screening where only a few contaminants or analytes are of concern or interest; and (3) field test kits are available for most heavy metals. Titrimetry and Colorimetry Disadvantages: (1) Are time consuming if a large number of samples must be analyzed; (2) each analyte of interest requires different reagents and test procedures making analysis of multiple analytes time consuming; (3) strict QA/QC procedures are more difficult to follow in the field using test kits; and (4) availability of colorimetric field test kits for specific toxic organics is still relatively limited.

<u>Frequency of Use</u>: Colorimetric techniques are commonly used for field analysis of nutrients in soil and in ground-water tracer studies. Use of colorimetric field test kits for field screening of contaminants is a relatively new and promising field screening technique.

Standard Methods/Guidelines: --

Sources for Additional Information: Davis et al. (1985-colorimetry, titrimetry), Fishman and Friedman (1989-colorimetry, titrimetry), U.S. EPA (1987).

CHROMIUM, TOTAL 1.5-Diphenylcarbohydrazide Method



1. Select the sample amount from tables below and digest according to procedure in Section II.

Note: If sample cannot be analyzed shortly after sampling. see Section IV for storage and preservation information.

Note: This is an EPA-approved method only if preceded by an EPA-approved nitric acid digestion. The Digesdahl digestion procedure is not EPA approved and cannot be used for permit reporting purposes. (See digestion information on page 22.)

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2. Use analysis volume 3. Fill a second 25-ml in the tables below that sample cell with corresponds to the deionized water to the sample amount selected 25-ml mark (the reagent in Step I. Pipet analysis blank).

volume into a 25-ml mixing graduated cylinder. If aliquot is more than 0.5 ml, pH adjust according to the last step in the digestion procedure in Section II. Dilute to the 25-ml mark with deionized water, if necessary. Pour contents of cylinder into a 25-ml sample cell.

Note: For proof of accuracy, use a 0.25 mg/l chromium standard solution (preparation given in the Accuracy Check) in place of the sample.

4. Add the contents of one Chromium 1 Reagent Powder Pillow to each

cell. Swirl to mix.

Method 8023

L	IQUID SAMPLES	;
Expected Conc.	Sample	Analysis
Chromium	Amount	Volume
(mg/l)	(ml)	(#21)
0.05-2.0	10.0	20.0
0.20-8	20.0	10.0
0.75-33	10.0	5.00
7.5-330	5.00	1.00
75-3300	1.00	0.500
	OIL SAMPLES	
Expected Conc.	Sample	Analysis
Chromium	Amount	Volume
(mg/kg)	(8)	(ml)
8-330	0.25	20
20-820	0.20	10
50-2200	0.15	5
350-16000	0.10	1
5	OLID SAMPLES	
Expected Conc.	Sample	Analysis
Chromium	Amount	Volume
(mg/kg)	(g)	(<u>m</u> l)
4.0-165	0.500	20.0
10-410	0.400	10.0
25-1100	0.300	5.00
190-8200	0.200	1.00
750-33000	0.100	0.500

CHROMIUM, continued



5. Place cells in a boiling water bath and wait for 5 minutes.

Notes If a precipitate forms while heating, add a second Chromium 1 Respent Powder Pillow and continue heating.



6. Remove cells from the water bath and cool to 25 °C under tap water. If necessary, add deionized water to the 25-ml mark of the sample cell.



cell. Swirl to mix.

Note: Add the contents of a

second Chromium 2 Respent

Chromium 1 Reagent Powder

ZERO

Pillow was added in Step 5.

Powder Pillow if a second

8. Add the contents of 7. Add contents of one Chromium 2 Reagent one Acid Reagent Powder Powder Pillow to each

Pillow to each cell. Swirl to mix. Note: Test results will not be

affected if a small portion of this reagent does not dissolve. Add contents of a second Acid Respect Powder Pillow if a second Chromium 1 Reagent Powder Pillow was added in Step 5.



9. Add contents of one ChromaVer 3 Chromium Reagent Powder Pillow to each cell. Swirl to mix.

Note: A purple color will develop if chromium is present.

The Pour-Thru Cell can be used.

10. Wait 5 minutes for

11. Zero instrument the color to develop. with reagent blank using settings below. Read the Note: Do not wait more than

mg/l chromium of other 20 minutes before completing cell. Steps II to 12.

> DR/3000 Program No. 13 Wavelength 540 nm



Wavelength 540 nm DR/700

Wavelength 550 nm Note: See Section I for information on instrument standardization.

Module No. 55.01



12. Calculate the total chromium (Cr) concentration of the sample using the following formula:

> total mg/l Cr = A × 2500 BxC

WHERE A = mg/l read, Step 11 B = ml (g) sample amount, Step 1 C = ml analysis volume, Step 2

Note: For solid and oil samples express the resulting concentration as mg/kg and not 25 mg/l.



10.5 WET CHEMISTRY ANALYTICAL TECHNIQUES

10.5.2 Immunochemical Techniques

Other Names Used to Describe Method: Enzyme immunoassay (EIA), enzyme linked immunosorbent assay (ELISA), radioimmunoassay (RIA), fluoroimmunoassay.

Uses at Contaminated Sites: EIA: Analyzing BTX (benzene, toluene, xylene), PCB, PCP (water, soil), cocaine, heroin, and pesticides.

<u>Method Description</u>: EIA techniques that involve the use of antibody reagents that react with the analyte of interest to produce reactions that can be analyzed colorimetrically are a recent development for trace organic analysis (see Section 10.5.1 for additional discussion of methods for colorimetric analysis). Figure 10.5.2 shows procedures for an EIA test for pentachlorophenol (PCP). Other types of immunoassay techniques include radioimmunoassay and fluoroimmunoassay. (See also, discussion of bioassays in Section 10.6.6.)

<u>Method Selection Considerations</u>: EIA Advantages: (1) Is the best suited technique for preliminary screening where only a few contaminants or analytes are of concern or interest; (2) EIA test kits are very simple, rapid (minutes), and inexpensive; and (3) have the potential for specific field tests for a large number of toxic organics with very low detection limits (ppb). EIA Disadvantages: (1) Is time consuming if a large number of samples must be analyzed; (2) each analyte of interest requires different reagents and test procedures making analysis of multiple analytes time consuming; (3) strict QA/QC procedures are more difficult to follow in the field using test kits; and (4) availability of kits for specific toxic organics is relatively limited at this time.

<u>Frequency of Use</u>: Enzyme immunoassays are a relatively new technique and have excellent potential for more extensive use in the future.

Standard Methods/Guidelines: --

Sources for Additional Information: U.S. EPA (1988b-immunoassays/FM-D4, 1991b, 1992). See also, Table 10-5.



Figure 10.5.2 Procedures for enzyme immunoassay test for PCPs: (A) Antibody disks, (B) read-out disks, (C) absorbent blotting reservoir, (D) crush vial containing lyophilized antibody (DuQuette et al., 1991).

10.5 WET CHEMISTRY ANALYTICAL TECHNIQUES

10.5.3 Liquid Chromatography

Other Names Used to Describe Method: High pressure/performance liquid chromatography (HPLC), thin-layer chromatography (TLC), ion (exchange) chromatography.

<u>Uses at Contaminated Sites</u>: HPLC: Analyzing PAHs and phenols; TLC: Analyzing nitrogen-containing aromatics; Ion chromatography: EPA Method 300.0 (Kopp and McKee, 1983) covers the following ions: Cl, F, nitrate-N, nitrite-N, orthophosphate-P, and sulfate; also can be used to analyze halide and fluorinated organic acid dyes in tracer studies.

<u>Method Description</u>: Liquid chromatography is a type of chromatography where the mobile liquid phase containing analytes of interest is injected into a stationary phase that is either liquid or solid. Numerous specific techniques, such as partition, adsorption, ion exchange, paper, and TLC, have been developed. Ion chromatography involves separation of ions (typically anions) on a column of ion exchange resin, which are detected conductimetrically (Figure 10.5.3). A TLC technique with potential for separation of nitrogen-containing compounds in the field has been developed. A field operable HPLC unit using UV/visible and fluorescence detectors (see Section 10.4.2) appears to be the best field screening technique for PAHs (see advantages below).

Method Selection Considerations: In general, liquid chromatography is able to detect more compounds than GC, but at generally higher detection limits. Ion Chromatography Advantages: (1) Is a well established technique for separation of both organic and inorganic species; (2) several ions can be measured in a single aqueous sample; (3) eliminates many of the interferences associated with other techniques, and is capable differentiating species of the same ion in some cases; and (4) sensitive and has a wider range of applicability so that accurate measurement can be made on samples containing moderate to substantial ionic concentrations. Ion Chromatography Disadvantages: (1) Very high concentrations of an ion relative to another ion of interest might interfere or preclude measurement of the ion present in lower concentrations; and (2) individual measurements have a relatively low dynamic range, so separate dilutions might be required to bring a sample concentration into the optimum analytical range. HPLC (for PAHs) Advantages: (1) Instrumentation requires fewer gases for field analyses; (3) larger sample volumes can be injected compared to GC, yielding lower method quantitation limits; and (4) provides better resolution (ppb) than GC for comparable analysis time. HPLC (for PAHs) Disadvantages: New method for which there has been relatively little actual field experience.

<u>Frequency of Use</u>: Ion chromatography is commonly used for laboratory analysis of major anions. Field application of other liquid chromatographic techniques is a new development that appears promising for specific applications, such as detection of PAHs (HPLC) and nitrogen-containing and other polynuclear aromatic compounds (TLC).

Standard Methods/Guidelines: --

Sources for Additional Information: Davis et al. (1985), Fishman and Friedman (1989), Hassett (1982-highpressure liquid chromatography), Thompson et al. (1989-ion chromatography). See also, Table 10-5.



Ion chromatography - anions

Figure 10.5.3 Ion chromatography system for anions (Fishman and Friedman, 1989).

10.5 WET CHEMISTRY ANALYTICAL TECHNIQUES

10.5.4 Electrochemical Techniques

Other Names Used to Describe Method: Coulometry, voltammetry, polarography. Techniques covered elsewhere: pH, Eh, DO, electrical conductance (Section 10.1); ion-selective electrodes (see Section 5.5.5); potentiometric/amperometric/conductometric electrochemical sensors (see Section 10.6.5).

Uses at Contaminated Sites: Coulometry: Detecting ionic tracers. Voltammetry/polarography: Determining if trace metals, ions, and organics are in soils, waters, and sediments.

<u>Method Description</u>: Coulometric methods of analysis measure the quantity of electricity (in coulombs, the amount of electricity flowing during the passage of a constant current of 1 ampere for 1 second) required to carry out a chemical reaction. Primary coulometric analysis involves direct reactions by oxidation or reduction at the proper electrode. Secondary coulometric analysis involves indirect reactions between the solution and a primary reactant produced at one of the electrodes. Voltammetry is the area of electroanalytical chemistry involved in measuring the current at an electrode as a function of potential or voltage. Numerous specific techniques have been developed and only a very general description is provided here. Polarography is a voltammetric method in which a dropping mercury electrode (DME), is used for very precise control of changes in currents applied to the electrode. Plots of current vs. potential allow identification of the analyte based on the shape of the curve and concentration based on wave height. Stripping voltammetry is a two-step process in which electrolytic deposition of the chemical species is followed by application of a voltage scan to cause electrolytic dissolution (stripping) of the species back into solution at characteristic potentials.

<u>Method Selection Considerations</u>: Coulometric Advantages: (1) Instrumentation is relatively simple; and (2) well suited for trace analysis of ionic tracers, such as chloride and bromide (see Section 4.3.1). Coulometric Disadvantages: Is not well suited for analysis of complex mixtures. Polarography Advantages: (1) Instrumentation is relatively simple; (2) depending on specific method, capable of sensitivity to sub-ppm; (3) good selectivity allows determination of many constituents without prior chemical separation; and (4) capable of measuring large ranges of concentration, ranging from concentrated extracts from solids to dilute natural waters. Polarography Disadvantages: (1) Is a relatively simple method requiring minimal sample preparation; and (2) is the most sensitive electroanalytical technique currently available (capable of metals analyses down to ppt level). Stripping Voltammetry Disadvantages: (1) Method is not likely to be useful for field or mobile laboratory; and (2) highest sensitivities are difficult to achieve for routine analysis.

<u>Frequency of Use</u>: Measurement of pH, Eh, and specific conductance (Section 10.1) are the most commonly used electrochemical techniques. Polarography and stripping voltammetry are not likely to be useful in field or mobile laboratory applications.

Standard Methods/Guidelines: --

Sources for Additional Information: Davis et al. (1985-coulometry), Fishman and Friedman (1989-voltammetry/polarography), Street and Peterson (1982-polarography and stripping voltammetry).

10.6 OTHER ANALYTICAL TECHNIQUES

10.6.1 Radiological Techniques

Other Names Used to Describe Method: Analytical techniques: X-ray diffraction (XRD), proton induced X-ray emission (PIXE), glancing incidence X-ray analysis (GIXA), (instrumental) neutron activation analysis([I])NAA); Techniques covered elsewhere: Nuclear borehole techniques (Section 3.3), radioisotope single-borehole tracer methods (3.5.4), radioisotope tracers (Section 4.4.5), X-ray fluorescence (Section 10.4.1), and electron microprobe analysis (Section 10.6.4).

<u>Uses at Contaminated Sites</u>: Detecting natural radioisotopes (i.e., gamma log, Section 3.3.1); performing elemental and mineralogical analysis (XRD, PIXE, INNA, gamma spectrometry, electron microprobe analysis, XRF); performing tracer studies (see Sections cited above).

<u>Method Description</u>: XRD involves the identification of minerals by directing a monochromatic x-ray beam at a powdered sample and using a scintillation, proportional, or geiger counter (see above) to detect the intensities and diffraction angles as the beam is rotated around the sample. Crystalline minerals can be identified by the characteristic position and intensities of the diffraction peaks. PIXE analysis uses a high-speed proton beam to displace inner-shell electrons of the sample elements. When the electrons return to their proper shells, x-rays are emitted that have energies characteristic of the elements and proportional to their mass. Computer processing provides data on all elements present in a given sample. In INNA, powdered samples are irradiated for specified times and neutron fluxes, depending on the elements of interest. Gamma-ray spectra of the irradiated samples are measured with Ge(Li) detectors coupled with multi-channel analyzers.

Method Selection Considerations: All radiological analytical methods have the disadvantage of requiring special health and safety precautions. XRD Advantages: (1) Is a relatively simple and inexpensive bulk sample method; (2) provides simultaneous multi-mineral characterization; and (3) is best used in conjunction with other more quantitative species-specific chemical methods. XRD Disadvantages: (1) Estimates of mineral percentages are only semi-quantitative; and (2) minerals present in small amounts often are difficult to discern in multicomponent mixtures. PIXE Advantages: (1) Provides simultaneous multi-element characterization; (2) is rapid (30 minutes/sample); and (3) is good for initial screening to identify presence of elements for which more precise analysis should be done. PIXE Disadvantages: (1) Instrumentation is expensive (but somewhat cheaper than ICP-AES); and (2) has relatively high detection limits (10s to 100s ppm). INAA Advantages: (1) Requires less sample preparation time compared to AAS and ICP-AES; and (2) sensitivity compares well with conventional spectrometric techniques for many elements. INAA Disadvantages: (1) Requires nuclear reactor for irradiation of samples; (2) sensitivity is highly dependent on the exact elements being measured and on the sample matrix; and (3) some elements, such as lead, cannot be measured.

Frequency of Use: Analytical techniques: XRD is a widely used method for mineral identification. PIXE and INAA are commonly used for analysis of coal fly ashes, but have received limited use for contaminated site characterization. All three methods are primarily laboratory methods, although XRD instrumentation probably could be used in a mobile laboratory. See neutron activation log (Section 3.3.5) and neutron-lifetime log (Section 3.3.6) for field applications using principles of neutron activation analysis.

Standard Methods/Guidelines: Analysis of radioactive substances: Thatcher et al. (1977); XRD of soil samples: ASTM (1985).

Sources for Additional Information: Analytical techniques: Davis et al. (1985-gamma, beta, NAA), Helmke (1986-neutron activation analysis), Thompson et al. (1989-INNA, PIXE, XRD), Whittig and Allardice (1986-XRD), Wong and Carlsen (1991-tritium field screening).

10.6 OTHER ANALYTICAL TECHNIQUES

10.6.2 Gravimetric/Volumetric Techniques

Other Names Used to Describe Method: --

<u>Uses at Contaminated Sites</u>: Characterizing particle size distribution; measuring bulk density; measuring dissolved/suspended solids; calculating contaminant concentrations/flux; measuring water flow during pumping tests (Section 4.2); measuring soil gravimetric moisture (Section 6.3.1).

Method Description: Gravimetric techniques involve measuring the mass of the material of interest. For chemical analyses, a mechanical or electronic analytical balance capable of measuring the mass of an object within 0.1 to 0.01 mg is used. Field applications require less sensitive devices, such as a hanging spring scale with a canvas sling or pail for weighing coarse fragments, and a scale or balance with 0.1 gram accuracy for weighing soil samples for field tests (Boulding, 1991). Volumetric techniques involve the measurement of volume. Volume of liquid samples for chemical analysis is easily measured by the use of graduated cylinders or sample containers of a known volume. For borehole aquifer characterization, volume is measured using flowmeters (Section 3.5.3), and in pumping tests, pumping rate can be determined in several ways: (1) Observing the time required to fill a container of known volume, (2) use of commercial water meters, (3) use of a circular orifice weir (Figure 10.6.2), or (4) channeling surface flow from the pump through flumes or weirs. For gases, volume typically is measured by using syringes of a known volume, or measuring the rate of gas flow through a tube of known diameter. Both gravimetric and volumetric measurements are required for soil characterization because the soils vary in bulk density (weight per unit volume), depending on the volume of pore space. There are four major methods for measuring bulk density: (1) The core method, which involves drying and weighing of an undisturbed core sample of known volume; (2) gamma-gamma logging (see Section 3.3.2); (3) the excavation method, which involves excavating an amount of soil (which is dried and weighed) and measuring the volume of sand required to fill the hole, or the volume of water required to fill a rubber-balloon; and (4) the clod method makes use of Archimedes' principle, and involves coating a clod of known weight with a water-repellent substance and weighing it first in air, then again while immersed in a liquid of known density.

Method Selection Considerations: A scale or balance of the required accuracy (see above) should be standard equipment for field investigations for soil characterization. Selection of appropriate volumetric measurement techniques for water and gases is straightforward. Bulk density is required for most vadose zone models (see Appendix C), and allows qualitative evaluation of the potential for transport of contaminants through the vadose zone. The core method is simple and accurate, but generally unsatisfactory in stony or very dry soils. Advantages and disadvantages of gamma-gamma logging are covered in Section 3.3.2. The excavation method is a simple and accurate field procedure, but lacks discrimination of localized horizons and is limited to around 12 centimeters below the surface. The clod method allows discrimination of localized horizons, but is more complex and usually gives higher bulk density values than other methods because interclod spaces are not taken into accurate scales for weighing samples are available (Section 6.3.1), and have the advantage of providing moisture content of the sample as well as bulk density.

<u>Frequency of Use</u>: Gravimetric and volumetric measurements are essential for the uses described above. The core method is probably the most commonly used method for measuring bulk density, followed by gamma-gamma logging.

Standard Methods/Guidelines: Gravimetric: Fishman and Friedman (1989). Soil bulk density: ASTM (1984rubber balloon method), Blake and Hartge (1986), Campbell (1991), Flint and Childs (1984).

Sources for Additional Information: Flow discharge measurement: Johnson (1964), Jorgensen (1969).



Figure 10.6.2 Volumetric techniques: Construction diagram of a circular orifice weir commonly used for measuring pumping rates of a high-capacity pump (Driscoll, 1986, by permission).

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10.6 OTHER ANALYTICAL TECHNIQUES

10.6.3 Magnetic Methods

Other Names Used to Describe Method: Magnetic susceptibility (MGS), electron spin resonance (ESR), nuclear magnetic resonance (NMR).

<u>Uses at Contaminated Sites</u>: MGS: Performing qualitative soil mineral characterization; detecting lateral changes in soil characteristics; ESR: Characterizing clay minerals and sorption of metals; NMR: Characterizing clay minerals and soil organic matter; borehole logging and soil moisture monitoring (see Sections 3.2.4 and 6.2.5).

<u>Method Description</u>: MGS is the tendency of atoms or ions in a sample to become aligned when placed in a magnetic field, and is obtained by measuring the magnetic moment per unit volume or mass induced in a sample by an applied magnetic field. The Gouy (Figure 10.6.3a) and Faraday (Figure 10.6.3b) susceptibility balances are two commonly used types. ESR measures the electron magnetic moment in solid, water, or air samples. The instrument consists of an electromagnet inducing a continuous magnetic field that can be varied in strength, a resonance cavity where the sample is positioned, a microwave source, and a detector that measures the sorption response of the sample (Figure 10.6.3c). NMR operates on the same principle as ESR, except that the nuclear magnetic moment (carbon or proton spectra) is measured instead of the electron magnetic moment. Figure 10.6.3d shows a NMR spectrometer using a radio frequency transmitter and receiver/detector that records proton or carbon spectra in response to variations in magnetic field. See also, section 3.2.4 for application of NMR as a borehole logging technique and for soil moisture monitoring.

<u>Method Selection Considerations</u>: MGS Advantages: Instrumentation and measurement procedures are relatively simple. MGS Disadvantages: Provides qualitative rather than quantitative information on mineralogy. ESR/NMR Advantages: (1) Are more amenable to quantitative interpretation than MGS; and (2) are well suited for controlled laboratory study of contaminant-soil interactions. ESR/NMR Disadvantages: (1) Are not well suited for the chemical characterization of complex soil chemistry; and (2) instrumentation is generally too bulky for use in mobile laboratories.

<u>Frequency of Use</u>: Laboratory applications for study of soil mineralogy and organic matter are relatively recent, but are becoming more widely used. Use for contaminated site characterization has been limited.

Standard Methods/Guidelines: McBride (1986).

Sources for Additional Information: MGS: Fine et al. (1992), Mullins (1977), Williams and Cooper (1990), Woolcock and Zafar (1992); NMR: Bleam (1991), Thorn (1987). See also, references for Section 3.2.4 and 6.2.5.


Figure 10.6.3 Magnetic methods: (a) Gouy magnetic susceptibility balances; (b) Faraday magnetic susceptibility balance; (c) Diagram of typical X-band ESR spectrometer; (d) Diagram of NMR spectrometer (McBride, 1986, by permission).

10. CHEMICAL FIELD SCREENING AND ANALYTICAL METHODS

10.6 OTHER ANALYTICAL TECHNIQUES

10.6.4 Microscopic Techniques

Other Names Used to Describe Method: Optical microscopy, scanning electron microscopy (SEM), electron microprobe analysis (EMPA).

<u>Uses at Contaminated Sites</u>: Optical microscopy: Observating soil morphologic features; identification of coarsegrained minerals. SEM and EMPA: Assessing morphology, composition, and identity of minerals.

Method Description: Microscopic techniques involve the visual identification and characterization of soil/solid waste morphologic features and minerals with instruments ranging from magnification of up to 20 times, using a simple hand lens, to magnifications of 50,000 times using an electron microscope. As magnification increases, resolution increases, but the area viewed decreases (Figure 10.6.4a). Optical microscopy: Stereoscopic microscopes can be used in the field for more detailed visual inspection of soil morphologic features at magnifications of 20 to 80 times. Petrographic microscopes for mineral identification require the preparation of thin sections by impregnating samples with epoxy resin and grinding the samples to a precise thickness. The thin sections are examined with magnifications ranging from 50 to 400 times. Minerals are distinguished by their color in polarized and nonpolarized light, refractive index, and crystal morphology. SEM involves the irradiation of a sample with a focused electron beam with very short wavelengths (about 100,000 times shorter than that for visible light) that can provide high image resolution. Secondary electrons emitted from the sample produce a topographical image of the sample, and backscattered electrons provide some qualitative information on elemental composition. EMPA is similar to SEM, except that the electron beam also produces X-ray fluorescence (see Section 10.4.1), which allows for quantitative interpretation of elemental concentration as well as topographic images (Figure 10.6.4b).

<u>Method Selection Considerations</u>: Optical Microscopy Advantages: (1) Is a simple, nondestructive technique that allows mineral identification without intermediate calculations or inferences; (2) sample preparation and examination are relatively quick, simple, and inexpensive; and (3) use of stereoscopic microscopes in field (5 to 6 inches working distance, 20 to 80 power) allows for observation of soil features that cannot be readily seen by eye or with a hand lens. Optical Microscopy Disadvantages: (1) Preparation of thin sections for accurate identification of minerals is not readily done in the field; (2) accurate mineral identification requires an experienced and skilled observer; and (3) identification of fine-grained material can be very difficult and might require other methods, such as X-ray diffraction (Section 10.6.1). SEM Advantages: (1) Has very high resolution (magnification from 20 to 50,000 times; and (2) can differentiate heterogeneity among fine-grained particles as well as heterogeneity within individual particles. SEM Disadvantages: (1) Equipment is nonportable and expensive; and (2) elemental information and topographic image interpretation is largely qualitative. EMPA Advantages: Able to produce images that depict elemental distribution; EMPA Disadvantages: (1) Analyses are expensive and relatively few instruments are available; (2) quantitative results for most elements are limited to concentrations of 50 to 100 ppm; and (3) has lower resolution than SEM (up to 2500 times).

<u>Frequency of Use:</u> Optical microscopy and SEM are commonly used in the laboratory for mineral identification and characterization. Hand lenses are standard equipment for observation of soils in the field; use of optical microscopes in the field is uncommon, but more widespread use for examination of soils would be beneficial. EMPA is most commonly used in the field of metallurgy, but could be used more widely in soil and waste studies if limitations of expense and limited instrument availability were reduced.

Standard Methods/Guidelines: --

Sources for Additional Information: Cady et al. (1986-optical microscope), Goldstein et al. (1981), Sawhney (1986-electron microprobe), Thompson et al. (1989-Chapter 16).



(b)

Figure 10.6.4 Microscopy: (a) Schematic illustration of the relationship between increasing levels of resolution and the area of the field under view (Cady et al., 1986, by permission); (b) Schematic diagram showing components of an electron microprobe and the signal produced from a specimen surface irradiated with an electron beam (Sawhney, 1986, by permission).

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10. CHEMICAL FIELD SCREENING AND ANALYTICAL METHODS

10.6 OTHER ANALYTICAL TECHNIQUES

10.6.5 Other Chemical Sensors

Other Names Used to Describe Method: Electrochemical sensors (amperometric/galvanic cell sensors, semiconductor sensors, spectroelectrochemical sensors), piezoelectric sensors (piezoelectric quartz microbalance, surface acoustic wave [SAW] sensor).

<u>Uses at Contaminated Sites</u>: Field screening of contaminants in air, soil, and ground-water samples; SAW: Screening for toxic/organophosphorus gases; Semiconductor: Screening for organochlorine; Pyrolysis-EC: Screening for alcohols, epoxide, formaldehyde, CO, and H2S.

Method Description: Electrochemical sensors: As the name implies, these sensors measure an electrochemical response when the sensor comes in contact with the analyte(s) of interest. Amperometric gas sensors are the best developed sensor of this type (see Section 10.3.2). These sensors typically consist of electrodes in contact with an electrolyte-saturated insulator. Selective membranes allow the gas of interest to enter the insulator and redox reactions on the sensing-electrode surface generate a current that is proportional to the analyte concentration. Figure 10.6.5 illustrates an exploded view of a typical amperometric sensor. Amperometric sensors are capable of detecting levels as low as ppb of many organic and inorganic air pollutants. Use of amperometric sensors for detecting contaminants in ground water is in developmental stages at this time. Semiconductor sensors are designed to respond electrically to the substance of interest. A semiconductor sensor designed to detect low concentrations of chlorinated and brominated organic compounds in vapor and water (using membrane extraction [see Section 10.2.5]) has recently been tested in the laboratory. Piezoelectric sensors: Several types of sensors using piezoelectric materials, which develop an electrical response to changes in pressure, have been developed. Typically, oscillating crystals are used as sensitive gravimetric detectors. Selective coatings allow specific organic solvent vapors to be sorbed on the crystal. The changed mass of the crystal resulting from sorption changes the frequency of oscillation, which can be correlated with concentration. SAW sensors also use piezoelectric materials and coatings that selectively sorb the vapor or gas of interest. Changes in the mass or mechanical modulus of the surface coating are measured by the change in velocity of electrically generated Rayleigh (surface) waves, as measured by travel time from the source of receiving electrodes in the sensor. Concentration with SAW sensors is related to changes in velocity.

Method Selection Considerations: Amperometric Sensor Advantages: (1) Is inexpensive; (2) is simple and reliable (no moving parts and sensor output is usually a linear function of concentration); and (3) is portable (units with sensor, electronics, battery, and readout device can easily fit in a shirt pocket). Amperometric Sensor Disadvantages: (1) Separate sensor required for each compound of interest; and (2) applications for groundwater monitoring are in early developmental stages. Piezoelectric and SAW Sensor Advantages: (1) Are portable; and (2) SAW vapor sensors have higher sensitivity than gravimetric piezoelectric sensors. Piezoelectric and SAW Sensor Disadvantages: (1) General difficulty in developing selective coatings that are not affected by complex mixtures; and (2) separate sensor required for each compound of interest.

<u>Frequency of Use</u>: Amperometric sensors: Commonly used in ambient air quality monitoring. Other sensors: Emerging technology area.

Standard Methods/Guidelines: --

Sources for Additional Information: See Table 10-5.



Figure 10.6.5 Parts of a typical amperometric sensor (Schmidt et al., 1988).

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10. CHEMICAL FIELD SCREENING AND ANALYTICAL METHODS

10.6 OTHER ANALYTICAL TECHNIQUES

10.6.6 Other Biological Techniques

Other Names Used to Describe Method: Field: Short term field bioassessments, biomonitoring, laser/microbe bioassay (LMB), immunochemical techniques (Section 10.5.2). Laboratory: Bioassays, toxicity tests, biomarkers.

<u>Uses at Contaminated Sites</u>: Assessing actual or potential biological impacts of contamination at a site; monitoring the effect of effluent on organisms; assessing the treatability of contaminated soil or ground-water for bioremediation.

Method Description: At the simplest level, a qualitative assessment of the presence or absence of contaminants at a site can be made by observing whether any vegetation appears to have been killed or growth inhibited by the presence of toxic contaminants. Short-term field bioassessments: Field screening studies include collection of small mammals, fish, benthic invertebrates, and plants for the purpose of evaluating alterations in community structure, population dynamics, bioaccumulation of toxicants, and histopathology. The LMB system is a recently developed technique that has potential for use in the field. Nineteen isogenic strains of Bacillus subtilis are used to characterize and quantify the toxicants present in an aqueous solution. The response of the bacteria to toxic substances in the solution is monitored by differential light scattering from a laser beam. The different strains respond differently to different toxicants and a computer analyzes the measured responses to the known response profiles to identify the type and concentration of toxicant. Figure 10.6.6 shows an example of the use of mussels for field biomonitoring of the effects of potentially toxic effluents. A series of field cages filled with mussels are placed along a gradient of contaminant concentrations. After a period of time (usually 7 to 30 days) the mussels are retrieved and taken to a laboratory for further testing and analysis. Numerous laboratory methods have been developed for biological assessment of toxicity, many of which can be run in mobile laboratories (see below). These can be broadly classified as: (1) Toxicity tests using specific aquatic and terrestrial organisms and/or microorganisms to measure biological response to specific contaminants or mixes of contaminants; and (2) the analysis of biomarkers, which are molecular biological indicators that can directly link specific chemicals or classes of chemicals to observed biological effects. The microtox bioassay, a colorimetric technique (see Section 10.5.1) that uses microorganisms, has been used to determine the appropriate range of waste application loading for soil-based waste treatment systems (see reference in Table 10-5).

<u>Method Selection Considerations</u>: General Advantages: (1) Qualitative observations of inhibition of vegetative growth can very easily be made; and (2) more sophisticated field bioassessment methods allow for correlation of contaminant levels to actual biological impacts. General Disadvantages: (1) Field techniques have not been widely used at contaminated sites so procedures are not well established; (2) personnel with specialized training are required; and (3) equipment for more sophisticated techniques might not be readily available. LMB Advantages: (1) Equipment is field portable and relatively fast (around 1 hour for a single sample); (2) can distinguish between substances with cytotoxic and genotoxic properties; (3) potential for both high sensitivity and high specificity for numerous toxic chemicals and chemical classes; and (4) computer processing and output speeds up and simplifies interpretation of results. LMB Disadvantages: (1) Is a new technique that has received limited field testing; and (2) ability to distinguish compounds in real-world complex mixtures has not yet been demonstrated.

<u>Frequency of Use</u>: EPA's Environmental Research Laboratories at Duluth, Minnesota, and Narragansett, Rhode Island, have mobile laboratories set up for ambient and effluent toxicity testing, which have been used primarily as part of NPDES programs. Use of short-term field bioassessment methods at contaminated sites has been fairly limited in the past, but these methods are being used with increasing frequency.

Standard Methods/Guidelines: Britton and Greeson (1989-algal growth potential bioassay). U.S. EPA (1986a) contains recommendations for use of bioassays for evaluation of hazardous waste land treatment demonstrations.

Sources for Additional Information: U.S. EPA (1987-Section 12.6), Warren-Hicks et al. (1989). See Table 10-5 for additional references.



Figure 10.6.6 Mussel field cages used to transplant mussels along transects (DiBona et al., 1989).

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Table 10-4 Reference Index for General Approaches to Field Screening/Analytical Methods and Extraction Procedures

Торіс	References
General Approaches to Field Scree	ening
Symposia	U.S. DOE (1988), U.S. EPA (1988a, 1991a)
Review Reports/Papers	Chudyk (1989), Coffey et al. (1988), Eastwood and Vo-Dinh (1991), Jenkins et al. (1988, 1989), Koglin and Poziomek (1990), Montgomery et al. (1985), National Institute for Petroleum and Energy Research (1990), Poziomek and Koglin (1991), Remata et al. (1990), U.S. EPA (1982, 1987, 1988a, 1991b)
Agency Research Programs	<u>U.S. EPA</u> : Chapman and Fredericks (1988-FASP), Fribush and Fisk (1991), Transue et al. (1991-FASP), Tuttle and Chapman (1989), U.S. EPA (1992); <u>Other</u> : Cornell (1991-New Jersey), Frank et al. (1991-DOE), Mackay (1991-U.S. Army), Madden and Johnson (1992-U.S. Army)
QA/QC	Mackiewicz (1990, 1991), Poziomek and Koglin (1991-cite 8 references from U.S. EPA, 1991, that are not included here)
Sample Extraction Procedures	
Headspace Analysis	Crockett and DeHaan (1991-soil VOCs), Golding et al. (1991-VOCs in soil/water), Hewitt et al. (1991), Ho et al. (1988), Hogan (1991), Holbrook (1987), Pankow (1986, 1991), Roe et al. (1989), Sims et al. (1991-soil), Spittler et al. (1988-soil, 1991-water), Stuart et al. (1991-BTEX in soil/water), Wylie (1988)
Soil Vacuum Extraction	Golding et al. (1991), Spittler (1991)
Purge and Trap	Chochran and Henson (1988), Hein (1988), Liebman et al. (1991), Linenberg and Robinson (1991), Sherman et al. (1988a-concentrator purge & trap), Turner et al. (1991), Wise et al. (1991a), Wylie (1988); <u>P/CCW</u> : Pankow (1991), Pankow and Rosen (1988)
Thermal Treatment	<u>Microwave-Assisted Digestion</u> : Grohse et al. (1988); <u>Thermal Desorption</u> : Pankow and Isabelle (1982), Pankow and Kristensen (1983), Pankow et al. (1988), Robbins et al. (1990), Schlesing et al. (1991), Vandegrift (1988), Wise et al. (1991b); <u>Thermal Extraction/Pyran Thermal Chromatograph</u> : Greenlaw et al. (1989), Henry et al. (1988), Junk et al. (1991a,b), Overton et al. (1988a,b); <u>XRF</u> <u>Sample Preparation</u> : Bernick et al. (1991), Harding (1991), Ramsey et al. (1991)
Soil (Micro)extraction	Semivolatile Organics: Kasper et al. (1991), Transue et al. (1991); <u>PCB extraction</u> : Keller and Ganapathi (1991), Twomey et al. (1990); <u>Tritium</u> : Wong and Carlsen (1991)
Other Methods	<u>Supercritical Fluid Extraction</u> : Liebman et al. (1991), Lopez-Avila et al. (1991), Schulten and Schnitzer (1991), Wright and Fruchter (1992); <u>Membrane extraction</u> : Melcher and Morabito (1991); <u>Extraction Disks</u> : Poziomek et al. (1991)

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Table 10-5 Reference Index for Screening/Analytical Methods

Торіс	References
Total/Specific Vapor	
Detectors	<u>Comparisons</u> : Clay and Spittler (1982), Gervasio and Davis (1989), Robbins et al. (1990), Smith and Jensen (1987), Spittler (1980, 1991); <u>Explosimeter</u> : Aller (1984), Robbins et al. (1990); <u>Flame Ionization Detector</u> : Gervasio and Davis (1989), Hein (1988), Robbins et al. (1990); <u>Organic Vapor Analyzer</u> : Barber and Braids (1982), Glaccum et al. (1983), Hogan (1991), Jermakian and Majika (1988), Robbins et al. (1989); <u>Photoionization Detector</u> : Brose and Gross (1988), Gervasio and Davis (1988), Hare (1987), Robbins et al. (1990); <u>P/T Argon Ionization Detector</u> : Linenberg and Robinson (1991); <u>Unspecified</u> : Stetter et al. (1984); <u>Mercury Vapor Analyzer</u> : Brass et al. (1991)
Portable Gas Chromatograph	<u>GC Comparisons/Validation</u> : Homsher et al. (1988), Spittler (1991); <u>Gas</u> <u>Chromatographs</u> : Baker et al. (1991-GC/FID), Berkely (1991-GC/PID), Buchmiller (1989), Carney et al. (1991-retention indices), Clay and Spittler (1982), Crockett and DeHaan (1991), Fowler and Bennett (1987), Golding et al. (1991- GC/FID), Hewitt et al. (1991), Ho et al. (1988), Kaelin and Prichett (1991- GC/argon ionization detector), Keller and Ganapathi (1991), Linenberg (1988), Moore (1991-GC/PID), Moreton et al. (1991-soil BTEX), Overton et al. (1988c), Quimby et al. (1982-GC/OVA), Reynolds et al. (1991), Robbat and Xyrafas (1988a), Shangraw (1988), Sherman et al. (1988b), Spittler (1980, 1984-PCBs), Spittler et al. (1982-GC/FID), Stuart et al. (1991-GC/PID), Turner et al. (1991- PT/GC), Wander et al. (1988), Wohltjen et al. (1991); <u>GC/AES</u> : Szelewski and Wilson (1988); <u>GC/FTIR</u> : Gurka et al. (1986)
Fieldable/Mobile Mass	
Spectrometer; GC/MS	<u>GC/MS</u> : Bruell and Hoag (1984), Gurka et al. (1986), McGinnis and Hafferty (1987-PCP), Moy (1989-PCB), Sinha (1991), Transue et al. (1991-GC/ECD, PAHs, PCP); <u>Mobile Mass Spectrometer</u> : Duret et al. (1991), Hadka and Dickinson (1988), Klainer et al. (1991), Trainor and Laukien (1988); <u>GC/ITMS or</u> <u>IID (Ion Trap Mobility Spectrometry or Ion Trap Detector</u>): Cispar et al. (1991), Cooks et al. (1991), Leibman et al. (1991), McClennen et al. (1991-MINIMASS), Wise et al. (1991a, 1991b); <u>Tandem MS (MS/MS</u>): Wise et al. (1991a; <u>GC/MS</u> : Meuzelaar et al. (1991), Robbat and Xyferas (1988b); <u>Thermal Desorption ITMS- MS/MS</u> : Wise et al. (1991b), <u>Thermal Desorption GC/MS</u> : Robbat et al. (1991); <u>Semivolatile Thermal Extraction GC/MS</u> : Henry et al. (1988), Junk et al. (1991a,b), Overton et al. (1988d)
Mobile Laboratories	Ben-Hur et al. (1984-mobile MS/MS), Burger (1991), Chapman et al. (1986), Engels et al. (1984), Franks et al. (1985), Greenlaw et al. (1989-thermal chromatograph/MS), McGinnis and Hafferty (1987-PCP), Moy (1989-PCB), Tuttle and Chapman (1989); <u>Cost Analysis</u> : Ganapathi et al. (1988); <u>Dedicated</u> <u>Laboratory</u> : Freeman and Karmazyn (1988)
Ion Mobility Spectrometry (Plasma Chromatography)	Bell and Eiceman (1991-GC/IMS), Burroughs et al. (1991), Clement et al. (1992), Davis (1991-data analysis), Hoffland and Shoff (1991), Reategui et al. (1988), Richter (1991), Snyder et al. (1991-microorganisms), Wise et al. (1990)

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Table 10-5 (cont.)

Торіс	References
Fluorescence/Luminescence/Spec	troscopic Techniques
X-Ray Fluorescence (XRF)	Ashe et al. (1991), Barich et al. (1988), Bernick et al. (1991), Carlson and Alexander (1991-QA/QC), Chappell et al. (1986), Coetzee et al. (1986-XRF, ICP- AES), Cole et al. (1991-XRF/CLP comparison), Everitt et al. (1988), Florkowski et al. (1971), Freiburg et al. (1987-XRF, AAS, AES), Furst et al. (1985), Gabry (1991-XRF vs CLP), Glanzman (1988), Grupp et al. (1988), Harding (1991- EDXRF), Harding and Walsh (1990-EDXRF), Jenkins (1984), Kendall (1991b), Meiri et al. (1990), Perlis and Chapin (1988), Piorek and Rhodes (1988- calibration), Raab et al. (1990), Ramsey et al. (1991-EDXRF calibration), Sackman et al. (1988), Smith and Lloyd (1986-XRF, AAS), Watson et al. (1989)
UV Fluorescence	<u>PAH Solvent Extract</u> : Popp (1989), Saenz et al. (1991), Theis et al. (1991); <u>With</u> <u>HPLC</u> : Mann and Vickers (1988), Riddell et al. (1991); <u>With Fiber Optics</u> : Chudyk et al. (1988), Gillispie and St. Germain (1988), Haas et al. (1988, 1991), Kenny et al. (1988), Lieberman et al. (1991), Smith et al. (1988), Taylor et al. (1991); <u>UV Surface Contamination Detector</u> : Richter (1991); <u>Airborne</u> : Guenneberg (1978)
Other Luminescence Methods	<u>Synchronous Fluorescence</u> : Gammage et al. (1988, 1991); <u>Spectrofluorometer/Solid State Fluorescence</u> : Poziomek et al. (1991); <u>Room</u> <u>Temperature Phosphorescence</u> : Vo-Dinh (1984), Vo-Dinh et al. (1991)
IR Spectroscopic Methods	<u>Review</u> : Kendall (1991a), Phelps and DeSha (1991-LIDAR, FITR); <u>LIDAR</u> : Mackay (1991); <u>Mobile FTTR</u> : Fateley et al. (1990), U.S. EPA (1991a-five papers, not indexed separately); <u>Other Infrared</u> : Gurka et al. (1986-GC/FTTR), Kasper et al. (1991-soil hydrocarbons), Richter (1991-IR laser absorption); <u>IR</u> <u>Reflectance/Transmission</u> : King and Clark (1988)
Other Spectroscopy	<u>Ultraviolet-Visible Absorption Spectroscopy</u> : Beemster and Schlager (1991-with fiber optics), Schlager and Beemster (1991), Thompson (1974); <u>UV Derivative Spectroscopy</u> : Hager and Jones (1990-BTEX); <u>Surface Enhanced Raman Spectroscopy</u> : Carrabba et al. (1988, 1991), Ferrell et al. (1988), Smith et al. (1988)
Wet Chemistry Analytical Techn	iques/Instrumentation
Immunochemical Methods	<u>Reports/Symposia</u> : Schnell and Chang (1990), Silverstein et al. (1992a, 1992b), Van Emmon and Mumma (1990), Vanderlaan et al. (1991); <u>Field Enzyme</u> <u>Immunoassay Test Kits</u> : Bushway et al. (1988), Chamerlik-Cooper et al. (1991- PCBs), Duquette et al. (1988, 1991-PCP), Ensys Inc. (1991), Harrison and Ferguson (1990), Ladouceur (1991), McMahon et al. (1988), Schmidt et al. (1988), Vanderlaan et al. (1988), Van Emon et al. (1991a-pesticies, 1991b-PCP kits); <u>Immunochemical Fiber Optic Sensors</u> : Bolts et al. (1988), Lin et al. (1988)

Table 10-5 (cont.)

Торіс	References			
Wet Chemistry Analytical Techniqu	Wet Chemistry Analytical Techniques/Instrumentation (cont.)			
Colorimetric Chemical Field Test Kits	<u>PCB</u> : Gabry (1987), Woolerton et al. (1988); <u>Other</u> : Hanby (1988-aromatic compounds), Jenkins et al. (1991-explosives), Lindsay and Baedecker (1988-aqueous sulfide), Schlesing et al. (1991-chlorinated organics), Silvestri et al. (1981), Stamnes et al. (1991-Cr Hach kit)			
Liquid Chromatography	<u>Thin-Layer Chromatography</u> : Brumley and Brownrigg (1991-PNAs), Silvestri et al. (1981); <u>High Performance Liquid Chromatography</u> : Betowski and Jones (1989), Ekambaram and Burch (1988-PAHs), Joseph (1992), Mann and Vickers (1988), Pace et al. (1992), Riddell et al. (1991-PAHs)			
Bioassessment Techniques				
Bioassays	Brown et al. (1984), Easterly et al. (1988), Felkner et al. (1988a,b-laser/microbe bioassay); <u>Microtox Assay</u> : Abbott and Sims (1989-PAHs), Bulich (1979), Matthews and Bulich (1984), Symons and Sims (1988)			
Bioassessments/Monitoring	Bohman et al. (1989), Charters (1988), Dermer et al. (1980-biochemical indicators), DiBona et al. (1989), Gardner et al. (1989), Gezo and Brusick (1987), Piekarz (1990), Steen (1987-toxicity testing), Warren-Hicks et al. (1989-field and laboratory methods)			
Other Chemical Sensors				
General	Adrian (1992), Edmonds (1981), Hollenberg and Sahn (1988-biosensors), Janata and Bezegh (1988), Wohltjen (1984)			
Electrochemical Sensors	Carrabba et al. (1991-SEFOS), Penrose et al. (1991-pyrolysis-EC), Schmidt et al. (1988)			
Piezoelectric Sensors	Alder and McCallum (1983), Guilbault and Jordan (1988), Hlalvay and Guilbault (1977), Mierzwinski and Witkiewicz (1989), Overton et al. (1988d)			
Surface Acoustic Wave (SAW)	Ballentine and Wohltjen (1989), Ballentine et al. (1986), Bartley (1991), Elton and Houle (1991-SAW/GC), Jarvis et al. (1991), Nieuwenhuizen and Barendsz (1987)			
Semiconductor	Penrose et al. (1991)			

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APPENDIX C

GUIDE TO MAJOR REFERENCES OF SUBSURFACE CHARACTERIZATION, MONITORING, AND ANALYTICAL METHODS

A very large technical literature has developed in the last 20 years on characterization and monitoring of contaminated sites. This appendix provides information on major documents published by EPA, other government organizations, universities, and commercial organizations, which provide information of one or more aspects of vadose zone and ground-water characterization and monitoring. Most of these documents relate wholly, or in part, to contaminated sites. Other documents that do not have this perspective are included only if they focus primarily on field methods that can be applicable to contaminated sites.

Table C-1 provides brief descriptive information on over 80 major references. These are categorized into the following groups in the table: (1) Soils and ground water, (2) vadose zone, (3) ground water, (4) soils and solid wastes, and (5) symposia proceedings. EPA publications that are available at no cost from the Center for Environmental Research Information in Cincinnati are indicated with an asterisk in the reference list at the end of this appendix. Wherever possible, the NTIS number of government publications available from the National Technical Information Service (NTIS) is provided. (The NTIS telephone number is 800-553-6847). Publications by the Electric Power Research Institute are available at no cost to government agencies (EPRI Research Reports Center, P.O. Box 50490, Palo Alto, CA 94303, telephone 415-965-4081).

There is a very large literature on subsurface site characterization and monitoring techniques scattered through various annual and intermittent conference series. The published proceedings of four regular conference series serve as an excellent source of information on the latest developments in field characterization and monitoring: (1) The annual National Outdoor Action Conference on Aquifer Restoration, Ground Water Monitoring and Geophysical Methods, (2) the annual Conference on Petroleum Hydrocarbons and Organic Chemicals in Ground Water: Prevention, Detection and Remediation, (3) the annual Conference on Hazardous Materials Control (formerly called Superfund), and (4) the annual Conference on Hazardous Wastes and Hazardous Materials. Proceedings of the first two series are published by the National Water Well Association (NWWA), which changed its name to the National Ground Water Association (NGWA) in 1991 (NGWA Bookstore, P.O. Box 182039, Columbus, OH 43218, telephone 614-761-1711), and the proceedings of the latter two series are published by the Hazardous Dept., 9300 Columbia Rd., Silver Spring, MD 20910-1702, telephone 301-587-9390).

In addition, the Association of Ground Water Scientists and Engineers of NWWA/NGWA has sponsored numerous conferences focussed on special topics or regional issues. Since 1990, NWWA/NGWA conferences have been published in a subscription series titled Ground Water Management in which six coupons are issued that can be redeemed for the publications in the series of interest to the subscription holder (\$140 members/\$192.50 nonmembers, see the NGWA address above).

Table C-2 lists the titles of more than 70 published conference/series proceedings focusing on ground water and/or contaminated sites. Many relevant papers in these proceedings are cited in earlier sections of this guide. EPA regional offices and laboratories have many of these documents, and EPA/NTIS numbers are indicated, where available. If a document of interest cannot be found in a nearby library, the indicated sponsor (NWWA/NGWA or HMCRI) should be contacted concerning its availability. Out-of-print NWWA/NGWA publications can be borrowed for a fee from the National Ground Water Information Center (6365 Riverside Drive, Dublin, OH 43017, telephone 614-761-1711).

Table C-3 provides information on major compilations of information on analytical procedures for constituents of geochemical interest at contaminated sites. Most of these books and reports focus on laboratory methods and procedures, but might be useful for additional information on basic analytical methods that can be used in mobile laboratories or adapted for more portable instrumentation.

References Topic Site Investigations Hydrologic Characterization Brakensiek et al. (1979), Brown et al. (1983), Bureau of Reclamation (1981), Dames & Moore (1974), Driscoll (1986), Nielsen and Johnson (1990), Nielsen and Sara (1992), Rehm et al. (1985), Thompson et al. (1989), UNESCO (1983), U.S. EPA (1991a,b), USGS (1977+), Waste Management of North America (1991), Zimmie and Riggs (1981); see also, Tables 4-5 and 4-6 Warren-Hicks et al. (1989); see also, Table 10-5 (bioassessment techniques) Ecological Assessment Specific Settings Surface Mining: Barrett et al. (1980); Hazardous Waste Sites: Cameron (1991), Cochran and Hodge (1985), Ford and Turina (1985), Lesage and Jackson (1992), Oudjik and Mujica (1989), Perket (1986), Sisk (1981), U.S. EPA (1987, 1989a); RCRA Facilities: U.S. EPA (1986d, 1989b,c); Low Level Radioactive Wastes: EG&G (1990); Remedial Operations: Ross and Keeley (1992), U.S. EPA (1988a, 1991a); Surface Impoundments: Silka and Swearingen (1978) Ground-Water Monitoring Collins and Johnson (1988), Crouch et al. (1976), Devinny et al. (1990), EG&G General Procedures (1990), Everett (1980), Everett et al. (1976), Fried (1975), Gillham et al. (1983), Keith (1992), Loftis and Ward (1979), Mooij and Rovers (1976), Morrison (1983), Nielsen (1991), Nielsen and Johnson (1990), Nielsen and Sara (1992), Ross and Keeley (1992), Todd et al. (1976), U.S. DOE (Various dates), U.S. EPA (1986b, 1990a,b, 1991a,b, 1993), van Duijvenbooden and van Waegeningh (1987) Aller et al. (1991), Barcelona et al. (1983), Driscoll (1986), Howsam (1990), Korte Monitoring Wells and Kearl (1985), Nielsen and Schalla (1991); see also, Tables 2-4 and B-2 See Tables 5-4 and 5-5 Sampling Equipment Sampling Procedures API (1987), Barcelona et al. (1983, 1985), Berg (1982), Classen (1982), Holden (1984), Keith (1988), Korte and Kearl (1985), Nash and Leslie (1991), Rainwater and Thatcher (1960), Scalf et al. (1981), Summers and Gherini (1987), Unwin (1982), Wood (1976); see also, Table B-4. Crouch et al. (1976), Everett et al. (1976), Loftis and Ward (1979) Costs Tinlin (1976); Solid Waste Disposal: Fenn et al. (1977), U.S. EPA (1981a,b, 1986f); Specific Settings RCRA Facilities: U.S. EPA (1983a,b, 1985, 1986a,d,e,f, 1989c); Enhanced Oil Recovery: Beck et al. (1981); Surface Mining: Everett (1979, 1983, 1985), Everett and Hoylman (1980a,b), Williams and Schuman (1987); Oil Shale: Everett (1985), Slawson (1979, 1980a,b); Electric Utilities: GeoTrans (1989), Redwine et al. (1985); Wastewater and Sludge Application: Ho et al. (1978); Waste Spills: Pilie et al. (1975), Yang and Bye (1979); Geothermal: Weiss et al. (1979) State/Local Guidance Documents* Connecticut Environmental Protection Agency (1983), Lindorff et al. (1987), NJDEP (1988), Santa Clara County Water District (1985), Stephens (1986)

Table C-1 Major Reference Sources on Subsurface Characterization and Monitoring Methods

Table C-1 (cont.)

Торіс	References
Microbiological Sampling	Bitton and Gerba (1984), Board and Lovelock (1973), Bordner et al. (1978), Britton and Greeson (1989), Costerton and Colwell (1979), Dunlap et al. (1977), Rosswall (1973), USGS (1977+); see also, references in Section 9.3.6
Vadose Zone Monitoring	
General	Everett et al. (1983), Nielsen and Johnson (1990), Nielsen and Sara (1992), Rehm et al. (1985), Rijtema and Wassink (1969), U.S. EPA (1986c), Wilson (1980)
Soil Solute	Devinny et al. (1990), Morrison (1983), Nash and Leslie (1991), Nielsen (1991), USGS (1977+); see also, Table 9-4
Soil Gas Soils	Devitt et al. (1987), Ford et al. (1984), Kerfoot and Barrows (1987), U.S. EPA (1988b); see also, references for Section 9.4.2
Field Characterization	Blume et al. (1991), Boulding (1991), Brakensiek et al. (1979), Bureau of Reclamation (1974, 1990), Cameron (1991), Hodgson (1978), SCS (1971)
General Sampling	Acker (1974), Barth et al. (1989), Cameron (1966), Corps of Engineers (1972), Keith (1992), Hodgson (1978), Hvorslev (1948, 1949), Mason (1992), McKeague (1978), Mooij and Rovers (1976), SCS (1984), U.S. DOE (Various dates); <u>Sediments</u> : Barth and Starks (1985), Edwards and Glysson (1988), Palmer (1985), Plumb (1981); see also, Table 2-5
Sampling for Soil Contaminants	API (1987, 1992), Boulding (1991), Brown et al. (1991), EG&G (1990), Ford et al. (1984), Goodwin et al. (1982), Keith (1988), Scalf et al. (1981), Schweitzer and Santolucito (1984), U.S. EPA (1986b, 1988b, 1989c, 1991a), van Duijvenbooden and van Waegeningh (1987).
Wastes	
Sampling	deVera (1980), Ford et al. (1984), Keith (1988, 1992), Simmons (1991), U.S. DOE (Various dates), U.S. EPA (1986b), Wolbach et al. (1984)
Agency/Organization Index	
U.S. EPA	Soils and Ground Water: Alier et al. (1991), Cochran and Hodge (1985), Dunlap et al. (1977), Everett et al. (1976), Fenn et al. (1977), Ford and Turina (1985), Ford et al. (1984), Ross and Keeley (1992), Scalf et al. (1981), Silka and Swearingen (1978), Sisk (1981), U.S. EPA (1986d, 1987, 1989a,b,c, 1990b, 1991a); <u>Vadose Zone</u> : Devitt et al. (1987), Everett et al. (1983), Kerfoot and Barrows (1987), U.S. EPA (1986c), Wilson (1980); <u>General Ground Water</u> : Barcelona et al. (1985), Berg (1982), Crouch et al. (1976), Loftis and Ward (1979), Tinlin (1976), Todd et al. (1976), U.S. EPA (1990a, 1991a), Yang and Bye (1979); <u>Ground-Water Guidance Documents</u> : U.S. EPA (1981a, 1981b, 1983a,b, 1985, 1986a,e,f, 1988a, 1993); <u>Soil and Solid/Liquid</u> <u>Waste</u> : Barth et al. (1989), Boulding (1991), Cameron (1991), deVera (1980), Hatayama et al. (1980), Mason (1992), Pilie et al. (1975), U.S. EPA (1986b), Yang and Bye (1979); <u>Energy Development Ground-Water Monitoring</u> : Beck et al. (1981), Everett (1979, 1983), Everett and Hoylman (1980a,b); Slawson (1979, 1980a,b), Weiss et al. (1979)

Table C-1 (cont.)

Торіс	References
Other Federal	Bureau of Reclamation: Bureau of Reclamation (1974, 1981, 1990); Department of Energy: EG&G (1990), U.S. DOE (Various dates); Fish and Wildlife Service: Brown et al. (1991); Forest Service: Barrett et al. (1980); NASA: Cameron et al. (1966); USATHAMA/Corps of Engineers: Corps of Engineers (1972), Goodwin et al. (1982), Hvorslev (1949), Plumb (1981); USDA/SCS: Brakensiek et al. (1979), SCS (1971, 1984); U.S. Geological Survey: Classen (1982), Edwards and Glysson (1988), Guy (1969), USGS (1977+), Wood (1976)
Other Government	<u>Canada</u> : McKeague (1978), Mooij and Rovers (1976); <u>States</u> ⁺ : Barcelona et al. (1983), Connecticut Environmental Protection Agency (1983), Lindorff et al. (1987), NJDEP (1988), Stephens (1986)
American Chemical Society (ACS)	Keith (1988, 1992), Nash and Leslie (1991), Schweitzer and Santolucito (1984)
Amercian Petroleum Institute (API)	API (1987, 1992), Gillham et al. (1983)
American Society for Testing and Materials (ASTM)	ASTM (Annual, 1992a,b); <u>Ground-Water and Vadose Zone STPs</u> : Collins and Johnson (1988), Nielsen and Johnson (1990), Nielsen and Sara (1992), Zimmie and Riggs (1980); <u>Hazardous Waste Solid Testing Conference Series</u> : (Papers in this series tend to focus on laboratory methods, but also include papers on field-oriented techniques): 1st (Conway and Mallow, 1981); 2nd (Conway and Gulledge, 1982); 3rd (Jackson et al., 1984); 4th (Petros et al., 1985); 5th (Perket, 1986); 6th (Lorenzen et al., 1986)
Consulting Firms	Dames & Moore (1974), Everett (1980), GeoTrans (1989), Waste Management of North America (1991)
Electric Power Research Institute (EPRI)	Redwine et al. (1985), Rehm et al. (1985), Summers and Gherini (1987), Thompson et al. (1989)
UNESCO	Brown et al. (1983); Symposia: Rijtema and Wassink (1969), UNESCO (1983)
Other	Devinny et al. (1990), Driscoll (1986), Everett (1985), Fried (1975), Ho et al. (1978), Holden (1984), Howsam (1990), Klute (1986), Lesage and Jackson (1992), Morrison (1983), Nielsen (1991), Oudjik and Mujica (1989), Simmons (1991), Unwin (1982), van Duijvenbooden and van Waegeningh (1987)

*The appropriate state regulatory agency should be contacted for the most current version of any guidance documents.

Sponsor	Year	Title
EPA/NWWA	1971	1st National Ground Water Quality Symposium (EPA-16060 GRB, NTIS PB214-614)
	1974	2nd (EPA-68-03-0367, NTIS PB257-312)
	1977	3rd (EPA/600/9-77/014, NTIS PB272-908)
	1979	4th (EPA/600/9-79/029, NTIS PB80-103476)
	1980	5th
	1983	6th (State, County, Regional, and Municipal Jurisdictions of Ground Water Protection)
	1984 1986	7th (Innovative Means of Dealing with Potential Sources of Ground Water Contamination) 8th (Anatomy of Superfund)
NWWA	1981	1st National Ground Water Quality Monitoring Symposium and Exposition
	1982	2nd National Symposium on Aquifer Restoration and Ground Water Monitoring
	1983	3rd
	1984	41h
	1985	5th
	1986	6th
	1987	1st National Outdoor Action Conference on Aquifer Restoration, Ground Water Monitoring, and Geophysical Methods
	1988	2nd
	1989	3rd
	1990	4th GWM 2
	1991	5th GWM 5
NGWA	1992	6th GWM 11
NWWA/API	1984	[1st] Conference on Petroleum Hydrocarbons and Organic Chemicals in Ground Water-Prevention, Detection, and Restoration
	1985	[2nd]
	1986	[3rd]
	1987	[4th]
	1988	[5th]
	1989	[6th]
	1990	[7th] GWM 4
	1991	[8th] GWM 8
NGWA/API	1992	[9th] GWM 14
Modeling		
NWWA/IGWMC	1984	1st Conference on Practical Applications of Ground Water Models
	1985	2nd
	1980	3rd Conterence on Solving Ground Water Problems with Models
	1988	Conterence on Geochemical Modeling of Ground Water Contamination
NGWA/IGWMC	1989	5th GWM 9
Geophysics		
NWWA/EPA	1984 1985 1986	[1st] Conference on Surface and Borehole Geophysical Methods in Ground Water Investigations [2nd] Surface and Borehole Geophysical Methods and Ground Water Instrumentation Conference and Exposition

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Table C-2 Conferences and Symposia with Papers Relevant to Subsurface Characterization and Monitoring

Table C-2 (Cont.)

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Sponsor	Year	Title
Vadose Zone		
NWWA/EPA	1983 1985 1986	[1st] Conference on Characterization and Monitoring in the Vadose (Unsaturated) Zone [2nd] 3rd
Karst		
NWWA	1986 [°] 1988 1991	[1st] Conference on Environmental Problems in Karst Terranes and Their Solutions 2nd 3rd Conference on Hydrogeology, Ecology, Monitoring and Management of Ground Water in Karst Terranes GWM 10
Misœllancous N	WWA Cor	iferences
NWWA/AGWSI	E 1988 1989	Ground Water Geochemistry Conference Conference on New Field Techniques for Quantifying Physical and Chemical Properties of Heterogeneous Aquifers
·	1990	Cluster of Conferences (Agricultural Impacts on Ground Water Quality; Ground Water Geochemistry Ground Water Management and Wellhead Protection; Environmental Site Assessments: Case Studies and Strategies) GWM 1
NGWA/AGWSE	1991 2 1992	Environmental Site Assessments Case Studies and Strategies: The Conference GWM 6 [2nd] Environmental Site Assessments Case Studies and Strategies: The Conference GWM 12
NWWA Eastern	Regional	Conferences
NWWA/AGWSI	2 1984 1985 1986 1987 1988 1989 1990 1991	 [1st] Eastern Regional Ground Water Conference [2nd] 3rd Annual Eastern Regional Ground Water Conference 4th [5th] Focus Conference on Eastern Regional Ground Water Issues [6th] [7th] GWM 3 [8th] GWM 7
NGWA/AGWSE	1992	[9th] GWM 13
Other NWWA R	legional Co	onferences
NWWA	1983	Eastern Regional Conference on Ground Water Management Western Regional Conference on Ground Water Management
	1984 1985	Conference on Ground Water Management Southern Regional Ground Water Conference Western Regional Ground Water Conference
	1986	Conference on Southwestern Ground Water Issues Focus Conference on Southeastern Ground Water Issues
	1987	Focus Conference on Midwestern Ground Water Issues Focus Conference on Northwestern Ground Water Issues [2nd] Focus Conference on Southwestern Ground Water Issues
	1900	Land Long conference on Southwestern Chound Water Issues

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Table C-2 (Cont.)

Sponsor Year Title

HMCRI

Hazardous Materials Control Research Institute Conferences

1980 1st National Conference on Management of Uncontrolled Hazardous Wastes Sites 1981 2nd 1982 3rd 4th 1983 1984 5th 1985 бth 1986 7th 1987 8th Superfund '87 1988 9th Superfund '88 10th Superfund '89 1989 1990 11th Superfund '90 1991 12th Hazardous Materials Control (HMC-Superfund '91) 1992 13th HMC-Superfund '92

Regional Hazardous Materials Control Conferences

1990	HMC—Great Lakes '90
1991	HMC-Northeast '91
1992	HMC—South '92
1984	1st National Conference on Hazardous Wastes and Environmental Emergencies
1985	2nd
1986	3rd National Conference on Hazardous Wastes and Hazardous Materials
1987	4th
1988	5th
1989	6th (HWHM '89)
1990	7th (HWHM '90)
	1990 1991 1992 1984 1985 1986 1987 1988 1989 1990

Miscellaneous Conferences

HMCRI	1992	National R&D Conference on the Control of Hazardous Materials
	1992	Federal Environmental Restoration '92

[]--Indicates that number is not included in the title of the published proceedings. GRM indicates that proceedings have been published in NWWA/NGWA's Ground Water Management Series.

Abbreviations:

AGWSE	Association of Ground Water Scientists and Engineers (NWWA/NGWA)
API	American Petroleum Institute
EPA	U.S. Environmental Protection Agency
HEW	U.S. Department of Health, Education and Welfare
HMCRI	Hazardous Materials Control Research Institute
IGWMC	International Ground Water Modeling Center
NWWA/NGWA	National Water Well Association (named changed to National Ground Water Association in 1992)

Table C-3 Major Compilations of Analytical Procedures for Constituents of Geochemical Interest

Reference	Description
Instrumentation Principles	Skoog (1985), Willard et al. (1988)
U.S. EPA Analytical Methods	Overviews: Mueller and Smith (1991), Wagner (1992); <u>General</u> : Kopp and McKee (1983); <u>Metals</u> : U.S. EPA (1991c); <u>Organics in Water</u> : Longbottom and Lichtenberg (1982); <u>Solid Waste (SW-846)</u> : U.S. EPA (1986b); <u>SW-846 Methods Studies</u> : Edgill (1989), Edgill and Wilburs (1989), Engel et al. (1988); <u>Drinking Water Analysis</u> : Long and Martin (1989), Pfaff (1981), U.S. EPA (1990c,d); <u>Pesticides</u> : Watts (1980); <u>Sediment</u> : Guy (1969), Plumb (1981), U.S. EPA (1989d); <u>Quality Control</u> : Booth (1979), Sharma (1979)
Other U.S. EPA Methods	Field Screening Methods: U.S. EPA (1988b); Ecological Assessment: Warren-Hicks et al. (1989)
U.S. Geological Survey TWRIs	The Techniques of Water Resource Investigation series includes manuals describing procedures for planning and conducting specialized work in water-resources investigations. Wood (1976) covers field analysis of unstable constituents; Fishman and Friedman (1989—supersedes Brown et al., 1970, Skougstad et al., 1979, and Fishman and Bradford, 1982) cover methods for analyzing inorganic constituents in water and fluvial sediment; Barnett and Mallory (1971) describe determination of minor elements in water by emission spectroscopy; Wershaw et al. (1987—supersedes Goerlizt and Brown, 1972) cover methods for determination of organic substances in water and fluvial sediments; Thatcher et al. (1977) cover methods for determination of radioactive substances in water and fluvial sediments; Brinton and Greeson (1989—supersedes Greeson et al., 1977) cover methods for collection and analysis of aquatic biological and microbiological samples; Friedman and Erdman (1982) cover quality assurance practices for the chemical and biological analyses of water and fluvial sediments; Guy (1969) covers laboratory methods for sediment analysis.
Other References	
Water Analysis	Standard Methods: APHA (1992), ASTM (Annual-Vols. ll.01 and 11.02), Hach (1991); Other Major References: Fresenius et al. (1988), Rainwater and Thatcher (1960), Thompson et al. (1989)
Soil Analysis	<u>Physical Properties</u> : ASTM (Annual-Vol. 4.08), Blume et al. (1991), Guy (1969- sediments), Klute (1986), SCS (1984), Smith and Mullins (1991), Topp et al. (1992); <u>Soil Chemistry</u> : Council on Soil Testing and Plant Analysis (1992), McKeague (1978), Westerman (1990)
Contaminants	Method Compilations: Plumb (1984), U.S. DOE (Various dates); Ground water: API (1987-petroleum hydrocarbons), Hach (1991—inorganics), Lesage and Jackson (1992); Soil: API (1987, 1992-petroleum hydrocarbons)
Solid Waste	<u>Hazardous Waste</u> : Silvestri et al. (1981), Wolbach et al. (1984); <u>Flue Gas</u> <u>Desulfurization Waste</u> : Noblett and Burke (1990), Radian Corporation (1988); <u>Oil</u> <u>Shale</u> : Wallace et al. (1984); <u>Mine Soils and Overburden</u> : Williams and Schuman (1987)
Some Suggestions Concerning Evaluation of Literature References

New developments in field techniques for ground-water and contaminated site investigations are changing so rapidly that care is required when evaluating the literature, especially when dealing with a method that is outside one's area of special expertise. Several factors affect the weight that should be given to conclusions or recommendations concerning a particular method: (1) Whether the information is from a peerreviewed or non-peer reviewed source; (2) where the authors come from; and (3) how recently it has been published.

Greatest weight should be given to the content of papers published in peer-reviewed scientific journals such as Ground Water and Ground Water Monitoring Review (retitled Ground Water Monitoring and Remediation beginning in 1993). Most conference proceedings (ASTM conferences being an exception) are not peer-reviewed, and consequently there is more likely to be diversity of opinion concerning conclusions or recommendations in individual papers. When non-peer-reviewed papers are considered, greater weight can be given to those written by individuals from academic institutions or research-oriented government agencies (U.S. Geological Survey, personnel from EPA research laboratories) vis-a-vis papers written by manufacturers or consultants who might have an interest in promoting a particular method. On the other hand, papers written by individuals from academic institutions and more research-oriented government agencies, such as the U.S. Geological Survey, that describe new techniques are more likely to be in developmental stages and not readily available for routine field use. Finally, more recently published papers can generally be given greater weight than earlier publications because they are more likely to address recent developments and advances in investigation techniques. As a general rule, review of multiple references from a variety of sources that deal with a specific method should help determine its appropriateness for a specific application or for specific site conditions. When in doubt, one or more experts should be consulted.

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APPENDIX D

GUIDE TO VADOSE ZONE MODELS FOR WATER BUDGET AND EXPOSURE MODELING

Preliminary site reconnaissance should provide sufficient information to develop a preliminary conceptual model (or models) of the site that has three main elements (see Figure D-1): (1) Characteristics of the waste source, (2) known or potential pathways for migration, and (3) receptors of exposure to the contaminant. Mathematical models or computer codes often are used to evaluate potential for transport of contaminants and to estimate exposure. The conceptual model, and computer modeling efforts are an iterative process in which: (1) The current model helps define the parameters that should be sampled or otherwise characterized; (2) data are collected; (3) the model is confirmed or modified to reflect the new information; (4) additional data are collected if necessary.

Depending on local conditions, the soil (weathered zone) can be separated from the water table (saturated zone) by an intervening zone, called the vadose zone where water flow occurs predominantly under unsaturated conditions. Modeling of contaminant transport in the vadose zone tends to be more complicated than in the saturated zone because variations on moisture content and gaseous phase transport also must be considered.

Computer codes that model the soil and vadose zone (also called variably saturated or saturated/unsaturated flow models) fall into three main groups: (1) Flow models, which deal only with the flow of water in variably saturated conditions; (2) transport models, which deal with the movement of contaminants or other chemicals under saturated/unsaturated conditions; and (3) geochemical models (also called distribution-of-species codes), which deal with reactions in the aqueous phase of the system.

This appendix focuses on vadose zone models that focus on the soil rooting zone, because they can be used for both water budget analysis, which is useful for characterization of the hydrologic system at the site, and evaluation of leaching potential to ground water, which is important for exposure assessment. U.S. EPA (1988b) provides guidance on the selection of vadose zone models for exposure assessment.

Table D-1 summarizes input data requirements for 10 near-surface vadose zone models, and Table D-2 indexes references where additional information can be found about these 10 models^1 . When a model has been selected, this table can be used to identify the soil parameters that must be estimated or measured during field investigation activities. The most important soil physical and hydrologic parameters, based on the frequency with which they are required in these 10 models, are: (1) Saturated hydraulic conductivity and saturated moisture content (90%), (2) soil bulk density and precipitation (80%), (3) unsaturated hydraulic conductivity, water retention, and soil porosity (70%); (4) soil texture and soil temperature (60%); (5) horizons or soil layering (50%); (6) air permeability and depth to ground water (20%). The other chemistry-related parameters in this table generally are obtained by laboratory analyses of soil or ground-water samples.

Numerous other computer codes have been developed for modeling flow and chemical transport in soil and the vadose zone. U.S. EPA (1988a) contains one or two page descriptions of 14 computer codes that can be used for modeling below the rooting zone (none of which are included in Table D-1). Information on 28 variably saturated flow models, and 26 variably saturated flow/solute transport models, are summarized in van der Heijde et al. (1988). Donigian and Rao (1986b) compare the capabilities of four of the models in Table D-1 (HELP, SESOIL, CREAMS, and PRZM), and six other models that deal with soil leaching. Sposito (1985) discusses a number of models that can be used for geochemical modeling of inorganic pollutants in soil system, without consideration of flow.

¹ Other models that might have value for application at contaminated sites for water budget analysis or contaminant transport assessment include: (1) CMLS (Chemical Movement in Layered Soils), (2) GLEAMS (Groundwater Loading Effects of Agricultural Management Systems), (3) HSWDS (Hydrologic Simulation of Waste Disposal Sites), (4) PESTAN (Pesticide Analytical Model), (5) RUSTIC (Risk of Unsaturated/Saturated Transport and Transport and Transformation of Chemical Concentrations), and (6) VLEACH (a one-dimensional finite difference vadose zone leaching model). The RETC code can be used to estimate unsaturated hydraulic conductivity as a function of moisture content based on other soil parameters, such as texture.



Figure D-1 Elements of a conceptual model for contaminated site characterization (Barth et al., 1989).

D-2

	Model Name [Reference(s)]										
Properties and Parameters	Help (A,B)	Sesoil (C,D)	Creams (E,F)	PRZM (G,H,I)	Vadoft (H,J)	Minteq (J)	Fowl™ (K)	Ritz (L)	Vip (M)	Chemflo (N)	
Soil bulk density	0	٠		•	٠	0	٠	•	۲	•	
Soil pH	0	•	0	0	0	٠	•	0	0	0	
Soil texture		0	•	•	•	0	0	٠	٠	0	
Depth to ground water	0	•	9	0	•	0	0	6	6	0	
Horizons (soil layering)	•	•	•	•	•	0	Ø	Ø	0	0	
Saturated hydraulic conducitivity	•	•	•	•	•	0	•	٠	٠	•	
Water retention	•	٠	•	•	•	0	•	0	0	•	
Air permeability	0	٠	0	0	٢	0	0	0	۲	0	
Climate (precipitation)	•	۲	•	•	0	0	•	•	٠	•	
Soil porosity	٠	•	٠	•	•	0	0	•	٠	0	
Soil organic content	0	٠	٠	•	•	٠	0	•	٠	6	
Cation Exchange Capacity (CEC)	0	٠	0	0	0	•	0	0	0	0	
Degradation parameters	•	•	٠	•	•	0	0	•	٠	•	
Soil grain size distribution	0	۲	Ø	0	۲	0	0	0	0	0	
Soil redox potential	0	0	0	0	0	•	0	0	0	(6)	
Soil/water partition coefficients	0	-	•	•	•	•	•	•	•	•	
Soil oxygen content	0	0	0	0	0	0	0	0	٠	0	
Soil temperature	0	•	۲	•	٠	٠	6	• •	٠	0	
Soil mineralogy	0	•	0	0	0	6	0	0	0	0	
Insaturated hydraulic conductivity	•	•	•	•	•	0	•	0	0	•	
Saturated soil moisture content	•	۲	•	•	•	0	•	•	٠	•	
Aicroorganism population	0	0	0	0	0	0	0	0	0	0	
Soil respiration	0	0	0	0	0	0	0	0	0	0	
Evaporation	•	•	•	•	6	0	0	•	•	•	
lir/water contaminant densities	6	0	0	0	0	0	•	•	•	0	
kir/water contaminant viscosities	0	0	0	0	0	0	0	0	۲	0	

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 A. Schroeder, et al., 1984.
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 K. Hostetler, Erickson, and Rai, 1988.
 Bequired ONot required Subsective of the models, the models, but not directly entered as input to models.

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 Bequired ONot required Subsective of the models, the models, but not directly entered as input to models.

 B. Schroeder, et al., 1984.
 I. Dean et al., 1989.
 M. Stevens et al., 1989.
 Used in the estimation of other required characteristics or the integretation of the models, but not directly entered as input to models.

 D. Chen, Wolfman, and Liu, 1987.
 J. Brown and Allison, 1987.
 N. Nofziger et al., 1989.
 N. Nofziger et al., 1989.

Source: Breckenridge et al. (1991)

Торіс	References	
General		
Vadose Zone Modeling	<u>Overviews</u> : Breckenridge et al. (1991), Donigan and Rao (1986a), El-Kadi an Beljin (1987), Hern and Melancon (1986a), Iskander (1981), Mangold and T (1987), Nielsen et al. (1990), Oster (1982), Thompson et al. (1989), U.S. EPA (1984, 1988a,b), van Genuchten (1987), Weaver et al. (1989), Whelan and Bu (1988); <u>Conference/Symposia</u> : Arnold et al. (1982), Evans and Nicholson (199 fractured rock), Wierenga and Bachelet (1988)	
Vadose Model Comparisons	Addiscott and Wagenet (1985), Donigian and Rao (1986b), Kincaid and Morrey (1984), Kincaid et al. (1984), Oster (1982), van der Heijde et al. (1988)	
Geochemical Modeling	Apps (1988), Jenne (1979), Jacobs and Whatley (1985), Jenne (1981), Melchior and Bassett (1990), NWWA (1988, 1990), U.S. EPA (1990a,b), Yeh and Tripathi (1989); <u>Model Comparisons</u> : Mangold and Tsang (1987), Nordstrom and Ball (1984), Nordstrom et al. (1979), Schechter et al. (1985), Sposito (1985)	
Field Testing	Hern and Melancon (1986b), Hern et al. (1986), van der Heijde et al. (1989), Weaver et al. (1989)	
Flow and Transport Models		
CHEMFLO*	Nofziger et al. (1989)	
CMLS	Ehteshami et al. (1991), Nofziger and Hornsby (1986, 1987)	
CREAMS	Barnes and Rodgers (1988), Beasley et al. (1991), Devaurs and Springer (1988), Donigian and Rao (1986b), Knisel (1980), Knisel and Leonard (1990), Knisel et al. (1985), Leonard and Ferreira (1984), van der Heijde et al. (1988)	
FOWL	Hostetler et al. (1988)	
GLEAMS	Beasley et al. (1991), Davis et al. (1990), Knisel and Leonard (1990), Knisel et al. (1991), Leonard et al. (1987)	
HELP	Barnes and Rodgers (1988), Donigian and Rao (1986b), Dwyer et al. (1988), Schroeder et al. (1984a,b), Schroeder and Peyton (1987a,b)	
HSWDS	Perrier and Gibson (1982)	
MINTEQ	Brown and Allison (1987), Loux et al. (1989, 1990), Morrey et al. (1986)	
PESTAN*	Hern and Melancon (1986b), Melancon et al. (1986), Ravi and Johnson (1992)	
PRZM	Banton and Villeneuve (1989), Carsel et al. (1984, 1985, 1988), Donigian and Rao (1986b), Heddon (1986), Hern and Melancon (1986b), Kincaid and Morrey (1984), Melancon et al. (1986), van der Heijde et al. (1988), Varshney et al. (1993), Whelan and Brown (1988)	

Table D-2 Reference Index for Near-Surface Vadose Zone and Geochemical Models

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Table D-2 (cont.)

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Topic	References
RETC*	van Genuchten et al. (1991)
RITZ*	Jury et al. (1983), Nofziger and Williams (1988), Sims et al. (1991), U.S. EPA (1986), van der Heijde et al. (1988)
RUSTIC (PRZM + VADOFT)	Dean et al. (1989), Varshney et al. (1993)
SESOIL	Bonazountas and Wagner (1984), Chen et al. (1987), Donigian and Rao (1986b), Hern and Melancon (1986b), Kincaid et al. (1984), Melancon et al. (1986), van der Heijde et al. (1988)
VADOFT	Dean et al. (1989), Huyakorn et al. (1988), Varshney et al. (1993)
VIP*	Grenney et al. (1987), McClean et al. (1988), Sims et al. (1991), Stevens et al. (1988, 1989), Symons et al. (1988)
VLEACH*	CH2M Hill (1990)
Unnamed	Tim and Mostaghimi (1989), Wagenet and Hutson (1986)

*Available from EPA Center for Subsurface Modeling Support, P.O. Box 1198, Ada OK 74820; (505-332-8800).

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