
Research and Development



Characterization of Hazardous Waste Sites—A Methods Manual:

Volume II. Available Sampling Methods, Second Edition



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CHARACTERIZATION OF HAZARDOUS
WASTE SITES--A METHODS MANUAL
VOLUME II
AVAILABLE SAMPLING METHODS
Second Edition

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NOTICE

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FOREWORD

Available Sampling Methods is part of a multivolume manual, entitled Characterization of Hazardous Waste Sites--A Methods Manual, developed by the U.S. Environmental Protection Agency. The sampling methods-document is dedicated to sampling procedures and sampling information only, and is meant to be used in conjunction with two companion documents that address general site characterization approaches, and available laboratory analytical methods for sample analysis. The sampling volume describes a collection of methods and materials sufficient to address most sampling situations that arise during routine waste site and hazardous spill investigations. The methods are compiled with detailed, practical information to provide field investigators with a set of functional operating procedures.

The first companion volume, Integrated Approach to Hazardous Waste Site Characterization, includes discussions on preliminary assessment, initial data evaluation, administrative procedures, offsite reconnaissance, site inspection, chain of custody, quality assurance, safety and training in addition to considerations concerning sampling strategy and methods selection. The second companion document, Available Laboratory Analytical Methods, outlines detailed methodology suitable for hazardous waste analysis and is organized by media and compound.

ABSTRACT

Investigations at hazardous waste sites and sites of chemical spills often require onsite measurements and sampling activities to assess the type and extent of contamination. This document is a compilation of sampling methods and materials suitable to address most needs that arise during routine waste site and hazardous spill investigations.

The sampling methods presented in this document are compiled by media, and were selected on the basis of practicality, economics, representativeness, comparability with analytical considerations, and safety, as well as other criteria. In addition to sampling procedures, sample handling and shipping, chain-of-custody procedures, instrument certification, equipment fabrication, and equipment decontamination procedures are described.

Sampling methods for soil, sludges, sediments, and bulk materials cover the solids medium. Ten methods are detailed for surface waters, groundwater and containerized liquids; twelve are presented for ambient air, soil gases and vapors, and headspace gases. A brief discussion of ionizing radiation survey instruments is also provided.

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SECTION 1.0

INTRODUCTION

1.1 GENERAL

Investigations at hazardous waste and environment-threatening spill sites place more restrictive demands on personnel, materials and methodologies than those usually found in routine environmental surveys. As a result, traditional procedures and protocols used for the acquisition of environmental samples often fail to meet the rigors and demands required for many hazardous waste sampling applications. Thus, the collection of hazardous waste samples will frequently require specialized equipment and protocols either developed specifically for such uses or modified from preexisting materials and/or techniques. Some important considerations are:

- Versatility--Methods and materials must be suitable to a wide range of situations and applications because of the unknown nature of many hazardous waste investigations and environmental spill responses.
- Safety--Hazardous wastes, by definition, are associated with both acute and chronic exposure to dangerous, toxic chemicals and this dictates that protective sample collection methods be used to minimize personnel exposure. In addition, instrumentation and equipment must be safe for use in the atmospheres in which they are being operated.
- Decontamination--Because of the nature of the materials being sampled, the option of using disposable sampling equipment must be considered since attempting field cleanup efforts may be impractical.
- Ease of Operation--Hazardous waste site investigations and response actions at environment-threatening spills generally require some level of hazard protection that may be cumbersome, limit the field of vision, or fatigue the sampler. Sample collection procedures must therefore be relatively simple to follow, expedite sample procurement and to reduce the chance of fatigue. Collection and monitoring equipment should be simple to operate, direct reading, and should not be unwieldy.

These and other factors associated with the procurement of hazardous waste samples need to be addressed in a compilation of practical, cost effective, and reliable methods and procedures capable of yielding representative samples for a diverse number of potential parameters and chemical matrices. These methods must be consonant with a variety of analytical considerations running the gamut from gross compatibility analyses (pH, flammability, water reactivity, etc.) to highly sophisticated techniques capable of resolution in the part per billion (ppb) range.

1.2 METHOD SELECTION CRITERIA

Major emphasis must be placed on the selection of sampling methods. Even a limited literature survey will disclose the existence of a great number of sampling methods, all of which have certain merits that warrant consideration. Therefore, selection criteria were chosen on which to base decisions for including the sampling methods found in this manual. The following is a listing, not necessarily in order of relative importance, of these criteria.

Practicality

The selected methods should stress the use of simple, pragmatic, proven procedures capable of being used or easily adapted to a variety of situations.

Representativeness

The essence of any sampling campaign is to collect samples that are representative of the material or medium under consideration. The selected methods, although strongly taking into consideration economics, simplicity, practicality, and portability, must also be capable of delivering a best practicable representation of the situation under investigation.

Economics

The costs of equipment, manpower and operational maintenance need to be considered in relation to overall benefit. Instrument durability, disposable equipment, cost of decontamination, and degree of precision and accuracy actually required are also factors to be considered.

Simplicity or Ease of Operation

Because of the nature of the material to be sampled, the hazards encountered during sampling, and the cumbersome safety equipment sometimes required, the sampling procedures selected must be relatively easy to follow and equipment simple to operate. Equipment should be portable, lightweight, rugged and, if possible, direct reading.

Comparability with Analytical Considerations

The uncertainty of sample integrity as it relates to the analytical techniques to be used should be reduced whenever possible. Errors induced by poorly selected sampling techniques, especially those used in uncontrolled situations, can be the weakest link in the quality of the generated data. Special consideration must therefore be given to the selection of sampling methods in relation to any adverse effects that might surface during analysis. Proper materials of construction, sample or species loss, and chemical reactivity are some of the factors that must receive attention.

Versatility

The diversity and sheer numbers of potential parameters and scenarios often preclude the use of novel approaches that are designed or better suited for classifying a small number of compounds in a limited, defined environment. The methods in question must be adaptable to a variety of sampling situations and chemical matrices. This factor should not, however, jeopardize sample integrity.

Safety

The risk to sampling personnel, intrinsic safety of instrumentation, and safety equipment required for conducting the sampling all need to be evaluated in relation to the selection of proper methods and procedures.

The above criteria were consulted during the selection of each of the methods listed in the following sections. Obviously, tradeoffs were necessary, and therefore, some methods may prove excellent for some situations and less satisfactory for others. This factor must be considered by any field investigator before using the procedures outlined here.

1.3 PURPOSE AND OBJECTIVE OF SAMPLING

The basic objective of any sampling campaign is to collect a sample which is representative of the media under investigation. More specifically, the purpose of sampling at hazardous waste sites is to acquire information that will aid investigators in determining the presence and identity of onsite contaminants and the extent to which these compounds have become integrated into the surrounding environment. This information can then be used as support for future litigations or as input to remedial investigations and risk assessments.

The term "sample" has already been defined as a representative part of the media under investigation. Representativeness, however, is a relative term and must be carefully considered, along with several other criteria, prior to the acquisition of samples. A list of the criteria is as follows.

- Representativeness--This sample possesses the same qualities or properties as the material under consideration. The degree of resemblance of the sample to the material in question is determined by the desired qualities under investigation and analytical techniques used.
- Sample size--This should be chosen carefully in respect to physical properties of the entire object and the requirements and/or limitations of both sampling and analytical techniques.
- Number and/or the frequency of subsample--Decisions on these considerations are based on what types of statistical information are desired and the nature of the material collected.
- Maintenance of sample integrity--The sample must retain the properties of the original medium conditions (at the time of sampling) through collection, transport, and delivery to the analyst.

1.4 TYPES OF SAMPLES

Before defining the general sample types, the nature of the media or materials under investigation must be discussed. Materials can be divided into three basic groups as outlined in Figure 1-1.¹

Of least concern to the sampler are homogeneous materials. These materials are generally defined as having uniform composition throughout. In this case, any sample increment can be considered representative of the material. On the other hand, heterogeneous samples present problems to the sampler because of changes in the quality of the material over distance.

When discussing types of samples, it is important to distinguish between the type of media to be sampled and the sampling technique that yields a specific type of sample. In relation to the media to be sampled, two basic types of samples can be considered: the environmental sample and the hazardous sample.

Environmental samples (ambient air, soils, rivers, streams, or biota) are generally dilute (in terms of pollutant concentration) and usually do not require the special handling procedures used for concentrated wastes. However, in certain instances, environmental samples can contain elevated concentrations of pollutants and in such cases would have to be handled as hazardous samples.

Hazardous or concentrated samples are those collected from drums, tanks, lagoons, pits, waste piles, fresh spills, etc., and require special handling procedures because of their potential toxicity or hazard. These samples can be further subdivided based on their degree of hazard; however, care should be taken when handling and shipping any wastes believed to be concentrated, regardless of the degree.

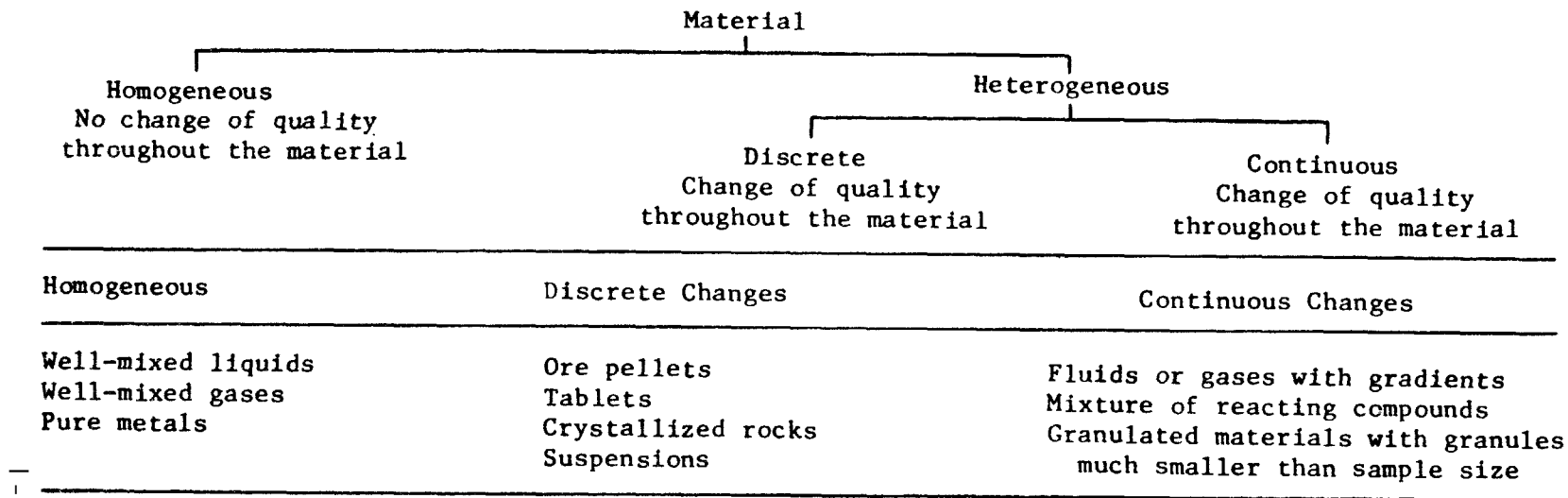
In general, two basic types of sampling techniques are recognized, both of which can be used for either environmental or concentrated samples.

Grab Samples

A grab sample is defined as a single sample representative of a specific location at a given point in time. The sample is collected all at once and at one particular point in the sample medium. The representativeness of such samples is defined by the nature of the materials being sampled. In general, as sources vary over time and distance, the representativeness of grab samples will decrease.

Composite Samples

Composites are combinations of more than one sample collected at various sampling locations and/or different points in time. Analysis of composite yields an average value and can, in certain instances, be used as an alternative to analyzing a number of individual grab samples and calculating an average



Source: Reference 1.

Figure 1-1. Types of material.

value. It should be noted, however, that compositing can mask problems by diluting isolated concentrations of some hazardous compounds below detection limits.

For sampling situations involving hazardous wastes, grab sampling techniques are generally preferred because grab sampling minimizes the amount of time sampling personnel must be in contact with the wastes, reduces risks associated with compositing unknowns, and eliminates chemical changes that might occur due to compositing. Compositing is still often used for environmental samples and may be used for hazardous samples under certain conditions. For example, compositing of hazardous waste is often performed (after compatibility tests have been completed) to determine an average value over a number of different locations (groups of drums). This procedure provides data that can be useful by providing an average concentration within a number of units, can serve to keep analytical costs down and can provide information useful to transporters and waste disposal operations.

1.5 SAMPLING PLAN

Before any sampling activities are begun, it is imperative that the purpose and goals of a program and the equipment, methodologies, and logistics to be used during the actual sampling be identified in the form of a work or sampling plan. This plan is developed when it becomes evident that a field investigation is necessary and should be initiated in conjunction with or immediately following the preliminary assessments. This plan should be concise, comprehensible and address the following items:

- Review existing work or background;
- Define goals and scope of work;
- Organization of the field teams;
- Statistical strategy;
- Quality assurance/Quality control procedures;
- Safety considerations; and
- Decontamination procedures.

Please note that this list of sampling plan components is not all inclusive. Additional elements may be inserted or altered depending on the needs of the project. It should be understood that in emergency situations personal judgement may have to be implemented. In any event, actions should be dictated by plan to maintain logical and consistent order to the task. (Additional details concerning the development of a Sampling Plan can be found in Volume I, Section 6.)

Reviewing Existing Work or Background Information

A synopsis of the site operational history as well as a review of previous study conclusions and recommendations are necessary in order to familiarize the field team members with the investigation.

In addition, this section should include regional or state maps locating the investigation area as well as detailed maps and photos of the local site. Of particular importance to the investigators is information pertaining to the following points: (1) the composition and characteristics of the wastes, (2) the adequate storage or destruction of wastes on the site, (3) the routes which the wastes could migrate off site, and (4) the effects that would occur (or might have occurred) through the discharge of waste.

Goals and Scope

A clear definition of the goals of the investigation and a detailed explanation of the tasks and phases designed to provide the information necessary to obtain the goals should be included.

The goals may be a general characterization of the site or a determination of offsite migration of contaminants or the collection of physical evidence for enforcement proceedings. Generally it is a combination of these or other potential goals which must be considered. Investigators must be aware that short-sighted goal assignment may limit utility of data for future applications.

The scope of the investigation should be outlined as discrete phases and tasks. The sequence and timeframe for each task should be delineated on a project time table or time line with key decision points and options clearly displayed.

Efficient arrangement of tasks to minimize onsite time will lead to reduced risks by reducing exposure times.

Organization of the Field Teams

Before sampling can commence, the following responsibilities must be delegated into the following roles:

- Project Team Leader--is primarily an administrator when not participating in the field investigation.
- Field Team Leader--is responsible for the overall operation and safety of the field team.
- Site Safety Officer--is primarily responsible for all safety procedures and operations.
- Command Post Supervisor--serves as a means of communication and never enters the exclusive area except for emergencies.
- Work Party--performs the onsite tasks necessary to fulfill the objectives.

Please note, that in many hazardous waste projects, one person may fulfill many roles.

Statistical Strategy

Implementation of the proper statistical strategy depends upon two essential points, the objectives or goals of the sampling plan and the amount of information available on the parameter or parameters of interest, i.e. time, spatial distribution, variability, etc. The following are among the different sampling schemes that could be chosen.

Random Sampling--

Random sampling uses the theory of random chance probabilities to choose representative sample locations. Random sampling is generally employed when little information exists concerning the material, location, etc. It is most effective when the population of available sampling locations is large enough

to lend statistical validity to the random selection process. Since one of the main difficulties with random sampling deals with achieving a truly random sample, it is advisable to use a table of random numbers to eliminate or reduce bias (Appendix G).

Systematic Sampling--

Systematic sampling involves the collection of samples at predetermined, regular intervals. It is the most often employed sampling scheme; however, care must be exercised to avoid bias. If, for example, there are periodic variations in the material to be sampled such that the systematic plan becomes partially phased with these variations, bias will result.

A systematic sampling plan is often the end result of an approach that was begun as random due to the tendency of investigators to subdivide large sample areas into smaller increments before randomizing.

Stratified Sampling--

Data and background information made available from the preliminary site survey, prior investigations conducted on site, and/or experience with similar situations can be useful in reducing the number of samples needed to attain a specified precision. Stratified sampling essentially involves the division of the sample population into groups based on knowledge of sample characteristics at these divisions. The purpose of the approach is to increase the precision of the estimates made by sampling. This objective should be met if the divisions are "selected in such a manner that the units within each division are more homogeneous than the total population."² The procedure used basically involves handling each division in a simple random approach.

Judgment Sampling--

A certain amount of judgment often enters into any sampling approach. In fact, a biased approach is the one most often employed when the intent is to document the presence of contamination (e.g., for enforcement purposes). Since judgment approaches tend to allow investigator bias to influence decisions, care must be exercised. Poor judgment can lead to poor quality data and improper conclusions. If judgment sampling is employed, it is generally advisable that enough samples be collected to lend credence to any conclusion drawn about the area under investigation because it is very difficult to actually measure sample accuracy. This is especially true for enforcement samples where the analytical results indicate no apparent sign of contamination. In such cases it is important to reduce the chance of committing a Type II statistical error. In such cases the inability to measure sample accuracy makes it difficult to rule out Type II errors (i.e., the likelihood that contaminants are present at the site even if not found in the samples).

Hybrid Sampling Schemes--

In reality, most sampling schemes consist of a combination or hybrid of the types previously described. For example, when selecting an appropriate plan for sampling drums at a hazardous waste site, the drums might be initially staged based on preliminary information concerning contents, program objectives, etc. (judgment, stratified sampling), and then sampled randomly within the

specified population groups (random sampling). Hybrid schemes are usually the method of choice as they can allow for greater diversity without compromising the objectives of the program.

For further details on this subject, please refer to Volume I, Section 6 of this series.

Quality Assurance/Quality Control

The adherence to a proper Quality Assurance--Quality Control plan is essential for a successful sampling effort. The two major concerns of a QA/QC plan are quality assurance samples and document control (chain of custody).

Quality Assurance Samples--

Quality assurance samples must be collected at any time legal action is anticipated. It is recommended that quality assurance samples be collected in all sampling surveys in order to know the quality of data collected. These additional samples are essential to any quality control aspects of the project and may also assist in reducing costs associated with resampling brought about by container breakage, errors in the analytical procedure, and data confirmation. The following is a list of the types of quality assurance samples required.

- Sample Blanks--Sample blanks are samples of deionized/distilled water, rinses of collection devices or containers, sampling media (e.g., sorbent), etc. that are handled in the same manner as the sample and subsequently analyzed to identify possible sources of contamination during collection, preservation, handling, or transport.
- Duplicates--Duplicates are essentially identical samples collected at the same time, in the same way, and contained, preserved, and transported in the same manner. These samples are often used to verify the reproducibility of the data.
- Split Samples--Split samples are duplicate samples given to the owner, operator, or person in charge for separate independent analysis.
- Spiked Samples--Spiked samples are duplicate samples that have a known amount of a substance of interest added to them. These samples are used to corroborate the accuracy of the analytical technique and could be used as an indicator of sample quality change during shipment to the laboratory.

Document Control /Chain-of-Custody--

Strict adherence to document and data control procedures is essential from the standpoint of good quality assurance/quality control and should be instituted as routine in any hazardous waste investigation. It becomes especially important when collected data is used to support enforcement litigations. All

collected information, data, calibration and maintenance records, samples, and documents, must therefore be accounted for and retrievable at any time during an investigation.

The purpose of document control is to ensure that all project documents be accounted for when the project is complete. Types of documents considered essential include maps, drawings, photographs, project work plans, quality assurance plans, serialized logbooks, data sheets, coding forms, confidential information, reports, etc.

Chain-of-custody procedures are necessary to document the sample identity, handling and shipping procedures, and in general to identify and assure the traceability of generated samples. Custody procedures trace the sample from collection, through any custody transfers, and finally to the analytical facility at which point internal laboratory procedures take over. Chain-of-custody is also necessary to document measures taken to prevent and/or detect tampering with samples, sampling equipment or the media to be sampled. A detailed description of Document Control/Chain-of-Custody Procedures can be found in Appendix D and in Volume I, Section 2.

Safety--

A more detailed discussion of safety considerations can be found in Volume I, Section 3 and Volume IV (pending release of Volume IV refer to Interim Standard Operating Safety Guides, Revised September 1982). These considerations should be carefully reviewed before engaging in any hazardous waste sampling endeavors. It is important, however, that safety be generally discussed at this time to provide a necessary reminder of the importance of taking proper, well developed precautions when dealing with hazardous materials.

Decontamination Procedures

Decontamination procedures are designed to provide two primary safeguards.

- Prevent the movement of contaminated materials into noncontaminated areas.
- Insure that samples collected during the investigation are not exposed to additional contamination from onsite materials or sampling equipment.

Proper decontamination is not only a health and safety concern but also an analytical and sampling consideration. The Sampling Plan will detail proper decontamination procedures to safeguard both the onsite personnel and the sample integrity. Appendix E provides generalized decontamination guidelines excerpted from Interim Standard Operating Safety Guides, September 1982, Office of Emergency and Remedial Response.

1.6 IMPLEMENTATION OF SAMPLING PLAN

It is the responsibility of the Field Team Leader to implement and obtain the goals of the Sampling Plan. This involves overseeing and coordinating five primary tasks:

- Management of the sampling team(s),
- Coordination with the analytical lab(s),
- Coordination with additional subcontractor efforts,
- Implementation of the Safety Plan or Coordination with the Safety Officer, and
- Implementation of the QA/QC Plan or Coordination with the Quality Assurance Officer.

Small scale efforts often utilize the Field Team Leader (FTL) as the Safety Officer and the Quality Assurance Officer. As the Sampling Plan increases in complexity, it becomes essential that the FTL designate Safety and Quality Assurance Officers in order to insure proper implementation. It then becomes the task of the FTL to coordinate their activities. Each of the five primary tasks are described below:

Management of the Sampling Team

The key task here is to insure that the goals of the sampling plan are obtained. In addition to the selection and proper implementation of methods, the FTL must continually adjust, and carefully document changes to the sampling plan to accommodate situations which may arise. This may involve, for example, relocating or adding sampling locations if the investigation uncovers new sources or should adverse weather make some locations inaccessible.

Thorough and detailed documentation of all onsite activities is also a critical responsibility. This includes records of all expenditures, manpower, and equipment uses and any changes of scope. These records are particularly important for Superfund investigation or any investigation where attempts will be made to recover costs from responsible parties. This aspect of the FTL's responsibilities is often overlooked or downplayed, but to do so is likely to later result in tremendous problems in cost recovery.

Coordination with the Analytical Lab

The sampling plan also serves to integrate the responsibilities of Sampling Teams and the analytical labs. It is critical that the sampling activities are coordinated with the laboratory. The following points illustrate the extent of this coordination and its importance to the project:

- Sampling schedules must be arranged with the laboratory manager to insure that the samples can be processed within the specified holding times.
- Labels and labeling information should be discussed. This will aid in the orderly transfer of field information to the proper laboratory personnel and insure that each sample receives only the specified analyses.
- Shipping arrangements, if necessary, must be agreed to in advance. Selection of a carrier, delivery times, and billing procedures must be mutually acceptable.
- Preservation requirements and equipment decontamination procedures should be specified. This would include specific recommendations reagents and cleaning solvents.
- A field and a laboratory coordinator should be designated from both groups to serve as points of communication. In most situations, the laboratory will specify the number and type of Quality Assurance/Quality Control samples. Should this decision be made by field personnel, information regarding these samples must be transmitted to the Laboratory's QA/QC director.

Failure to properly coordinate these activities can result in complete data loss, or at a minimum a reduction in its quality and overall reliability. Either of these outcomes translates into potentially significant waste of time and money.

Coordination with Other Subcontractor Efforts

Many investigations require the services of a team of subcontractors. These include Drilling Contractors, Geophysical Investigation Teams, Aerial Photographic Contractors, and Trade Contractors (electricians, plumbers, carpenters and fencing contractors). It is the FTL's responsibility to coordinate their activities, insure adherence to the sampling plan, or contractual requirements.

Here again, thorough recordkeeping and documentation is critical.

Implementation of the Safety Plan

The Safety Plan must be implemented prior to full scale mobilization of onsite activities. This would include establishing decontamination stations, command posts, first aid stations, etc. If the scope of the investigation is large, the FTL should designate a Site Safety Officer to implement the safety plan.

Volume I, Section 3 discusses the basic aspects of a Safety Plan. Volume IV will provide details on the preparation and provisions of safety plans, however until release of this document, this information is contained in the Interim Standard Operation Safety Guides, September 1982, U.S. EPA, OERR.

Implementation of the QA/QC Plan.

The QA/QC Plan must address all phases of the investigation including field measurements, laboratory analysis, subcontractor activities, documentation/recordkeeping and report preparation. As with the Safety Plan, the complexity and time demands of this task increase with the overall project complexity. Therefore on larger projects, the FTL may designate a QA/QC officer.

The FTL must be familiar with the terms of the QA/QC Plan and insure that they are implemented by all field and laboratory personnel. Of particular concern to the FTL are document control and chain of custody procedures. As stated earlier, and cannot be over emphasized, are the demands on the FTL for documentation and recordkeeping, these tasks must also closely adhere to document control procedures. Without these records and documented assurance of their completeness and validity litigation and cost recovery efforts will be severely handicapped.

1.7 REFERENCES

1. Kateman, G. and F. W. Pijpers. Quality Control in Analytical Chemistry. John Wiley and Sons, New York, 1981.
2. Smith, R. and G. James. The Sampling of Bulk Materials. Analytical Sciences Monographs, Volume 8. The Royal Society of Chemistry, London, 1981.

SECTION 2.0

SOLIDS

2.1 GENERAL

The sampling of solid or semi-solid materials is complicated by the structural properties of the material. For example, the presence of entrapped gases and fluids is often an integral part of the substance and may be of consequence in the analytical techniques for which the sample was collected. It is necessary in most cases to collect a sample which does not alter this balance. In addition, physical strength and density of the material demand sampling devices of significant rigidity and strength. As a result a great deal of disturbance will occur at the sample-sampler interface. These effects can be reduced by careful sampling and by collecting aliquots with a high volume to surface area ratio.

A solid does not necessarily have uniform characteristics with respect to distance or depth. Those portions which form boundaries with the container, define the edges of a pile, or contact the atmosphere do not necessarily represent the material as a whole. Care must be exercised in order to prevent aeration or significant changes in moisture content. Samples should be tightly capped and protected from direct light.¹

Most commercially available solids sampling devices are steel, brass or plastic. In general, use of stainless steel is the most practical and several manufacturers will fabricate their equipment with all stainless steel parts on a special order basis. Another alternative is to have sampler contact surfaces Teflon coated. This can be accomplished by either sending the device to a commercial coater or by in-house application of spray-on Teflon coatings. Some devices, especially those for soil sampling, have traditionally been chrome- or nickel-plated steel. These should be particularly avoided, or the plating should be removed because scratches and flaking of the plating material can drastically effect the results of trace element analysis. Plated or painted surfaces, can be used in many cases if the outside coating is first removed by using abrasives. Such practice can yield a significant cost savings over more expensive materials, so long as the exposed material will affect the sample.

This section is divided into three subsections which address the sampling of soils, sludge and sediments, and bulk materials.

2.2 SOILS

Soil sampling is an important adjunct to groundwater monitoring. Sampling of the soil horizons above the groundwater table can detect contaminants before they have migrated into the water table, and can help establish the amount of contamination sorbed on aquifer solids that have the potential of contributing to the groundwater contamination.

Soil types can vary considerably on a hazardous waste site. These variations, along with vegetation, can effect the rate of contaminant migration through the soil. It is important, therefore, that a detailed record be maintained during sampling operations, particularly of location, depth, and such characteristics as grain size, color and odor, and/or readings obtained on field monitoring equipment. Subsurface conditions are often stable on a daily basis and may demonstrate only slight seasonal variation especially with respect to temperature, available oxygen, and light penetration. Changes in any of these conditions can radically alter the rate of chemical reactions or the activity of associated microbiological community. As a result samples should be kept at their at-depth temperature or lower, protected from direct light, sealed tightly in glass bottles, and analyzed as soon as possible.

The physical properties of the soil, its grain size, cohesiveness, associated moisture, and such factors as depth to bedrock and water table will limit the depth from which samples can be collected and the method required to collect them. Often this information on soil properties can be acquired from published soil surveys obtainable through the U.S. Geological Survey (USGS) and other government and farm agencies. A comprehensive listing of these offices and currently available soil surveys is included in the "NEIC Manual for Groundwater/Subsurface Investigations at Hazardous Waste Sites."² Most of the methods employed for soil sampling at hazardous waste sites are adaptations of techniques long employed by foundation engineers and geologists. This section presents those methods which can be employed with a minimum of special training, equipment or cost. More detailed methods capable of sampling to greater depths in more difficult soil conditions, or that can simultaneously install groundwater monitor wells, usually require professional assistance. These techniques are discussed more fully in the "Manual for Ground-water Sampling Procedures."³

Collection of samples from near the soil surface can be accomplished with tools such as spades, shovels, and scoops. With this type of readily available equipment the soil cover can be removed to the required depth; then a stainless steel scoop can be used to collect the sample. An undisturbed sample can be collected from this excavation by employing a thin wall tube sampler. This device is, as the name implies, a metal tube generally 2.5 to 7.5 cm in diameter and 30.5 to 61.0 cm long. The tube is forced into the soil, then extracted. Friction will usually hold the sample material in the tube during the extraction. The construction material is generally steel, and some samplers can utilize plastic liners and interchangeable cutting tips. The liners are useful for trace element sampling but are generally not suitable for organic analysis due to the possibility that materials in the liner will

leach out and become incorporated as part of the sample. The liner tubes can further be capped off and used as sample containers for transport to the lab.

Interchangeable cutting tips facilitate smoother penetration with reduced sample disturbance. They are available in various styles and construction suitable for moist, dry, sandy or heavy-duty applications. The design of these cutting tips will further aid in maintaining the sample in the tube during sample extraction.

Augers are also very effective for soil sampling. Bucket type augers can be used directly for soil sample collection or to advance a borehole to the desired depth so then a thin wall tube can be employed.

Kits are available that include, in conjunction with the tube sampler and cutting tips, an auger point and a series of extension rods. These kits allow for hand augering a borehole. The auger can then be removed and a tube sampler lowered and forced into the soil at the completion depth. Though kits are available with sufficient tools to reach depths in excess of 7 meters, soil structure, impenetrable rock, and water levels usually prevent reaching such completion depths. Kits that include 1 meter of drill rod and the ability to order additional extensions will in practice prove satisfactory. The need for soil information at greater depths will normally require professional assistance. Consideration should be given to supplementing this information with groundwater monitoring since soil sampling can be conducted in conjunction with well completion.

For those wishing a more in-depth discussion of soils and soil sampling, refer to the Preparation of Soil Sampling Protocol: Techniques and Strategies, (EPA 600/4-83-020) by Dr. Benjamin J. Mason, prepared under contract to the U.S. Environmental Protection Agency, Environmental Monitoring Systems Laboratory--Las Vegas, August 1983. This report discusses in detail the factors that influence the selection of a particular sampling scheme or the use of a particular sampling method with a strong emphasis on statistical design and data analysis. Another document, Soil Sampling Quality Assurance User's Guide, (EPA 600/4-84-043) by Dr. Delbert S. Barth and Dr. Benjamin J. Mason, prepared by the Environmental Research Center, University of Nevada-Las Vegas under a cooperative agreement with the Environmental Protection Agency (May 1984) will also be helpful.

2.2.1 METHOD II-1: SOIL SAMPLING WITH A SPADE AND SCOOP

Discussion

The simplest, most direct method of collecting soil samples for subsequent analysis is with the use of a spade and scoop. A normal lawn or garden spade can be utilized to remove the top cover of soil to the required depth and then a smaller stainless steel scoop can be used to collect the sample.

Uses

This method can be used in most soil types but is limited somewhat to sampling the near surface. Samples from depths greater than 50 cm become extremely labor intensive in most soil types. Very accurate, representative samples can be collected with this procedure depending on the care and precision demonstrated by the technician. The use of a flat, pointed mason trowel to cut a block of the desired soil will be of aid when undisturbed profiles are required. A stainless steel scoop or lab spoon will suffice in most other applications. Care should be exercised to avoid the use of devices plated with chrome or other materials. Plating is particularly common with garden implements such as potting trowels.

Procedures for Use

1. Carefully remove the top layer of soil to the desired sample depth with a precleaned spade.
2. Using a precleaned stainless steel scoop or trowel, remove and discard a thin layer of soil from the area which comes in contact with the shovel.
3. Transfer sample into an appropriate sample bottle with a stainless steel lab spoon or equivalent.
4. Check that a Teflon liner is present in the cap if required. Secure the cap tightly. The chemical preservation of solids is generally not recommended. Refrigeration is usually the best approach supplemented by a minimal holding time. For specific containerization and preservation requirements consult Appendix A.
5. Label the sample bottle with the appropriate sample tag. Be sure to label the tag "carefully and clearly", addressing all the categories or parameters. Complete all chain-of-custody documents and record in the field log book.
6. Decontaminate equipment after use and between sample locations. For specific decontamination guidelines, consult Appendix E.

2.2.2 METHOD 11-2: SUBSURFACE SOLID SAMPLING WITH AUGER
AND THIN-WALL TUBE SAMPLER

Discussion

This system consists of an auger bit, a series of drill rods, a "T" handle, and a thin-wall tube corer (see Figure 2-1). The auger bit is used to bore a hole to the desired sampling depth and then withdrawn. The auger tip is then replaced with the tube corer, lowered down the borehole, and forced into the soil at the completion depth. The corer is then withdrawn and the sample collected.

Alternately the sample can be recovered directly from the auger. This technique however, does not provide an "undisturbed" sample as would be collected with a thin tube sampler. In situations where the soil is rocky, it may not be possible to force a thin tube sampler through the soil or sample recovery may be poor. Sampling directly from the auger may be the only viable method. Several auger types are available which include Bucket type, continuous flight (screw) and posthole augers. Bucket types are good for direct sample recovery and are fast and provide a large volume of sample. When continuous flight (screw) augers are utilized, the sample can be collected directly off the flights, however, this technique will provide a somewhat unrepresentative sample as the exact sample depth will not be known. The continuous flights auger are, however, satisfactory for use when a composite of the entire soil column is desired. Posthole augers have limited utility for sample acquisition as they are designed more for their ability to cut through fibrous, heavily rooted, swampy areas. In soils where the borehole will not remain open when the tool is removed, a temporary casing can be used until the desired sampling depth is reached.

Uses

This system can be used in a wide variety of soil conditions. It can be used to sample both from the surface, by simply driving the corer without preliminary boring, or to depths in excess of 6 meters. The presence of rock layers and the collapse of the borehole, however, usually prohibit sampling at depths in excess of 2 meters. Interchangeable cutting tips on the corer reduce the disturbance to the soil during sampling and aid in maintaining the core in the device during removal from the borehole.

Procedures for Use

1. Attach the auger bit to a drill rod extension and further attach the "T" handle to the drill rod.
2. Clear the area to be sampled of any surface debris (twigs, rocks, litter). It may be advisable to remove the first 8 to 5 cm of surface soil for an area approximately 15 cm in radius around the drilling location.

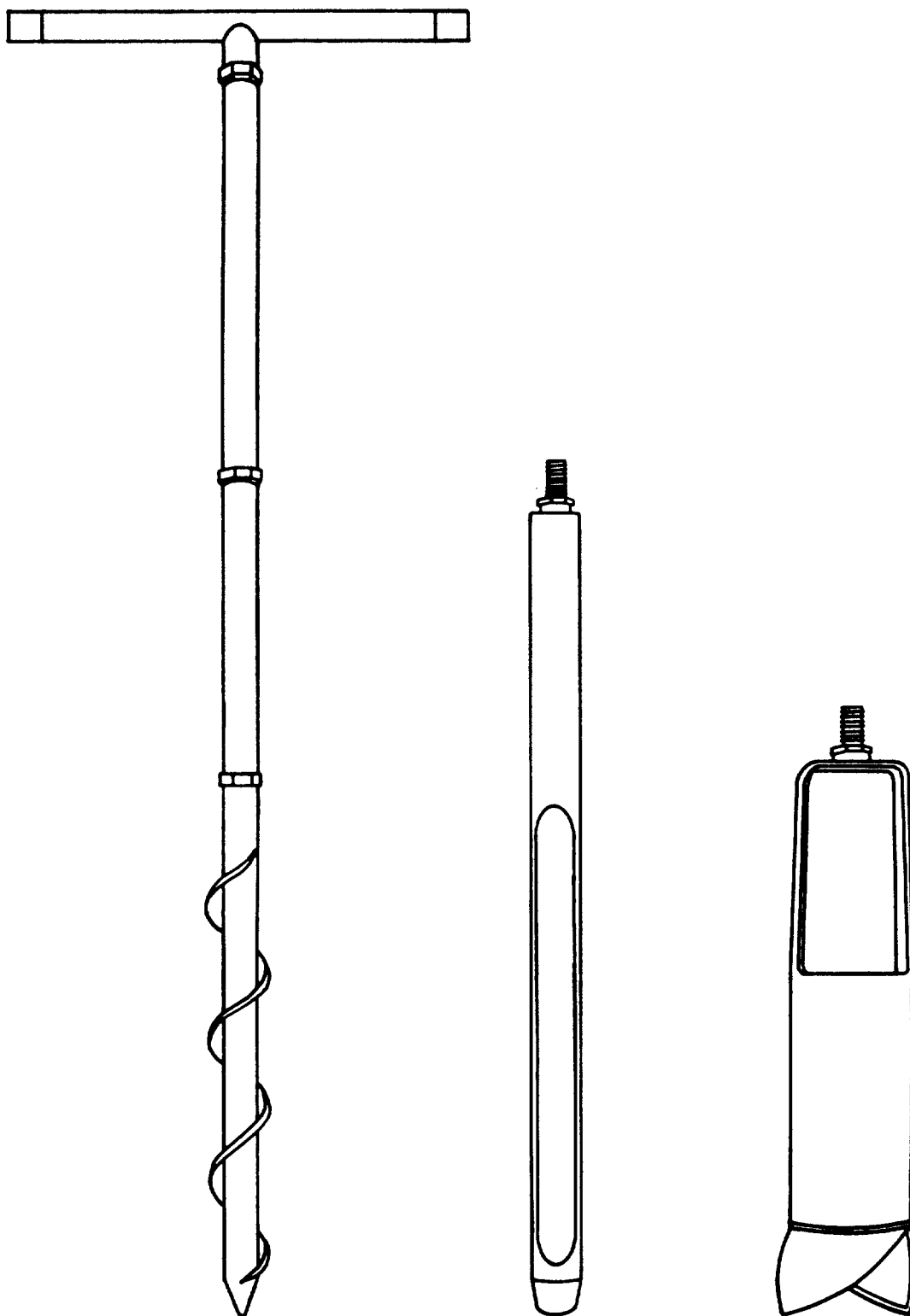


Figure 2-1. Augers and thin-wall tube sampler.

3. Begin drilling, periodically removing accumulated soils. This prevents accidentally brushing loose material back down the borehole when removing the auger or adding drill rods.
4. After reaching desired depth, slowly and carefully remove auger from boring. (Note: When sampling directly from auger, collect sample after auger is removed from boring and proceed to Step 10.)
5. Remove auger tip from drill rods and replace with a precleaned thin-wall tube sampler. Install proper cutting tip.
6. Carefully lower corer down borehole. Gradually force corer into soil. Care should be taken to avoid scraping the borehole sides. Hammering of the drill rods to facilitate coring should be avoided as the vibrations may cause the boring walls to collapse.
7. Remove corer and unscrew drill rods.
8. Remove cutting tip and remove core from device.
9. Discard top of core (approximately 2.5 cm), which represents any material collected by the corer before penetration of the layer in question. Place remaining core into sample container.
10. Check that a Teflon liner is present in the cap if required. Secure the cap tightly. The chemical preservation of solids is generally not recommended. Refrigeration is usually the best approach supplemented by a minimal holding time. Consult Appendix A for containerization and preservation recommendations.
11. Label the sample bottle with the appropriate sample tag. Be sure to label the tag carefully and clearly, addressing all the categories or parameters. Complete all chain-of-custody documents and record in the field logbook.
12. Decontaminate sampling equipment after use and between sampling locations. Refer to Appendix E for decontamination requirements.

Sources

deVera, E. R., Simmons, B. P., Stephens, R. D., and Storm, D. L. "Samplers and Sampling Procedures for Hazardous Waste Streams." EPA 600/2-80-018, January 1980.

2.3 SLUDGES AND SEDIMENTS

In general and for the purpose of this manual, sludges will be defined as semi-dry materials ranging from dewatered solids to high viscosity liquids. Sediments are the deposited material underlying a body of water. On occasion they are exposed by evaporation, stream rerouting, or other means of water loss. In these instances they can be readily collected by soil or sludge collection methods.

Sludges can often be sampled by the use of a stainless steel scoop or trier. Frequently sludges form as a result of settling of the higher density components of a liquid. In this instance the sludge may still have a liquid layer above it. When the liquid layer is sufficiently shallow, the sludge may be scooped up by a device such as the pond sampler described in Section III, Method III-2, or preferably by using a thin-tube sampler as described in this section (see Method II-4). The latter is preferable as it results in less sample disturbance and will also collect an aliquot of the overlying liquid, thus preventing drying or excessive sample oxidation before analysis. Sludges which develop in 55-gallon drums can usually be collected by employing the glass tubes used for the liquid portion sample (Method III-5) as a thin-tube sampler. The frictional forces which hold the sludge in the tube can be supplemented by maintaining a seal above the tube. When the overlying layer is deep, a small gravity corer such as those used in limnological studies will be useful. Gravity corers, such as Phlegers, are easier to preclean and decontaminate than piston type corers.

If the sludge layer is shallow, less than 30 centimeters, corer penetration may damage the container liner or bottom. In this instance a Ponar or Eckman grab may be applicable, as grab samplers are generally capable of only a few centimeters of penetration. Of the two, Ponar grab samplers are more applicable to a wider range of sediments and sludges. They penetrate deeper and seal better than the spring-activated Eckman dredges, especially in granular substrates.

In many instances sediments and sludges can be collected with a peristaltic pump as described in Method III-3. This method is limited to slurried samples less than approximately 20 percent solid. The weight of the material will also greatly reduce the lift capacity of the pump, however, it may still be useful in extending the reach of the sampler laterally toward the center of a vessel. In slurries not fully agitated, a bias may also be introduced toward the liquid portion of the material.

Sediments can be collected in much the same manner as described above for sludges; however, a number of additional factors may be considered. Streams, lakes, and impoundments, for instance, will likely demonstrate significant variations in sediment composition with respect to distance from inflows, discharges, or other disturbances. It is important, therefore, to document exact sampling location by means of triangulation with stable references on the banks of the stream or lake. In addition, the presence of rocks, debris, and organic material may complicate sampling and preclude the use of or require modification to some devices. Sampling of sediments should therefore be conducted to reflect these and other variants.

2.3.1 METHOD 11-3: COLLECTION OF SLUDGE OR SEDIMENT SAMPLES WITH A SCOOP

Discussion

Sludge and sediment samples are collected using the simple laboratory scoop or garden type trowel specified in Method 11-1. This method is more applicable to sludges but it can be used for sediments provided the water depth is very shallow (a few centimeters). It should be noted, however, that this method can be disruptive to the water/sediment interface and might cause substantial alterations in sample integrity if extreme care is not exercised. The stainless steel laboratory scoop is generally recommended due to its noncorrosive nature. Single grab samples may be collected or, if the area in question is large, it can be divided into grids and multiple samples can be collected and composite.

Uses

This method provides for a simple, quick, and easy means of collecting a disturbed sample of a sludge or sediment.

Procedures for Use

1. Sketch the sample area or note recognizable features for future reference.
2. Insert scoop or trowel into material and remove sample. In the case of sludges exposed to air, it may be desirable to remove the first 1-2 cm of material prior to collecting sample.
3. If compositing a series of grab samples, use a stainless steel mixing bowl or Teflon tray for mixing.
4. Transfer sample into an appropriate sample bottle with a stainless steel lab spoon or equivalent.
5. Check that a Teflon liner is present in cap if required. Secure the cap tightly. The chemical preservation of solids is generally not recommended. Refrigeration is usually the best approach supplemented by a minimal holding time. Containerization and preservation requirements are detailed in Appendix A.
6. Label the sample bottle with the appropriate sample tag. Be sure to label the tag carefully and clearly, addressing all the categories or parameters. Complete all chain-of-custody documents and record in the field logbook.
7. Decontaminate sampling equipment after use and between sample locations according to the guidelines presented in Appendix E.

2.3.2 METHOD 11-4: SAMPLING SLUDGE OR SEDIMENTS WITH A HAND CORER

Discussion

This device is essentially the same type of thin-wall corer described for collecting soil samples (Method 11-2). It is modified by the addition of a handle to facilitate driving the corer (see Figure 2-2) and a check valve on top to prevent washout during retrieval through an overlying water layer.

Uses

Hand corers are applicable to the same situations and materials as the scoop described in Method 11-3. It has the advantage of collecting an undisturbed sample which can profile any stratification in the sample as a result of changes in the deposition.

Some hand corers can be fitted with extension handles which will allow the collection of samples underlying a shallow layer of liquid. Most corers can also be adapted to hold liners generally available in brass, polycarbonate plastic or Teflon. Care should be taken to choose a material which will not compromise the intended analytical procedures.

Procedures for Use

1. Inspect the corer for proper precleaning, and select sample location.
2. Force corer in with smooth continuous motion.
3. Twist corer then withdraw in a single smooth motion.
4. Remove nosepiece and withdraw sample into a stainless steel or Teflon tray.
5. Transfer sample into an appropriate sample bottle with a stainless steel lab spoon or equivalent.
6. Check that a Teflon liner is present in cap if required. Secure the cap tightly. The chemical preservation of solids is generally not recommended. Refrigeration is usually the best approach supplemented by a minimal holding time. Appendix A, Sample Containerization and Preservation should be consulted for specific requirements.
7. Label the sample bottle with the appropriate sample tag. Be sure to label the tag carefully and clearly, addressing all the categories or parameters. Complete all chain-of-custody documents and record in the field logbook.
8. Decontaminate sampling equipment after use and between sample locations as required by procedures in Appendix E, Decontamination.

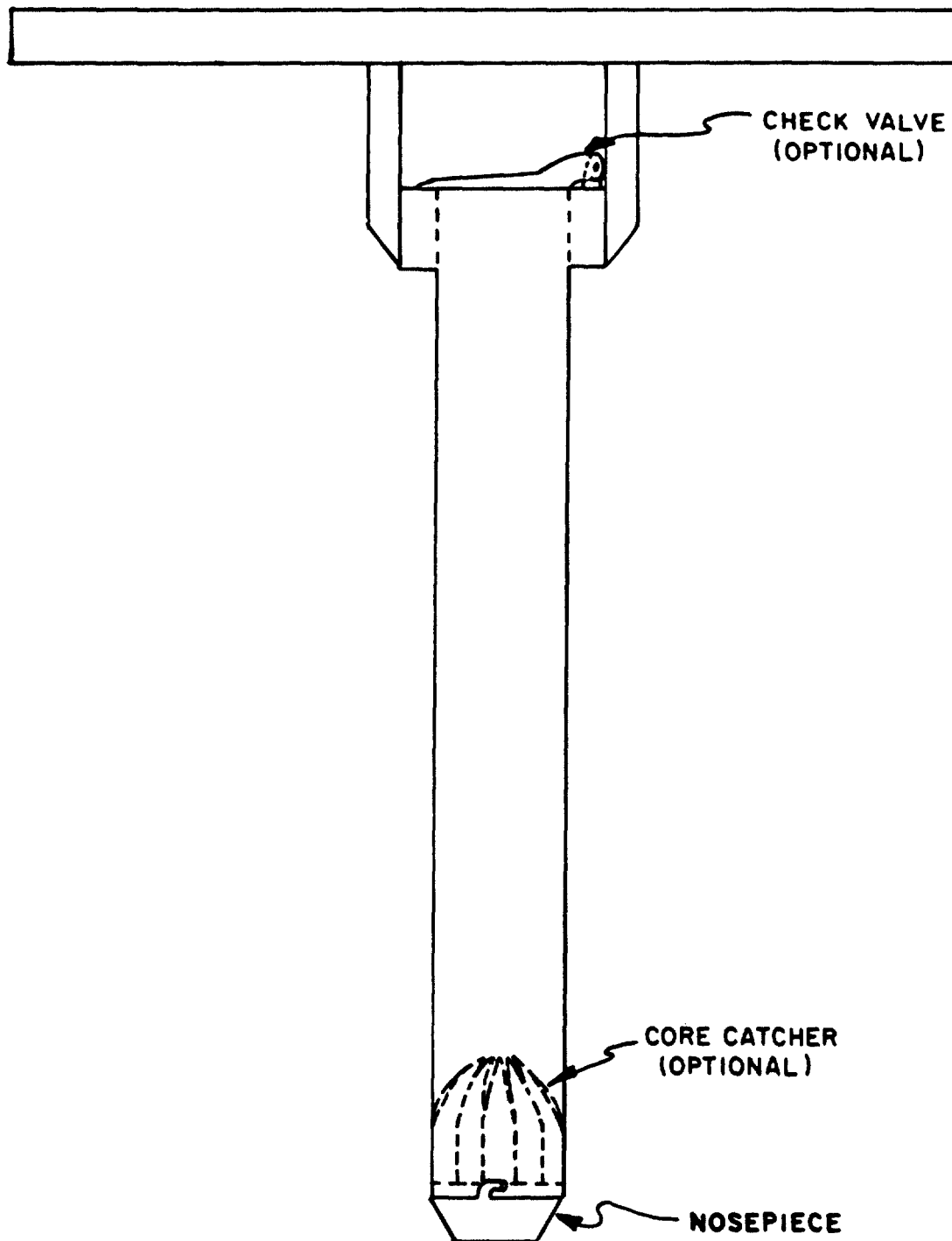


Figure 2-2. Hand corer.

2.3.3 METHOD 11-5: SAMPLING BOTTOM SLUDGES OR SEDIMENTS WITH A GRAVITY CORER

Discussion

A gravity corer is a metal tube with a replacement tapered nosepiece on the bottom and a ball or other type of check valve on the top. The check valve allows water to pass through the corer on descent but prevents a washout during recovery. The tapered nosepiece facilitates cutting and reduces core disturbance during penetration.

Most corers are constructed of brass or steel and many can accept plastic liners and additional weights (see Figure 2-3).

Uses

Corers are capable of collecting samples of most sludges and sediments. They collect essentially undisturbed samples which represent the profile of strata which may develop in sediments and sludges during variations in the deposition process. Depending on the density of the substrate and the weight of the corer, penetration to depths of 75 cm (30 inches) can be attained.

Care should be exercised when using gravity corers in vessels or lagoons that have liners because penetration depths could exceed that of the substrate and result in damage to the liner material.

Procedures for Use

1. Attach a precleaned corer to the required length of sample line. Solid braided 5 mm (3/16 inch) nylon line is sufficient; 20 mm (3/4 inch) nylon, however, is easier to grasp during hand hoisting.
2. Secure the free end of the line to a fixed support to prevent accidental loss of the corer.
3. Allow corer to free fall through liquid to bottom.
4. Retrieve corer with a smooth, continuous lifting motion. Do not bump corer as this may result in some sample loss.
5. Remove nosepiece from corer and slide sample out of corer into stainless steel or Teflon pan.
6. Transfer sample into appropriate sample bottle with a stainless steel lab spoon or equivalent.
7. Check that a Teflon liner is present in cap if required. Secure the cap tightly. The chemical preservation of solids is generally not recommended. Refrigeration is usually the best approach supplemented by a minimal holding time. Refer to Appendix A for sample containerization and preservation guidelines.

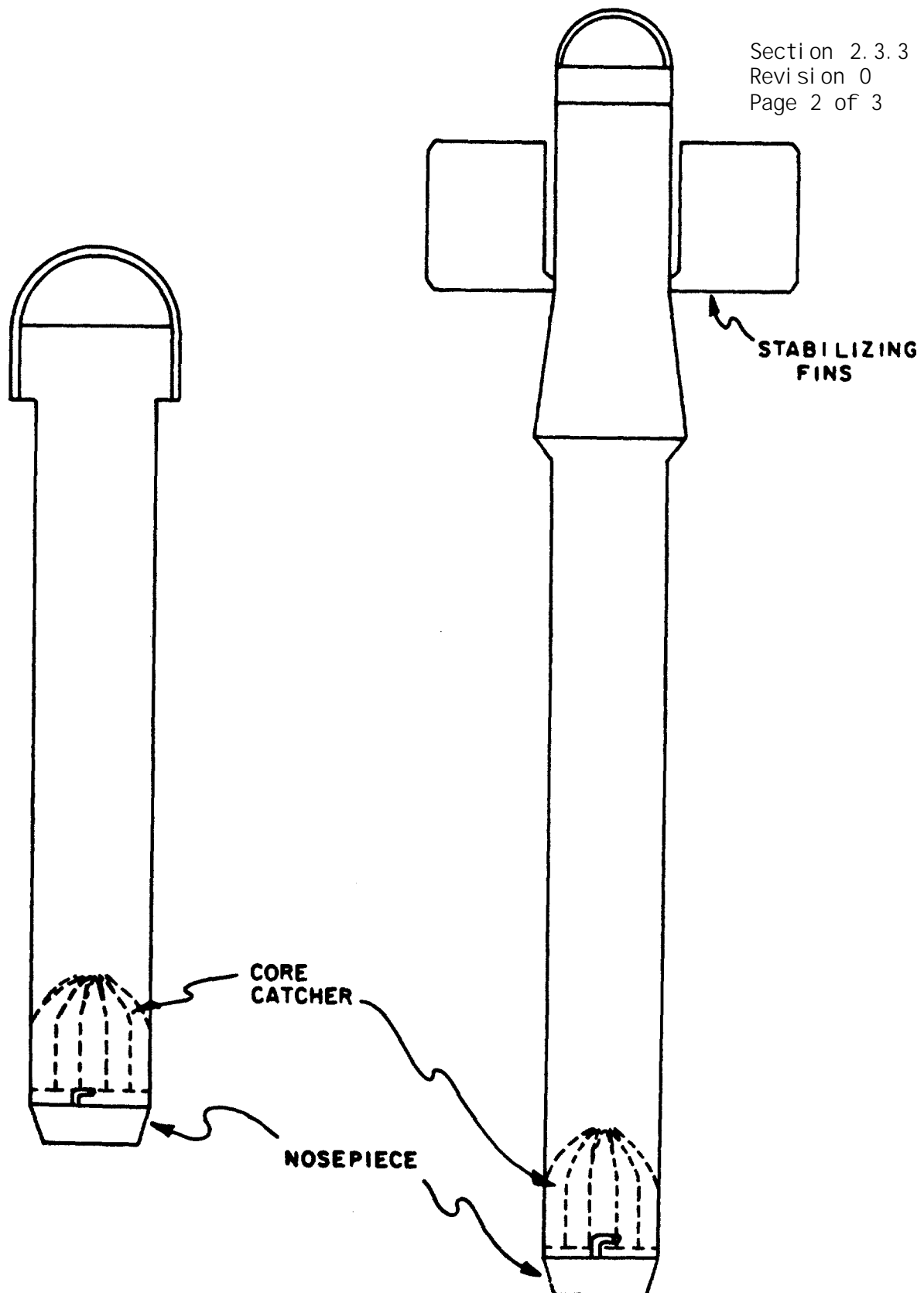


Figure 2-3. Gravity corers.

8. Label the sample bottle with the appropriate sample tag. Be sure to label the tag carefully and clearly, addressing all the categories or parameters. Complete all chain-of-custody documents and record in the field logbook.
9. Consult Appendix E for decontamination requirements and decontaminate sampling equipment after use and between sampling locations.

Sources

American Public Health Association. "Standard Methods for the Examination of Water and Wastewater" 14th Edition, Washington, D.C. 1975.

2.3.4 METHOD 11-6: SAMPLING BOTTOM SLUDGES OR SEDIMENTS WITH A PONAR GRAB

Discussion

The Ponar grab is a clamshell type scoop activated by a counter lever system. The shell is opened and latched in place and slowly lowered to the bottom. When tension is released on the lowering cable the latch releases and the lifting action of the cable on the lever system closes the clamshell (see Figure 2-4).

Uses

Ponars are capable of sampling most types of sludges and sediments from silts to granular materials. They are available in a "Petite" version with a 232 square centimeter sample area that is light enough to be operated without a winch or crane. Penetration depths will usually not exceed several centimeters. Grab samplers, unlike the corers described in Method 11-5, are not capable of collecting undisturbed samples. As a result, material in the first centimeter of sludge cannot be separated from that at lower depths. The sampling action of these devices causes agitation currents which may temporarily resuspend some settled solids. This disturbance can be minimized by slowly lowering the sampler the last half meter and allowing a very slow contact with the bottom. It is advisable, however, to only collect sludge or sediment samples after all overlying water samples have been obtained.

Procedures for Use

1. Attach a precleaned Ponar to the necessary length of sample line. Solid braided 5 mm (3/16 inch) nylon line is usually of sufficient strength; however, 20 mm (3/4 inch) or greater nylon line allows for easier hand hoisting.
2. Measure and mark the distance to bottom on the sample line. A secondary mark, 1 meter shallower, will indicate proximity so that lowering rate can be reduced, thus preventing unnecessary bottom disturbance.
3. Open sampler jaws until latched. From this point on, support sampler by its lift line or the sampler will be tripped and the jaws will close.
4. Tie free end of sample line to fixed support to prevent accidental loss of sampler.
5. Begin lowering the sampler until the proximity mark is reached.
6. Slow rate of descent through last meter until contact is felt.
7. Allow sample line to slack several centimeters. In strong currents more slack may be necessary to release mechanism.

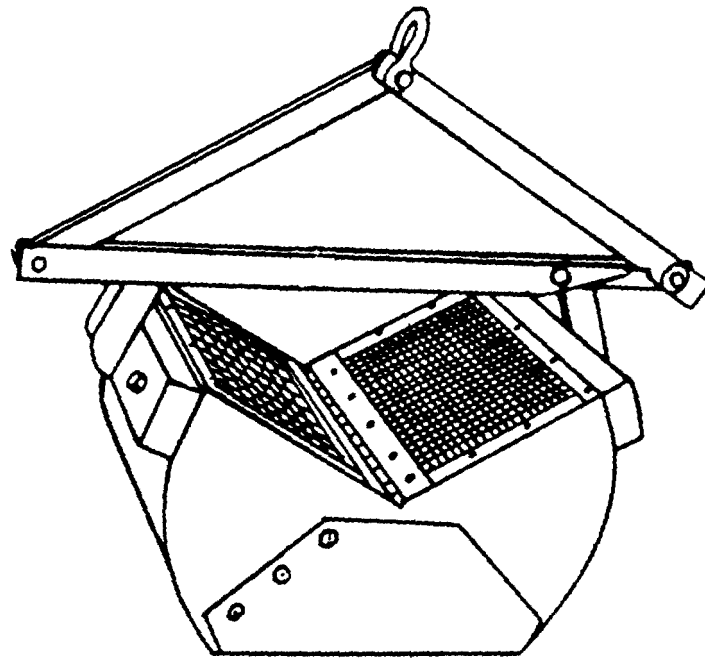


Figure 2-4. Ponar grab

8. Slowly raise dredge clear of water surface.
9. Place Ponar into a stainless steel or Teflon tray and open. Lift Ponar clear of the tray.
10. Collect a suitable aliquot with a stainless steel lab spoon or equivalent and place sample into appropriate sample bottle. Appendix A contains containerization and preservation requirements.
11. Check for a Teflon liner in cap if required and secure cap tightly. The chemical preservation of solids is generally not recommended. Refrigeration is usually the best approach supplemented by a minimal holding time.
12. Label the sample bottle with the appropriate sample tag. Be sure to label the tag carefully and clearly, addressing all the categories or parameters. Complete all chain-of-custody documents and record in the field logbook.
13. Consult Appendix E, Decontamination for appropriate decontamination procedures to be used on sampling equipment after use and between sampling locations.

Sources

American Public Health Association. "Standard Methods for the Examination of Water and Wastewater" 14th Edition, American Public Health Association, Washington, D.C. 1975.

Lind, Owen T. "Handbook of Common Methods in Limnology." C.V. Mosby Company, St. Louis, 1974.

2.4 BULK MATERIALS

Unlike soils which are heterogeneous associations of earthen and manufactured substances, bulk materials are generally a homogeneous collection of a single identifiable product. They are usually contained in bags, drums or hoppers although on occasion large amounts of the material may be piled directly on the ground, either deliberately or as the result of a spill.

Those surfaces exposed to the atmosphere may undergo some chemical alteration or degradation and should be avoided during sample collection. Since the process producing the bulk material may demonstrate some variation with respect to time, it is advisable to collect a series of samples as one composite to represent the material.

Bulk materials in an unconsolidated state may be readily collected by a stainless steel scoop. When the amount of the material is large, a composite can be collected by the use of a grain thief (see Figure 2-6). This device is essentially a long hollow tube with evenly spaced openings along its length. This tube is placed inside an outer sleeve with similar openings and forced into the material. The inner sleeve is rotated until its openings align with those on the outer sleeve, thus allowing the material to enter. The inner sleeve is then further rotated sealing the openings, the device is withdrawn, and the sample recovered.

Grain thieves are available in many materials including brass and various plastics. As with other sampling devices, care should be taken to choose a construction material which will not compromise the desired analytical results.

A more detailed treatment of this subject (Bulk Materials) can be found in The Sampling of Bulk Materials by R. Smith and G. V. James, The Royal Society of Chemistry, London (1981). Although this book does not deal specifically with hazardous waste sampling, the concepts discussed, especially on the subject of the establishment of a sampling scheme, are readily applicable.

2.4.1 METHOD 11-7: SAMPLING OF BULK MATERIAL WITH A SCOOP OR TRIER

Discussion

A typical sampling trier (Figure 2-5) is a long tube with a slot that extends almost its entire length. The tip and edges of the tube slot are sharpened to allow the trier to cut a core of the material to be sampled when rotated after insertion into the material. Sampling triers are usually made of stainless steel with wooden handles. They are about 61 to 100 cm long and 1.27 to 2.54 cm in diameter. They can be purchased readily from laboratory supply houses.

A laboratory scoop or garden variety trowel can also be used to sample bulk material. The trowel looks like a small shovel. The blade is usually about 7 by 13 cm with a sharp tip. A laboratory scoop is similar to the trowel, but the blade is usually more curved and has a closed upper end to permit the containment of material. Scoops come in different sizes and shapes. Stainless steel or polypropylene scoops with 7 by 15 cm blades are preferred. A trowel can be bought from hardware stores; the scoop can be bought from laboratory supply houses.

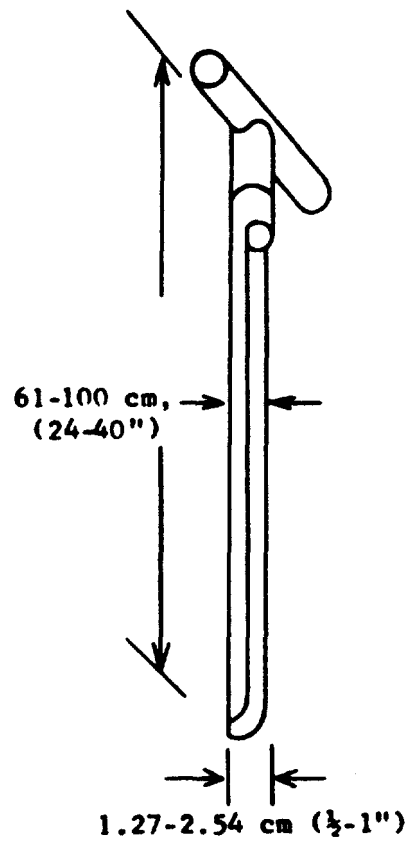
Uses

The use of the trier is similar to that of the grain sampler discussed in Method 11-8. It is preferred over the grain sampler when the powdered or granular material to be sampled is moist or sticky.

The trowel or lab scoop can be used in some cases for sampling dry, granular or powdered material in bins or other shallow containers. The lab scoop is a superior choice since it is usually made of materials less subject to corrosion or chemical reactions.

Procedures for Use

1. Insert the precleaned trier into the waste material at a 0 to 45° angle from horizontal. This orientation minimizes the spillage of sample from the sampler. Extraction of samples might require tilting of the containers.
2. Rotate the trier once or twice to cut a core of material.
3. Slowly withdraw the trier, making sure that the slot is facing upward.
4. Transfer the sample into a suitable container with the aid of a spatula and/or brush.
5. If composite sampling is desired, repeat the sampling at different points two or more times and combine the samples in the same sample container.



Source: Reference 4.

Figure 2-5. Sampling trier.

6. Check that a Teflon liner is present in the cap if required. Secure the cap tightly. The chemical preservation of solids is generally not recommended. Refrigeration is usually the best approach supplemented by a minimal holding time. Consult Appendix A for sample containerization and preservation requirements.
7. Label the sample bottle with the appropriate sample tag. Be sure to label the tag carefully and clearly, addressing all the categories or parameters. Complete all chain-of-custody documents and record in the field logbook.
8. Clean and decontaminate sampler after use and between sampling locations as per guidelines presented in Appendix E, Decontamination.

Sources

deVera, E.R., Simmons, B.P., Stephens, R.D., and Storm, D.L. "Samplers and Sampling Procedures for Hazardous Waste Streams." EPA-600/2-80-018. January 1980.

2.4.2 METHOD 11-8: SAMPLING BULK MATERIALS WITH A GRAIN THIEF

Discussion

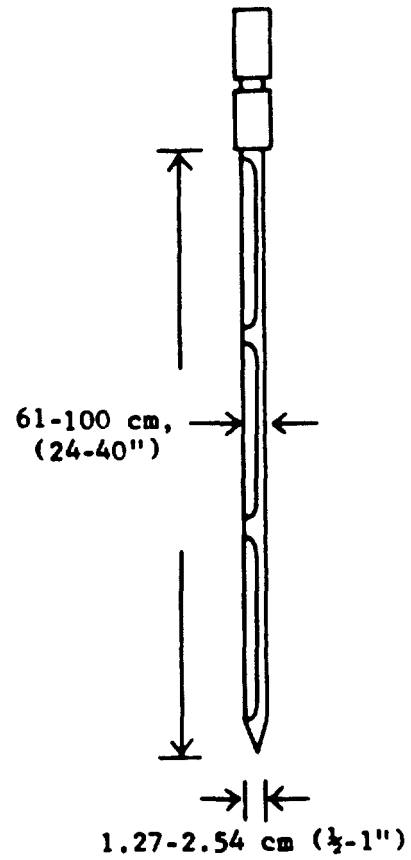
The grain thief (Figure 2-6) consists of two slotted telescoping tubes, usually made of brass or stainless steel. The outer tube has a conical, pointed tip on one end that permits the sampler to penetrate the material being sampled. The sampler is opened and closed by rotating the inner tube. Grain thieves are generally 61 to 100 cm long by 1.27 to 2.54 cm in diameter, and they are commercially available at laboratory supply houses.

Uses

The grain thief is used for sampling powdered or granular wastes or materials in bags, fiberdrums, sacks or similar containers. This sampler is most useful when the solids are no greater than 0.6 cm in diameter.

Procedures for Use

1. While the precleaned sampler is in the closed position, insert it into the granular or powdered material or waste being sampled from a point near a top edge or corner, through the center, and to a point diagonally opposite the point of entry.
2. Rotate the inner tube of the sampler into the open position.
3. Wiggle the sampler a few times to allow materials to enter the open slots.
4. Place the sampler in the closed position and withdraw from the material being sampled.
5. Place the sampler in a horizontal position with the slots facing upward.
6. Rotate and slide away the outer tube from the inner tube.
7. Transfer the collected sample in the inner tube into a suitable sample container.
8. If composite sampling is desired, collect two or more core samples at different points, and combine the samples in the same container.
9. Check that the Teflon liner is present in the cap if required. Secure the cap tightly. The chemical preservation of solids is generally not recommended. Refrigeration is usually the best approach supplemented by a minimal holding time. Appendix A should be consulted for containerization and preservation requirements.



Source: Reference 4.

Figure 2-6. Grain thief.

10. Label the sample bottle with the appropriate sample tag. Be sure to label the tag carefully and clearly, addressing all the categories or parameters. Complete all chain-of-custody documents and record in the field logbook.
11. Decontaminate equipment after use and between sampling locations using recommended techniques of Appendix E.

Sources

deVera, E.R., Simmons, B.P., Stephens, R.D., and Storm, D.L. "Samplers and Sampling Procedures for Hazardous Waste Streams." EPA-600/2-80-018. January 1980.

Horwitz, W., Sensel, A., Reynolds, H., and Parks, D.L., editors. Animal Feed: Sampling Procedure. In: Official Methods of Analysis. The Association of Official Analytical Chemists. 12th Edition. Washington, D.C. 1979.

2.5 REFERENCES

1. Smith, R. and G. James. The Sampling of Bulk Materials. Analytical Sciences Monograph, Volume 8. The Royal Society of Chemistry, London. 1981.
2. Sisk, S. W. NEIC Manual for Groundwater/Subsurface Investigations at Hazardous Waste Sites. EPA-330/9-81-002. 1981.
3. Scalf, M., J. McNabb, W. Dunlap, R. Crosby, and J. Fryberger. Manual for Groundwater Sampling Procedures. R. S. Kerr Environmental Research Laboratory, Office of Research and Development, Ada, OK. 1980.
4. deVera, E. R., B. P. Simmons, R. D. Stephen, and D. L. Storm. Samplers and Sampling Procedures for Hazardous Waste Streams. EPA-600/2-80-018. January 1980.

SECTION 3.0

LIQUIDS

3.1 GENERAL

Liquids by their nature are a relatively easy substance to collect. Obtaining representative samples, however, is more difficult. Density, volatility, temperature, currents, and a wealth of other mechanisms cause changes in the composition of a liquid with respect to both time and distance. Accurate sampling must be responsive to these dynamics and reflect their actions.

For the purpose of this manual liquids will include both aqueous and nonaqueous solutions and will be subdivided as surface waters, containerized liquids, and ground waters. Surface waters will be considered as any fluid body, flowing or otherwise, whose surface is open to the atmosphere. This will include rivers, streams, discharges, ponds, and impoundments, both aqueous and nonaqueous. The containerized liquid section will address sampling of both sealed and unsealed containers of sizes varying from drums to large tanks. Some overlap may occur between these two sections; when in doubt, both sections should be consulted. The groundwater section will be concerned with obtaining samples from subsurface waters but will not include methods for well construction.

3.2 SURFACE WATERS

The choice of sample locations in surface waters is an important consideration which must be addressed prior to sample acquisition, since it will often effect the selection of sampling equipment. Selection of representative locations will depend on many factors including stream dimensions, shape, flow rate (velocity), inputs and discharges. The USGS publishes the national Handbook of Recommended Methods for Water Data Acquisition which addresses this problem in detail, in addition to several other concepts including flow measurement.

Generally the selection of sample locations will be detailed in the sampling plan, however familiarity with the concepts in the USGS Handbook will assist the samplers in accommodating in-field adjustments. Most often depth integrated and/or cross-sectional composite samples are preferable to single-point grabs. In practice safe access and handling as well as other physical limitations will be influential factors during sample acquisition at hazardous waste contaminated sites.

Samples from shallow depths can be readily collected by merely submerging the sample container. The method is advantageous when the sample might be significantly altered during transfer from a collection vessel into another container. This is the case with samples collected for oil and grease analysis since considerable material may adhere to the sample transfer container and as a result produce inaccurately low analytical results. Similarly the transfer of a liquid into a small sample container for volatile organic analysis, if not done carefully, could result in significant aeration and resultant loss of volatile species. Though simple, representative, and generally free from substantial material disturbances, it has significant shortcomings when applied to a hazardous waste, since the external surface of each container would then need to be decontaminated.

In general the use of a sampling device, either disposable or constructed of a nonreactive material such as glass, stainless steel, or Teflon, is the most prudent method. The device should have a capacity of at least 500 ml, if possible, to minimize the number of times the liquid must be disturbed, thus reducing agitation of any sediment layers.

A 1-liter stainless steel beaker with pour spout and handle works well. It is easily cleaned and considerably less expensive than Teflon. Though still more expensive than other plastics it is more durable and generally more inert under field conditions. Also useful are large stainless steel ice scoops and ladles available from commercial kitchen and laboratory supply houses.

It is often necessary to collect liquid samples at some distance from shore or the edge of the containment. In this instance an adaptation which extends the reach of the technician is advantageous. Such a device is the pond sampler as devised by the California Department of Health.² It incorporates a telescoping heavy-duty aluminum pole with an adjustable beaker clamp attached to the end (see Method III-2). The beaker previously

described, a disposable glass or plastic container, or the actual sample container itself, can be fitted into the clamp. In situations where cross contamination is of concern, use of a disposable container or the actual sample container is always advantageous. The cost of properly cleaning usually outweighs the cost of disposal of otherwise reusable glassware or bottles. This is especially true when the cleanup must be done in the field. The potential contamination of samples for volatile organic analysis by the mere presence of organic solvents necessary for proper field cleaning is usually too great to risk.

Another method of extending the reach of sampling efforts is the use of a small peristaltic pump (see Method III-3). In this method the sample is drawn in through heavy-wall Teflon tubing and pumped directly into the sample container. This system allows the operator to reach out into the liquid body, sample from depth, or sweep the width of narrow streams.

If a medical grade silicone tubing is used in the peristaltic pump, the system is suitable for sampling almost any parameter including most organics.^{3,4} Some volatile stripping, however, may occur, and though the system may have a high flow rate, some material may be lost on the tubing. Therefore, pumping methods should be avoided for sampling volatile organics or oil and grease. Battery-operated pumps of this type are available and can be easily hand-carried or carried with a shoulder sling. It is necessary in most situations to change both the Teflon suction line as well as the silicon pump tubing between sample locations to avoid cross-contamination. This requires maintaining a sufficiently large stock of material to avoid having to clean the tubing in the field.

These tubings are quite expensive but their relatively inert nature makes thorough decontamination in the lab both practical and simple thus allowing reuse. It should be noted that the Teflon suction tubing is an effective substitute for that supplied with the sophisticated automatic liquid waste samplers such as the ISCO Model 2100 and Manning Models S-3000 and S-4040.

When medical grade silicon tubing is not available or the analytical requirements are particularly strict, the system can be altered as described in Method III-3, Figure 3-3. In this configuration the sample volume accumulates in the vacuum flask and does not enter the pump. The integrity of the collection system can now be maintained with only the most nonreactive material contacting the sample. Some loss in lift ability will result since the pump is now moving air, a compressible gas rather than an essentially noncompressible liquid.

It may on occasion be necessary to sample large bodies of water where a near surface sample will not sufficiently characterize the body as a whole. In this instance again the above-mentioned pump is quite serviceable. It is capable of lifting water from depths in excess of 6 meters. Since the lift capacity is actually measured as the distance above the hydrostatic surface, it is possible to withdraw samples from depths significantly below the water surface. It should be noted that this lift ability decreases somewhat with higher density fluids and with increased wear on the silicone pump tubing.

Similarly increases in altitude will decrease the pumps ability to lift from depth. When sampling a liquid stream which exhibits a considerable flow rate, it may be necessary to weight the bottom of the suction line. The stainless steel strainer suction weight supplied with the ISCO and Manning samplers usually works well. A heavier weight can be constructed by filling a short (7.5 cm to 10 cm) length of Teflon tubing with lead and plugging both ends with tight-fitting Teflon plugs. This weight can then be clamped with stainless steel band clamps to the suction tubing.

Situations may still arise where a sample must be collected from depths beyond the capabilities of a peristaltic pump. In this instance an at-depth sampler may be required, such as a Kemmerer, ASTM Bomb (Bacon Bomb) or Van Dorn sampler. These devices work well; however, care must be utilized in selecting devices that are made of materials that will not contaminate the sample. Van Dorn samplers are not generally recommended for organics as they rely on an elastic closing mechanism that can effect samples. They are readily available in a totally nonmetallic design which is very useful for sample collection for trace metal analysis.

Kemmerer samplers are available on special order or adaptable for sample collection for organic analysis by substituting Teflon for the rubber or plastic stoppers. If the device is further ordered with stainless steel metallic parts in addition to Teflon stoppers it becomes a very versatile sampler.

The submersible pumps discussed in conjunction with groundwater sampling (Section 3.4) may also be useful in this application.

3.2.1 METHOD III-1: SAMPLING SURFACE WATERS USING A DIPPER OR OTHER TRANSFER DEVICE

Discussion

A dipper or other container constructed of inert material, such as stainless steel or Teflon, can be used to transfer liquid wastes from their source to a sample bottle. This prevents unnecessary contamination of the outer surface of the sample bottle that would otherwise result from direct immersion in the liquid. Use of this device also prevents the technician from having to physically contact the waste stream. Depending upon the sampling application, the transfer vessel can be either disposed of or reused. If reused, the vessel should be thoroughly rinsed and/or decontaminated prior to sampling a different source.

Uses

A transfer device can be utilized in most sampling situations except where aeration must be eliminated (samples for volatile organic analysis) or where significant material may be lost due to adhesion to the transfer container.

Procedures for Use

1. Submerge a precleaned stainless steel dipper or other suitable device with minimal surface disturbance.
2. Allow the device to fill slowly and continuously.
3. Retrieve the dipper/device from the surface water with minimal disturbance.
4. Remove the cap from the sample bottle and slightly tilt the mouth of the bottle below the dipper/device edge.
5. Empty the dipper/device slowly, allowing the sample stream to flow gently down the side of the bottle with minimal entry turbulence.
6. Continue delivery of the sample until the bottle is almost completely filled. Leave adequate ullage to allow for expansion.
7. Select appropriate bottles and preserve the sample if necessary as per guidelines in Appendix A.
8. Check that a Teflon liner is present in the cap if required. Secure the cap tightly.
9. Label the sample bottle with an appropriate sample tag. Be sure to label the tag carefully and clearly, addressing all the categories or parameters. Record the information in the field logbook and complete the chain-of-custody form.

10. Properly clean and decontaminate the equipment prior to reuse or storage (Appendix E).

Sources

GCA Corporation, "Quality Assurance Plan, Love Canal Study - Appendix A, Sampling Procedures," EPA Contract 68-02-3168.

3.2.2 METHOD III-2: USE OF POND SAMPLER FOR THE COLLECTION OF SURFACE WATER SAMPLES

Discussion

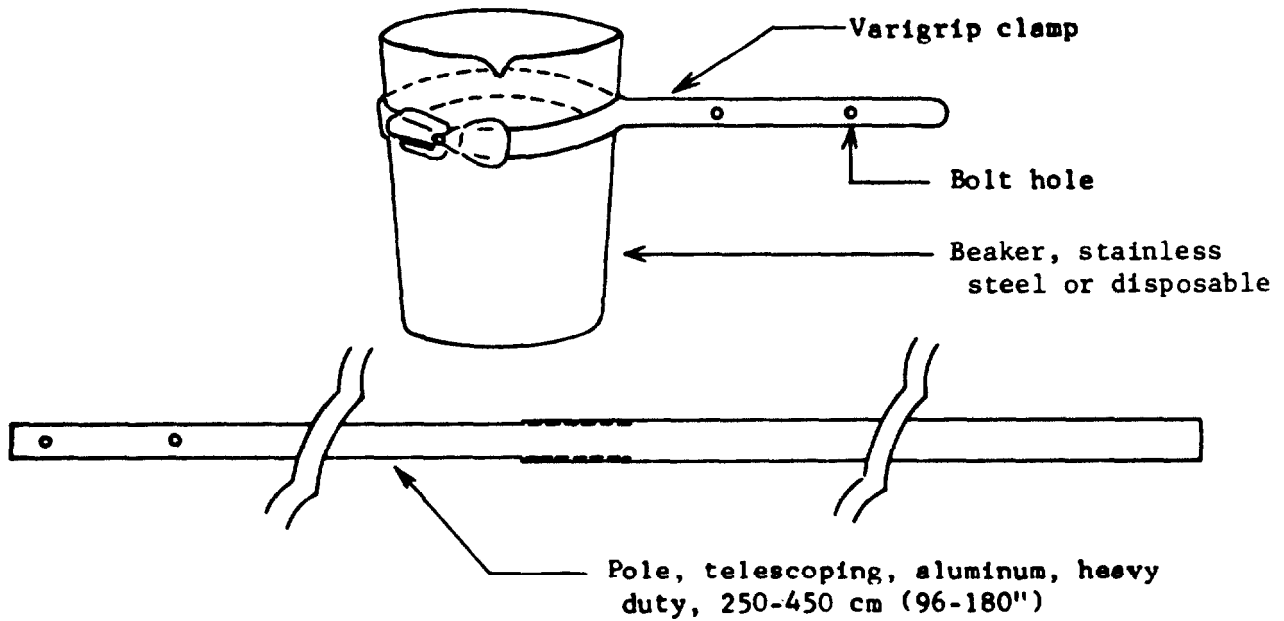
The pond sampler consists of an adjustable clamp attached to the end of a two- or three-piece telescoping aluminum tube that serves as the handle. The clamp is used to secure a sampling beaker (see Figure 3-1). The sampler is not commercially available, but it is easily and inexpensively fabricated. The tubes can be readily purchased from most hardware or swimming pool supply stores. The adjustable clamp and sampling beaker can be obtained from most laboratory supply houses. The materials required to fabricate the sampler are given in Appendix B.

Uses

The pond sampler is used to collect liquid waste samples from disposal ponds, pits, lagoons, and similar reservoirs. Grab samples can be obtained at distances as far as 3.5 m from the edge of the ponds. The tubular aluminum handle may bow when sampling very viscous liquids if sampling is not done slowly.

Procedures for Use

1. Assemble the pond sampler. Make sure that the sampling beaker and the bolts and-nuts that secure the clamp to the pole are tightened properly.
2. With proper protective garment and gear, take grab samples by slowly submerging the precleaned beaker with minimal surface disturbance.
3. Retrieve the pond sampler from the surface water with minimal disturbance.
4. Remove the cap from the sample bottle and slightly tilt the mouth of the bottle below the dipper/device edge.
5. Empty the sampler slowly, allowing the sample stream to flow gently down the side of the bottle with minimal entry turbulence.
6. Continue delivery of the sample until the bottle is almost completely filled.
7. Select appropriate sample bottles and preserve the sample if necessary as per guidelines in Appendix A.
8. Check that a Teflon liner is present in the cap if required. Secure the cap tightly.



Source: Reference 1.

Figure 3-1. Pond sampler.

9. Label the sample bottle with an appropriate sample tag. Be sure to label the tag carefully and clearly, addressing all the categories or parameters. Record the information in the field logbook and complete the chain-of-custody documents.
10. Properly clean and decontaminate the equipment prior to reuse or storage using recommended guidelines of Appendix E.

Sources

deVera, E.R., Simmons, B.P., Stephens, R.D., and Storm, D.L. "Samplers and Sampling Procedures for Hazardous Waste Streams," EPA-600/2-80-018, January 1980.

GCA Corporation, "Quality Assurance Plan, Love Canal Study - Appendix A, Sampling Procedures," EPA Contract 68-02-3168.

3.2.3 METHOD III-3: PERISTALTIC PUMP FOR SAMPLING SURFACE WATER BODIES

Discussion

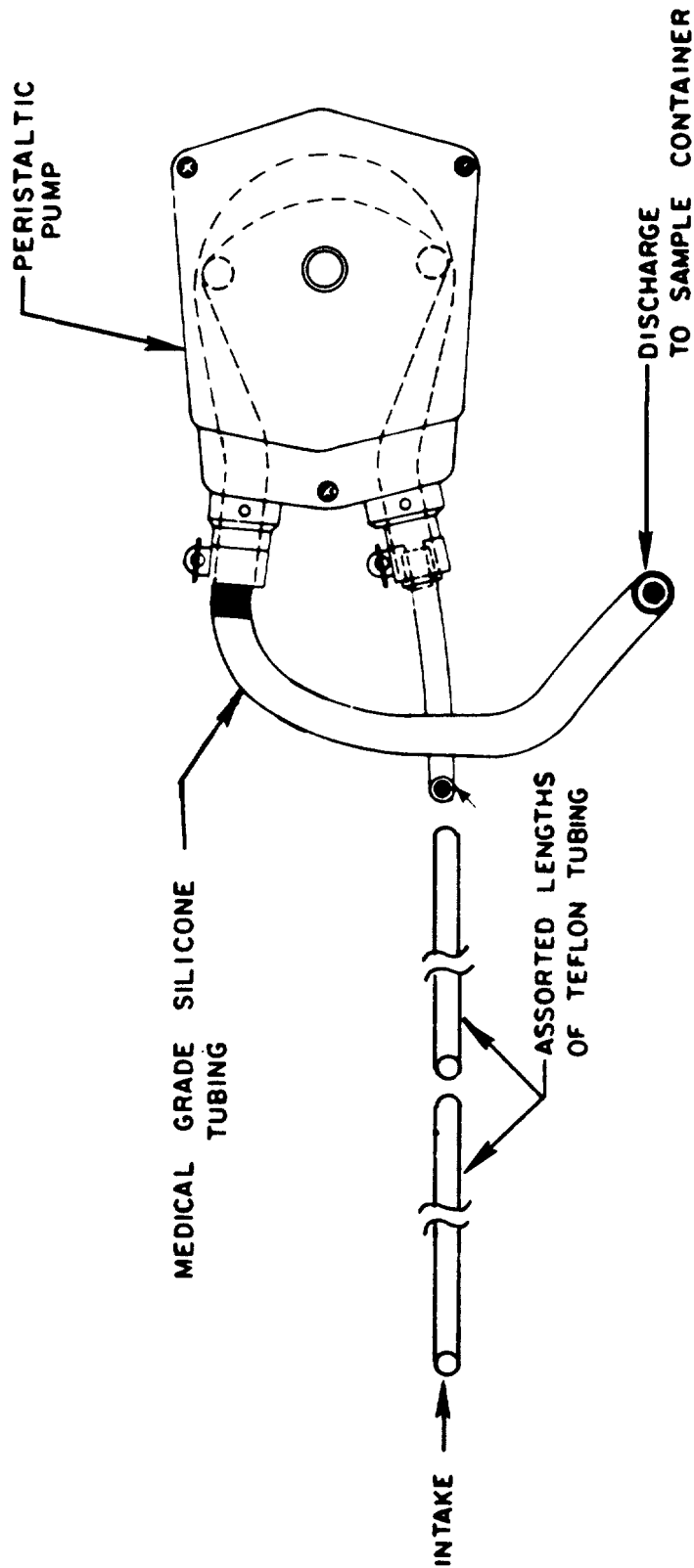
This collection system consists of a peristaltic pump capable of achieving a pump rate of 1 to 3 lpm, and an assortment of Teflon tubing for extending the suction intake. A battery operated pump is preferable as it eliminates the need for DC generators or AC inverters.

Uses

The system, as shown in Figures 3-2 and 3-3, is highly versatile. It is portable and the sample collection is conducted through essentially chemically nonreactive material. It is practical for a wide range of applications including streams, ponds, and containers. This procedure can both extend the lateral reach of the sampler and allow sampling from depth. Likewise, it can function both as a well purge and a sample collection system. The chief disadvantage of this method is the limited lift capacity of the pump, approximately 8 meters.

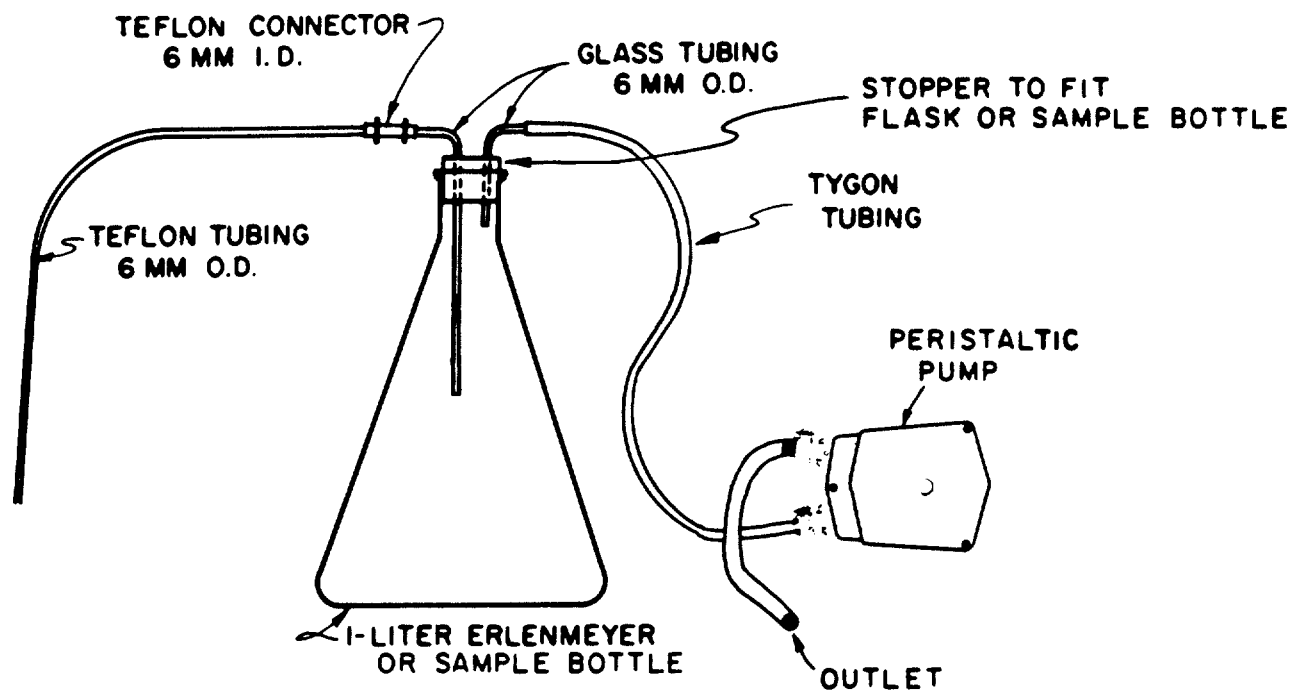
Procedures for Use

1. Install clean, medical-grade silicone tubing in the pump head, as per the manufacturer's instructions. Allow sufficient tubing on discharge side to facilitate convenient dispensation of liquid into sample bottles and only enough on the suction end for attachment to the intake line. This practice will minimize sample contact with the silicone pump tubing.
2. Select the length of suction intake tubing necessary to reach the required sample depth and attach to intake side of pump tubing. Heavy-wall Teflon, of a diameter equal to the required pump tubing, suits most applications. (Heavier wall will allow for a slightly greater lateral reach.)
3. If possible, allow several liters of sample to pass through system, before actual sample collection. Collect this purge volume and then return to source after the sample aliquot has been withdrawn.
4. Fill necessary sample bottles by allowing pump discharge to flow gently down the side of bottle with minimal entry turbulence. Cap each bottle as filled.
5. Select appropriate bottles and preserve the sample if necessary as per guidelines in Appendix A.
6. Check that a Teflon liner is present in the cap if required. Secure the cap tightly.



Source: Reference 2.

Figure 3-2. Peristaltic pump for liquid sampling.



Source Reference 4.

Figure 3-3. Peristaltic pump for liquid sampling (modified).

7. Label the sample bottle with an appropriate tag. Be sure to complete the tag with all necessary information. Record the information in the field logbook and complete the chain-of-custody documents.
8. Allow system to drain, then disassemble. Return tubing to lab for decontamination (if feasible). See Appendix E for general decontamination procedures.

Sources

U.S. Environmental Protection Agency. "Procedures Manual for Ground Water Monitoring at Solid Waste Disposal Facilities." EPA-530/SW-611. August 1977.

3.2.4 METHOD III-4: COLLECTION OF WATER SAMPLES FROM DEPTH WITH A KEMMERER BOTTLE

Discussion

The Kemmerer bottle is a messenger-activated water sampling device (see Figure 3-4). In the open position water flows easily through the device. Once lowered to the desired depth a messenger is dropped down the sample line tripping the release mechanism and closing the bottle. In the closed position the bottle is sealed, both on top and bottom, from any additional contact with the water column and can be retrieved.

Most commercially available Kemmerer bottles are of brass or plastic construction. Modification of existing systems with nonreactive materials such as Teflon, glass or stainless steel would be only partially successful due to the complicated machining necessary for the release mechanism. Other modifications such as a stoppered bottom drain are simpler and useful in minimizing sample disturbance during transfer to the appropriate containers.

Uses

The Kemmerer bottle is currently the most practical method of collecting discrete, at-depth samples from surface waters or vessels where the collection depth exceeds the lift capacity of pumps. The application is limited however by the incomparability of various construction materials with some analytical techniques. Proper selection, i.e., all metal assemblies for organic analysis or all plastic assemblies for trace element analysis, will overcome this deficiency.

Procedures for Use

1. Inspect Kemmerer bottle for thorough cleaning and insure that sample drain valve is closed (if bottle is so equipped).
2. Measure and then mark sample line at desired sampling depth.
3. Open bottle by lifting top stopper-trip head assembly.
4. Gradually lower bottle until desired level is reached (predesignated mark from Step 2).
5. Place messenger on sample line and release.
6. Retrieve sampler; hold sampler by center stem to prevent accidental opening of bottom stopper.
7. Rinse or wipe off exterior of sampler body (wear proper gloves and protective clothing).

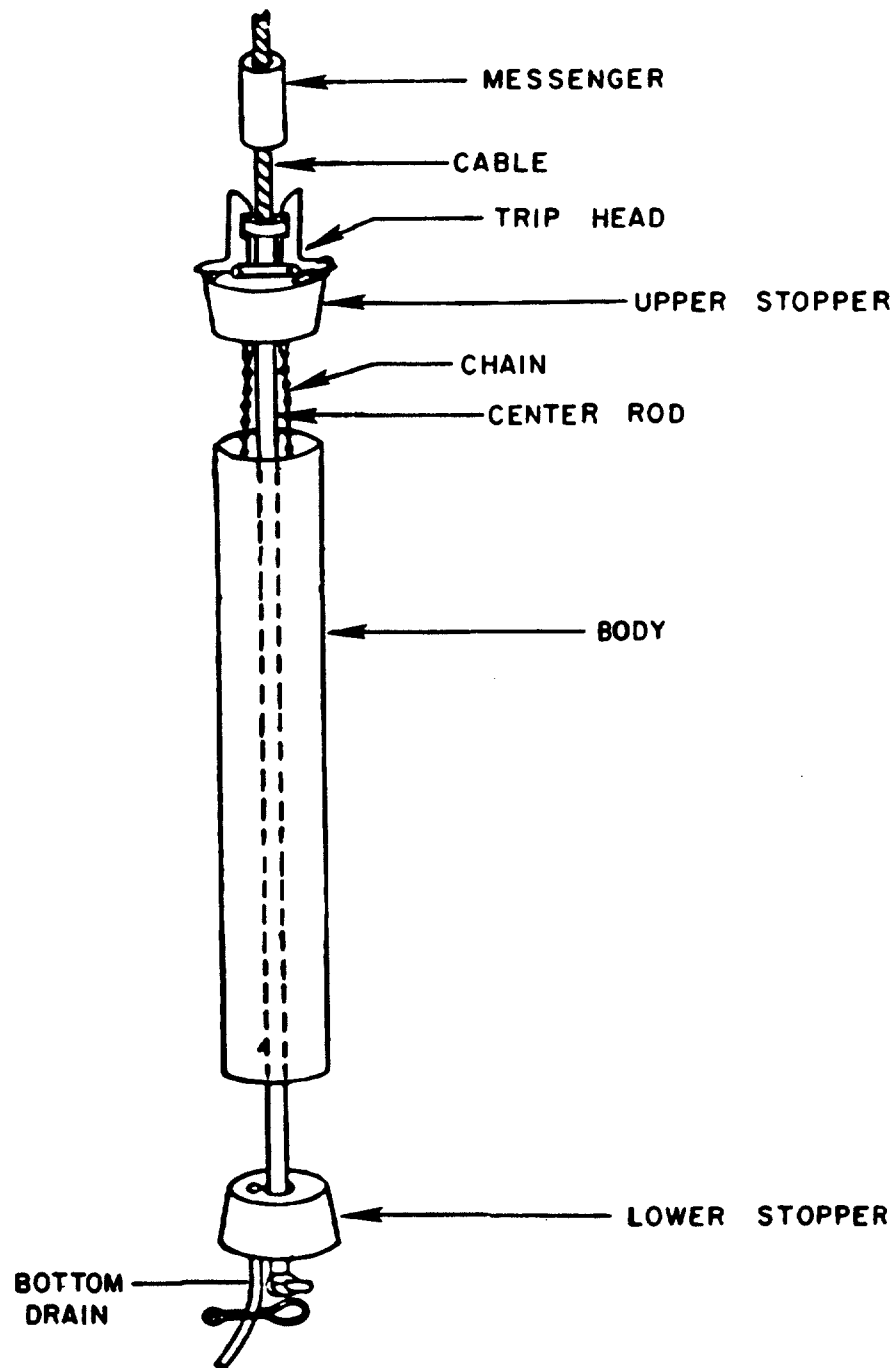


Figure 3-4. Modified Kemmerer sampler.

8. Recover sample by grasping lower stopper and sampler body with one hand (gloved), and transfer sample by either (a) lifting top stopper with other hand and carefully pouring contents into sample bottles, or (b) holding drain valve (if present) over sample bottle and opening valve.
9. Allow sample to flow slowly down side of sample bottle with minimal disturbance.
10. Select sample bottles and preserve the sample if necessary as per guidelines in Appendix A.
11. Check that a Teflon liner is present in the cap if required. Secure the cap tightly.
12. Label the sample bottle with an appropriate tag. Be sure to complete the tag with all necessary information. Record the information in the field logbook and complete all chain-of-custody records.
13. Decontaminate sampler and messenger or place in plastic bag for return to lab. See Appendix E for general decontamination procedures.

Sources

U.S. Environmental Protection Agency, "Procedures Manual for Ground Water Monitoring at Solid Waste Disposal Facilities." EPA-530/SW-611, August 1977.

3.3 CONTAINERIZED LIQUIDS

The sampling of tanks, containers, and drums present unique problems not associated with natural water bodies. Containers of this sort are generally closed except for small access ports, manways, or hatches on the larger vessels or taps and bungs on smaller drums. The physical size, shape, construction material, and location of access will limit the types of equipment and methods of collection.

When liquids are contained in sealed vessels, gas vapor pressures build up, sludges settle out, and density layerings develop. The potential for explosive reactions or the release of noxious gases when containers are opened requires considerable safeguards. The vessels should be opened with extreme caution. Preliminary sampling of any headspace gases may be warranted. Section 4.4 details procedures for sampling headspace gases. As a minimum, a preliminary check with an organic vapor analyzer may help determine needed levels of personnel protection and may be of aid in selecting a sampling method.

In most cases it is impossible to observe the contents of these sealed or partially sealed vessels. Since some layering or stratification is likely in any solution left undisturbed over time, a sample must be taken that represents the entire depth of the vessel.

Agitation to disrupt the layers and rehomogenize the sample is physically difficult and almost always undesirable. In vessels greater than 1 meter in depth the method of choice is to slowly, in known increments of length, lower the suction line from a peristaltic pump. Discrete samples can be collected from various depths then combined or analyzed separately. If the depth of the vessel is greater than the lift capacity of the pump, an at-depth water sampler, such as the Kemmerer type discussed in Method III-4, or the ASTM Bomb (Bacon Bomb) may be required. In situations where the reactive nature of the contents are known, a small submersible pump may be used.

When sampling a previously sealed vessel, a check should be made for the presence of a bottom sludge. This is easily accomplished by measuring the depth to apparent bottom then comparing it to the known interior depth. Methods for sampling a bottom sludge are found in Section 2.3.

The sampling of drums for hazardous liquid wastes is a very taxing situation with present equipment. The most widely used method is a glass tube, 6 mm to 16 mm I.D., that is lowered into the drum. The top of the tube is sealed with a stopper or the thumb and the tube withdrawn. The bottom of the tube is then placed over a glass jar, the stopper removed from the top and the contents drained into the containers. After collection of sufficient sample the tube is then broken up into the drum. This method is simple, relatively inexpensive, and quick and collects a sample without having to decontaminate equipment. It does, however, have serious drawbacks. Most low density fluids do not hold well in the glass tubes. A great deal of the potential sample flows out of the bottom of the tube as it is raised from the drum, thereby reducing the representativeness of collected material. Many

3.3.1 METHOD III-5: COLLECTION OF LIQUID CONTAINERIZED WASTES USING GLASS TUBES

Description

Liquid samples from opened containers (55-gallon drums) are collected using lengths of glass tubing. The glass tubes are normally 122 cm in length and 6 to 16 mm inside diameter. Larger diameter tubes may be used for more viscous fluids if sampling with the small diameter tube is not adequate. The tubing is broken up and discarded in the container after the sample has been collected, eliminating difficult cleanup and disposal problems. This method should not be attempted with less than a two-man sampling team.

Uses

This method provides for a quick, relatively inexpensive means of collecting concentrated containerized wastes. The major disadvantage is from potential sample loss which is especially prevalent when sampling less viscous fluids. Splashing can also be a problem and proper protective clothing (e.g., butyl rubber apron, face shields, boot covers) should always be worn.

Procedures for Use

1. Remove cover from sample container opening.
2. Insert glass tubing slowly to almost the bottom of the container. Try to keep at least 30 cm of tubing above the top of the container.
3. Allow the waste in the drum to reach its natural level in the tube.
4. Cap the top of the tube with a safety-gloved thumb or a rubber stopper.
5. Carefully remove the capped tube from the drum and insert the uncapped end in the sample container.
6. Release the thumb or stopper on the tube and allow the sample container to fill to approximately 90 percent of its capacity.
7. Repeat steps 2 through 6 if more volume is needed to fill the sample container.
8. Remove the tube from the sample container and replace the tube in the drum.
9. Cap the sample container tightly with a Teflon-lined cap and affix the sample identification tag.
10. Break the glass sampling tube in such a way that all parts of it are discarded inside the drum. (Note: see the initial discussion to this section for exceptions.)

variations to this technique have been reported. These include the incorporation of a small suction device (i.e., pipette bulb) to the top of the tube as well as the use of various tube sizes. Some success has been reported with tubes that have been heated at one end then drawn to form a much smaller orifice. This allows the use of larger diameter tubing, therefore a greater volume of sample per attempt, while reducing the material loss from the tube bottom normally associated with larger diameter tubes.

It should be noted that in some instances disposal of the tube by breaking it into the drum may interfere with eventual plans for the removal of its contents. The use of this technique should therefore be cleared with the project officer, or other disposal techniques evaluated.

In many instances a drum containing waste material will have a sludge layer on the bottom (Method III-5). Slow insertion of the sample tube down into this layer and then a gradual withdrawal will allow the sludge to act as a bottom plug to maintain the fluid in the tube. The plug can be gently removed and placed into the sample container by the use of a stainless steel lab spoon. These spoons are relatively inexpensive and can be disposed of in the original waste container with the glass transfer tube.

Designs exist for equipment that will collect a sample from the full depth of a drum and maintain it in the transfer tube until delivery to the sample bottle. These designs include primarily the Composite Liquid Waste Sampler (COLIWASA) and modifications thereof.² The COLIWASA is difficult to properly decontaminate in the field; its applicability is therefore limited to those cases when a sample of the full depth of the drum is absolutely necessary. The COLIWASA can be somewhat modified for this task by making the lift rod of stainless steel, the bottom stopper of Teflon, and the body of glass tubing. In this configuration the glass tube can be broken into the drum leaving only the center rod and the stopper to be decontaminated. In a preliminary investigation where the total number of drums to be sampled is small an equal number of both the center rods and bottom stoppers could be made in advance thus eliminating the time involved for onsite cleanup. Heat shrinkable Teflon tubing or other types of Teflon coating can also be used to cover the stainless steel rod if contact of the stainless steel with the waste is undesirable.

3. When a solid is encountered in a drum (either layer or bottom sludge) the optional method described above may be used to collect a core of the material, or the material may be collected with a disposable scoop attached to a length of wooden or plastic rod.
4. If analysis is to be performed onsite, packing steps 12 and 13 may be deleted. These steps are necessary for transporting and/or shipping samples.

Sources

American Society for Testing and Materials. "Standard Recommended Practices for Sampling Industrial Chemicals," ASTM E-300-73.

U.S. Environmental Protection Agency, "Technical Methods for Investigating Sites Containing Hazardous Substances, Technical Monograph 1-29, Draft," Ecology and the Environment, June 1981.

11. Replace the bung or place plastic over the drum.
12. Place sample container in a Ziplock plastic bag (one per bag).
13. Place each bagged container in a 1-gallon metal paint can (or appropriate sized container) and pack in vermiculite packing material. Place lid on the can.
14. Mark the sample identification number on the outside of each paint can and complete chain-of-custody log and the field logbook.

Optional Method (if sample of bottom sludge is desired)

1. Remove cover from container opening.
2. Insert glass tubing slowly almost to the bottom of the container. Try to keep at least 30 cm of tubing above the top of the container.
3. Allow the waste in the drum to reach its natural level in the tube.
4. Gently push the tube towards the bottom of the drum into the sludge layer. Do not force it.
5. Cap the top of the tube with a safely-gloved thumb or rubber stopper.
6. Carefully remove the capped tube from the drum and insert the uncapped end in the sample container.
7. Release the thumb or stopper on the tube and allow the sample container to fill to approximately 90 percent of its capacity. If necessary, the sludge plug in the bottom of the tube can be dislodged with the aid of a stainless steel laboratory spatula.
8. Repeat if more volume is needed to fill sample container and recap the tube.
9. Proceed as in Steps 9 through 14 above.

Note:

1. If a reaction is observed when the glass tube is inserted (violent agitation, smoke, light, heat, etc.) the investigator should leave the area immediately.
2. If the glass tube becomes cloudy or smokey after insertion into the drum, the presence of hydrofluoric acid is indicated and a comparable length of rigid plastic tubing should be used to collect the sample.

3.3.2 METHOD III-6: SAMPLING CONTAINERIZED WASTES USING THE COMPOSITE LIQUID WASTE SAMPLER (COLIWASA)

Discussion

The COLIWASA is a much cited sampler designed to permit representative sampling of multiphase wastes from drums and other containerized wastes. The sampler is commercially available or can be easily fabricated from a variety of materials including PVC, glass, or Teflon. In its usual configuration it consists of a 152 cm by 4 cm (inside diameter) section of tubing with a neoprene stopper at one end attached by a rod running the length of the tube to a locking mechanism at the other end. Manipulation of the locking mechanism opens and closes the sampler by raising and lowering the neoprene stopper. A current recommended model of the COLIWASA is shown in Figure 3-5; however, the design can be modified and/or adapted somewhat to meet the needs of the sampler.

Uses

The COLIWASA is primarily used to sample most containerized liquids. The plastic COLIWASA is reported to be able to sample most containerized liquid wastes except for those containing ketones, nitrobenzene, dimethylformamide, mesityl oxide and tetrahydrofuran. A glass COLIWASA is able to handle all wastes unable to be sampled with the plastic unit except strong alkali and hydrofluoric acid solution. Due to the unknown nature of most containerized waste, it would therefore be advisable to eliminate the use of PVC materials and use samplers composed of glass or Teflon.

The major drawbacks associated with using a COLIWASA concern decontamination and costs. The sampler is difficult if not impossible to decontaminate in the field and its high cost in relation to alternative procedures (glass tubes) make it an impractical throwaway item. It still has applications, however, especially in instances where a true representation of a multiphase waste is absolutely necessary. For this reason, the procedure for its use is included.

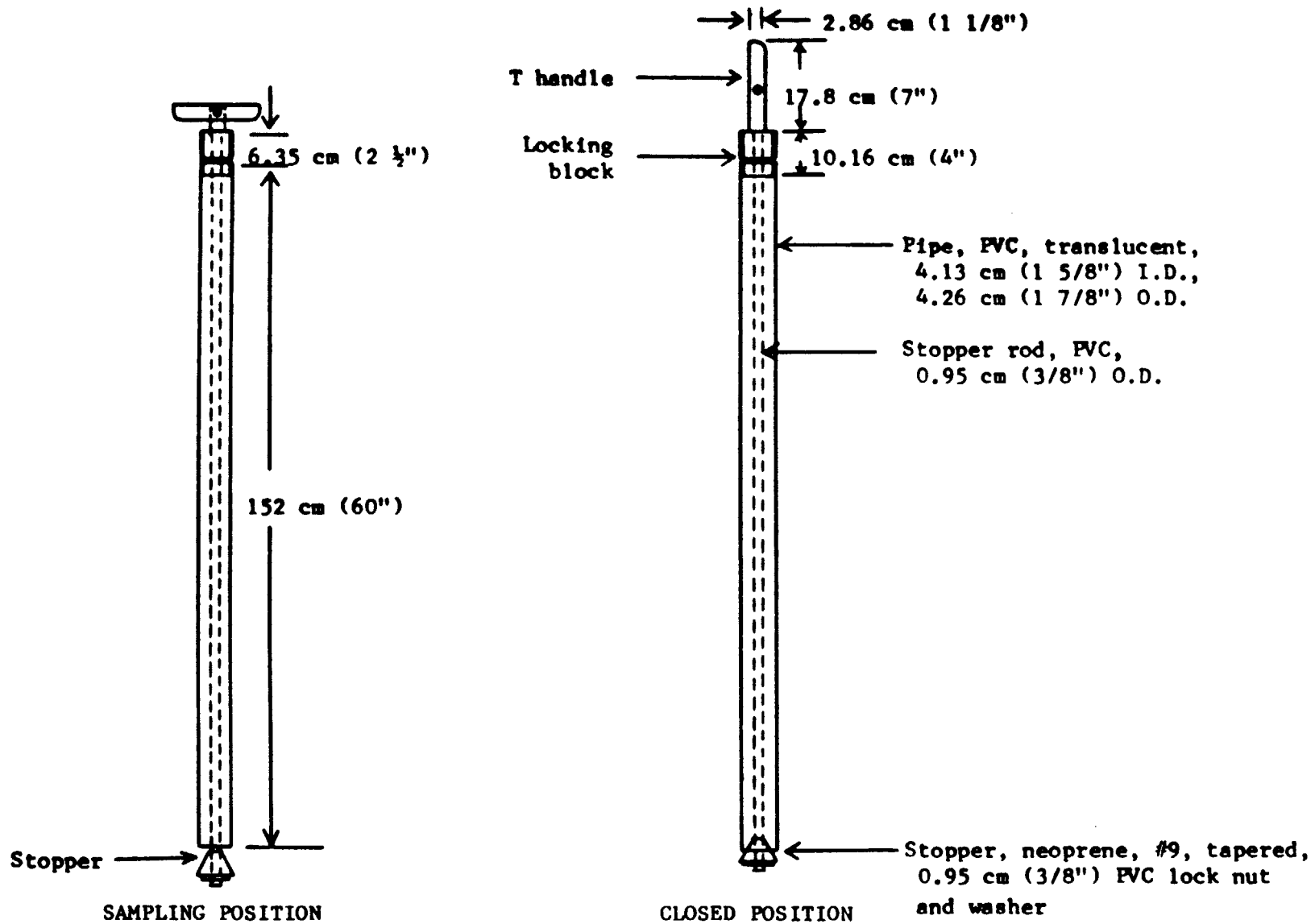
Procedures for Use

1. Choose the material (see Appendix B) to be used to fabricate the COLIWASA and assemble the sampler as shown in Figure 3-5.
2. Make sure that the sampler is clean.
3. Check to make sure the sampler is functioning properly. Adjust the locking mechanism if necessary to make sure the neoprene rubber stopper provides a tight closure.
4. Wear necessary protective clothing and gear and observe required sampling precautions.

5. Put the sampler in the open position by placing the stopper rod handle in the T-position and pushing the rod down until the handle sits against the sampler's locking block.
6. Slowly lower the sampler into the liquid waste. (Lower the sampler at a rate that permits the levels of the liquid inside and outside the sampler tube to be about the same. If the level of the liquid in the sample tube is lower than that outside the sampler, the sampling rate is too fast and will result in a nonrepresentative sample).
7. When the sampler stopper hits the bottom of the waste container, push the sampler tube downward against the stopper to close the sampler. Lock the sampler in the closed position by turning the T handle until it is upright and one end rests tightly on the locking block.
8. Slowly withdraw the sampler from the waste container with one hand while wiping the sampler tube with a disposable cloth or rag with the other hand.
9. Carefully discharge the sample into a suitable sample container by slowly pulling the lower end of the T handle away from the locking block while the lower end of the sampler is positioned in a sample container.
10. Cap the sample container with a Teflon-lined cap; attach label and seal; record in field logbook; and complete sample analysis request sheet and chain-of-custody record.
11. Unscrew the T handle of the sampler and disengage the locking block. Clean sampler onsite or store the contaminated parts of the sampler in a plastic storage tube for subsequent cleaning. Store used rags in plastic bags for subsequent disposal. See Appendix E for general decontamination procedures.

Sources

deVera, E. R., Simmons, B. P., Stephens, R. D., and Storm, D. L. "Samplers and Sampling Procedures for Hazardous Waste Streams." EPA 600/2-80-019, January 1980.



Source: Reference 6.

Figure 3-5. Composite liquid waste sampler (Coliwsa).

3.4 GROUNDWATER

Groundwater sampling suffers from many of the same difficulties as closed containers, such as the inability to observe what is being collected or what disturbances are resulting from that collection.

There are essentially two sources from which to collect groundwater, either from wells or from seeps and springs. The former is more complex and a discussion of its intricacies will follow later. The sampling of seeps and springs is considerably easier, but it may be less indicative of the actual groundwater quality than well sampling.

Seeps and springs are generally areas where the surface contour intersects the water table. These areas usually have well established microbiological populations evidenced by extensive moss and algal growths. These microbiological populations usually extend for some distance into the water-bearing formation (aquifer) and are generally more populous and of different species than those associated with the bulk of the aquifer. Their effect on the oxygen content, pH, nutrient and metals concentrations in the groundwater can be extensive. The water, therefore, that seeps from these areas may be substantially altered, and not representative of the conditions deeper in the subsurface. They can, however, yield some information if properly interpreted. If the area in question is without developed wells they are certainly worth consideration, especially for the ease with which they can be sampled.

A stainless steel scoop of the type found in ice machines is ideal for collecting samples from seeps. The flat bottom can be pressed against the bank and the water will flow with very little additional disturbance into the scoop, for transfer to the sample bottles. It is important to collect the sample as close to the actual seep as possible to reduce contact time with the atmosphere and potential for surface contamination.

For the purposes of this document, groundwater monitoring via wells will include only the actual sampling of existing wells. The methods and techniques for placement, construction, and development of wells for groundwater monitoring are varied and complicated. The "Manual for Ground-Water Sampling Procedures"⁶ and "NEIC Manual for Groundwater/Subsurface Investigations at Hazardous Waste Sites"⁷ provide considerable information for establishing a full groundwater monitoring program including the completion of monitor wells. It is, however, necessary to know the well depth, diameter, construction material, type and size of the well screen if used, vertical position of the well screen or slotted section of casing, and type of annular packing if any. This information will aid in evaluating the suitability of the well for sampling for a particular analysis. For instance, if the well has a galvanized steel casing with a brass well screen, it would not be suitable for trace element analysis. Similarly, if the well is located in a swampy area, the type and amount of grout or fill around the well casing would determine the degree of surface water inflow to the well that might be expected. Most of the information necessary is available on the well drillers log. An example of a completed drillers log is included as Figure 3-6. It should be noted,

OBSERVATION WELL CONSTRUCTION SUMMARY

PROJECT _____		WELL NO. <u>PB137</u>	
SITE _____		AQUIFER <u>Glacial Drift</u>	
COORDINATES _____		DATE COMPLETED <u>10/27/82</u>	
SUPERVISED BY <u>P. Huidobro</u>		_____	

<p>GROUND ELEVATION 909.14</p> <p>Topsol</p> <p>Peat</p> <p>Sand</p> <p>Peat</p> <p>Alternating layers of gray till and outwash</p> <p>Outwash</p> <p>Gray till</p> <p>Medium sand with pebbly gravel</p> <p>Limestone bedrock</p> <p>GENERALIZED STRATIGRAPHY (Assumed to be similar to PB136)</p>		<p>Elevation of reference point <u>910.9</u></p> <p>Height of reference point above ground surface <u>1.76</u></p> <p>Depth of surface seal <u>1'</u></p> <p>Type of surface seal: <u>concrete</u></p> <p>I.D. of surface casing <u>4"</u></p> <p>Type of surface casing: <u>steel with vented locking cap</u></p> <p>Depth of surface casing <u>5.24'</u></p> <p>I.D. of riser pipe <u>2"</u></p> <p>Type of riser pipe: <u>galvanized steel</u></p> <p>Diameter of borehole <u>4"</u></p> <p>Type of filler: <u>1/1 cement/bentonite</u></p> <p>Elevation / depth of top of seal <u>834.54/74.6'</u></p> <p>Type of seal: <u>1/1 cement/bentonite</u></p> <p>Type of gravel pack <u>silica sand No. 20</u></p> <p>Elev./depth of top of gravel pack <u>833.54'/75.6'</u></p> <p>Elevation / depth of top of screen <u>826.84'/82.3'</u></p> <p>Description of screen <u>No. 10 slot welded galvanized steel</u></p> <p>I.D. of screen section _____</p> <p>Elevation / depth of bottom of screen <u>822.64'/86.5'</u></p> <p>Elev./depth of bottom of gravel pack <u>822.64'/86.5'</u></p> <p>Elev./depth of bottom of plugged blank section <u>822.64'/86.5'</u></p> <p>Type of filler below plugged section _____</p> <p>Elevation of bottom of borehole <u>822.64'/86.5'</u></p>
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Form 1002

Figure 3-6. Sample well construction form.

however, that the actual well depth may be somewhat less than the completion depth listed on the log as a result of aquifer invasion through the screen or open-hole sloughing below the casing. This may be particularly noticable in wells that have had only sporadic use or have been idle long. It is recommended then that actual well depth be checked by field measurement whenever possible.

Measurement of the well depth can be accomplished by sounding the well with a reusable weight attached to a disposable line. Slowly lower the weight into the well until the bottom is detected. With the line taut, mark the top of casing level on the line with waterproof ink. Recover the line and weight from the well and accurately measure the length of line below the mark. Discard the line and thoroughly clean the weight before reuse. Next, measure the casing length above (or below) ground level and subtract (or add) to obtain well depth.⁶ When measuring potentially contaminated wells, wear appropriate safety gear to avoid skin contact with well water.

The depth to the water level in the well must be measured in order to calculate the liquid bore volume for prepurging and is also important to any hydrological interpretations of the analytical results. Depths to water are normally measured with respect to the top of casing, as in well-depth determinations. Several methods are available including: (1) the electric sounder, (2) the chalked steel tape, and (3) the popper.⁶

The electric sounder, although not the most accurate, is recommended for initial site work because of the minimal potential for equipment contamination and simplicity of use. Sounders usually consist of a conductivity cell at the end of a graduated wire, and a battery powered buzzer. When the cell contacts the water the increased conductivity completes the circuit and allows current to flow to the alarm buzzer. The depth to water can then be read from the graduations on the wire or the wire can be measured directly. This device may not be suitable for use if a potentially flammable or explosive layer is present in the well, unless it is an intrinsically safe version. A discussion of electrical product certification is presented in Appendix F.

The chalked steel tape is a more accurate device for measuring static water levels. Coat the lower 0.5 to 1.0 meters of a steel measuring tape on either side with either carpenter's chalk or any of the various indicating pastes. Attach a weight to the lower end to keep the tape taut and lower it into the center of the well (condensate on the casing wall may prematurely wet the tape). Listen for a hallow "popping" sound when the weight reaches water. Then lower the tape very slowly for at least another 15 cm, preferably to an even increment. Next, carefully withdraw the tape from the well; determine water depth by subtracting the wetted length of tape from the total length of tape in the well. In small-diameter wells, the volume of the weight may cause the water to rise by displacement. In general the use of indicating paste or chalk should be discouraged although they may not present a significant problem if water samples are not collected. As with all depth measurement devices, thoroughly clean the wetted section of the tape and the weight before reuse to avoid cross contamination.

The metal tape and popper is another simple and reliable method for measuring depth to water in wells more than 3.8 cm (1.5 in.) in diameter. The popper is a metal cylinder with a concave undersurface fastened to the end of the metal tape. Raise and drop the popper until it hits the water surface and makes a distinct "popping" sound. Adjust the tape length so that the popper just hits the water surface. Read the depth to water from the tape measure.

To obtain a representative sample of the groundwater it must be understood that the composition of the water within the well casing and in close proximity to the well is probably not representative of the overall groundwater quality at that sampling site. This is due to the possible presence of drilling contaminants near the well and because important environmental conditions such as the oxidation-reduction potential may differ drastically near the well from the conditions in the surrounding water-bearing materials. For these reasons it is highly desirable that a well be pumped or bailed until the well is thoroughly flushed of standing water and contains fresh water from the aquifer. The recommended amount of purging before sampling is dependent on many factors including the characteristics of the well, the hydrogeological nature of the aquifer, the type of sampling equipment being used, and the parameters being sampled. A common procedure is to pump or bail the well until a minimum of two (2) to ten (10) bore-volumes have been removed.

Gibb⁸ notes that removing all water from the well bore is only possible if the well is pumped dry and suggests two alternative approaches: (a) monitor the water level in the well while pumping. When the water level has "stabilized" most if not all of the water being pumped is coming from the aquifer. (b) monitor the temperature, conductivity, or pH of the water while pumping. When these parameters "stabilize" it is probable that little or no water from casing storage is being pumped.⁵

The use of an indicating analysis such as pH, temperature, redox potential, or, most commonly, conductivity, may be the most accurate and reliable method of assuring complete well purging and it also reduces the likelihood of over purging. The technique is easily implemented in the field and gives a rapid and positive indication of changes in the well bore water. This change in the water character and subsequent stabilization can normally be interpreted as evidence that sufficient purging has occurred. It should be noted that the sensitivity of these parameters to changes as a result of exposure of groundwater to surface level conditions (i.e., changes in the partial pressure of dissolved gases or the conditions of the purging system) make in-situ monitoring desirable. An alternative to this would be to conduct these measurements in a closed cell attached to the discharge side of the pump system.

Other factors which will influence the amount of purging required before sampling include the pumping rate and the placement of the pumping equipment within the column of water in the well bore. For example, recent studies have shown that if a pump is lowered immediately to the bottom of a well before pumping, it may take some time for the column of water above it to be exchanged if the transmissivity of the aquifer is high and the well screen

is at the bottom of the casing.^{5,8} In such cases the pump will be drawing water primarily from the aquifer.

This has been further documented in studies conducted by the National Council of the Paper Industry for Air and Stream Improvement (NCASI)⁴ on a full-scale model of a 2-inch PVC well. They found that purging from just below the water surface insured a more complete removal of the casing water than by withdrawal from well below the surface. It was also evident that when purging did occur from just below the surface, satisfactory results could be obtained at any of a wide range of pumping rates with either a peristaltic or a submersible pump.

Because of the potential for further environmental contamination, planning for purge water disposal is a necessary part of well monitoring. Alternatives range from dumping it on the ground (not back down the well) to full containment, treatment, and disposal. If the well is believed to be contaminated, the best practice is to contain the purge water and store it until the water samples have been analyzed. Once the contaminants are identified, appropriate treatment requirements can be determined.

There are many methods available for well purging. In some cases bailing will suffice, however it can become tedious and labor intensive in deep or large diameter wells. In some situations, an inflatable packer can be utilized above and/or below the pump to reduce the casing volume requiring purging. This technique is particularly adaptable to wells with more than one screened interval to isolate the aquifer of interest. The size and weight of a pump and packer assembly usually require tripods/derricks and hoist equipment which are not easily implemented. Additionally the packer may be constructed of rubber material which may effect some analysis although viton packers are feasible.

Gas pressure lift systems are useful in many instances. They are usually light, easy to install, and can be powered by several different pressure systems, usually compressed nitrogen or air. The effect of the contact between the pressure gas and the groundwater usually results in changes in the dissolved gas content.⁵ As a result pH, conductivity, or other analysis used to determine purge completion must be conducted down hole.

Peristaltic pumps are widely used for purging of wells with water levels close to the surface (less than 8 meters). They are reasonably portable, light, and easily adaptable to ground level monitoring of purge indicator parameters by attaching a flow-through cell. These pumps require a minimum of down hole equipment and can easily be cleaned in the field; or the entire tubing assembly can be changed for each well.

Several manufacturers are marketing submersible pumps specifically designed for groundwater monitoring. They are generally capable of fitting down 2-inch ID wells. Most of these pumps have effective depth limitations of less than 150 feet. Although some can operate to depths in excess of 300 ft., they usually have substantially reduced discharge flows and significant power consumption. All electrically powered equipment should be checked for

electrical safety certification (UL, FM, NFPA, NEC). Appendix F discusses the various certifications in detail.

Three basic designs are currently available:

- Eductors--A pump is used to circulate water through a venturi, the resultant pressure drop across the venturi is used to draw sample into the recirculating stream. A split stream is drawn off the recirculating stream equal to the flow at the intake. These systems require priming water and must be run long enough to insure complete removal of the priming water from the recirculation loop. They readily fit into a 2-inch diameter well and will recover samples from as deep as 100 ft or 50 ft in a 1-1/2 inch ID well.
- Submersible Motor--A small submersible electric motor is used to drive a common stator/rotor pump. They will fit inside a 2-inch ID well and operate at depths to approximately 150 ft. The discharge flow varies with depth" from 1.2 gpm at 10 ft to 0.6 gpm at 125 ft.
- Bladder Compression--A flexible bladder with a check valve at either end is suspended inside a rigid chamber. Hydrostatic pressure forces water into the bladder. The chamber is then pressurized which squeezes the bladder and forces the water out the other check valve and into the discharge tubing. This cycle is then repeated until the sample is recovered.

The pump is operated by a compressed air source, either bottled gas or a small DC powered compressor. Pumps are available that can sample from depths of 250 ft. Flow rates vary with models but range from about 2.0 gpm at 25 ft to 0.5 gpm at 150 ft. (At least one model is capable of 0.78 gpm at 250 ft.)

Once the well has been sufficiently purged, the actual sampling should begin as soon as the water level begins to approach its pre-purge level. Sampling for volatile organics may begin even sooner, before substantial volatilization begins. If recovery is very slow, it may be necessary to wait several hours or even until the following day before sufficient volume is available for all the necessary analyses. In this instance a volatile organics sample set may be collected soon after completion of the purging process and a second set with the remaining samples. When a pump is used for sample collection, its rate should be controlled, if possible to closely match the transmissivity of the formation. Excessive draw down of the well during sampling may result in nonrepresentative samples due to changes in groundwater flow.⁹

Bailers are probably the simplest means of collecting groundwater samples. They result in a minimum of sample disturbance if carefully handled. They can be constructed of noncontaminating materials, and their low relative cost makes the use of a separate device for each well practical, thus eliminating in-field

cleaning and cross contamination. Peristaltic pumps can be used for sampling in most shallow wells. They require a minimum of down-hole equipment and cross contamination can be eliminated by replacement of the suction tubing between wells. Gibb¹⁰ as well as NCASI⁴ found little difference between samples withdrawn by a peristaltic pump and those taken by a bailer. These pumps however may not be suitable for the collection of volatile organics due to possible gas stripping; therefore, their use should be supplemented by a bailer when sampling includes volatile organic species.

The use of submersible pumps as described previously for sample collection is possible provided they are constructed of suitably noncontaminating materials. They can operate at depths beyond the capabilities of peristaltic pumps and at which depths bailing becomes tedious. The chief drawback, however, is the difficulty of avoiding cross contamination between wells. These systems are generally too expensive to allow for several separate units. Though some units can be easily disassembled and allow for replacement of most sample contacted surfaces, field decontamination still may be difficult and should properly require solvents that may lead to sample contamination. Their use therefore, in multiple well programs, should be carefully considered against bailers.

In general, gas pressure displacement systems where gas interfaces with the liquid should not be used for sample collection as they have been shown to cause considerable changes in the groundwater character.¹⁰

3.4.1 METHOD III-7: PURGING WITH A PERISTALTIC PUMP

Discussion

The peristaltic pump as described in the surface water sampling section Method III-3 can be implemented for the presample purging of groundwater monitor wells.

Uses

The use of a peristaltic pump for well purging is particularly advantageous since the same system can later be used for sample collection (see Method III-10). The application, however, is limited to wells with a depth of less than approximately 8 meters, due to the limited lift capabilities of peristaltic action.

Procedures for Use

1. Using clean equipment, sound well for total depth and water level, then calculate the fluid volume in the casing ("casing volume").
2. Determine depth from casing top to mid-point of screen or well section open to aquifer. (Consult drillers log, or sound for bottom.)
3. If depth to mid-point of screen is in excess of 8 meters, choose alternate system.
4. Lower intake into the well to a short distance below the water level and begin water removal. Collect or dispose of purged water in an acceptable manner. Lower suction intake, as required, to maintain submergence.
5. Measure rate of discharge frequently. A bucket and stopwatch are most commonly used.
6. Purge a minimum of four casing volumes or until discharge, pH, temperature, or conductivity stabilize. See discussion on well purging in Section 3.4, Groundwater.
7. After pumping, monitor water level recovery. Recovery rate may be useful in determining sample rate.

3.4.2 METHOD III-8: PURGING WITH A GAS PRESSURE DISPLACEMENT SYSTEM

Discussion

A pressure displacement system consists of a chamber equipped with a gas inlet line, a water discharge line and two check valves (see Figure 3-7). When the chamber is lowered into the casing, water floods it from the bottom through the check valve. Once full, a gas (i.e., nitrogen or air) is forced into the top of the chamber sufficient to result in the upward displacement of the water out the discharge tube. The check valve in the bottom prevents water from being forced back into the casing, and the upper check valve prevents water from flowing back into the chamber when the gas pressure is released. This cycle can be repeated as necessary until purging is complete.

Uses

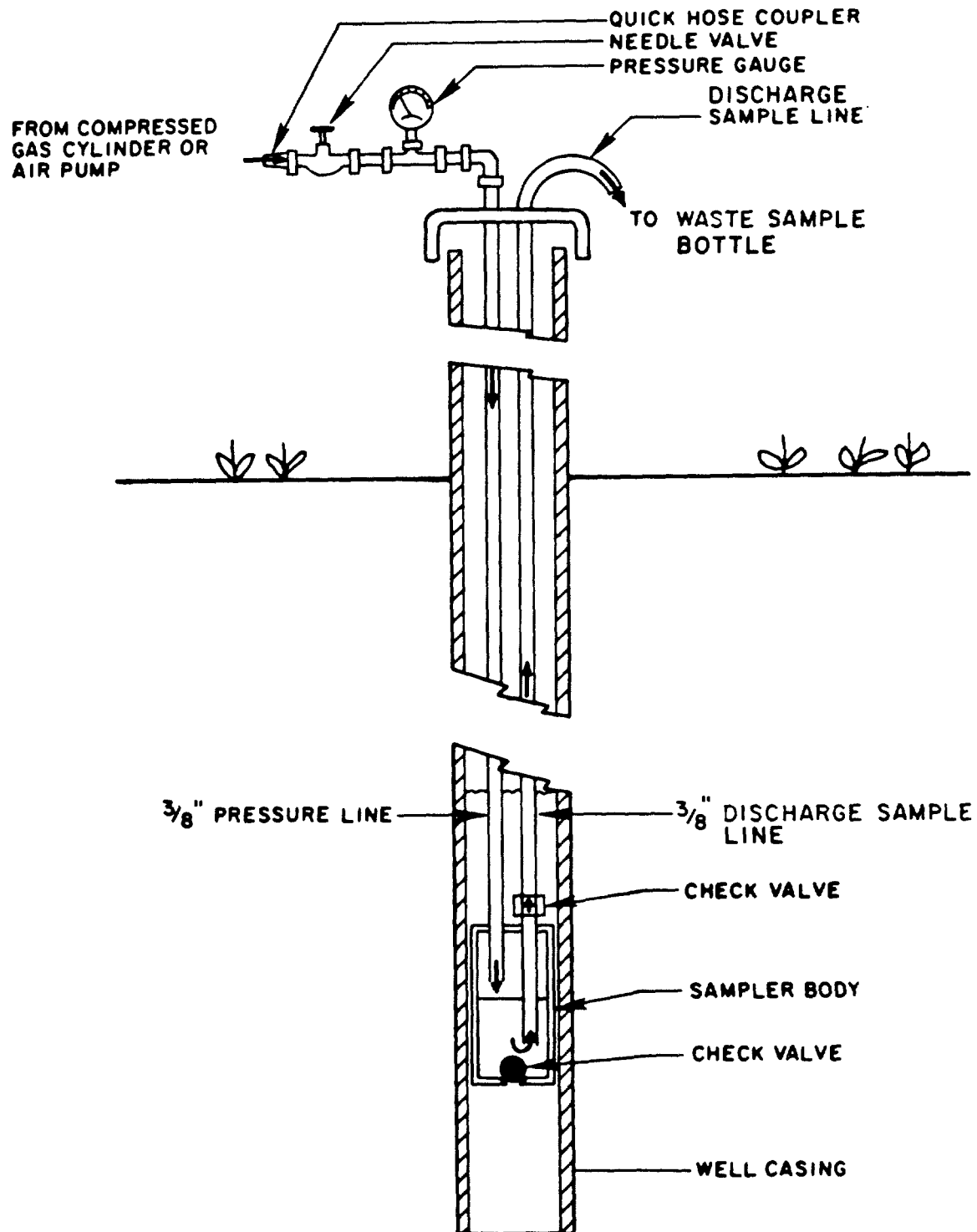
The pressure lift system is particularly useful when the well depth is beyond the capability of a peristaltic pump. The water is displaced up the discharge tube by the increased gas pressure above the water level. The potential for increased gas diffusion into the water makes this system unsuitable for sampling for volatile organic or most pH critical parameters.¹⁰

Procedures for Use

1. Using clean noncontaminating equipment, i.e., an electronic level indicator (avoid indicating pastes) determine the water level in the well, then calculate the fluid volume in the casing.
2. Determine depth to midpoint of screen or well section open to aquifer (consult drillers log).
3. Lower displacement chamber until top is just below water level.
4. Attach gas supply line to pressure adjustment valve on cap.
5. Gradually increase gas pressure to maintain discharge flow rate.
6. Measure rate of discharge frequently. A bucket and stopwatch are usually sufficient.
7. Purge a minimum of four casing volumes or until discharge characteristics stabilize (see discussion on well purging in Section 3.4, Groundwater).
8. After pumping, monitor water level recovery. Recovery rate may be useful in determining sample rate.

Source

U. S. Environmental Protection Agency. "Procedures Manual for Ground Water Monitoring at Solid Waste Disposal Facilities," EPA-530/SW-611, August 1977.



Source: Reference 6.

Figure 3-7. Gas pressure displacement system.

3.4.3 METHOD III-9: SAMPLING MONITOR WELLS WITH A BUCKET TYPE BAILER

Discussion

Bucket type bailers are tall narrow buckets equipped with a check valve on the bottom. This valve allows water to enter from the bottom as the bailer is lowered, then prevents its release as the bailer is raised (see Figure 3-8). Top filling bailers are also available and may be useful for well purging but generally result in increased sample turbulence and are not recommended for sample acquisition.

Uses

This device is particularly useful when samples must be recovered from depths greater than the range (or capability) of suction lift pumps, when volatile stripping is of concern, or when well casing diameters are too narrow to accept submersible pumps. It is the method of choice for the collection of samples which are susceptible to volatile component stripping or degradation due to the aeration associated with most other recovery systems. Samples can be recovered with a minimum of aeration if care is taken to gradually lower the bailer until it contacts the water surface and is then allowed to sink as it fills. Teflon is generally the best construction material but other materials (PVC, stainless steel, etc.) are acceptable if compatible with designated sample analysis. The primary disadvantages of bailers are their limited sample volume and inability to collect discrete samples from a depth below the water surface.

Procedures for Use

1. Using clean, noncontaminating equipment, i.e., an electronic level indicator (avoid indicating paste), determine the water level in the well, then calculate the fluid volume in the casing.
2. Purge well as per Methods III-7 or III-8.
3. Attach precleaned bailer to cable or line for lowering.
4. Lower bailer slowly until it contacts water surface.
5. Allow bailer to sink and fill with a minimum of surface disturbance.
6. Slowly raise bailer to surface. Do not allow bailer line to contact ground.
7. Tip bailer to allow slow discharge from top to flow gently down the side of the sample bottle with minimum entry turbulence.
8. Repeat steps 2-5 as needed to acquire sufficient volume.
9. Select sample bottles and preserve the sample, if necessary, according to the guidelines in Appendix A.

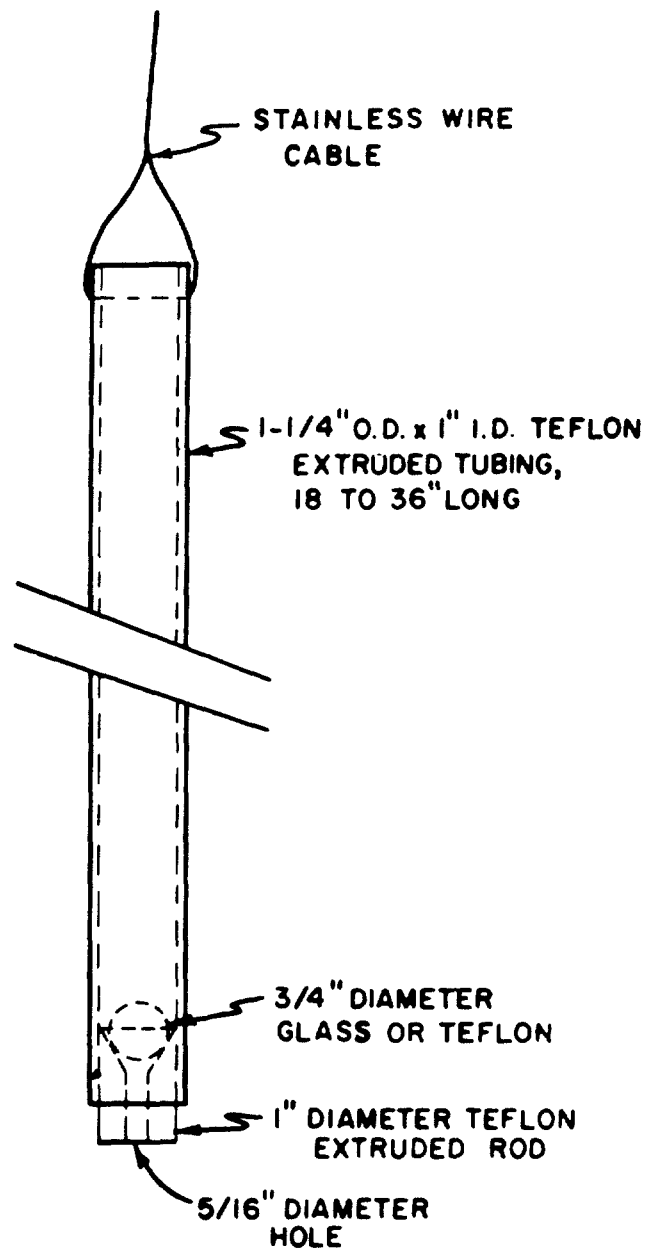


Figure 3-8. Teflon bailer.

10. Check that a Teflon-liner is present in cap if required. Secure the cap tightly.
11. Label the sample bottle with an appropriate tag. Be sure to complete the tag with all necessary information. Record the information in the field logbook and complete all chain-of-custody documents.
12. Thoroughly decontaminate the bailer after each use according to specific laboratory instructions, or the general guidelines in Appendix E. In some cases, especially where trace analysis is desired, it may be prudent to use a separate bailer for each well.

Sources

Dunlap, W. J., McNabb, J. F., Scalf, M. R. and Crosby, R. L., "Sampling for Organic Chemicals and Microorganism in the Subsurface." EPA-600/2-77-176, August 1977.

3.4.4 METHOD III-10: SAMPLING MONITOR WELLS WITH A PERISTALTIC PUMP

Discussion

A pump system is considerably advantageous when analytical requirements demand sample volumes in excess of several liters. The major drawback of a pump system is the potential for increased volatile component stripping as a result of the required lift vacuum. Samples for volatile organic analysis should be collected with a bailer as described in Method III-9 and should precede any sample collection which may further disturb the well bore content.

Uses

The peristaltic pump system can be used for monitor well sampling whenever the lift requirements do not exceed 8 meters (for deeper wells see Method III-11). It becomes particularly important to use a heavy wall tubing in this application in order to prevent tubing collapse under the high vacuums needed for lifting from depth.

Procedures for Use

1. Using clean, noncontaminating equipment, i.e., an electronic level indicator (avoid indicating paste), determine the water level in the well, then calculate the fluid volume in the casing.
2. Purge well as per Methods III-7 or III-8.
3. If soundings show sufficient level of recovery, prepare pump system. If insufficient recovery is noted allow additional time to collect samples on a periodic schedule which will allow recovery between samplings.
4. Collect volatile organic analysis samples if required with bucket type bailer (Method III-9).
5. Install clean medical grade silicon tubing in peristaltic pump head.
6. Attach pump to required length of precleaned Teflon suction line and lower to midpoint of well screen if known or slightly below existing water level.
7. Consider the first liter of liquid collected as a system purge/rinse.
NOTE : If well yield is insufficient for required analysis this purge volume may be suitable for some less critical analysis.
8. Fill necessary sample bottles by allowing pump discharge to flow gently down the side of bottle with minimal entry turbulence. Cap each bottle as filled.
9. Select sample bottles and preserve the sample if necessary as per guidelines in Appendix A.

10. Check that a Teflon-liner is present in cap if required. Secure the cap tightly.
11. Label the sample bottle with an appropriate tag. Be sure to complete the tag with all necessary information. Complete chain-of-custody documents and field logbook.
12. Allow system to drain then disassemble. Return tubing to lab for decontamination. See Appendix E for general decontamination procedures.

Sources

Dunlap, W. J., McNabb, J. F., Scalf, M. R. and Crosby, R. L. "Sampling for Organic Chemicals and Microorganisms in the Subsurface," EPA-600/2-77-176, August 1977.

3.4.5 METHOD III-11: SAMPLING MONITOR WELLS WITH A SUBMERSIBLE PUMP

Discussion

Several types of submersible pumps are available for groundwater monitoring and offer considerable advantages over other systems. They are able to operate from depths beyond the capabilities of peristaltic pumps and save significant time and effort relative to hand bailing. Further, if constructed of suitable materials and properly used, they can both purge and adequately sample the well.

Uses

Submersible pumps generally use one of two types of power supplies, either electric or compressed gas. Electric powered pumps generally run off a 12 VDC rechargeable battery from an automotive electrical system. Those units powered by compressed gas normally use a small electric compressor which also needs 12 VDC power. They may also utilize compressed gas from bottles or even high performance hand pumps.

These pumps are generally constructed of "more or less" noncontaminating materials "suitable for Priority Pollutant Sampling". They often contain plastics, rubber or metal parts which may contribute or otherwise effect the analysis of samples for certain trace components. Investigations requiring samples for a wide range of trace analysis may preclude their use for sample acquisition; however, they may still be useful for purging. In any case, when doubt remains, bailers are the best choice for actual sample acquisition.

Procedures for Use

1. Using clean, noncontaminating equipment, i.e., an electronic level indicator (avoid indicating paste or chalk), determine the water level in the well, then calculate the fluid volume in the casing
2. Lower the precleaned pump to just below the water level and begin pumping. Collect or dispose of purged water in an acceptable manner. Lower the pump as required to maintain submergence.
3. Measure rate of discharge frequently. A bucket and stopwatch are commonly used.
4. Purge a minimum of four casing volumes or until discharge pH, conductivity, or temperature stabilize. See discussion on well purging in Section 3.4, Groundwater. (Note: If the pump is constructed of materials compatible with the required sample analysis and if the well has recovered sufficiently (resound water level) sample acquisition can proceed as follows. It should be cautioned that all down hole and potentially wetted surfaces must also be noncontaminating/noncontributing. This includes power and suspension cables and compressed gas or sample tubing.)

5. Fill necessary sample bottles by allowing pump discharge to flow gently down the side of bottle with minimal entry turbulence. Cap each bottle as filled.
6. Select appropriate sample bottles and preserve the sample if necessary as per guidelines in Appendix A.
7. Check that a Teflon-liner is present in cap if required. Secure the cap tightly.
8. Label the sample bottle with an appropriate tag. Be sure to complete the tag with all necessary information. Complete chain-of-custody documents and field logbook.
9. Allow system to drain then disassemble. Return tubing to lab for decontamination. See Appendix E for general decontamination procedures.

3.5 REFERENCES

1. United States Department of the Interior National Handbook of Recommended Methods for Water--Data Acquisition. Reston, Virginia. 1977.
2. deVera, E. R., B. P. Simmons, N. D. Stephen, and D. L. Storm. Samplers and Sampling Procedures for Hazardous Waste Streams. EPA-600/2-80-018.
3. Instrument Specialties Company, Instruction Manual, Model 2100 Wastewater Sampler. Lincoln, Nebraska. January 1980.
4. National Council of the Paper Industry for Air and Stream Improvement, Inc. A Guide to Groundwater Sampling. Technical Bulletin No. 362. Madison, New York. January 1982.
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9. U.S. Environmental Protection Agency. Procedures Manual for Groundwater Monitoring at Solid Waste Disposal Facilities. EPA-530/SW-611. August 1977.
10. Gibb, J. P., R. M. Schuller, and R. A. Griffin. Collection of Representative Water Quality Data From Monitoring Wells in Land Disposal.

SECTION 4.0

GASES, VAPORS, AND AEROSOLS

4.1 GENERAL

Air monitoring at hazardous waste sites and environmental spills can be quite useful as an indicator of potential safety problems and as a means of screening for the presence of possible airborne contaminants. Monitoring is also important as a means of determining the specific identity and concentration of airborne toxic and hazardous pollutants onsite and the extent of their migration offsite for both worker and public health risk assessments. For the purpose of this document, sampling for gases, vapors and aerosols at hazardous waste sites and environmental spills falls into three general categories: the ambient atmosphere, soil gases, and container headspace gases.

4.2 AMBIENT

Ambient concentrations of airborne contaminants are greatly affected by the topography and meteorology of the surrounding area, and the investigator must be cognizant of this when choosing monitoring methods and equipment. Besides the obvious effects of temperature, wind, and precipitation in relation to dispersion and deposition of atmospheric pollutants, heat and sunlight can dramatically increase rates of volatilization, and cold and calm may cause stagnant conditions to prevail tending to reduce migration and to concentrate pollutants in low-lying areas. Accurate detection of atmospheric pollutants must take into account these and other factors if a successful sampling effort is desired.

Of major importance when discussing the sampling of ambient atmospheres is the use of portable analytical instrumentation. In addition to being portable, these devices need to be rugged and easy to operate and need to provide real time data in order to best meet the requirements inherent to field applications. They must also be proven safe when used in hazardous waste environments. Electrical devices and instruments which use flame or combustion principles must be of a type that eliminate the possibility of igniting combustible atmospheres. All instruments used should be "approved" or "certified" by Underwriters Laboratory (UL) or Factory Mutual Systems (FM) according to provisions set forth by the National Electrical Code (NEC). A detailed discussion of the various electrical product certification programs is presented in Appendix F. In addition, this appendix contains an explanation of the various atmosphere Classes, Divisions, and Groups for which these devices are approved.

In order to insure safe operation, the user must also become familiar with the detailed operation and maintenance procedures found only in the operating manual of each specific instrument. The investigator should keep in mind that the procedures outlined here are necessarily general and intended only to supplement the instrument operating manual. Investigators must also familiarize themselves with the limitations of each instrument. Inability to detect certain compounds, insensitivity (e.g., contaminants in the solid phase), slow response time, pump rate capacity, etc. are all factors which may affect the safety of the operator and/or quality of the data.

Field instrumentation is invaluable during initial site surveys for assessing the potential hazards that exist. Information of this nature is needed in order to determine the degree of protection required for personnel or to provide direction for further quantification of specific parameters.

Instruments such as portable oxygen indicators and combustible gas detectors would be the instruments of choice when a general safety assessment of an unknown atmosphere is necessary. Such atmospheres present many hazards including oxygen deficiency, explosivity, flammability, etc., and data obtained with these instruments can be used by the onsite safety officer to generally assess the presence of these dangers and dictate precautionary measures to be taken. They can be used to screen pockets or depressions in

the land contour, areas in close proximity to drums or spills, or closed in, unventilated rooms which may not have enough oxygen to support life or which allow combustible vapors to concentrate.

Other instruments that may be required for evaluating the hazard potential of ambient or workplace atmospheres are those which utilize flame ionization (FID) and photoionization (PID) detectors. These detectors are important due to the increased levels of sensitivity they can provide (for specific compound classes) and when used in conjunction with chromatographic columns, can specifically characterize and/or identify hazardous materials at spills or dump sites.

The Century OVA and AID Model 550 represent a type of instrument which uses a flame ionization detector. In its simplest form this type is used to determine the presence of gaseous and/or vapor phase hydrocarbons. These instruments responded to most gaseous/vapor phase organics present. The readings are referenced to a single component standard gas (usually methane). The response of such instruments is often termed "total hydrocarbons;" however, this is misleading since not all hydrocarbons are detected, specifically, important particulate hydrocarbons (i.e., pesticides and polynuclear aromatics), and polychlorinated biphenyls. In addition, the response to mixtures of vapor phase hydrocarbons depends upon the ratios and the types of organic compounds present and cannot be related to a specific vapor concentration. FIDs do, nonetheless, provide a useful and reliable tool for general assessment purposes.

Photoionization analyzers such as the portable HNU Model P1-101 are also capable of detecting the presence of a wide variety of chemical species, both organic and inorganic. As with FID's, photoionization detectors suffer similar limitations of detector response to component mixtures. The inability to respond to certain compounds must be recognized; however, PID's can provide important information for evaluation purposes.

As stated previously, the usefulness of both portable FID's and PID's can be expanded when used in conjunction with gas chromatography. The Century units offer a chromatography option which, when used properly, can be quite a valuable tool for aiding in specific compound identification. At present, the HNU P1-101 is not available with a chromatography option; however Spittler and Orl report success with a portable photoionization detector/gas chromatography (Photovac 10A10, Thornhill, Ontario) capable of sensitivity in the 0.1 to 10 ppb range. In all cases it should be realized that chromatography can be quite complex and demands the skills of an experienced operator to obtain valid and meaningful results.

Additional useful instruments and devices include those adapted from industrial hygiene practices and/or techniques. These include stain detector tubes and personnel collection devices. Detection by these methods is the most specific of all of the devices thus far described. These methods are therefore extremely useful for compound identification and quantification.

Stain detector tubes such as manufactured by National Drager, Matheson-Kitagawa, Bendix Corporation, and MSA provide an immediate indicator of a specific chemical or species of interest. They are somewhat limited due to small sample volume, interferences, degree of accuracy, operator judgement, etc.; however, they are valuable as a quick, relatively simple, direct-reading method of determining specific gas concentrations.

Collection devices such as solid sorbents, chemical absorbing solutions and filters are the most accurate of the methods used for properly identifying and quantifying species of interest. Use of these methods requires adherence to very specific procedures and conditions of the type found in the "NIOSH Manual of Analytical Methods,"² EPA Federal Reference Methods, or specific papers documenting procedures and characteristics of sorbent resins. Collected samples are subsequently analyzed at an offsite analytical laboratory that usually yields an analytical precision and accuracy presently unavailable in most field applications.

It should be noted, at this point, that ambient monitoring, within the context of this section, deals with area monitoring and not personnel monitoring. Although ambient methods can provide information on the types of contaminants present and the relative magnitude of contamination, it is not a substitute for personnel monitoring when worker exposure is the prime concern. In such cases, NIOSH methodologies should be consulted and appropriate methods chosen dependent upon specific monitoring requirements.

4.2.1 METHOD IV-1: DETERMINING OXYGEN CONTENT IN AMBIENT AND WORKPLACE ENVIRONMENTS WITH A PORTABLE OXYGEN MONITOR

Discussion

A portable oxygen monitor has three principle components for operation: the airflow system, the oxygen sensing device, and the microamp meter. Typically the air is drawn through the oxygen sensor with a built-in pump or aspirator bulb, although some instruments use passive cells. The sensor indicates the oxygen content and the information is translated electrochemically to the meter.

Most monitors have meters which indicate the oxygen content from 0-25 percent. There are also oxygen monitors available which indicate concentrations on scales from 0-5 percent and 0-100 percent. The most useful for ambient measurements is the 0-25 percent oxygen content readout. Many instruments also have alarm modes which can be set to activate at a specified oxygen concentration.

Uses

Portable oxygen monitors are invaluable when initially responding to hazardous material spills or waste site situations. They are useful in screening depressions in the land, unventilated rooms, or other areas that may not contain enough oxygen to support life. When used properly the portable oxygen monitor will indicate the percent oxygen in the test atmosphere. Normal oxygen concentration required for respiration is 20.9 percent.

Procedures for Use

1. Make sure instrument is clean and serviceable, especially sample lines and detector surfaces.
2. Consult records on instrument maintenance to determine if detector solution should be changed. Some instruments will need this service after as little as 1-2 weeks of use.
3. Check battery charge level. If in doubt, charge battery as detailed in operating manual. Some units have charge level indicators while others have alarms that will indicate a low charge.
4. Verify that sample pump is operable (if so equipped) when analyzer is on.
5. Turn instrument on and, using calibration knob on instrument, calibrate against fresh air (20.9 percent O₂) by aligning meter needle at 20.9 percent.
6. If unit is equipped with alarm mode, set alarm at desired level.

7. A quick field check can be accomplished by exhaling into the sensor, this should cause a definite drop in O₂ readings and activate any alarms. Allow for instrument warmup, if necessary, before entering site to take readings.
8. Position intake assembly or sensor in close proximity to area in question to get accurate reading.
9. If alarm occurs, personnel should evacuate area, unless equipped with supplied air equipment suitable for use in an IDLH atmosphere.
10. Some important factors to keep in mind during use are:
 - Slow sweeping motions may assist in the prevention of bypassing problem areas.
 - Operation of instrument in temperatures outside of manufacturer specified operating range may compromise accuracy of readings or damage unit. The instrument should always be calibrated at the temperature of intended use.
 - Presence of known or unknown interfering gases, especially oxidants, can affect readings (for example the Edmont Model 60-400 Oxygen Monitor has interferences of the following gases in concentrations greater than 0.25 percent or 2500 ppm: SO₂, fluorine, chlorine, bromine, iodine and nitrogen oxides). See the operating manual for unit being used.
 - The oxygen detector can also be poisoned (decrease in sensitivity) by exposure to various gases. Some detectors are poisoned by concentration of mercaptans and hydrogen sulfide greater than or equal to 1 percent. See operating manual for unit being used.
 - When relying on alarm mode for warnings of oxygen deficient atmospheres, a manual check of the alarm function at regular intervals is recommended.
 - Wherever applicable, protect instrument with a disposable cover to prevent contamination.
 - Most units will have rechargeable battery packs that provide continuous operation for 8-12 hours. Recharging batteries prior to expiration of the specified interval will insure operation while on a site.
 - More than any other factor, effective utilization of unit requires operator with full understanding of operating principles and procedures for the specific instrument in use.

Sources

Edmont Model 60-400 Combustible Gas/Oxygen Monitor Instruction Manual.
Manufactured by Energetic Science, Elmsford, NY 10523.

U.S. Environmental Protection Agency. "Hazardous Materials Response
Operations Training Manual." National Training and Operational Center,
Cincinnati, OH.

4.2.2 METHOD IV-2: DETERMINATION OF COMBUSTIBLE GAS LEVELS USING A PORTABLE COMBUSTIBLE GAS INDICATOR

Discussion

A combustible gas indicator consists of three primary components: the sensor (hotwire, catalytic, solid state, etc.), signal processor and readout display. Sample is introduced to the sensor either by diffusion into a passive sensor or by pumping. The sensor produces a signal which is processed and displayed as the ratio of the combustible gas present to the total required to reach the lower explosion limit (LEL).

The lower explosive limit (also LFL, lower flammability limit) is defined as the lowest concentration of gas or vapor in air which can be ignited by an ignition source and cause an explosion or flame propagation. Conversely, the upper explosive limit or UEL (also UFL, upper flammability limit) is the concentration of gas in air above which there is insufficient oxygen available to support combustion, and an explosion is unlikely. A flame, however, may burn at the gas-air interface or, should additional air enter the mixture, a very explosive atmosphere may develop. In general, the instruments respond in the following manner.

- The meter indicates 0.5 LEL (50 percent). This means that 50 percent of the concentration of combustible gas needed to reach an unstable combustible situation is present. If the LEL of the gas is 5 percent in air, then the instrument indicates a 2.5 percent mixture is present.
- The meter needle stays above 1.0 LEL (100 percent). This means that the concentration of combustible gas is greater than the LEL and less than the UEL and, therefore, immediately combustible and explosive.
- The meter needle rises above the 1.0 (100 percent) mark and then returns to zero. This response indicates the ambient atmosphere has a combustible gas concentration greater than the UEL.

Of the many instruments commercially available for detecting combustible or explosive gas, some are not certified safe for operation in the atmospheres they can detect. It is important to use only those monitors that are certified safe for use in atmospheres greater than 25 percent of the LEL. Appendix F discusses the electrical product certification programs and details the various atmosphere divisions, classes and groups for which these products are tested.

Some combustible gas monitors provide readouts in units of percent LEL, some in percent combustible gases by volume, and some have scales for both. Many situations may occur where the types of combustible gases to be encountered are unknown. In such instances the more explosive the calibration gas (the lower the LEL) the more sensitive the indication of explosivity and thus the greater the margin of safety. The operator should be familiar with

the LEL concentrations for specific gases to effectively use instruments that provide data in percent combustible (by volume) only.

Although monitors can be purchased that are factory calibrated using gases such as butane, pentane, natural gas, or petroleum vapors, methane calibration is the most common. The LEL of methane is 5 percent by volume in air, therefore, an air mixture containing 5 percent methane will be read as 100 percent LEL and will be explosive if a source of ignition is present. When combustible gases other than methane are sampled, the relative response of the detector for these other gases must be considered. Recalibration to other gases may be possible; see manufacturers recommendations. The relative sensitivity of the detector and the differences in LEL for different gases will produce varying meter responses equal concentrations of different gases. Actual correlation equations that will convert the percent LEL (based on methane) read by the unit to a percent LEL for another combustible gas can usually be found in the operating manual.

Many units also have alarm systems which can be adjusted for various LEL's and several are available that incorporate oxygen analyzers.

Uses

In general, combustible gas detectors are used to determine the potential for combustion or explosion of unknown atmospheres. These instruments, in combination with oxygen detectors and radiation survey instrumentation, should be the first monitors used when entering a hazardous area. In this sense they provide a general indication of the degree of immediate hazard to personnel and can be used to assist the safety officer in making decisions on levels of protection required at the site. However, they provide little or no information about the presence of compounds hazardous or toxic at trace level concentrations.

Procedure for Use

1. Make sure instrument is clean and serviceable, especially sample lines and detector surfaces.
2. Check battery charge level. If in doubt, charge battery as described in operating manual. Some units have charge level meters, while others have only low charge alarms.
3. Turn unit to ON position, and allow instrument sufficient warmup time.
4. Verify that sample pump is operable (if so equipped) when analyzer is ON.
5. With the intake assembly in combustible gas-free ambient air, zero the meter by rotating the zero control until the meter reads 0 percent LEL.

6. Calibrate unit against known concentration of a calibration gas by rotating the calibration control (span or gain) until the meter reads the same concentration as the known standard. For those instruments with internal or nonadjustable span, a calibration curve should be prepared, using concentrations in the range expected to be encountered.
7. If necessary, adjust alarm setting to appropriate combustibility setting.
8. Position intake assembly or cell in close proximity to area in question to get accurate reading.
9. If alarm occurs, or if readings reach the action levels designated in the safety plan, personnel should evacuate area.
10. If instrument malfunction occurs, personnel should evacuate area.
11. Some important factors to keep in mind during use are:
 - Slow sweeping motions of intake or cell assembly will help assure that problem atmospheres are not bypassed. Cover an area from floor (ground) to ceiling, or above breathing zone.
 - Operation of unit in temperatures outside of recommended operating range may compromise accuracy of readings or damage the instrument.
 - Platinum filament detectors may be poisoned (reduced in sensitivity) by gases such as leaded gasoline vapors (tetraethyl lead), sulfur compounds (mercaptans and hydrogen sulfide) and silicon compounds.
 - Many combustible gas detectors are not designed for use in oxygen-enriched or depleted atmospheres. If this condition is encountered or suspected, personnel should evacuate the area. Specially designed units are available for operation in such atmospheres.
 - An oxygen detector should always be used in conjunction with explosimeters.
 - Accurate data depends on regular calibration and battery charging. See operating manual.
 - More than any other factor, effective utilization of unit requires operator with full understanding of operating principles and procedures for the specific instrument in use.

Sources

Edmont Model 60-400 Combustible Gas/Oxygen Monitor Instruction Manual.
Manufactured by Energetic Science, Elmsford, NY 10523.

U.S. Environmental Protection Agency. "Hazardous Materials Incident
Response Operators Training Manual." National Training and Operational
Training Center, Cincinnati, Ohio.

4.2.3 METHOD IV-3: MONITORING ORGANIC VAPORS USING A PORTABLE FLAME IONIZATION DETECTOR

Discussion

A flame ionization detector (FID) will respond to most organic vapors as they form positively charged ions when combusted in a hydrogen flame. The magnitude of the response is a function of the detector sensitivity and the ionization properties of the particular compound as well as its concentration. As a result, this signal must be compared to that generated by calibration with a known concentration of a standard gas. The sample concentration is then reported as the ppm equivalent of the calibration compound. Most units are calibrated with a known concentration of methane; however, almost any gaseous hydrocarbon that produces a response can be used. Many models also have built-in calibration circuits which can insure that the electronic response to a known signal remains constant.

Some models can be equipped with an option that provides chromatographic separation of the sample gas constituents. This permits a tentative qualification and quantification to be made of the resultant peaks which have retention times equal to those of known standards. This option requires the use of a chart recorder for recording the peak areas and retention times, and in such a mode, prevents the instrument from providing a continuous readout. Use of a chromatographic option also requires additional expertise if reliable, consistent results are desired.

Most portable FID's rely on the sample gas to supply the combustion air to the detector flame, so they are designed to operate in ambient atmospheres with relatively normal oxygen concentrations (21 percent). This design precludes the sampling of process vents, poorly ventilated or sealed containers, or any sample gas hydrocarbon concentration sufficient to reduce the available oxygen or otherwise saturate the detector. In such instances adaptations are usually available to supply a source of oxygen from a compressed gas bottle or introduce the gas through a dilution system with a known (calibrated) dilution factor.

Uses

A portable FID is useful as a general screening tool to detect the presence of most organic vapors. It will not, however, respond to particulate hydrocarbons such as pesticides, PNAs, and PCBs. It can be used to detect pockets of gaseous hydrocarbons in depressions or confined spaces, to screen drums or other containers for the presence of entrapped vapors, or generally to assess an area for the presence of elevated levels of vapor phase organics.

Procedure for Use

The procedures presented in this section are intended to apply to any portable FID; therefore, detailed operating instructions must be obtained from the operating manual of the specific unit to be used.

1. Check battery charge level indicator; if in doubt, recharge battery as described in manual.
2. Turn instrument on and allow adequate warmup time.
3. If equipped with internal calibration capability, perform instrument calibration. Perform zero and other calibration procedures as described in operating manual.
4. If equipped with an alarm mode, set alarm at desired concentration.
5. Turn on pump and check for leaks by covering sample inlet and observing rotameter. Indicator ball should drop to zero level.
6. With pump operating, open hydrogen gas storage tank valve and open supply regulator to allow fuel gas flow to detector chamber.
7. Depress igniter switch, observe indicator needle for positive response and listen for a "pop." If flame fails to light, depress igniter switch again.
8. Once detector flame is lit, unit is ready for use.
9. If calibration to a specific hydrocarbon species is desired, complete this procedure according to the manufacturers instructions.
10. Hold sample probe in close proximity to area in question as low sample rate allows for only very localized readings.
11. Slow sweeping motion will help prevent the bypassing of problem areas. Make sure batteries are recharged within time frame specified in operator manual. Usual length of operating time between charges is 8-12 hours.
12. Some units have alarms that signal operator if detector flame goes out. If this alarm sounds, evacuate all personnel and relight flame in known safe area then reenter site.
13. Monitor fuel and/or combustion air supply gauges regularly to insure sufficient gas supplies.
14. High background readings after prolonged use may indicate sample probe and/or in-line filters (in front of detector) need to be cleaned. Use of pipe cleaners or clean air blown backwards through filters is adequate. Do not use organic solvents as detector will respond to solvent as well.
15. Representative readings will also depend on performance of routine maintenance as described in detail in operating manual. Also, since unit contains pressurized gas supplies, perform leak check procedures regularly, as leaking hydrogen gas is explosive.

16. As with any field instrument, accurate results depend on the operator being completely familiar with the operator's manual for the particular unit.
17. Concentrations beyond the greatest scale factor of the instrument or in excess of 30 percent (0.3) LEL of the sample component require system modification. Similar modification may be necessary for sampling in oxygen-deficient atmospheres. This usually entails increasing the combustion air to the detector by sample dilution or by an independent air supply. A dilution system is simply the apparatus required to supply a filtered, controlled air supply for analyzers that use the sample gas stream as the source of combustion air. A dilution system can, by selection of various critical orifices, dilute a gas stream by ratios up to 100:1.
18. Always be sure that carrier gas flow (usually sample gas) is initiated prior to lighting the detector flame.

Sources

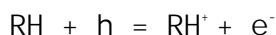
Analabs, A Unit of Foxboro Analytical. "Operating and Service Manual for Century Systems' Portable Organic Vapor Analyzer (OVA) Model OVA-108 and Optional Accessories, Revision C," North Haven, Connecticut.

4.2.4 METHOD IV-4: MONITORING TOXIC GASES AND VAPORS USING A PHOTOIONIZATION DETECTION

Discussion

This method is designed to detect, measure and record real-time levels of many organic and inorganic vapors in air. A photoionization detector (PID) will respond to most vaporous compounds in air that have an ionization potential less than or equal to that supplied by the ionizing source in the detector, an ultraviolet lamp. The magnitude of this response is a function of the detector sensitivity and the concentration and ionization properties of the individual compound. Though it can be calibrated to a particular compound, the instrument cannot distinguish between detectable compounds in a mixture of gases, and therefore indicates an integrated response which is a function of the response factors and concentrations of all ionizable species present.

The analyzer employs the principle of photoionization for detection. This process is termed photoionization since the absorption of ultraviolet light (a photon) by a molecule leads to ionization as shown in the equation:



where RH = trace gas,

h = a photon with an energy \geq Ionization Potential of RH

The sensor consists of a sealed ultraviolet light source that emits photons which are energetic enough to ionize many trace species (particularly organics) but do not ionize the major components of air such as O₂, N₂, CO, CO₂, or H₂O. A chamber adjacent to the ultraviolet source contains a pair of electrodes. When a positive potential is applied to one electrode, the field created drives any-ions formed by absorption of UV light to the collector electrode where the current (proportional to concentration) is measured. This signal is amplified and conditioned and then sent to the output display.

To minimize adsorption of various sample gases, the ion chamber is usually made of an inert fluorocarbon material. The sample line is kept as short as possible, and a rapid flow of sample gas is maintained through the ion chamber volume.

Uses

The portable photoionization detector is useful as a general survey instrument at waste sites and hazardous material spills. As such, it is similar to an FID in application; however, its capabilities are somewhat broader in that it can detect certain inorganic vapors. Conversely, the PID is unable to respond to certain low molecular weight hydrocarbons (e.g., methane and ethane) that are readily detected by FID. In addition, certain toxic gases and vapors (e.g., carbon tetrachloride, HCN) have high ionization potentials and cannot be detected with a PID.

Procedure for Use

The procedural steps delineated herein are intentionally general. The operating manual for the unit being used should be consulted for specific instructions.

1. Check battery charge level. If in doubt, charge battery as described in manual.
2. Turn unit on. Verification of UV lamp operation can be made by looking into sensor for purple glow of the lamp.
3. Perform zero and calibration procedure as described in operating manual. Calibration for specific compounds can be performed so that instrument response is proportional to the calibration gas concentration.
4. If so equipped, set alarm at desired level.
5. Once calibrated, unit is ready for use.
6. Position intake assembly in close proximity to area in question as the low sampling rate allows for only very localized readings.
7. A slow sweeping motion of intake assembly will help prevent the by-passing of problem areas.
8. Be prepared to evacuate the area if preset alarm sounds. Operators utilizing supplied air systems may not need to consider this action.
9. Static voltage sources such as AC power lines, radio transmissions, or transformers may interfere with measurements. See operating manual for discussion of necessary considerations.
10. Regular cleaning and maintenance of instrument and accessories will assure representative readings.
11. As with any field instrument, accurate results depend on the operator being completely familiar with the operator's manual for the unit in use.

Sources

HNU Systems Inc., "Instruction Manual for Model PI 101 Photoionization Analyzer." 1975.

4.2.5 METHOD IV-5: USE OF PORTABLE, FIELD-OPERABLE GAS CHROMATOGRAPHS

Discussion

The use of field portable gas chromatography (GC) for obtaining data on concentrations of certain volatile organic compounds in ambient air in and around waste sites has been demonstrated.³ While their ability to provide unambiguous identification and quantitation of target compounds may be limited by such factors as interferences, ambient conditions, and operator experience, the data supplied in preliminary assessments using these instruments may be used in determining air sampler placement and approximate compound concentrations.

Gas chromatography is a technique in which components of a mixture are separated in the gas phase using a solid phase sorbent. The mixture is placed on the front end of GC column (generally a 1/8" stainless steel tube packed with the appropriate sorbent) and flushed through the column with an inert carrier gas. Compounds are eluted from the column according to such factors as their affinity for the sorbent and volatility, and routed into a detector, which may be designed to detect compounds having specific properties or may respond to a more general class.

Identification of compounds is generally based on elution time from the column. This retention time is dependent upon a number of factors; however if these factors are held constant, retention times for individual compounds will hold fairly steady. Standard mixtures of the compounds of interest are run to determine retention times for the target compounds, and sample runs are compared to identify specific eluting peaks.

While factors affecting retention time can be held constant in laboratory settings, this may not be possible under field conditions, where lack of power and variable environmental conditions may force compromises in the analysis. A major factor affecting retention time, for instance, is temperature of the GC column. Since the field portable GC's described here are designed to work from battery power, sufficient energy may not be available to maintain the column at constant temperature. Variations in ambient temperature will then make retention times shift, making identifications ambiguous. Calibrations can be run at several column (ambient) temperatures to provide a family of curves, thus, reducing bias from temperature changes.

Another element contributing to ambiguous identification is complexity of the sample. Compounds eluting with close or similar retention times may give "false positive" identifications or false high quantitations. Quantitations are based on comparison of response of sample components to response of standards of known concentrations. Again, these values may be inaccurate due to uncontrollable variables such as environmental conditions or sample complexity.

Several detectors are available for gas chromatography, ranging from the very simple (such as thermal conductivity) to the more complex (such as mass spectrometry). Due to power restrictions and other restraints, such as size

and reagent availability, field instruments have been limited to about three or four detectors. The most common are:

- Flame Ionization Detectors (FID). The FID will respond to most organic compounds as they form positively charged ions when combusted in a hydrogen flame. The magnitude of the response is a function of the detector sensitivity and the ionization properties of the particular compound as well as its concentration. As a result, this signal must be compared to that generated by calibration with a known concentration of a standard. The output of the detector is generally recorded on a strip chart recorder as intensity versus retention time, producing a GC "peak". The area under the peak (using an integrator) or the peak height at maximum intensity can be used for quantitation. Maximum sensitivity is generally in the mg/m^3 (ppm) range.
- Photo-Ionization Detector (PID). The PID also ionizes sample introduced into it and responds to positive ions produced by an ultraviolet light source. Again, the magnitude of response is dependent on concentration and ionization properties of the compound. Response is measured as with the FID.

The PID offers two advantages over the FID. First, it is sensitive to some compounds to the $\mu\text{g/m}^3$ (ppb) range, especially light aromatics such as benzene, toluene, and xylene. Secondly, at least one model (the HNu Model 301 Portable GC) can be equipped with lamps of differing ionization potential, providing some degree of selectivity. Compounds with ionization potentials above that of the energy of the lamp being used exhibit vastly reduced response as compared to compounds with lower ionization potential. For instance, toluene (ionization potential 8.8 eV) will respond strongly when ionized by a 9.5 eV lamp, while n-heptane (10.08 eV) will exhibit a greatly reduced response. To assist in lamp selection, a listing of ionization potentials for various compounds is typically included in the owner's manual.

The instruments listed below have been designed to be field portable or are easily adaptable to field use, e.g. by addition of a battery pack or small gas cylinders.

- Century Systems Model OVA-138 Organic Vapor Analyzer (OVA).⁴ The OVA-128 has two models: a "survey mode" to provide nonspecific quantitation and/or detection of organic vapors (described more fully in Method IV-3) and a GC mode, in which a gas chromatography column is attached to the detector to possibly provide identification of the vapor constituents. The OVA utilizes a flame ionization detector (FID) which, in the survey mode, will yield sensitivities to 0.1 ppm (methane). The instrument maintains its own power and gas supplies.
- Model 10A10 Photovac. This system uses a PID with a fixed ionization potential. Sensitivities to 0.1 ppb for certain compounds have been

reported.^{3,5} The system is almost completely self-contained, with internal power and gas supplies, with only a strip chart recorder external to the main body of the instrument.

- HNu Model 301 Gas Chromatography. This unit is essentially a compact laboratory instrument which is made field portable with the addition of a "field pack", containing battery and gas supplies. Field-usable detectors include an FID and a PID with a selection of lamps ranging from 9.5-11.7 eV.
- Sentex Sensing Technology Scentor Automated Gas Chromatography. This is a relative new instrument utilizing an Argon Ionization Electron Capture or Flame Ionization Detector. Sampling and analysis are completely automated. Samples are collected on a sorbent cartridge, then thermally desorbed into the GC column. The instrument maintains a known standard gas internally for use in quantification. Sensitivities for most organic compounds are reported at the low ppb range and low part-per-trillion range for polar compounds.⁶

Uses

In theory, any compound which can pass through a gas chromatographic column as a discrete "peak" and is capable of being detected by the detector, is amenable to this method. In practice, this may not always be the case. A partial list of compounds measured in the field with portable GCs is given in Table 4-1. Some of the factors which could be considered before using a field portable GC in a field survey are as follows:

1. Column Selection. Selection of appropriate gas chromatography column packing and column length is necessary. A critical element in this selection is the ability of the column to elute the compound(s) of interest as a discrete "plug" at the temperature at which the column will be operated.

Successful use of two-column packings has been reported for general purpose use:⁴ a 10 percent OV-101 on 60/80 mesh Chromasorb-W/AW-DCMS treated and a 1 percent TCEP on 60/80 mesh Chromasorb W-HP. Spittler³ has reported use of a 12-inch carbowax column for rapid determination of volatile loading and a 4-ft SE30 column for more efficient separation and quantitation.

2. Compound Volatility. Generally, compounds exhibiting a vapor pressure of less than 1 mm (Hg) at 20°C will be troublesome to measure with a field-portable GC. Reasons for this include adsorption of the components on unheated syringe walls, the inability to elute the compound from the GC column at ambient temperatures, and the GC's inability to measure particulate-bound organic matter. Compounds in this class include PCBs, PAH, and most pesticides.

TABLE 4-1. COMPOUNDS SHOWN TO BE AMENABLE TO FIELD GC ANALYSIS⁴

Di chl oromethane (methy l ene chl ori de)
Tri chl oromethane (chl oroform)
Carbon tetrachl ori de
Di chl oroethane (ethy l ene di chl ori de)
1,2-trans di chl oroethy l ene
Tri chl oroethy l ene (TCE)
1,1,1-tri chl oroethane
Tetrachl oroethy l ene
Di methyl ketone (acetone)
Methy l ethy l ketone (MEK)
Methy l isobuty l ketone (MI BK)
Benzene
Methy l benzene (tol uene)
Chl orobenzene
Ethy l benzene
Ni tro benzene
1,2-di methyl benzene (O-xyl ene)
1,3-di methyl benzene (m-xyl ene)
1,4-di methyl benzene (p-xyl ene)
Ethy l acetate

The ability of a portable GC to provide unambiguous data may be limited. It can, however, with proper use and data interpretation, be used to detect and identify components of pockets of gaseous hydrocarbons in depressions or confined spaces, vapors in drums or other containers, or in ambient air.

Procedures for Use

The procedures presented in this section are intended to apply to any portable GC; therefore, detailed operating instructions must be obtained from the operating manual of the specific unit to be used. Some procedures, such as the preparation of standards, can be used with any instrument, and these are described here.

1. Standard Preparation

GC standards in air are prepared by using quantities of pure solvent which are small enough to vaporize completely in a 40 ml VOA vial at ambient pressure and temperature. To do this, 1 μ l of the pure standard is injected through the septum into the 40 ml vial. The resultant concentration is calculated by:

$$C = \frac{A \times D}{V + A}$$

where: C = compound concentration (in μ g/cc)

A = amount injected (in μ l)

D = density of the standard (in g/ml)

V = volume of the vial and the air in it (in liters)

For example, 1 μ l of toluene is injected into a clean 40 ml VOA vial and allowed to vaporize. The resultant concentration is then calculated as:

$$\frac{(1 \mu\text{l}) \times (0.866 \text{ g/ml})}{(0.4 \text{ l})} = 2.2 \mu\text{g/cc}$$

This standard can then be used to prepare standards of lesser concentration by further dilution with air on a volume/volume basis.

Several standards can be injected into one vial to make a multi-component standard to save calibration time. Care should be taken in component selection to prevent the resultant chromatograms from being too complex to determine individual compound responses. To extract a portion of the vapor, penetrate the septum with a clean syringe and fill to the desired volume. Remove the syringe and the standard is ready for injection into the chromatography. Total volume removed should not exceed 1 ml. Volumes in excess of 1 ml will detrimentally effect reproducibility of standards.

2. Sample Collection

Samples can be collected and introduced into the GC by either of two basic techniques: direct injection with a gas-tight syringe or, when so equipped, through the gas sampling loop.

An air sample can be obtained with a gas-tight syringe at the sampling site. The syringe is filled and emptied several times in the contaminated atmosphere. The syringe is filled once again and the sample is carried off site to an area where the GC is set up for field analysis. The sample can then be injected into the GC column, a chromatogram produced, and further qualitative/quantitative analysis performed. Sample volumes of 10 μ l to 1 ml can be used.

Syringes may become contaminated if high concentrations of organics are encountered. If a gas-tight syringe becomes contaminated, the easiest method of cleaning is to bake it in the oven at 105°C overnight. It may be possible to clean the syringe by removing the plunger, inserting the needle into the injection port, and allowing carrier gas to flow through it for several minutes.

Some instruments are equipped with a gas sampling loop to facilitate sample collection and injection. This device is essentially a length of steel tubing with a known volume that is fitted to the head of the chromatographic column with a two-way valve. With the valve in the "load" position sample gas can be drawn through the loop with a pump, then when the valve is switched to the "inject" position, the loop is isolated from the pump system and the carrier gas is diverted into the loop to sweep its contents onto the column. The sample loop improves the consistency of the injections by assuring a constant volume (provided temperature and pressure are constant) and a consistent injection speed.

3. Operation

- a. Check battery charger level indicator; if in doubt, recharge battery as described in the manual.
- b. Turn instrument on and allow adequate warmup time.
- c. Follow operating procedures for lighting FID flame (if used), lighting PID lamp (if used), establishing carrier gas flow, zeroing recorder response, etc.
- d. Using the procedure described below, inject an appropriate amount of the standard described in Section 1. Hold the syringe in two hands, using one to guide the needle into the septum and the other to provide force to pierce the septum and to prevent the plunger from being forced out by the pressure from the GC. Insert the needle through the septum as far into the injection port as possible, swiftly and smoothly depress the plunger, hesitate one second, and withdraw the needle.

- e. Mark injection time and sample I.D. on output recorder strip chart, and start timer for monitoring retention times. Record pertinent parameters in analytical logbook for documentation of analytical conditions.
 - f. Record retention times and peak heights (or integrated peak areas if an integrator is available) of each eluting peak.
 - g. Chromatograms of blank injections should be made at frequent intervals, especially after running a sample with a high contaminant loading, to guard against syringe contamination.
 - h. At least one standard run should be made on each day of analysis, or when conditions change sufficiently to warrant re-calibration.
4. Compound Identification

Injection of standard mixtures of compounds will allow determination of retention time for each of the compounds. The elution order of the compounds in these mixtures will remain constant for a given GC column and should be determined prior to field use. Tables 4-2 through 4-4 give elution orders and approximate retention times for several compounds on several columns.

After both standards and samples have been run, comparison of the samples to standards should be made. If retention time matches are noted, another column is installed in the GC and standards and samples re-run, with subsequent comparisons made. A retention-time match on two different columns provide a fairly reliable identification, however, given the purpose of this screening, the data should be labeled "tentative" or "preliminary".

To save analytical time, the following strategy has been used. Since shorter GC columns result in shorter retention times at the cost of lower resolution between peaks, a short column is used to do the first set of analyses, and longer columns are used to verify compound identity and quantitation.

5. Quantitation

Following tentative identification of compounds in the sample, a determination of quantitation can be made. This can be based either on peak height or on the integrated area under the peak if an integrator is used.

First, using the response to the standard, calculate a response factor as:

$$RF = \frac{R}{V \times C}$$

TABLE 4-2. SELECTED RETENTION TIMES 8-INCH 3 PERCENT DIISODECYL PHTHALATE ON CHROMASORB W

0°C		40°C	
Compound	Time, Seconds	Compound	Time, Seconds
Freon 113	15.3	Pentane	11.7
Pentane	18.0	Vinylidene Chloride	12.6
Ethhanethiol	18.0	Hexane	12.6
Vinylidene Chloride	19.8	Freon 113	12.6
Hexane	21.6	Ethrane	12.6
Methylene Chloride	26.1	Ethanethiol	12.6
Methyl Acetate	30.6	Methylene Chloride	14.4
Acrylonitrile	31.5	Vinyl Acetate	15.3
Vinyl Acetate	32.4	Trichloroethane 1,1,1	15.3
Ethrane	36.0	Methyl Acetate	16.2
Methyl Alcohol	36.9	Halothane	17.1
Halothane	36.9	Chloroform	17.1
Acetonitrile	36.9	Acetonitrile	17.1
Acetone	36.9	Heptane	18.9
Trichloroethane 1,1,1	40.5	Ethyl Acetate	18.9
Heptane	45.9	Acrylonitrile	18.9
Methyl Acrylate	48.6	Methyl Acrylate	19.8
Ethyl Acetate	49.5	Ethylene Dichloride	19.8
Benzene	49.5	Acetone	19.8
Chloroform	51.3	Methyl Ethyl Ketone	21.6

(continued)

TABLE 4-2 (continued)

0°C		40°C	
Compound	Time, Seconds	Compound	Time, Seconds
Tetrahydrofuran	65.7	Methyl Alcohol	21.6
Ethylene Dichloride	66.6	Benzene	21.6
Trichloroethylene	76.5	Tetrahydrofuran	22.5
Methyl Ethyl Ketone	76.5	Trichloroethylene	23.4
Ethyl Acrylate	95.4	Ethyl Acrylate	26.1
Methyl Methacrylate	100.8	Methyl Methacrylate	27.0
Toluene	133.2	Toluene	37.8
Tetrachloroethylene	171.0	Propanol, N	38.7
Propanol, N	182.7	Methyl Isobutyl Ketone	39.6
Methyl Isobutyl Ketone	278.1	Tetrachloroethylene	43.2
Trichloroethane 1,1,2	313.2	Trichloroethane 1,1,2	54.0
Ethylene Dibromide	320.4	Ethylene Dibromide	59.4
Ethyl Benzene	369.9	Ethyl Benzene	67.5
Styrene	677.7	Butyl Acrylate	108.0
Pyridine	1000.0	Styrene	117.0
Pentanol	1000.0	Pentanol	163.8
Butyl Acrylate	1000.0	Pyridine	1000.0

(continued)

TABLE 4-3. SELECTED RETENTION TIMES 8-INCH 10 PERCENT OV-101 ON CHROMASORB W GC COLUMN

0°C		40°C	
Compound	Time, Seconds	Compound	Time, Seconds
Methyl Alcohol	21.3	Pentane	11.5
Ethanethiol	23.9	Ethrane	12.4
Vinylidene Chloride	25.2	Ethanethiol	12.4
Pentane	25.2	Methyl Acetate	13.3
Ethrane	25.7	Acetone	13.3
Acetone	27.5	Methylene Chloride	14.2
Methylene Chloride	30.1	Methyl Alcohol	14.2
Methyl Acetate	30.1	Halothane	14.2
Freon 113	30.1	Freon 113	14.2
Acetonitrile	30.1	Vinylidene Chloride	14.4
Acrylonitrile	31.0	Acetonitrile	15.9
Halothane	35.4	Vinyl Acetate	16.9
Vinyl Acetate	43.2	Acrylonitrile	16.8
Methyl Ethyl Ketone	48.7	Methyl Ethyl Ketone	18.6
Propanol, N	49.5	Hexane	18.6
Hexane	49.6	Ethyl Acetate	18.6
Chloroform	55.8	Chloroform	18.6
Methyl Acrylate	56.7	Propanol, N	19.8
Ethyl Acetate	59.4	Ethylene Dichloride	21.3
Tetrahydrofuran	64.8	Methyl Acrylate	21.3

(continued)

TABLE 4-3 (continued)

0°C		40°C	
Compound	Time, Seconds	Compound	Time, Seconds
Ethylene Dichloride	68.2	Tetrahydrofuran	22.5
Trichloroethane 1,1,1	70.2	Trichloroethane 1,1,1	23.0
Benzene	78.8	Benzene	24.8
Trichloroethylene	121.5	Ethyl Acrylate	25.7
Ethyl Acetate	124.9	Trichloroethylene	27.5
Heptane	134.6	Heptane	30.1
Methyl Methacrylate	141.7	Methyl Methacrylate	31.0
Methyl Isobutyl Ketone	182.5	Methyl Isobutyl Ketone	37.2
Trichloroethane 1,1,2	225.0	Trichloroethane 1,1,2	41.6
Toluene	241.2	Toluene	45.0
Pyridine	254.7	Methyl Acrylate	56.7
Ethylene Dibromide	291.4	Ethylene Dibromide	56.7
Pentanol	360.0	Tetrachloroethylene	65.7
Tetrachloroethylene	367.2	Pentanol	77.4
Ethyl Benzene	571.4	Pyridine	85.5
Styrene	916.2	Ethyl Benzene	94.8
Butyl Acrylate	1000.0	Styrene	118.8

TABLE 4-4. SELECTED RETENTION TIMES 8-INCH 1 PERCENT TCEP ON CHROMASORB W-HP

0°C		40°C	
Compound	Time, Seconds	Compound	Time, Seconds
Pentane	13.5	Pentane	11.7
Freon 113	16.2	Hexane	12.6
Hexane	19.8	Freon 113	12.6
Vinylidene Chloride	21.6	Vinylidene Chloride	14.1
Heptane	26.1	Ethanethiol	14.4
Ethanethiol	26.1	Heptane	16.2
Trichloroethane 1,1,1	51.3	Halothane	18.9
Halothane	57.6	Ethrane	18.9
Methylene Chloride	63.0	Trichloroethane 1,1,1	21.6
Methyl Acetate	79.2	Methylene Chloride	23.4
Ethrane	81.9	Methyl Acetate	23.4
Trichloroethylene	84.6	Trichloroethylene	27.9
Tetrahydrofuran	97.2	Acetone	27.9
Acetone	98.1	Ethyl Acetate	28.8
Vinyl Acetate	99.0	Vinyl Acetate	29.7
Benzene	102.6	Tetrahydrofuran	29.7
Chloroform	107.1	Chloroform	30.6
Ethyl Acetate	126.0	Benzene	32.4
Methyl Acrylate	138.6	Tetrachloroethylene	36.0
Methyl Alcohol	143.1	Methyl Ethyl Ketone	39.6

(continued)

TABLE 4-4 (continued)

0°C		40°C	
Compound	Time, Seconds	Compound	Time, Seconds
Tetrachloroethylene	152.1	Methyl Acrylate	41.4
Methyl Ethyl Ketone	179.1	Methyl Alcohol	45.0
Acrylonitrile	202.5	Toluene	48.6
Ethylene Dichloride	225.0	Acrylonitrile	49.5
Toluene	232.2	Ethylene Dichloride	50.4
Methyl Methacrylate	247.5	Ethyl Acrylate	50.4
Ethyl Acrylate	247.5	Methyl Methacrylate	54.9
Acetonitrile	324.0	Propanol, N	61.2
Propanol, N	340.2	Methyl Isobutyl Ketone	69.3
Methyl Isobutyl Ketone	450.0	Acetonitrile	69.3
Ethyl Benzene	464.4	Ethyl Benzene	87.3
Trichloroethane 1,1,2	1000.0	Ethylene Dibromide	162.9
Styrene	1000.0	Butyl Acrylate	165.6
Pyridine	1000.0	Trichloroethane 1,1,2	167.4
Pentanol	1000.0	Styrene	208.8
Ethylene Dibromide	1000.0	Pentanol	216.0
Butyl Acrylate	1000.0	Pyridine	1000.0

where: RF = response factor

C = concentration of the standard (µg/ml)

V = volume of standard injected (ml)

R = response to the standard (in integrated counts or measured linear dimensions).

Sample concentrations can then be estimated by:

$$C^1 = \frac{R^1}{RF \times V^1}$$

where: C¹ = sample concentration (µg/ml)

R¹ = response of sample peak (in the same units as used for the standard)

V¹ = volume of sample injected (ml)

RF = calculated response factor.

4.2.6 METHOD IV-6: STAIN DETECTOR TUBE METHOD FOR SAMPLING GASEOUS COMPOUNDS

Discussion

A relatively simple method for determining concentrations of specific gaseous pollutants is through the use of stain detector tubes. They are usually calibrated in ppm for easy interpretation and are either direct reading or referenced to a supplied concentration scale or color change chart. The limiting factors in the application of this methodology are the small number of compounds for which detector tubes are available, interfering agents and cross-sensitivities, short sampling time, and the extremely small sample volume used. Most detector tubes are species specific; however, some detect groups of compounds, e.g., "total hydrocarbons."

The detector tubes are specific for individual compounds and require specific sampling techniques. This information is supplied with the tubes and details the required sample volume, proper tube preparation and insertion into the pump, and a discussion of the applicability and limitations of the tube. In general the tubes are opened by snapping off the tips on either end and inserting them into the pump so that the arrow on the tube indicates flow toward the pump. The required sample volume is then pulled through the tube. An indicator chemical in the tube will demonstrate a color change, the length of which is proportional to the concentration of the compound in question.

The detector tube and pump are the two major components of the system. Pumps used for drawing air through the tubes come in two basic forms: bellows pump and piston-type (syringe). These pumps are manufactured under strict specifications so as to draw only a specified volume of gas and are designed to be used with tubes of the same manufacturer.

Uses

Stain detector tubes are useful for screening sources to verify the Presence of suspected compounds and to provide some degree of quantification. They are generally inadequate for ambient air sampling applications due to the low sample volumes collected. They are more useful for detection of compounds at higher levels such as in drums, confined work areas, pockets or depressions, etc.

Procedure for Use

1. Perform necessary pump leak check procedures. This is usually accomplished by plugging pump inlet, drawing a vacuum on the pump, holding it for at least 1 minute and determining visually if leak allows bellows to inflate or piston fails to return completely into pump. The pump can be plugged using a sealed detector tube.

2. Break open both ends of detector tube, insert correct end into pump, and sample according to instructions. Most tubes have some kind of indicator (i.e., arrow, prefilter) that helps determine which end of tube is the inlet. The direction of the concentration scale is also a guide.
3. Visually inspect tube for color changes and record corresponding gas concentration.
4. Additional Notes
 - Prior to use, check tube expiration date, because most have a defined shelf life.
 - Some tube manufacturers advise that tubes showing negative results can be reused before they are rendered useless. The error potential and risk associated with reusing a previously opened tube is not advisable when working with hazardous materials.
 - Some types of detector tubes have reagent ampules which must be broken to activate the indicator. Also, some procedures call for use of multiple tubes, in series for multiple parameter detection, or specific interference removal.
 - The standard range of measurement or the detector sensitivity can sometimes be extended by changing the number of pump volumes pulled through the tube. The upper range limit can be extended by decreasing the number of pump volumes, and the lower range limit can be extended by increasing the number of pump volumes.
 - Tubes and pumps of different manufacturers should not be used interchangeably. For example, Drager tubes should be used only with Drager pumps.

Sources

Dragerwerk Ag Lubeck. "Detector Tube Handbook, Air Investigations and Technical Gas Analysis with Drager Tubes." 4th Edition, August 1979.

Matheson Safety Products. Operating Instructions for Matheson-Kitagawa Detector Tubes, Matheson Gas Products Model 8014 - Toxic Gas Detector.

4.2.7 METHOD IV-7: SAMPLING FOR VOLATILE ORGANICS IN AMBIENT AIR USING SOLID SORBENTS

Discussion

Solid sorbent cartridges can be used quite successfully to collect samples of volatile organics in ambient air and workplace environments. The sample apparatus consists of a sampling cartridge packed with a solid sorbent of desirable characteristics (e.g., Tenax-GC, activated charcoal, XAD-2) and a pump system capable of maintaining a constant flow rate across the collection media for a specified period of time.

In principle, organic vapors present in the air are adsorbed on the collection media and subsequently desorbed, thermally or chemically, in the laboratory. An aliquot of the desorbed sample is then subjected to chromatographic analysis (either capillary or packed column) followed by flame ionization or mass spectrometric detection.

Although several sorbents or sorbent combinations have been utilized for collection and concentration of volatile organic species, at present the porous polymer, Tenax-GC, is the most widely studied for a wide variety of compounds at concentrations typically found in ambient air. Tenax-GC is hydrophobic, thermally stable up to 360°C, and permits thermal desorption of organic species with volatility greater than n-eicosane at temperatures of 280°C.⁷ Glass or glass-lined stainless-steel sampling cartridges of various sizes and configurations are available and can be purchased prepacked or packed to specifications in the laboratory. In any case, the sorbent and/or prepacked tubes must be thoroughly precleaned, conditioned, and checked for freedom from interferences prior to use.

Other sorbents or combination of sorbents may be applied with equal success depending upon the nature of the ambient environment and the specific species of volatiles under investigation. Monsanto Research Corporation reports success with a combination sorbent system based on Tenax-GC, Porapak R, and Amborsorb XE-340 which has been used to collect a broad range of organic compounds.⁸ NIOSH procedures may also be used and the "NIOSH Manual of Analytical Methods"⁹ should be consulted where applicable. Finally, if the detection of specific organics is desired, the characteristics of the compound and sorbent of interest should be researched^{9,10,11} and all sampling parameters adjusted to meet these criteria.

The recommended procedure^{13,14,15} involves precleaning a batch of Tenax by Soxhlet extraction in methanol first, and then pentane for 24 hours each. The sorbent is then oven dried, packed in tubes and conditioned under carrier gas flow at 270°C (desorption conditions for 4 hours). The conditioning can be performed a final time (1 hour run) just prior to use. Cartridges are then stored in Teflon-capped culture tubes packed in aluminum foil, and then in 1 gallon "paint cans," for shipment to and from the field. Culture tubes should be wrapped in foil to limit exposure of sampling cartridges to UV light. Analysis should be instituted as quickly as possible in order to prevent sample

degradation. Schlitt, et al.,⁷ recommend a maximum storage time of 48 hours; however, this is often impractical, and a maximum storage period of 30 days has been used successfully in a previous study.^{15, 16} In any event, sorbent cartridges should be transported in solvent free coolers packed with "blue ice" and stored at 4°C while awaiting analysis.

The outlined procedure utilizes a borosilicate glass tube, outside diameter 16 mm (5/8") by 10 cm in length. The tube is packed with 1.2 grams of Tenax-GC sorbent with a plug of glass wool at each end (double plug at inlet). The personal monitoring pumps can be any low-flow model capable of maintaining consistent flows at the rates prescribed. Personnel monitoring pumps are available from a number of vendors and range in sophistication (and, accordingly, price) from very simple models to programmable ones capable of compensating for increasing pressure differential in addition to other features. Care should be taken to select a pump capable of operating in the desired flow rate range and which has features most likely to be used by the investigator.

Due to the wide range of volatility and breakthrough volumes of compounds amenable to this method, it is recommended that samples be collected at widely different flow rates and therefore different volumes. This approach will guard against analytical system overload and breakthrough of the more volatile organics while at the same time maintaining maximum sensitivity for all compounds. To achieve this, four (4) samplers are placed at each desired location and run for a specified time period (usually 4-8 hours). As a general guideline, one sampler should be operated within each of the following ranges.

<u>Flow Rate Range</u>	<u>Total Volume</u>
10- 20 cc/min	5-10 liters
20- 40 cc/min	10-20 liters
40- 80 cc/min	20-40 liters
80-160 cc/min	40-80 liters

This approach is based upon Standard Operating Procedures (SOPS) in use at EPA's Environmental Monitoring Systems Laboratory (EMSL) at Research Triangle Park (RTP), North Carolina. It is highly recommended for those cases where little is known about the compounds and concentrations present at a given site. The procedure can be costly, however, due to the potential number of samples that could be generated for subsequent analysis. Time and cost constraints may therefore dictate a compromise (collection of fewer samples/location) which may unfortunately jeopardize the quality of the resultant data. It is therefore recommended that as much background information concerning suspected compounds and approximate concentrations be collected as possible before deciding on a compromised approach. It is also recommended that at a minimum, the delineated procedure be adhered to for undefined situations.

Uses

The method outlined below has been successfully used for quantitative analysis of the compounds listed in Table 4-5 in ambient air and is based on procedures used successfully for qualitative and quantitative analysis during several programs. A brief review of the literature^{7, 17, 18, 19, 20, 21} reveals that a number of additional compounds (Table 4-6) have been analyzed either qualitatively or quantitatively using modifications of the procedure described herein. Other additional compounds may be amenable to the method, and to aid in determining both applicability and appropriate sampling volumes, a list of retention volumes for the described sorbent tube has been included as Table 4-7. It should be noted, however, that since many of the compounds included in these lists have not been analyzed using this specific method, testing or further review should be performed to perfect and prove the method for these compounds prior to actual sampling and analysis.

Procedures for Use

1. Calibration of Sampling Pumps Equipped with Rotameter, Needle Valve Combination--
 - a. Select a set of sampling pumps and assemble necessary equipment (see Figure 4-1). Calibrate each pump as follows:
 - b. Measure ambient air temperature, barometric pressure and relative humidity. Determine water vapor pressure from tables.
 - c. If battery test is available, check battery.
 - d. Place "calibrator" sorbent cartridge in line and start pump. Allow pump to stabilize. Do not use this "calibrator" cartridge for actual sampling, however, it can be reused for additional calibration runs.
 - e. Determine actual flow rate, f , with bubble tube flowmeter.

$$f = \frac{\text{Distance traveled (ml)}}{\text{Travel time (min)}}$$

- f. Adjust flow rate of the sampler to the desired rate by adjusting the needle valve. Verify that the flow rate has been achieved by checking against bubble tube three times. Calculate a mean value by summing the values of the three individual readings and dividing by three. The deviation of the individual flow rates from the mean flow rate should not exceed ± 5 percent.

TABLE 4-5. COMPOUNDS SUCCESSFULLY MONITORED USING
TENAX SAMPLING PROTOCOLS²²

2-Chl oropropane	1-Bromo-3-chl oropropane
1, 1-Di chl oroethene	Ethyl benzene
Bromoethane	Bromoform
1-Chl oropropane	Ethenyl benzene
Bromochl oromethane	o-Xyl ene
Chl oroform	1, 1, 2, 2-Tetrachl oroethane
Tetrahydrofuran	Bromobenzene
1, 2-Di chl oroethane	Benzal dehyde
1, 1, 1-Tri chl oroethane	Pentachl oroethane
Benzene	4-Chl orostyrene
Carbon tetrachl ori de	3-Chl oro-1-propene
Di bromethane	1, 4-Di chl orobutone
1, 2-Di chl oropropane	1, 2, 3-Tri chl oropropane
Tri chl oroethene	1, 1-Di chl oroethane
1, 1, 2-Tri chl oroethane	2-Chl orobutane
2, 3-Di chl orobutane	2-Chl oroethyl vi nyl ether
Bromotri chl oromethane	1, 1, 1, 2-Tetrachl oroethane
Tol uene	p-Di oxane
1, 3-Di chl oropropane	Epi chl orohydron
1, 2-Di bromethane	1, 3-Di chl orobutane
Tetrachl oroethene	p-Di chl orobenzene
Chl orobenzene	ci s-1, 4-Di chl oro-2-butene
1, 2-Di bromopropane	n-Butyl benzene
Ni trobenzene	3, 4-Di chl oro-1-butene
Acetophenone	1, 3, 5-Tri methyl benzene
Benzoni tri te	
I so-propyl benzene	
p-I sopropyl tol uene	

TABLE 4-6. LITERATURE SUMMARY - VOLATILE ORGANICS AMENABLE
TO COLLECTION BY TENAX SORBENT CARTRIDGES

Component	Reference(s)
N-Nitroso dimethyl amine	17
β-Propiolactone	17
Ethyl methanesulfonate	17
Nitromethane	17
Glycidaldehyde	17
Butadiene diepoxide	17
Styrene Epoxide	17
Aniline	17
Bis (chloromethyl) ether	17
Bis (2-chloromethyl) ether	17
Diethyl Sulfate	18
Acrolein	18
Propylene Oxide	18
Cyclohexene Oxide	18
Styrene Oxide	18
Acetophenone	18, 7
Methanol	20
Ethanol	20
Propanol	20
Ethyl Acetate	20
Acetone	20, 7
1, 2, 4-Tri chlorobenzene	21
1, 2, 3, 5-Tetrachlorobenzene	21
Hexachlorobenzene	21
p-Chlorophenol	21
2, 4, 6-Tri chlorophenol	21
Diphenyl Oxide	21
o-Phenyl phenol	21
Pentachlorophenol	21
n-Pentane	7
2-Methyl pentane	7
3-Methyl pentane	7
n-Hexane	7
Heptane	7
n-Octane	7
n-Nonane	7
Propyl benzene	7
Trimethyl benzene	7
o-Ethyl toluene	7

TABLE 4-7. APPROXIMATE RETENTION VOLUMES AT 38°C (100°F)
(liters/gram of Tenax)

=====		
Amines	dimethylamine	0.8
	isobutylamine	9
	t-butylamine	0.8
	di-(n-butyl)amine	1200
	pyridine	56
Ethers	diethyl ether	4
	propylene oxide	3
Esters	ethyl acetate	20
	methyl acrylate	20
	methyl methacrylate	70
Ketones	acetone	3
	methyl ethyl ketone	10
	methyl vinyl ketone	10
	acetophenone	860
Aldehydes	acetaldehyde	0.6
	benzaldehyde	920
Alcohols	methanol	0.3
	n-propanol	4
	allyl alcohol	5
Aromatics	benzene	19
	toluene	97
	ethyl benzene	200
	cumene	440
Hydrocarbons	n-hexane	5
	n-heptane	20
	2,2-dimethylbutane	0.1
	2,4-dimethylpentane	20
	4-methyl-1-pentane	3
	cyclohexane	8
	methyl chloride	2
	methyl bromide	0.8
Halogenated Hydrocarbons	vinyl chloride	0.5
=====		

(continued)

TABLE 4-7 (continued)

=====		
	methylene chloride	2
	chloroform	8
	carbon tetrachloride	8
	1,2-dichloroethane	10
	1,1,1-trichloroethane	6
	tetrachloroethene	80
	trichloroethene	20
	1, chloro-2-methyl propene	6
	3-chloro-2-methyl propene	7
	1,2-dichloropropane	30
	1,3-dichloropropane	90
	epichlorohydrin (1-chloro-2,3-epoxy propane)	30
	3-chloro-1-butane	5
	allyl chloride	4
	4-chloro-1-butene	10
	1-chloro-2-butene	20
	chlorobenzene	150
	o-dichlorobenzene	300
	m-dichlorobenzene	400
	benzyl chloride	500
	bromoform	100
	ethylene dibromide	60
	bromobenzene	300
Halogenated	2-chloroethyl ethyl ether	70
Ethers	Bis-(chloromethyl) ether	120
Nitrosamines	N-nitrosodimethylamine	90
	N-nitrosodiethylamine	420
Oxygenated	acrolein	3
Hydrocarbons	glycidaldehyde	40
	propylene oxide	4
	butadiene diepoxide	210
	cyclohexene oxide	330
	styrene oxide	930
=====		

(continued)

TABLE 4-7 (continued)

=====		
	phenol	330
	acetophenone	600
	B-propyl acetone	100
Nitrogenous	nitromethane	9
Hydrocarbons	aniline	680
Sulfur	diethyl sulfate	7
Compounds	ethyl methane sulfate	830
=====		

4-41

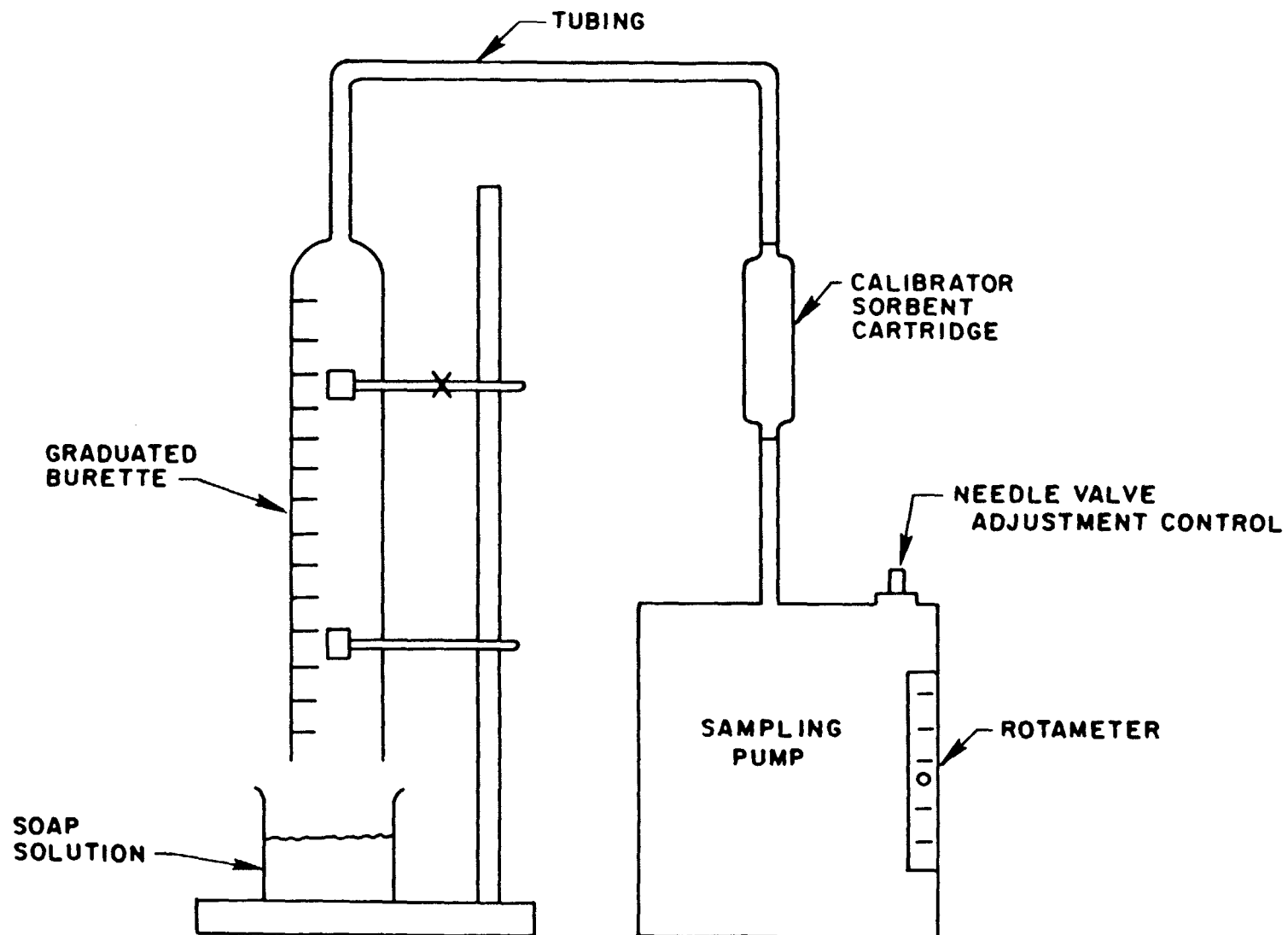


Figure 4-1. Calibration schematic for rotameter and needle valve combination.

- g. Calculate flow rate at standard conditions as follows:

$$F = f \left(\frac{T_{std}}{T_c} \right) \left(\frac{P_c - P_{H_2O}}{P_{std}} \right)$$

Nomenclature--

- F = flow rate at standard conditions, liters/min (ft³/min)
 f = actual flow rate at calibration conditions, liters/min (ft³/min)
 T_c = temperature of air during calibration, °K (°R)
 P_c = pressure of air during calibration, mm Hg (in. Hg)
 T_{std} = standard absolute temperature, 298°K (537°R)
 P_{std} = standard absolute pressure, 760 mm Hg (29.92 in. Hg)
 P_{H_2O} = vapor pressure of water at T_c , mm Hg (in. Hg)

- h. Mark level achieved during flow rate setting on rotameter and record on sampling data sheets for reference. To ensure acceptable flow rate precision, this reference setting on the rotameter should be maintained during sampling. The rotameter can either be integral with the pump (as in the MSA Monitaire Model S) or separate.
- i. Note that, in this case, the rotameter is used only as a visual reference. To ensure correctness of the reference point, calibration conditions should not deviate from sampling conditions by more than the following:

Temperature $\pm 15^\circ\text{C}$
Barometric pressure ± 10 mm Hg.

2. Calibration of Constant Flow Pump Systems--

- a. Select a set of sample pumps and assemble necessary equipment. Figure 4-2 depicts a calibration set-up developed by E.I. DuPont de Nemours and Company for use with their constant flow sampling pumps. This configuration or equivalent is acceptable.
- b. Calibrate each pump according to the following basic steps. In addition, consult the manufacturer's instructions for specific details unique to the model in use.

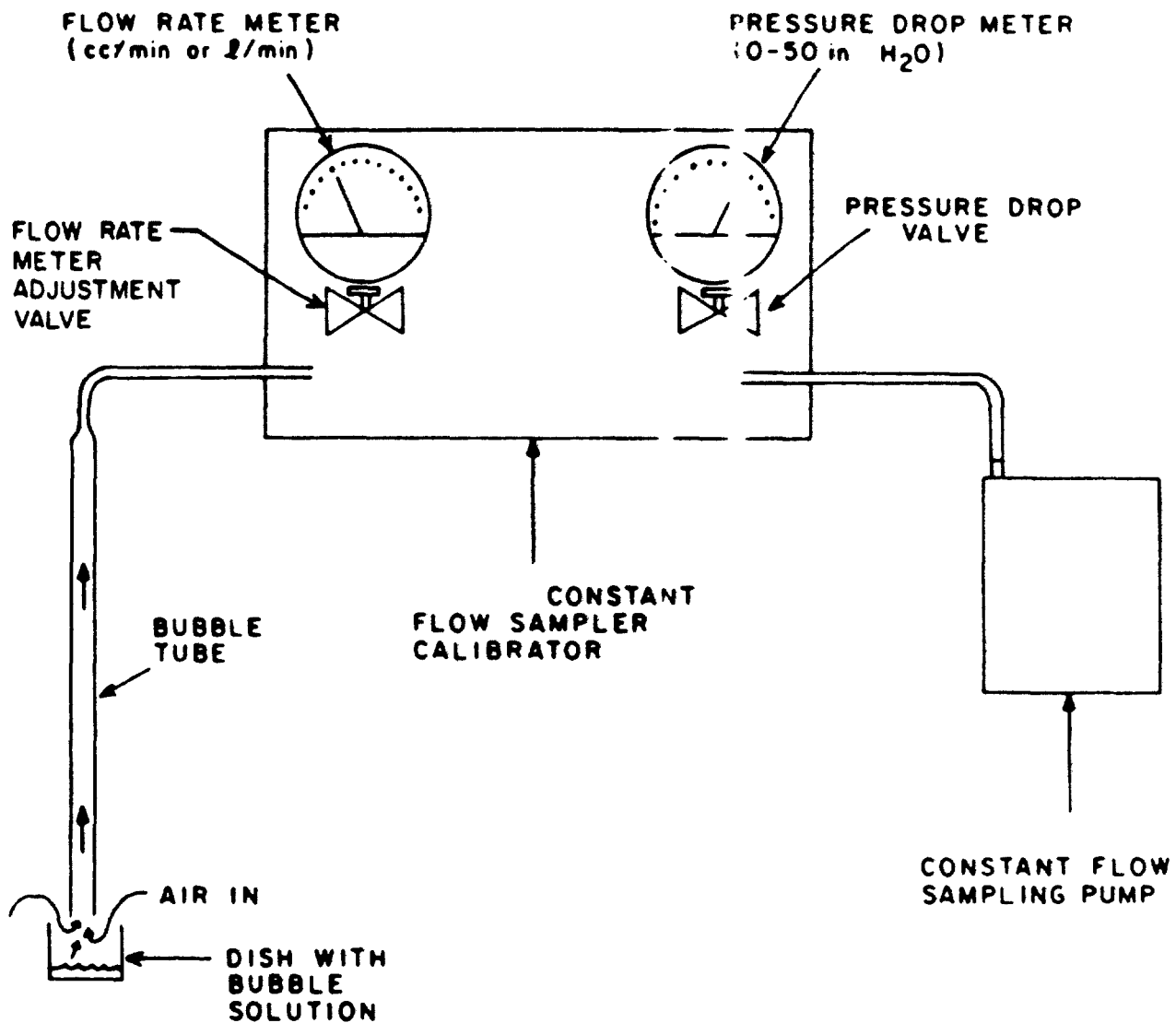


Figure 4-2. Calibration configuration for constant flow samplers.

- c. Measure ambient air temperature, relative humidity, and barometric pressure. Determine water vapor pressure from tables.
- d. If battery test is available, check battery.
- e. Connect pump inlet to bubble tube flowmeter* and start pump. Allow pump to stabilize before taking readings.
- f. Adjust to desired flow rate. With the calibration configuration depicted in Figure 4-2, the flow rate can be adjusted to the target range with the magnetic flow rate meter.
- g. If a flow rate meter is not available, determine the initial flow rate using the bubble tube flow meter and record the time required for the bubble to travel between the appropriate volume markings on the tube.
- h. Calculate the actual flow rate as follows:

$$f = \frac{\text{Distance traveled (ml)}}{\text{Travel time (min)}}$$

- i. Once the flow rate of each sampler has been adjusted to the appropriate setting, verify three times with the bubble tube and record the results. Calculate the mean flow rate by summing the values of the three individual readings and dividing by three. The deviation of the individual flow rates from the mean flow rate should not exceed ± 5 percent.
- j. Calculate flow rate and total sample volume at standard conditions as follows:

$$F = f \left(\frac{T_{std}}{T_c} \right) \left(\frac{P_c - P_{H_2O}}{P_{std}} \right)$$

Nomenclature--

F = flow rate at standard conditions, liters/min (ft³/min)

*Constant-flow type pumps such as the DuPont P-4000 have the ability to maintain the flow rate with ± 5 percent at pressure drops as high as 25 inches of water. For this reason, it is unnecessary to place a "calibrator" sorbent cartridge in line as long as the expected pressure drop does not exceed that specified by the manufacturer.

f = actual flow rate at calibration conditions, liters/min
(ft³/min)

T_c = temperature of air during calibration, °K (°R)

P_c = pressure of air during calibration, mm Hg (in. Hg)

T_{std} = standard absolute temperature, 298°K (537°R)

P_{std} = standard absolute pressure, 760 mm Hg (29.92 in. Hg)

P_{H_2O} = vapor pressure of water at T_c , mm Hg (in. Hg)

- k. When using the calibration configuration depicted in Figure 4-2, the flow maintenance feature of the pump can be checked by inducing a pressure drop across the system using the pressure drop valve and determining the subsequent flow rate with the bubble tube. The maximum difference between the flow rate determined in step i and the flow rate measured at the maximum expected pressure drop should not exceed ± 5 percent.

3. Sampling Procedure--

- a. Assemble sampling train (Figure 4-3). The general procedure requires four trains operated within the flow rate ranges listed in the preceding discussion be set up at each location. Duplicate samplers operating at the same flow rate as one or more of the listed ranges can be set up to collect duplicate samples for backup or quality assurance purposes. Set trains up at desired location and hang samplers on a tripod, music stand, or similar device. To insure stability in wind gusts, weighting of these devices or some method of anchoring is advisable. Nonsparking wooden stakes or fence post can be used but are more work, more difficult to move (which is often necessary), and may incur more risk where buried hazards exist.
- b. Record all initial information (time, counter reading, cartridge number, pump number, sampler, blank number, barometric pressure, ambient temperature, relative humidity, etc.).
- c. Start pump and observe system to determine if appropriate flow rate is being maintained. For systems utilizing rotameters, the calibrated rotameter setting should be maintained during the entire run and should be monitored regularly. Any adjustments to the rotameter are made by opening or closing the needle valve and are noted in the field log. Most constant flow sampling pumps have low flow indicators and/or an automatic shut-off feature at low flow conditions. These should be initially observed and periodically monitored during the course of sampling.

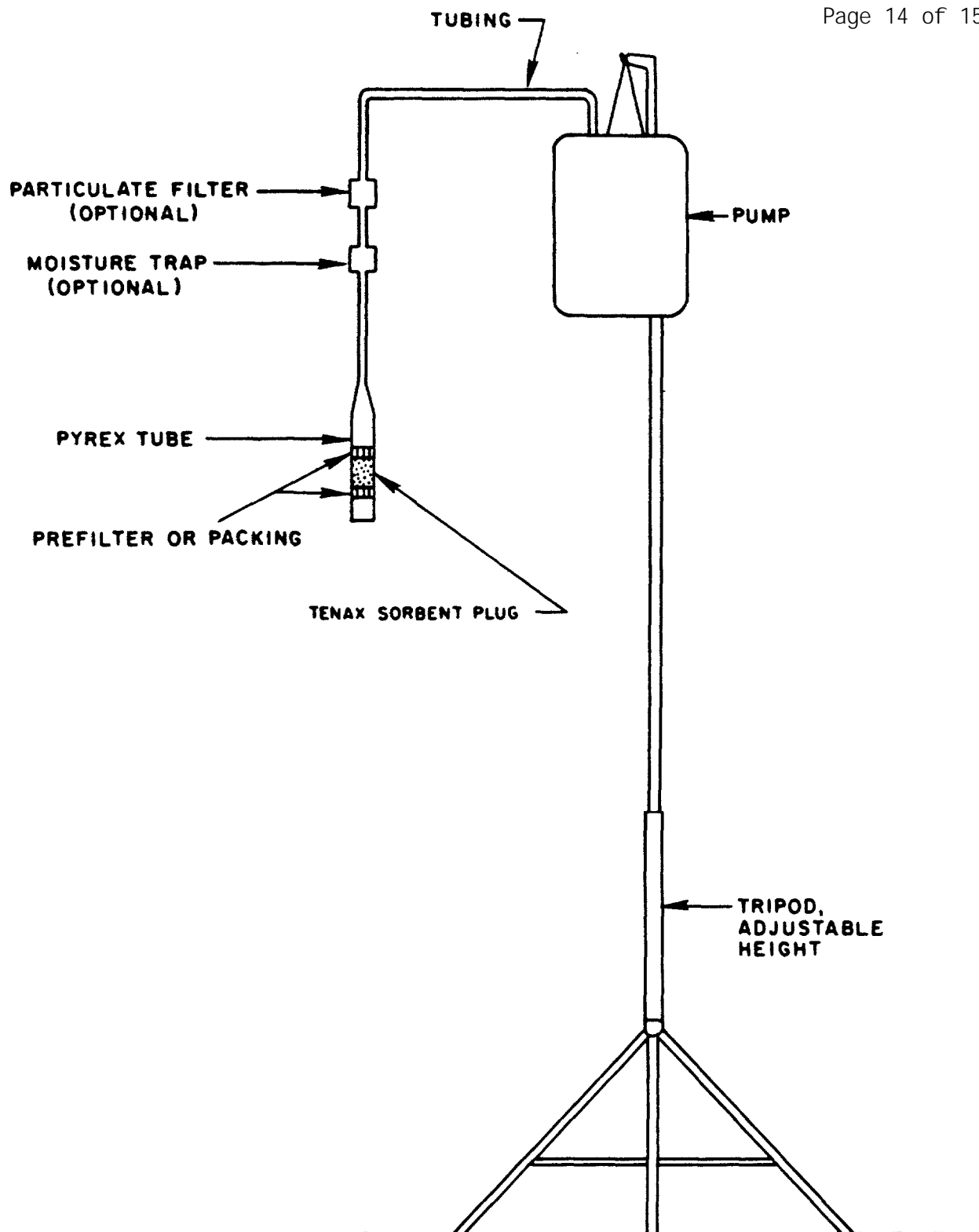


Figure 4-3. Tenax sampler.

- d. Allow pump to run for desired sampling time.
- e. At the end of the sampling period, observe rotameter level or low flow indicator to determine if flow rate has been maintained. Shut down sampling pump and record all pertinent information (counter reading, time, barometric pressure, relative humidity, ambient temperature, problems, comments, etc.). The final flow rate should not deviate from the initial flow rate by more than ± 5 percent.
- f. Remove sorbent cartridge (use clean lint-free gloves) and place in culture tube. Place cartridge in culture tube with the inlet facing up and mark accordingly on the outside of the culture tube. Do not put any marking on Tenax cartridges.
- g. Place plugs of glass wool below and above the cartridge and tightly cap and label the culture tube. Before proceeding with the packing, the tube should be shaken to insure that the cartridge does not rattle inside the culture tube.
- h. Place sample identification tag on culture tube and fill out chain-of-custody form.
- i. Calculate total sample volume at standard conditions

$$V_{std} = F \times t$$

where

F = flow rate at standard conditions, liters/min (ft^3/min)

t = total sample time (min)

V_{std} = sample gas volume at dry standard conditions, dsl (dscf)

Sources

GCA Corporation. "Quality Assurance Plan, Love Canal Study, Appendix A, Sampling Procedures." EPA Contract 68-02-3168.

GCA Corporation. "Guidelines for Air Monitoring at Hazardous Waste Sites for Volatile and Semivolatile Organic Compounds Using Tenax and Polyurethane Foam Sorbents." EPA Contract 68-02-3168. April 1983.

4.2.8 METHOD IV-8: COLLECTING SEMI-VOLATILE ORGANIC COMPOUNDS IN AMBIENT AIR USING POLYURETHANE FOAM (LOW VOLUME SAMPLERS)

Description

Polyurethane foam (PUF) has been shown to be an excellent collection media for trapping and concentrating a variety of semi-volatile organic compounds (defined here as exhibiting a vapor pressure less than or equal to 1 mm (Hg) at 20°C). Foams plugs are cut from the type of PUF used for furniture upholstery, pillows, and mattresses and is Soxhlet extracted with high grade hexane (pesticide quality or equivalent) prior to being fitted into specialized sampling cartridges. To sample airborne organics, a known volume of air is drawn through the collection media.

Cylindrical polyurethane foam plugs (polyether type, 0.021 gm/cm³) are cut from 3-inch stock using a 25 mm circular template, then cleaned in a soxhlet extractor to remove potential interferences. After drying to remove excess solvent and analyzing of one plug from each batch to verify freedom from contamination, the plugs are placed (under slight compression) in 22 mm (inside diameter) by 10 cm long hexane-rinsed glass tubes. The glass tubes are constructed from 22 mm (inside diameter) stock which has been tapered at one end to facilitate attachment to the sampling pump. A teflon reducing adaptor can also be fabricated which permits attachment to the sampling pump with no modification to the glass tube. The cartridges are then placed in teflon sealed 38 mm x 200 mm culture tubes wrapped with aluminum foil to protect the sampling cartridges from ultraviolet light.

Any high-volume personnel sampling pump capable of maintaining a constant flow rate of 3 to 4 liter/minute can be used. Samples are collected at this nominal flow rate for between 8 to 12 hours allowing a total sample volume of between 1 to 4 cubic meters (m³).

Polyurethane foam has been shown to be excellent for trapping a wide variety of semivolatile organic compounds in ambient air including numerous chlorinated pesticides,^{23, 24, 25, 26} polychlorinated biphenyls (PCBs),²⁴ polychlorinated naphthalenes,²⁹ herbicides and their corresponding methyl esters,^{25, 27} organophosphorus pesticides,^{24, 25} chlorinated benzenes,²⁵ chlorinated phenols,²⁵ and polynuclear aromatic hydrocarbons.^{23, 28} Table 4-8 lists the representative components of the above compound classes that have been collected in ambient air using this technique.

Uses

This procedure and modifications of this procedure have been used successfully to collect airborne chlorinated organics including pesticides, PCBs, and a variety of chlorinated benzenes and phenols and is generally applicable to the measurement of such compounds in the ng/m³ to µg/m³ range when sensitive analytical techniques are employed (GC/Electron Capture). These methods are generally not applicable for the more volatile organic compounds (those exhibiting a vapor pressure of greater than 1 mm (Hg)

TABLE 4-8. ORGANIC COMPOUNDS COLLECTED IN AMBIENT AIR USING LOW VOLUME OR HIGH VOLUME POLYURETHANE FOAM SAMPLERS

Polychlorinated biphenyls ^{24, 25, 26S, 31, 32, 33}

Aroclor 1221
Aroclor 1232
Aroclor 1242
Aroclor 1016
Aroclor 1248
Aroclor 1254
Aroclor 1260

Chlorinated pesticides ^{32, 34, 33, 25, 50, 35}

α chlordane	p, p'-DDT
γ chlordane	Endosulfan
Chlordane (technical)	Heptachlor
Mirex	Aldrin
α BHC	
β BHC	
γ BHC (lindane)	
δ BHC	
p, p'-DDD	
p, p'-DDE	

Polychlorinated naphthalenes ³⁴

Halowax 1001
Halowax 1013

Chlorinated Benzenes ^{25, 35, 26, 33}

1, 2, 3-trichlorobenzene
1, 2, 3, 4-tetrachlorobenzene
Pentachlorobenzene
Hexachlorobenzene
Pentachloronitrobenzene

Chlorinated phenols ²⁵

2, 4, 5-trichlorophenol
Pentachlorophenol

(continued)

TABLE 4-8 (continued)

=====

Herbicide Esters ²⁵

2,4-D Esters, isopropyl
2,4-D Esters, butyl
2,4-D Esters, isobutyl
2,4-D Esters, isooethyl

Organophosphorus pesticides ^{32, 25, 24, 32, 34}

Mevinphos
Dichlorvos
Ronnell
Chlorpyrifos
Diazinon
Methyl parathion

Ethyl parathion
Parathion
Malathion

Carbamate pesticides ^{25, 32}

Propoxur
Carbofuran
Bendiocarb
Mexacarbate
Carbaryl

Urea pesticides ²⁵

Monuron
Duron
Luron
Terbuturon
Fluometuron
Chlorotoluron

Triazine pesticides ^{25, 32}

Simazine
Atrazine
Propazine

=====

(continued)

TABLE 4-8 (continued)

=====

Pyrethrin pesticides ²⁵

Pyrethrin I
Pyrethrin II
Allethrin
d-Trans-allethrin
Dicrotophos
Resmethrin
Fenvalerate

Polynuclear aromatic hydrocarbons ^{34, 28, 36, 30, 29}

Naphthalene	Benzo (a) anthracene
Biphenyl	Chrysene/triphenylene
Fluorene	Benzo (b) fluoranthene
Dibenzothiophene	Benzo (e) pyrene
Phenanthrene	Benzo (a) pyrene
Anthracene	Perylene
Carbazole	o-Phenylene pyrene
2-Methyl anthracene	Dibenzo (at/ah) anthracene
1-Methyl phenanthrene	Benzo (g, h, i) perylene
Fluoranthene	Coronene
Pyrene	
Benzo (a) fluorene	
Benzo (b) fluorene	

=====

at 20°C) nor are they applicable for differentiating between vapor phase organics and those adsorbed on particulate matter. When collection of such compounds is desired, it will be necessary to utilize separate collection media (Tenax-GC, filters, etc.) or combination cartridges.

Although sampling trains consisting of a particulate pre-filter followed by a backup sorbent cartridge have been described,³⁰ indications are that large portions of the particulate associated organic compounds may volatilize off the particulate and onto the backup sorbent during collection, giving higher than actual vapor base measurements and lower than actual particulate associated measurements.^{50, 45}

The listing supplied in Table 4-8 will aid investigators in determining the applicability of this sampling method to their particular application. Potential users are cautioned that method validation studies for many of the compounds listed, including determination of collection efficiencies, resorption recoveries, etc., have not been conducted.

The investigator should keep in mind that the procedure described herein is meant, in its broadest application, to be a screening technique and is therefore necessarily general. If specific conditions, compounds of interest, concentrations, detection requirements, etc. are known, such factors should be carefully considered and the appropriate literature sources reviewed to optimize procedures relevant to specific needs. For instance, use of a cartridge composed of a PUF/adsorbent resin "sandwich" has been described by Lewis and McLeod which may be useful in collecting compounds with low PUF breakthrough volumes.²⁵

Procedures for Use

1. Calibrate the sampling pump as per the procedure outlined in Method IV-7. Adjust pumps to a target flow rate of 3 to 4 liters/minute.
2. Sampling procedures.
 - a. Assemble sampling train (see Figure 4-4). Set train up at desired location and hang sampler on a tripod, music stand, or similar device. To insure stability in wind gusts, weighting of these devices or some method of anchoring is advisable. The use of wooden or other nonsparking stakes can be used, but experience has shown these to be less convenient and often more work than weighted stands.
 - b. Record all initial information (time, counter reading, cartridge number, pump number, sampler, blank number, barometric pressure, ambient temperature, relative humidity, etc.).
 - c. Start pump and observe system to determine if appropriate flow rate is being maintained. For systems utilizing rotameters the calibrated rotameter setting should be maintained during the

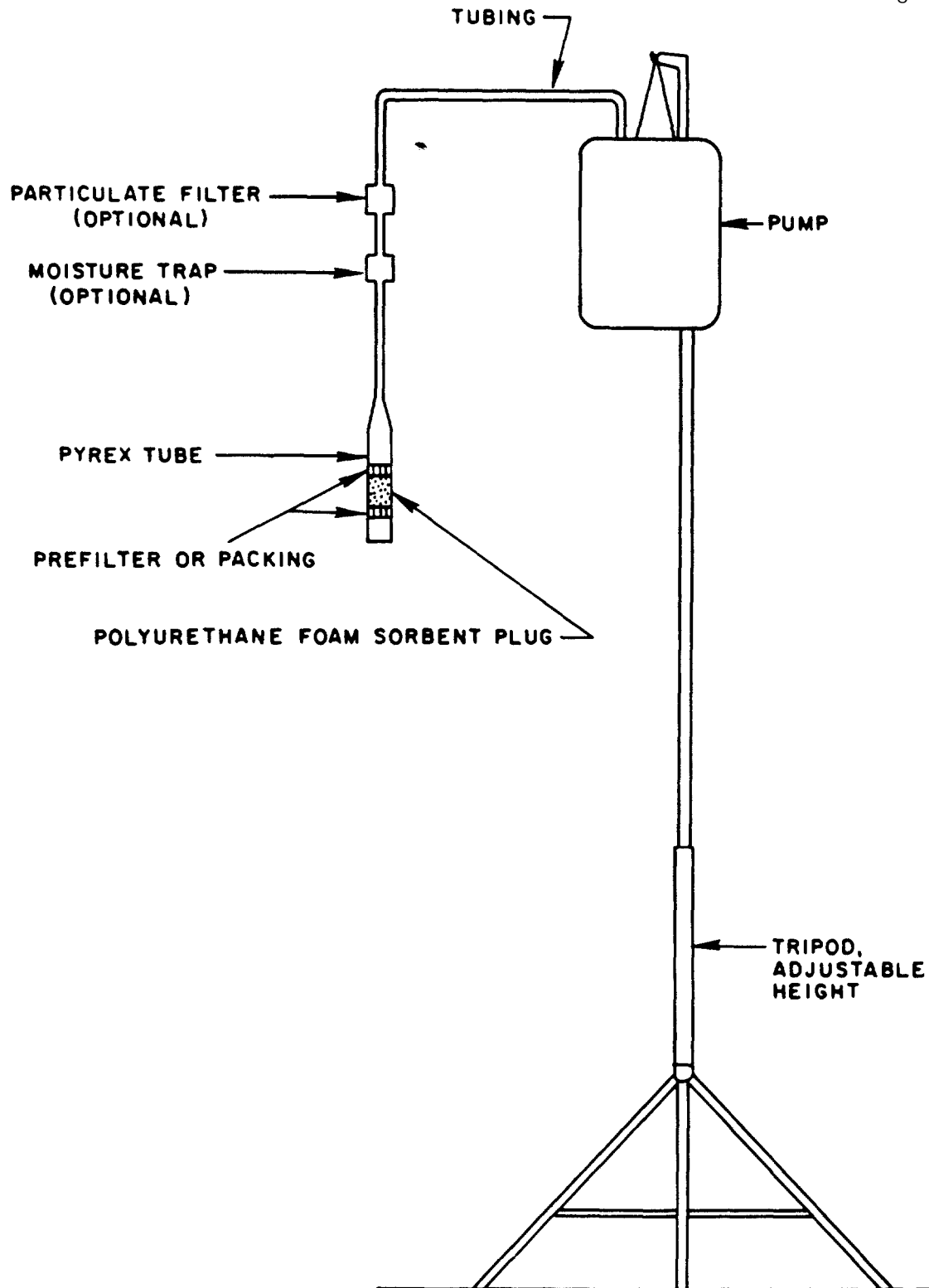


Figure 4-4. PUF sampling train schematic.

entire run and should be monitored regularly. Any adjustments to the rotameter are made by opening or closing the needle valve and should be noted in the field log. Most constant flow sampling pumps have low-flow indicators and/or an automatic shut-off feature at low-flow conditions. These should be initially observed and periodically monitored during the course of sampling.

- d. Allow pump to run for desired sampling time.
- e. At end of sampling period, observe rotameter level or low flow indicator to determine if flow rate has been maintained. Shut down sampling pump and record all pertinent information (counter reading, time, barometric pressure, relative humidity, ambient temperature, problems, comments, etc.). The final flow rate should not deviate from the initial flow rate by more than $\pm 5\%$.
- f. Remove PUF cartridge (use clean gloves) and wrap it with hexane-rinsed aluminum foil.
- g. Place foil-covered cartridge in a hexane-rinsed glass bottle or culture tube that has been properly labeled. Plugs of glass wool are placed below and above the cartridge and the tube is tightly capped. The tube should be gently shaken to insure that the cartridge does not rattle inside the culture tube.
- h. Place sample identification tag on sample bottle or culture tube and fill out chain of custody form.
- i. Calculate total sample volume at standard conditions as in Method IV-7.

Sources

GCA Corporation. "Quality Assurance Plan, Love Canal Study, Appendix A, Sampling Procedures." EPA Contract 68-02-3168.

Lewis, Robert G. and MacLeod, Kathryn E. "Portable Sampler for Pesticides and Semivolatile Industrial Organic Chemicals in Air." Analytical Chemistry, Volume 54, pp. 310-315, 1982.

GCA Corporation. "Guidelines for Air Monitoring at Hazardous Waste Sites for Volatile and Semivolatile Organic Compounds Using Tenax and Polyurethane Foam Sorbents." EPA Contract 68-02-3168, April 1983.

4.2.9 METHOD IV-9: DETERMINATION OF TOTAL SUSPENDED PARTICULATE IN AMBIENT AIR USING HIGH-VOLUME SAMPLING TECHNIQUE

Description

Ambient air is drawn into a covered housing and through a filter by means of a high-volume blower at flow rates between 1.13 to 1.70 m³/min (40 to 60 ft³/min). Particles within the size range of 100 to 0.1 µm diameter are collected on the filter although sampler flow rate and geometry tends to favor particles less than 60 µm aerodynamic diameter. The mass concentration of suspended particulate is computed by measuring the mass of collected particulate (gravimetric analysis) and the volume of air sampled.

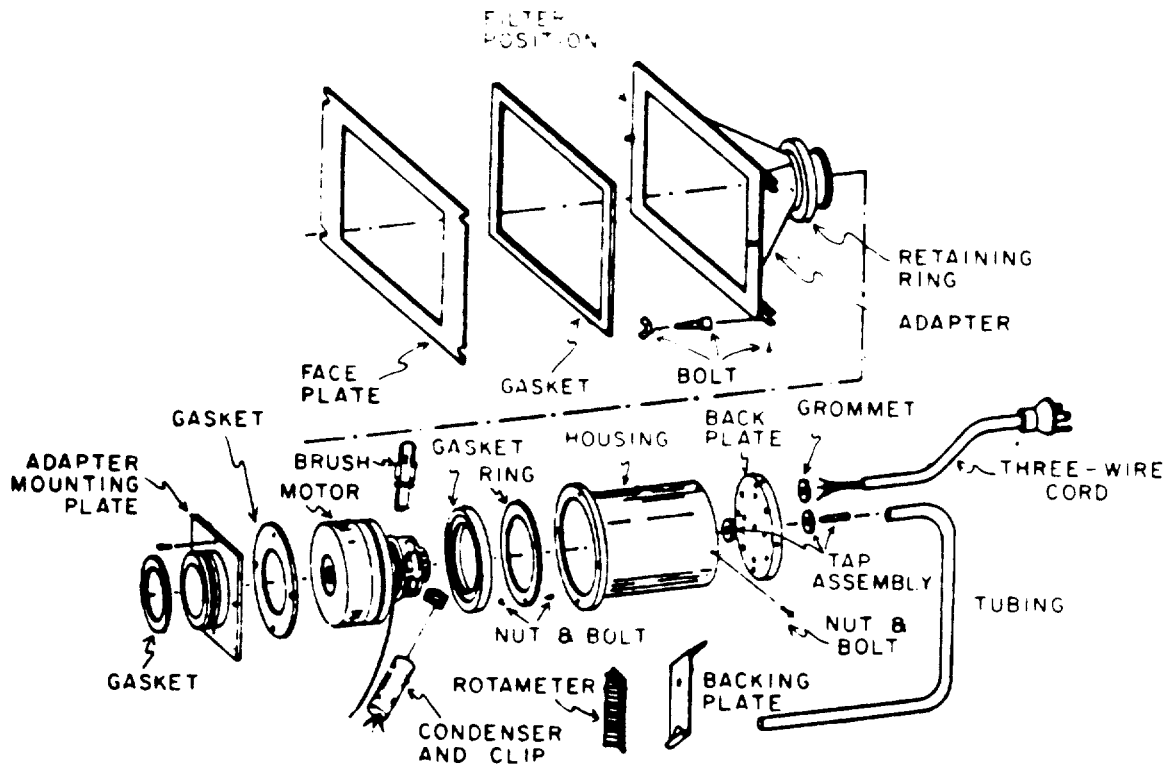
High volume ambient air samplers (Figures 4-5 and 4-6) are readily available from a number of vendors and should meet the specifications described in 40 CFR Part 50 Appendix B--Reference Method for the Determination of Suspended Particulate in the Atmosphere (High Volume Method).³⁷ Filter media (glass fiber filters) with a collection efficiency of at least 99 percent for particles of 0.3 µm diameter are also specified for use. Other types of filter media (e.g., paper) or specially prepared filters may be desired in instances where specific analysis is contemplated or low background levels of certain pollutants is desired.

After sample collection, pretared filters are analyzed gravimetrically to determine the total particulate loading. Trace metal analyses may be accomplished by extracting all or part of the filter and analyzing the extract accordingly (i.e., atomic absorption, ICP). It should be noted that when trace metal analysis is desired, it is extremely important to submit blank filters from each lot to the laboratory to determine background concentrations.

Modified high-volume sampling techniques have also been used to efficiently collect certain organic compounds. Stratton, et al.,³¹ and Jackson and Lewis³⁸ describe samplers modified to include a throat extension between the filter housing and blower that contains polyurethane foam sorbent. This arrangement can also be used to trap polynuclear aromatic hydrocarbon (PNAs). Additional sorbents or combinations can be used dependent upon specific collection requirements. As with trace metal analysis, it is important that blank filters and sorbents be submitted to the laboratory to determine the existence of background concentrations.

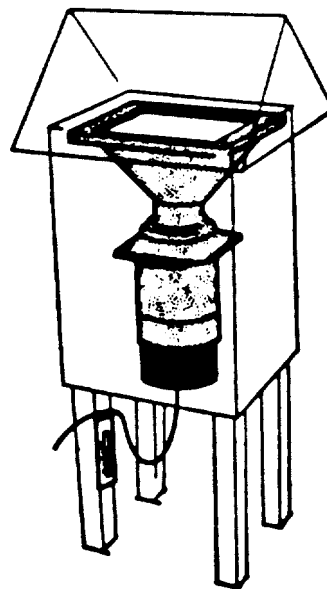
Uses

The described procedures can be used to collect Total Suspended Particulate (TSP) matter in ambient air. The collected material may be extracted and analyzed for trace metals or particulate related organics of low volatility. In the latter case, backup collection techniques (PUF) would be advisable.



Source: Reference 37.

Figure 4-5. Exploded view of typical high-volume air sampler parts.



Source: Reference 37.

Figure 4-6. Assembled sampler and shelter.

Procedure for Use

1. Calibration

Refer to 40 CFR 50, Appendix B, Part 8.0--Calibration as amended³⁷ and the EPA Proposed Changes to Ambient Measurement Methodology for Carbon Monoxide, Particulate, Sulfur Dioxide, 47 FR 2341, January 15, 1982.35

Essentially, samplers must be calibrated when first purchased, after major maintenance on the sampler (e.g., replacement of motor or brushes), any time the flow measuring device (rotameter or recorder) has been replaced or repaired, or any time a one-point calibration check deviates from the calibration curve by more than ± 6 percent.

The following procedure is based on the use of a certified variable resistance orifice as the sampler calibration device and a continuous flow-rate recorder (Dickson recorder) used to ensure the accuracy of air volume measurements. Samplers may also be equipped with an electronic flow controlling mechanism to perform the same function. Flow-rate controllers and recorders are not as yet required; however, errors resulting from nonconstant flow rates can be greatly reduced by using such devices. In addition, the currently approved flow indicators (rotameters) have been shown to be subject to a variety of errors caused by physical damage, dirt deposition, and flow restrictions in connecting tubing.

- a. Remove filter retaining plate from the sampler to be calibrated and place a clean filter in the filter holder.
- b. Attach the variable resistance orifice (VRO) to the sampler and position the orifice setting to full open. Secure the VRO fall plate to insure an air tight seal with the orifice gasket. Attach a slack tube manometer to the sampler unit.
- c. Plug sampler into 120-volt source, while checking manometer to insure that the orifice pressure drop does not exceed the range of the manometer. Let the sampler run for about 5 minutes.
- d. Turn motor off and place a fresh chart on the unit. The chart should include the following information: high-volume sampler identification, date and time of calibration, and operator's name. The chart should be labeled "Calibration Data."
- e. Check the recorder for proper operation, and zero the pen if necessary.
- f. Determine five approximately equally spaced intermediate points which provide pressure drops between the desired maximum and minimum operating point and record the following data on the calibration sheet:

- pressure drop from the manometer (in. H₂O)
- flow rate indicated on Dickson recorder, liters/mm (ft³/min).

Repeat three points centralized in the vicinity of the expected sampler flow rate to insure accuracy in the field.

(The Dickson Recorder should be tapped gently prior to reading, to insure that the recorder pen is in its final position.)

- g. Record the airflow rate from the VR0 high-volume calibration curve for each flow recorder reading.

$$\text{ACCEPTABILITY} = 100 \left| \frac{(Q_o - Q_c)}{Q_c} \right| \text{ within } 5\%$$

where:

Q_o = observed flow rate

Q_c = flow rate from calibrated curve

- h. If the air flow rate exceeds the acceptable limits, rerun points for which percent deviation exceeds 5 percent until acceptance limits are attained.
- i. Correct the sample flow rate to standard conditions using the following formula:

$$Q_2 = Q_1 \left[\frac{T_2 P_1}{T_1 P_2} \right]$$

where: Q₂ = corrected flow rate std. liters/min (std. ft³/min)

Q₁ = recorded flow rate from chart, liters/min (ft³/min).

T₁ = temperature during calibration, °K (°R).

T₂ = standard temperature, 298°K (537°R).

P₁ = atmospheric pressure during calibration, mmHg (in. Hg).

P₂ = standard absolute pressure, 760 mmHg 29.92 in. Hg).

2. Sample Collection

Total suspended particulate measurements are normally collected over a 24-hour sampling period; however, this requirement may be altered for hazardous waste sampling applications. Monitoring objectives may require sampling at specific time intervals only (e.g., during drum excavations), and high particulate loadings due to heavy equipment traffic may also require shortened sampling periods. Sampling time selection will therefore be site specific and obviously dependent upon a number of unique factors.

a. Installation of Clean Filter

- (1) Remove faceplate by loosening the four wing nuts and rotating the bolts outward.
- (2) Obtain a clean, weighed filter and record the filter number, high-volume sampler serial number, flowmeter serial number, location, run date, and start time on the data sheet.
- (3) Carefully place the clean filter rough side up, on the wire screen, and center the filter so that when the faceplate is in position, the gasket will form a tight seal on the outside edge of the filter.
- (4) Replace faceplate, being careful not to move the filter, and tighten the wing nuts evenly until the gasket forms an airtight seal against the filter.

b. Operation Checks

- (1) Allow sampler motor to warm up at least 5 minutes to reach normal operating temperature.
- (2) Assure that the flow recorder is connected to the sampler using the same tubing as was used to calibrate the sampler.
- (3) Place a new chart on the recorder and set at correct start time.
- (4) Record "Run Start" time and date, site identification, and sampler number on the chart.
- (5) Turn sampler off and set clock switch to desired setting. Total suspended particulate samples are normally collected over a 24-hour period; however, this requirement may be altered depending on monitoring applications.

c. Removing Exposed Filter

- (1) Turn sampler "on" and allow to warm up at least 5 minutes.
- (2) Check flow recorder chart for proper operation.
- (3) Turn sampler "off" and record elapsed time in logbook and on the data sheet.
- (4) Remove chart and place in envelope.
- (5) Carefully loosen wing nuts and remove faceplate gasket.
- (6) Remove the exposed filter by gently grasping the ends of the filter and lifting it from the screen. Fold the filter lengthwise at the middle with the exposed side "in." If the collected sample is not centered on the filter, fold the filter accordingly so that sample touches sample only.
- (7) Place the filter in a glassine envelope, and place glassine envelope with data sheet in a folder for return to sample bank.
- (8) Visually inspect for signs of leakage, damage, etc., to the sampler and repair if necessary.

Sources

United States Environmental Protection Agency. "Appendix B--Reference Method for the Determination of Suspended Particulate in the Atmosphere (High Volume Method)". 40 CFR Part 50. November 25, 1971.

United States Environmental Protection Agency. "Proposed Changes to Ambient Measurement Methodology for Carbon Monoxide, Particulate and Sulfur Dioxide." 47 CFR 2341. January 15, 1982.

4.3 SOIL GASES AND VAPORS

Monitoring of soil gases can often serve as a quick method of determining the extent of pollutant migration or establishing perimeters of a site containing buried wastes. Soil-gas exchange with the ambient atmosphere greatly dilutes gaseous components making them difficult to detect. Therefore, sampling in the soil can provide a more concentrated source for underground waste detection. Soil-gas sampling also has particular applicability to the identification of methane fluxes at sanitary landfills.

4.3.1 METHOD IV-10: MONITORING GAS AND VAPORS FROM TEST HOLE

Discussion

Gas samples can be withdrawn from test holes by using a nonsparking probe, brass and Teflon being the most suitable. The probe is then attached to the gas inlet of the desired gas monitor such as those described in the ambient gases section and Method IV-1 through IV-8. The test holes are easily prepared by driving a metal rod (approximately 1 in. diameter) into the soil with a drive weight. Commercial bar hole-makers are available that combine the steel hole-making bar and drive weight into one unit (see Figure 4-7).³⁹

Uses

This system is particularly adapted for rapid evaluation of waste sites for soil gas generation. When used in conjunction with a hydrocarbon analyzer or an explosimeter it can rapidly determine the areal extent of a waste site or the location of a particular emission source. It is recommended that the test area be screened with a metal detector before sampling.

Procedures for Use

1. Select location free from rocks and debris. Screen location with metal detector to verify absence of drums and pipes.
2. Place bar point on ground and raise drive weight, then allow weight to fall on bar. It is only necessary to guide the weight in its vertical travel.
3. Continue until desired depth or any penetration resistance is reached.
4. Remove bar hole-maker.
5. Attach suitable length of Teflon tubing (stainless steel or brass may be used in some instances but may result in some gas adsorption/absorption) to monitor instrument gas inlet.
6. Lower tubing into test hole and operate monitor or gas sampling device as listed in Methods IV-1 through IV-8.
7. Record results.
8. Remove sample tubing and observe that instrument readings return to background. If not, change tubing before proceeding to next test location.
9. Tramp over and recover test hole.

Sources

Flower, F.B. "Case History of Landfill Gas Movement Through Soils."
Rutgers University, New Brunswick, New Jersey.

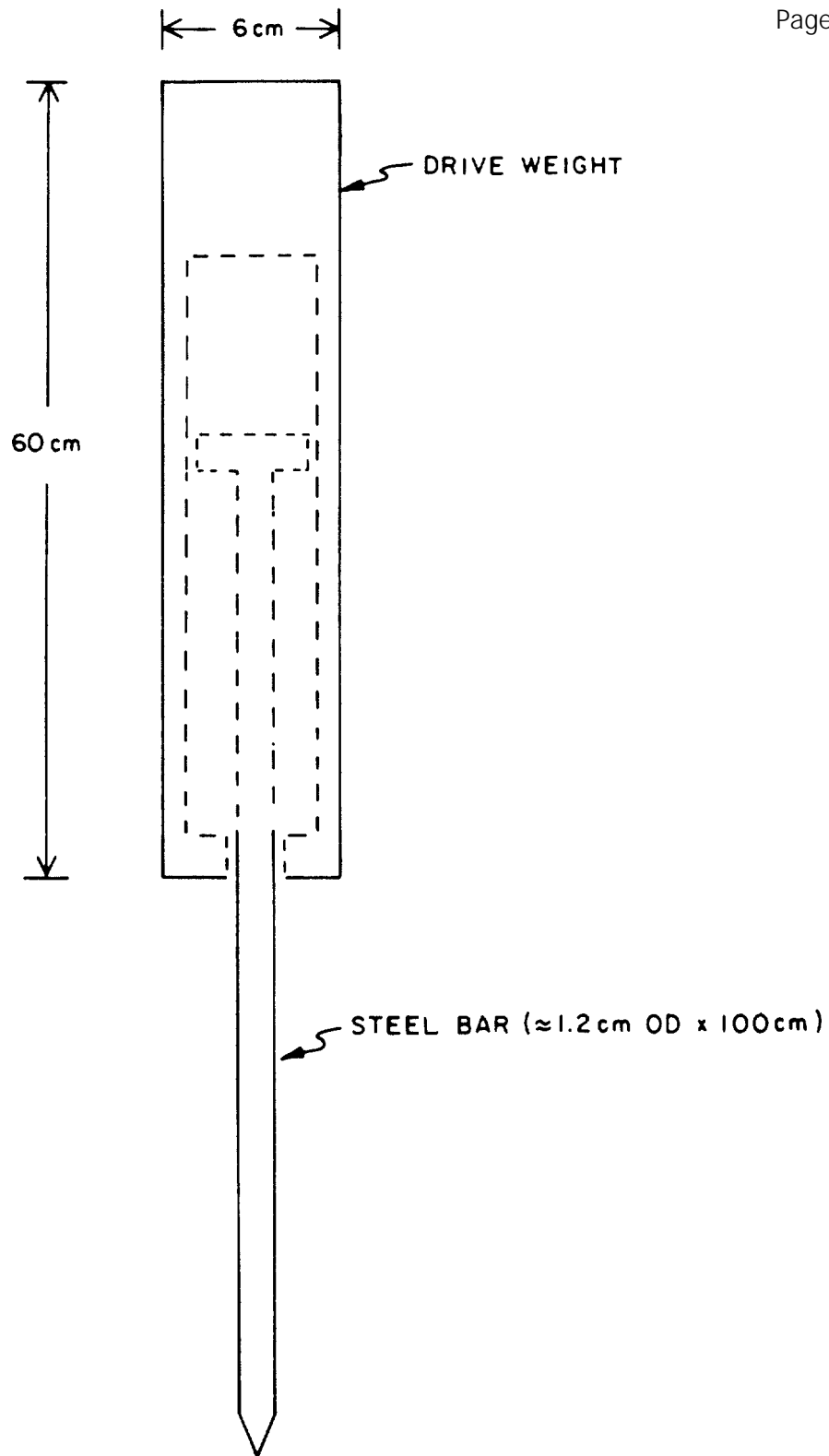


Figure 4-7. Bar hole-maker.

4.3.2 METHOD IV-11: MONITORING GAS AND VAPORS FROM WELLS

Discussion

The sampling of wells for gases and vapors can be accomplished by lowering an intake probe through a sealed cap on the top of the well, (Figure 4-8). The intake probe should be of a nonsparking material that will further minimize adsorption or desorption effects. Teflon or glass are preferable to steel or brass in this application. The intake probe is then connected to the desired gas monitor such as those described in the ambient gases section and Methods IV-1 through IV-8.

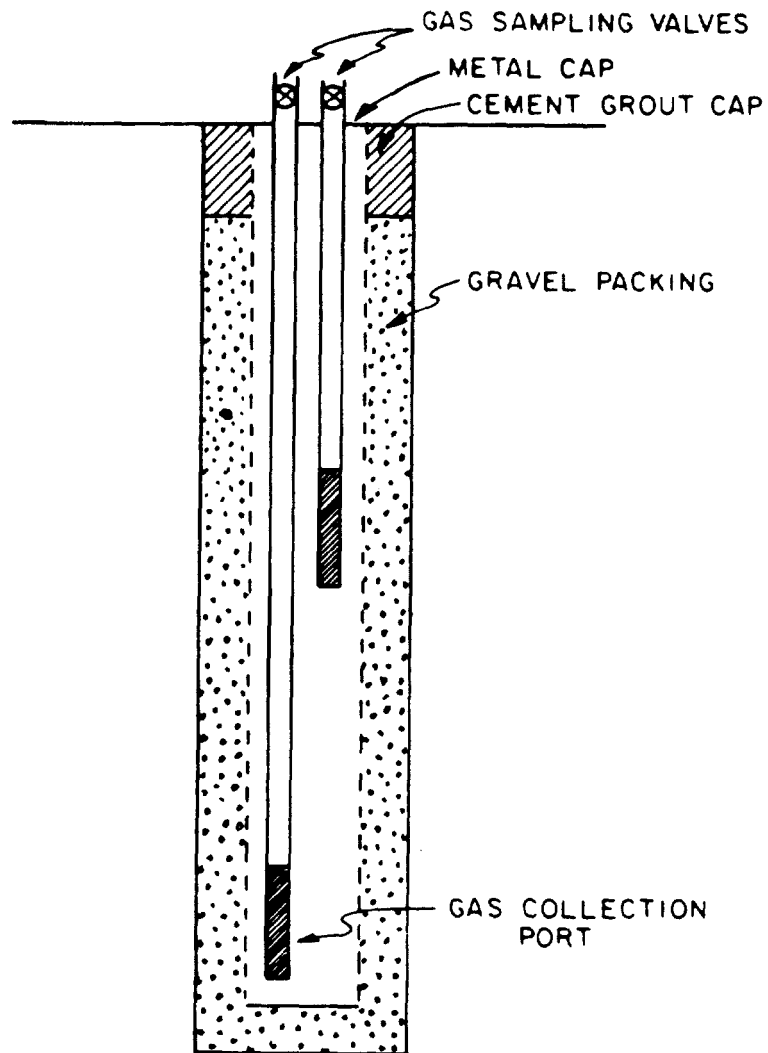
Uses

Existing groundwater monitoring wells can be used to check for the presence of those gases volatilized or otherwise liberated from the groundwater. In some cases, the groundwater level will be below the top of the screened portion of the well allowing free soil gases to enter the well casing.

Wells especially designed for soil-gas monitoring can also be placed by conventional well placement techniques. The well casing, however, is perforated the entire distance, the annular space is packed with gravel, and the top is sealed with a grout cap.⁴⁰ The top of the casing can even be equipped with a sampling valve to allow easy coupling to the monitoring instruments.

Procedures for Use

1. Sound the well for water level or bottom.
2. Select the required length of Teflon tubing. It should be of sufficient length to approach the water level or well bottom, but not so long as to allow water or bottom sediments to enter probe inlet. An inside diameter of 1/8 inch is usually sufficient. However, because this size lacks rigidity, a small weight can be secured to the inlet end to facilitate placement.
3. Lower the tubing through an appropriate sized stopper on the top of the well casing. A wooden plug serves well. It is not critical to maintain an effective seal around the tubing.
4. Lower intake to near bottom and attach outlet to monitor inlet.
5. Proceed with instrument operation according to Methods IV-1 through IV-8 or the instrument operator's manual. Note: When using an adsorption technique for qualification/quantification, Sisk⁴ recommends a sample rate of 1 lpm for 5 to 30 minutes through Tenax GC (see Method IV-7).



Source: Reference 40.

Figure 4-8. Gas sampling well.

6. Gradually raise the intake tubing while observing the instrument readings.
7. Record readings, then remove probe and close casing.
8. If instrument fails to return to background level, replace sample inlet tube before Proceeding to next well. Note: Sometimes vapors may condense on the lower portion of the sample tube, merely cutting off the bottom several centimeters of the intake tube may remove the source of contamination and allow reuse of the remaining sample tube.

Sources

Hatayama, H.R. "Special Sampling Techniques Used for Investigating Uncontrolled Hazardous Waste Sites in California." In: National Conference on Management of Uncontrolled Hazardous Waste Sites. Hazardous Material Control Research Institute, Silver Springs, Maryland. 1981.

4.4 HEADSPACE GASES

Headspace gases are the accumulated gaseous components found above solid or liquid layers in closed vessels. These gases may be the result of volatilization, degradation, or chemical reaction. Poorly ventilated or partially sealed areas can also act to concentrate gas vapors. Component concentrations normally exceed those found in ambient measurements. Therefore, the previously described ambient methods must be modified for handling these higher concentrations and for the remote sensing of container contents. The anticipated higher concentrations can be dealt with by altering the instrument detector range, reducing the sample gas flow rate into the instrument, or utilizing a sample dilution system. These techniques are necessary for the prevention of saturation, poisoning, and/or gross deterioration of the detector element. When lengthy extensions are used, one must also take into account increased time lags for instrument response.

Most ambient measurement devices have sample intakes which are highly directional and localized. The use of an extension will allow the operator to obtain samples from varying depths and distances within containers while maintaining a safe position.

Headspace gases are often found in certain types of containers. Bulging, stainless steel, lined, or other special designated drums are more likely to contain hazardous headspace gases. A preliminary scan of the external seams, edges, or any corroded areas with a vapor analyzer may indicate the nature of the contents.

Poorly ventilated vessels can usually be sampled for headspace gases through small hatches or openings. Fully sealed vessels must be approached more cautiously since breaching may result in the uncontrolled release of pressurized gases or the potential for violent reactions with the ambient atmosphere. Any decision to open a sealed vessel should be based on sound need and the investigator must be cognizant of the inherent dangers, and take appropriate safety precautions.

4.4.1 METHOD IV-12: SAMPLING OF HEADSPACE GASES IN SEMI SEALED VESSELS

Discussion

Sampling of headspace gases involves merely extending the intake or otherwise conducting the contained gas to the detection device. Any of the procedures discussed in the ambient section (Methods IV-1 through IV-8) can be employed. The use of Teflon tubing of approximately 4.8 or 6.4 mm (3/16 or 1/4 inch) inside diameter works well as a probe extension.

Uses

This system is viable in a wide variety of applications. It is simple, and only requires some adaption to match the extension tubing to the instrument intake. The likelihood of high concentrations of contaminants is, however, greater in contained vessels and, as a result, there is the potential for detector saturation and fouling. It is advisable to place any instruments used in this role in their highest operating range. Flame ionization detectors that utilize the sample gas stream as their combustion air may have insufficient oxygen for combustion and will likely require use of a dilution probe. The introduction of entrained droplets from the container contents should also be avoided. Careful handling of the extension tube to avoid close contact with the materials surface and in some instances the use of a glass wool filter plug will prevent material buildup in the probe and detector.

Procedures for Use

1. Select an appropriate monitoring instrument or device that will characterize the gas if present. A combustible gas detector, hydrocarbon vapor analyzer or stain detector tube is normally used. Be particularly aware of the limitations of the instrument in use.
2. Attach the proper size and length tubing that will reach into the container. The tubing seal with the monitoring instrument should be leak tight.
3. Insert tubing into container or vessel opening and operate instrument as per Methods IV-1 through IV-8 and the appropriate operators manual.

4.4.2 METHOD IV-13: SAMPLING OF HEADSPACE GASES IN SEALED VESSELS

Discussion

Sealed vessels, especially 55-gallon drums, present problems when sampling for entrapped gases. The container must be opened to accept a sample probe while still preventing uncontrolled release of its potentially hazardous contents. Further, this must be accomplished while still protecting the safety of the inspector.

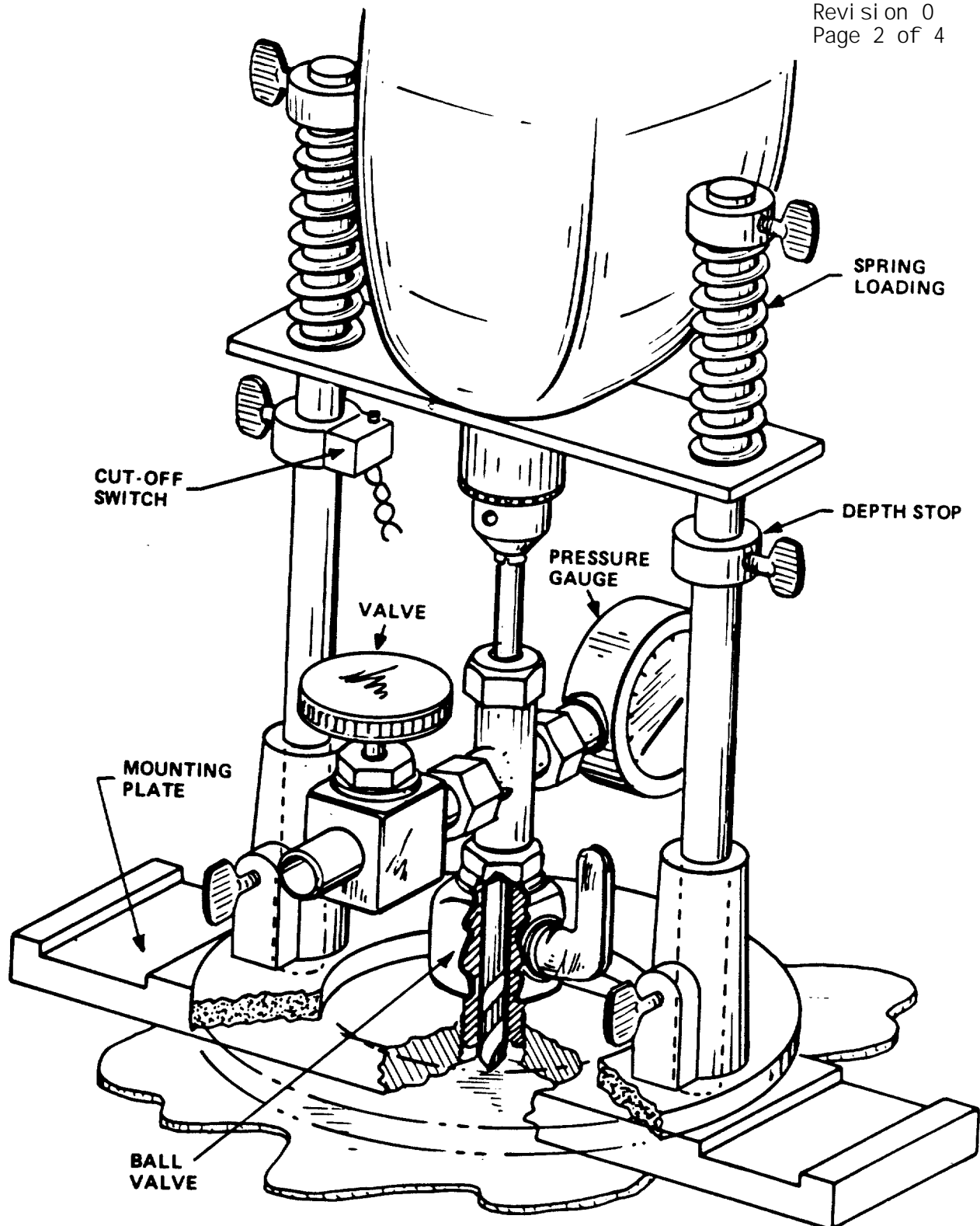
On large vessels and tanks inspection valves and petcocks are normally available. Sealed drums, however, are not designed to contain gases that often develop as reaction products of the contents and have no such provisions.

Leak-free sample tops can be installed on these drums by attaching a mechanism that will drill through a leak-tight fitting strapped to the drum (Figure 4-9). The system consists of a battery operated drill with a remote control switch. The drill is mounted on a simple spring-controlled frame which guides the drill bit through a Swagelok cross fitting. The Swagelok cross is attached to a ball valve which, in turn, is attached to a mounting plate. The mounting plate underside is gasketed with closed cell Neoprene foam. The mounting plate is held against the container using standard steel packaging straps. The cross fitting contains a Teflon seal which allows the drill bit to rotate without allowing gases from the container to escape during drum penetration. A pressure gauge is attached to one side of the Swagelok cross while a needle valve is attached to the side opposite the gauge. The pressure gauge permits the waste handler to observe the internal pressure of the container while the valve permits the removal of sample gas for analysis. The valve and pressure gauge can also be used to insure pressure equalization prior to further opening of the container. A light is located on the remote control switch which indicates when the drum has been pierced. The electrical control system is interlocked so that drill operation automatically stops upon penetration of the container by the drill bit. The whole assembly is activated remotely. Once the bit has penetrated the drum, contained gases flow between the drill bit and the inside of the fittings. Release of the gases is controlled by a needle valve. After sampling, the drill mechanism is pulled away from the container until the drill bit clears the ball valve. The ball valve is then closed, and the piercing mechanism up to the ball valve is removed from the container. The ball valve and mounting plate are left intact to serve as a permanent seal for the opening.

The monitors and detectors described in the Ambient Section (Methods IV-1 through IV-8) can then be adapted to the needle valve and the gas directed to the instrument.

Uses

This device has been used on 55-gallon drums but would also be applicable to other size drums and vessels. Fabrication specifications for this device are found in Appendix B.



Source: Reference 41.

Figure 4-9. Drilling mechanism.

Procedures for Use

1. Assemble the drill assembly as per Appendix B - Equipment Availability and Fabrication.
2. Brush clear any loose rust or dirt to assure a leak-free seal. Seat assembly against the drum side. Tighten mounting straps using portable packaging equipment.
3. Assure that all fittings are snug and needle valve is fully closed.
4. Deploy remote control cable to full extent and stand behind safety screen.
5. Activate drill.
6. After penetration is indicated by light on remote control unit, approach container while monitoring internal drum pressure with pressure gauge on sampler.
7. Attach desired monitor instrument for container content characterization. Any device listed in the ambient section can be employed (Method IV-1 through IV-8). The instrument can be attached by using an appropriate size Teflon tubing (see Method IV-11). After sampling, close needle valve. Extra caution is necessary for highly pressurized containers, as most monitors are designed to accept ambient pressure gases.
8. After proper quantification and/or identification of the contained gas, the safety officer should decide whether the gas can be vented or should be properly contained for later disposal.
9. The full assembly can be removed if the gas has been properly vented or disposed of; otherwise the drill can be loosened from the bit and removed from the guide assembly as outlined below.
 - a. Pull drilling mechanism away from container until the drill bit clears the ball valve. Close ball valve.
 - b. Loosen nut containing Teflon seal.
 - c. Unscrew bolts, holding drill assembly to mounting plate.
 - d. Remove drill assembly from mounting plate pulling drill bit through Teflon seal.
 - e. Remove cross fitting as unit from ball valve.

The remaining mounting plate and ball valve serve as a permanent seal until the container can be disposed of properly.

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SECTION 5.0

IONIZING RADIATION

5.1 GENERAL

Radiation monitoring should be one of the first tasks performed when initially approaching a waste site or hazardous material spill. This requirement is dictated by the potential risk to human health on contact with a radioactive source as exposure to even small amounts of energy may result in marked biological damage.

Radiation monitoring for hazardous waste situations essentially involves two approaches: personnel monitoring and survey monitoring. Personnel monitoring uses instruments designed to measure total cumulative radiation exposure which can be used to estimate the absorbed dose (in units of rad or rem). The instruments are worn or carried directly by the personnel being monitored and consist of such devices as film badges, thermoluminescent dosimeters, self-reading dosimeters, and pocket chambers. Survey instruments are meant to measure ionizing radiation -- expressed as an air exposure rate (in units of milliroentgens/hr) or activity of the source expressed as a disintegration rate (counts/minute). As do personnel monitors, these devices rely on the ability of radiation to cause ionizations and consist of ionization chambers, proportional counters, Geiger-Mueller instruments, and scintillation devices. They are particularly useful in performing initial field surveys to detect and locate the presence of radioactive sources and in drum screening procedures performed prior to further drum handling (i.e., staging, sampling, compositing, etc.).

Although all of these detection instruments rely on the ability of radiation to cause ionization, each differs in its sensitivity, i.e., its ability to detect different types and varying intensities of radiation. Basically there are four main groups of ionizing radiation types. These include:

- heavy, positively charged particles such as alpha particles, protons, deuterons, tritons, and possibly mesons each of which exhibit similar mechanisms of interaction with matter;
- beta particles including both positrons and electrons;
- electromagnetic radiation including x-ray and gamma radiation; and
- neutrons.

For the purposes of this section, however, only alpha, beta and gamma radiation will be discussed, as they are the types most likely to be encountered at a hazardous waste site.

Alpha particles are characterized as a charged particle having two protons and two neutrons and, due to this large mass and charge (in addition to high velocity), have a high probability of interacting or colliding with orbital electrons and atomic nuclei. They have a limited tissue penetration ability, however, since this type of radiation tends to lose its energy over short distances. It is therefore easy to shield against and poses little threat outside of the human body. However, due to its high specific ionization, alpha radiation is capable of totally destroying cellular material if it is able to locate within the body (e.g., by ingestion, inhalation, etc.).

Beta particles are negatively charged particles that can be construed as high-speed electrons. In contrast with electrons, however, beta particles originate in the nucleus. They exhibit medium specific ionization and penetration when compared to alpha particles. Although they pose a greater external body threat than alpha, beta particles of low energy are usually stopped by the horny dead layers of the skin. Beta particles with enough energy to penetrate the basal layer of the epidermis, however, still pose an external threat. They can be shielded by a few millimeters of aluminum and, like alpha particles, generally present a greater threat if their source is located inside the body.

Gamma radiation is a type of electromagnetic radiation of nuclear origin with a zero rest mass and no charge. It has the lowest specific ionization of the three classifications and possesses the ability to penetrate tissue for great distances. It therefore constitutes the greatest external radiation hazard (in comparison to alpha and beta) as it is capable of deep penetration within the body and is a threat to all organs. For this reason gamma radiation is the most routinely monitored radiation type at hazardous waste sites and environmental spills.

5.2 PERSONNEL MONITORS

Although no specific method is outlined in this manual for personnel radiation monitors (this is best covered by individual manufacturer instructions), it is important that their existence and basic characteristics be mentioned in this section. For this reason, the three basic types of personnel monitors, namely film badges, thermoluminescent dosimeters, and ion chambers, will be discussed as to the specific characteristics and relative advantages of each.

Film Badge

The use of films for monitoring personnel exposure is considered to be the most practical, although least accurate, of the existing methods. The method employs a gelatin base with a silver halide spread on film or glass. Radiation interacts within the silver halide in the emulsion by means of ionizations, thereby causing the formation of a latent image which, upon development, is converted into a black deposit of metallic silver. This darkening can then be related to the type, energy, and quantity of radiation received by the film badge. It is capable of recording a permanent record of personnel exposure.

Thermoluminescent Dosimeters

Thermoluminescent dosimeters (TLD) can replace film badges for most applications. In general they are more sensitive and more accurate than film badges and can be processed more quickly and less expensively. These devices detect radiation by storing ionization energy in defects of the crystal lattice of certain doped solids, such as LiF (Mn) and $\text{Ca F}_2 (\text{Mn})$. The altered energy levels are read out by heating the solid which then releases visible light. The light output is proportional to the absorbed radiation energy and can be related to exposure or dose units. TLD's can be reused but do not provide a permanent record of exposure because the information is erased upon readout. A permanent record is kept in the form of the original glow curve (light output vs. time (or temperature)) trace which can be stored on paper or in electronic memory.

Self-Reading Dosimeter

A self-reading dosimeter is essentially an ion chamber containing two electrodes, one being a thin quartz loop free to move with respect to its mounting and the other a fixed heavy quartz fiber. Like charges are placed on both loops causing the movable one to be repelled from the fixed loop. Ionization entering the chamber reduces the charge on the loops allowing the movable one to return towards its neutral position, the distance being proportional to the dose received in the chamber. The device also includes an optical system and transparent scale which permits instant results at any time without external readers. They are rugged, sensitive instruments small enough to be worn comfortably and extremely useful for measuring integrated exposure levels.

Pocket Chambers

A pocket ion chamber is basically a cylindrical electrode and a coaxial collecting rod which is insulated from the rest of the device. A charge is placed on the collecting rod, and this charge is subsequently reduced when ions formed upon exposure to radiation collect on the rod. The main disadvantage of the pocket chamber is that the collecting-rod charging procedure and the determination of exposure must be accomplished externally on a unit called a "charger-reader." The main advantages of pocket chambers, in comparison to the direct-reading dosimeter, are the low cost and simplicity.

5.3 SURVEY INSTRUMENTS

Radiation survey instruments must meet the same criteria as previously outlined for other monitors used at hazardous waste sites. They should be portable, rugged, sensitive, simple in design and operation, reliable, and intrinsically safe for use in explosive atmospheres. No one survey instrument or type of instrument can be expected to totally meet all of these criteria and the investigator must be aware of the characteristics (and limitations) of each type of detector.

It is of primary concern that the proper instruments are chosen for the particular survey requirements. Radiation survey instruments are designed to detect only certain types of radiation and only operate within certain exposure rate ranges. In most cases, more than one kind of instrument will be needed to insure that an area is free of radioactive sources or contamination. An instrument sensitive to background levels of gamma radiation should be the first one used. Scintillation detectors meet this requirement. Geiger-Mueller detectors also meet this need and have the additional advantages of being sensitive to beta radiation. Detection of alpha radiation requires another instrument having a thin window detector. Either gas ionization or scintillation principles may be employed in alpha detectors. This discussion will be limited to ionization chambers, proportional counters, Geiger-Mueller counters, and scintillation detectors.

Ionization Chambers

Ionization chambers are instruments in which the ionization initially produced within the chamber by radiation is measured without further gas amplification. It consists of a gas-filled envelope (usually air at atmospheric pressure) with two electrodes at different electrical potentials. The walls of the tube generally serve as the cathode and a wire mounted down the center of the tube serves as the anode. Ionizing radiation entering the chamber produces ions which migrate towards the electrode due to the applied potential, producing a current. This current requires amplification to a measurable level before it can be recorded on a meter. These are high-range instruments (low sensitivity) and are used extensively for measuring high intensity beta, gamma, or x-radiation. No aural indication is possible with these instruments and operators must be constantly aware of the meter to determine radiation intensity. Ionization chambers do not record individual radiation particles but integrate all signals produced as an electric current to drive the meter. They should be calibrated to the type and intensity of radiation desired to be measured in milliroentgens/hr (or roentgens/hr).

Proportional Counter

Instruments of this type derive their name due to their operation in the proportional region of the gas ionization detector response curve. Instrument probes have an extremely thin window that allows alpha particles to enter and as such are used extensively for this purpose by adjusting instrument operating parameters to discriminate against beta and gamma radiation. The meter is read in counts per minute and usually has several sensitivity

scales. It should be noted that due to the nature of alpha particles, it is important to hold the probe as close as possible to (though not in contact with) the surface being monitored. The window of the proportional counter is delicate in construction therefore requires care when using as a field instrument.

Geiger-Mueller Counter

These instruments operate principally in the same manner as ionization chambers except that secondary electrons are formed allowing for greater sensitivity. The chambers are filled with an inert gas such as argon, helium, or neon (below atmospheric pressure) and a quenching-gas which functions to control the secondary electron formation. These instruments are very sensitive and are commonly used to detect low level gamma and/or beta radiation. Meters are read in counts/minute or milliroentgens/hour. The gas amplification process inherent in this type of detector allows a single beta particle or gamma photon to be detected. It should be noted that these devices are sensitive instruments and care should be taken not to exceed their maximum capacity to prevent damage to the GM tube.

Scintillation Detectors

These devices depend upon light produced when ionizing radiation interacts with a media (solid crystal used in survey instruments). The produced flashes of light or scintillations fall onto a photomultiplier tube which converts them to electrical impulses. These impulses are amplified and subsequently measured to give an indication of the level of radiation present. These are extremely sensitive instruments used to detect alpha, beta, or gamma radiation simply by choosing the correct crystal. Alpha particles are detected with a silver activated zinc sulfide screen, beta radiation with an anthracene crystal (covered with a thin metal foil to screen alpha particles), and gamma or x-ray with a sodium iodide crystal. The instrument can be calibrated in the same manner as for ion chambers and Geiger-Mueller instruments. The operator should keep in mind that in older models the photomultiplier tube may be damaged if directly exposed to light without first disconnecting the voltage.

5.3.1 METHOD V-1: RADIATION SURVEY INSTRUMENTS

Discussion

As previously noted, a variety of radiation survey instrumentation exists, each capable of responding to different types and levels of ionizing radiation. The procedure delineated below is therefore purposely general and simply outlines common instrument features and operational steps. It is by no means meant to replace the instrument instruction manual but is only meant to serve as a supplemental guide.

Uses

Radiation survey instruments are used to detect the presence of radioactive sources. They are useful when making decisions concerning personal safety, determining levels of contamination, and meeting transportation and disposal requirements.

Procedures for Use

1. Choose an instrument or interchangeable detector tube which is consistent with the investigative requirements.
2. Turn selector switch to the standby or the warm-up position and allow instrument to warm-up for 1-2 minutes.
3. Turn instrument selector switch to battery check position and check battery strength.
4. Turn range selector switch to appropriate scale factor (e.g., 100X, 10X, 1X, 0.1X) and check or calibrate instrument with a radioactive check source (if available). Note: At a minimum, Coleman-type lantern mantles may be used as a check source. Lantern mantles are treated with a substance containing radioactive Thorium oxide.
5. Turn audio switch on if desired.
6. Choose needle response (fast/slow response).
7. Turn range selector to most sensitive setting and determine natural background radiation (0.01-0.02 mR/hr).
8. Scan suspected surfaces or areas. When in doubt, use most sensitive ranges first. Read scale in mR/hr or counts/minutes.

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SECTION 6.0

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APPENDIX A
SAMPLE CONTAINERIZATION
AND PRESERVATION

Acidity and Alkalinity

Apparatus and Materials:

- Polyethylene or borosilicate glass (Pyrex or equivalent) bottles.

Sample Collection, Preservation, and Handling:

- Fill sample bottles completely and cap tightly.
- Store samples at 4°C.
- All samples should be analyzed within 14 days of collection.

Quality Control:

- Dissolved gases contributing to acidity or alkalinity, such as carbon dioxide, hydrogen sulfide, or ammonia, may be lost or gained during sampling or storage. Sample bottles must be capped and sealed tightly, avoiding sample agitation or prolonged exposure to air.

Asbestos

Apparatus and Materials:

- 1-liter Polyethylene bottles

Sample Collection, Preservation and Handling:

- Leave air space at the top of the sample container to allow for shaking the sample.
- Avoid contacting the sample with acid
- If the sample cannot be filtered within 48 hours of collection, add 1 ml of a 2.71 percent solution of mercuric chloride per liter of sample to prevent bacterial growth.
- Store at 4°C

Quality Control:

The sample bottle should be rinsed at least twice with the water that is being sampled.

Bacteria

Apparatus and Materials:

- Polypropylene or glass bottles. Samples for bacteriological examination must be collected in bottles that have been cleansed and rinsed with great care, given a final rinse with distilled water, and sterilized.

Bottles of glass capable of being sterilized and of any suitable size and shape may be used for samples intended for bacteriological examination. Bottles shall hold a sufficient volume of sample for all the required tests, permit proper washing, and maintain the samples uncontaminated until the examinations are completed. Ground glass stoppered bottles, preferably wide-mouth and of break-resistant glass, are recommended. Polypropylene bottles of suitable size, wide-mouth, and capable of being sterilized are also satisfactory.

Metal or plastic screw cap closures may be used on sample bottles provided that no volatile compounds are produced on sterilization, and that they are equipped with liners that do not produce toxic or bacteriostatic compounds on sterilization.

Before sterilization, cover the tops and necks of sample bottles having glass closures with metal foil, rubberized cloth, heavy impermeable paper, or milk bottle cover caps.

Glassware shall be sterilized for not less than 60 minutes at a temperature of 170°C.

For plastic bottles that distort on autoclaving, low temperature ethylene oxide gas sterilization should be used.

- Sodium thiosulfate (ACS), 10 percent solution. When sampling water containing residual chlorine, sodium thiosulfate should be added to the clean sample bottle before sterilization in an amount sufficient to provide an approximate concentration of 100 mg/l in the sample. This can be accomplished by adding to a 500 ml bottle, 0.4 ml of a 10 percent solution of sodium thiosulfate (this will neutralize a sample containing about 15 mg/l of residual chlorine). The bottle is then stoppered, capped, and sterilized.
- Water samples high in copper or zinc and wastewater samples high in heavy metals should be collected in sample bottles containing a chelating agent that will reduce metal toxicity. This is particularly significant when such samples are in transit for 24 hours or more. Ethylenediaminetetraacetic acid (EDTA) is a satisfactory chelating agent. A concentration of 372 mg/l should be

added separately to the sample bottle before sterilization (0.3 ml of a 15 percent solution in a 500 ml bottle) or it may be combined with the sodium thiosulfate solution before addition.

Sample Collection, Preservation, and Handling:

- When the sample is collected, leave ample air space in the bottle (at least 2.5 cm or 1 in.) to facilitate mixing of the sample by shaking, preparatory to examination. Care must be exercised to take samples that will be representative of the water being tested and to avoid contamination of the sample at the time of collection or in the period before examination.

The sampling bottle shall be kept unopened until the moment it is to be filled. Remove the stopper and hood or cap as a unit, taking care to avoid soiling. During sampling, do not handle the stopper or cap and neck of the bottle and protect them from contamination. Hold the bottle near the base, fill it without rinsing, replace the stopper or cap immediately, and secure the hood around the neck of the bottle.

- Store samples at 4°C.
- All samples should be analyzed within 6 hours of collection.

Quality Control:

- The bacteriological examination of a water sample should be started promptly after collection to avoid unpredictable changes. The time and temperature of storage of all samples should be recorded and should be considered in the interpretation of data.

Bicarbonate/Carbonate

Apparatus and Materials:

- Polyethylene or glass bottles

Sample Collection, Preservation and Handling:

- Bicarbonate/Carbonate analysis should be performed onsite. If onsite determination is not possible, completely fill the sample bottle, leaving no headspace, and return it to the laboratory as quickly as possible for analysis.
- Store sample at 4°C until analyzed.

Quality Control:

- Carbon dioxide may be lost or gained during sampling and storage. Sample bottles must be capped and sealed tightly, avoiding sample agitation or prolonged exposure to air.

Biochemical-Oxygen Demand (BOD)

Apparatus and Materials:

- Polyethylene or glass bottles.

Sample Collection, Preservation, and Handling:

- If possible, avoid samples containing residual chlorine by sampling before chlorination. Notify laboratory if sample is from a chlorinated effluent.
- Store sample at 4° until analyzed.
- All samples should be analyzed within 48 hours of collection.

Quality Control:

- Samples for BOD analysis may undergo significant degradation during storage between collection and analysis, resulting in a low BOD value. Minimize reduction of BOD by promptly analyzing the sample.

Bromide

Apparatus and Materials:

- Polyethylene or glass bottles.

Sample Collection, Preservation, and Handling:

- There are no required preservation techniques, although storage at 4°C is recommended.
- All samples must be analyzed within 28 days of collection.

Quality Control:

- No special precautions.

Carbonate

See Bicarbonate/Carbonate

Chloride

Apparatus and Materials:

- Polyethylene or glass bottles.

Sample Collection, Preservation, and Handling:

- No preservative necessary.
- All samples must be analyzed within 28 days of collection.

Quality Control:

- No special precautions.

Chlorine Demand

Apparatus and Materials:

- Polyethylene or glass bottles.
- Testing apparatus and reagents, if analysis is to be performed onsite.

Sample Collection, Preservation, and Handling:

- Chlorine in aqueous solution is unstable, and the chlorine content of samples or solutions, particularly weak solutions, will decrease rapidly. Exposure to sunlight or other strong light or agitation will accelerate the reduction of chlorine. Therefore, sample must be analyzed onsite or brought immediately to the laboratory. The maximum holding time is 2 hours.

Quality Control:

- Chlorine determinations must begin immediately after sampling. Excessive light and agitation should be avoided.

Chromium VI

Apparatus and Materials:

- Polyethylene or glass bottles.

Sample Collection, Preservation, and Handling:

- Store samples at 4°C.
- All samples must be analyzed within 24 hours of collection.
- Do not contact sample with acid

Chemical Oxygen Demand (COD)

Apparatus and Materials:

- Polyethylene or glass bottles.
- Cone. sulfuric acid, H_2SO_4 (ACS).

Sample Collection, Preservation, and Handling:

- Preserve the sample by acidification with cone. sulfuric acid to a pH less than 2.
- Store samples at 4°C.
- All samples must be analyzed within 28 days of collection.

Quality Control:

- Unstable samples should be tested without delay.

Color

Apparatus and Materials:

- Polyethylene or glass bottles.

Sample Collection, Preservation, and Handling:

- Store samples at 4°C.
- All samples must be analyzed within 48 hours of collection.

Quality Control:

- No special precautions.

Conductance

Apparatus and Materials:

- Polyethylene or glass bottles.

Sample Collection, Preservation, and Handling:

- Store samples at 4°C.
- All samples must be analyzed within 28 days.

Quality Control:

- No special precautions.

Cyanide, Total and Amenable to Chlorination

Apparatus and Materials:

- Polyethylene or glass bottles.
- Sodium hydroxide solution (ACS).
- Ascorbic acid.

Sample Collection, Preservation, and Handling:

- Because most cyanides are highly reactive and unstable, analyze samples as soon as possible. Preserve the sample by addition of 2 ml of 10 N NaOH to raise the pH of the sample to 12 or above and store in a closed, dark bottle at 4°C.
- If residual chlorine is present in the sample, add 0.6 g ascorbic acid.
- All samples should be analyzed within 14 days of collection.

Quality Control:

- Maximum holding time is 24 hours when sulfide is present. Optionally, all samples may be tested with lead acetate paper before the pH adjustment in order to determine if sulfide is present. If sulfide is present, it can be removed by the addition of cadmium nitrate powder until a negative spot test is obtained. The sample is filtered and then NaOH is added to pH 12.

Fluoride

Apparatus and Materials:

- Polyethylene bottles.

Sample Collection, Preservation, and Handling:

- Polyethylene bottles are required for collecting and storing samples for fluoride analysis. Always rinse the bottle with a portion of the sample.
- All samples must be analyzed within 28 days of collection.

Quality Control:

- No special precautions.

Hardness

Apparatus and Materials:

- Polyethylene or glass bottles.

Sample Collection, Preservation, and Handling:

- Acidify with HNO_3 to pH 2, store samples at 4°C.
- Samples should be analyzed within 6 months of collection.

Quality Control:

- Serious errors may be introduced during sampling and storage by failure to remove residues of previous samples from the sample container; therefore all containers and sampling equipment should be thoroughly cleaned before use.

Hydrazine

Apparatus and Materials:

- Polyethylene or glass bottles.

Sample Collection, Preservation, and Storage:

- If the sample cannot be analyzed immediately, collect it under acid. Add 90 ml of sample to 100 ml of (1 + 9) HCl: one volume conc. HCl mixed with nine volumes **H₂O**.

Quality Control:

- Avoid contacting the sample with oxidizing agents which may diminish the hydrazine content.

Iodide

Apparatus and Materials:

- Polyethylene or glass containers.

Sample Collection, Preservation, and Handling:

- Store samples at 4°C, analyze within 24 hours of collection.

Quality Control:

- No special precautions.

Iodine

Apparatus and Materials:

- Polyethylene or glass containers

Sample Collection, Preservation, and Handling:

- The samples must be analyzed onsite or brought immediately to the laboratory. The maximum holding time is 2 hours.

Quality Control:

- Iodine determinations must begin immediately after sampling.

Metals - Except Chromium VI

Apparatus and Materials:

- Polyethylene or glass bottles.
- Nitric acid (1 + 1): Mix equal volumes of conc. nitric acid, HNO_3 (ACS), with deionized water.
- Deionized water.

Sample Collection, Preservation, and Handling:

- Wash and rinse sample container thoroughly with 1 + 1 nitric acid, then with deionized water before use.
- Acidify the sample with 1 + 1 nitric acid to a pH of 2.0 or less. Normally, 3 ml of 1 + 1 nitric acid per liter should be sufficient to preserve the samples. This will keep the metals in solution and minimize their adsorption on the container wall.
- All samples should be analyzed within 6 months of collection. An exception is mercury analysis, which must be completed within 28 days.

Quality Control:

- Serious errors may be introduced during sampling and storage by failure to remove residues of previous samples from the sample container; therefore, follow the described rinsing procedure for all containers and sampling equipment.

Nitrogen

Ammonia
Nitrate-Nitrite
Kjeldahl Nitrogen

Ammonia

Apparatus and Materials:

- Polyethylene or glass bottles.
- conc. sulfuric acid, H₂SO₄ (ACS).

Sample Collection, Preservation, and Handling:

- In the event that a prompt analysis is impossible, add conc. sulfuric acid to lower sample pH to less than 2.
- All samples should be analyzed within 28 days of collection.
- Store samples at 4°C.

Quality Control:

- The most reliable results are obtained from fresh samples.

Kjeldahl Nitrogen

Apparatus and Materials:

- Polyethylene or glass bottles.
- conc. sulfuric acid (H₂SO₄) (ACS).

Sample Collection, Preservation, and Handling:

- Acidify samples with conc. sulfuric acid to a pH of 2.0 or less.
- Store samples at 4°C.
- All samples should be analyzed within 28 days of collection.

Quality Control:

- The most reliable results are obtained in fresh samples. If prompt analysis is impossible, retard biological activity with the above preservation method.

Nitrate and Nitrite

Apparatus and Materials:

- Polyethylene or glass bottles.
- conc. sulfuric acid, H_2SO_4 (ACS)

Sample Collection, Preservation, and Handling:

- Store samples at 4°C.
- All samples should be analyzed within 48 hours of collection.
- If nitrate or nitrate plus nitrite are to be determined, preserve the sample by addition of H_2SO_4 to a pH of 2.0 or less.
- Sulfuric acid should not be added to samples requiring analysis for nitrite only.

Quality Control:

- Nitrate and nitrite determinations should be made promptly after sampling.

Oil and Grease

Apparatus and Materials:

- Glass bottles.
- Conc. sulfuric acid (**H₂SO₄**) (ACS).

Sample Collection, Preservation, and Handling:

- Collect a representative sample in a wide-mouth glass bottle and acidify in the sample bottle with conc. sulfuric acid to a pH of 2.0 or less. If other parameters are to be analyzed for, collect a separate sample for the oil and grease determination to avoid subdividing the sample in the laboratory.
- Store samples at 4°C.
- All samples should be analyzed within 28 days of collection.

Quality Control:

- Because losses of grease will occur on sampling equipment, the collection of a composite sample is impractical. Individual portions collected at prescribed intervals must be analyzed separately to obtain the average concentration over an extended period of time.

Organics

Purgeables - Method 624
Extractables - Method 625
Pesticides/PCBs - Method 608

Method 624 Purgeables

Apparatus and Materials:

- The water sample is to be collected in two (2) 40 ml vials with Teflon-faced silicone septa and screw caps and maintained at 4°C until the sampler's responsibility has been relieved at the Sample Bank.
- Container Preparation
 1. Wash 40 ml vials with screw caps (Pierce No. 13075 or equivalent) and Teflon-faced silicone septa (Pierce No. 12722 or equivalent) separately, utilizing a solution of Alconox detergent or equivalent, and hot tap water.
 2. Rinse thoroughly with deionized water.
 3. Place vials, caps, and septa on precleaned aluminum foil (as described above) and bake in an oven for one hour at 105°C.
 4. Allow the vials to cool with the septa properly inserted and the caps screwed on loosely. Tighten down caps when cool.
 5. Store vials in an area not subject to contamination by air or other sources.

Sample Collection, Preservation, and Handling

- If the sample contains residual chlorine, add sodium thiosulfate as a preservative (10 mg/40 ml is sufficient for up to 5 ppm Cl_2) to the empty sample bottles just prior to shipping to the sampling site.
- If aromatic compounds such as benzene, toluene and ethylbenzene are to be determined one of the following procedures should be used to minimize degradation of these compounds by microbial action.
 - Collect about 500 ml of sample in a clean container. Adjust the pH of the sample to about 2 by addition of 1+1 HCl. Cap the container and invert once to mix; check the pH with narrow range (1.4 to 2.8) pH paper. Transfer the sample to a 40 ml vial as described below. If residual chlorine is present, add sodium thiosulfate to another sample container and fill as described below.
 - Alternatively, the addition of the HgCl_2 to the sampling vial (approximately 12 mg per 40 ml vial) has been found effective for inhibiting microbial action.

- The following procedures apply to sampling directly with the sample vial.
 1. Collect a single undisturbed sample of water for the analysis of volatile organics. Submerge the sample vial just below the surface upside down and slowly invert. Accomplish this task creating as little disturbance as possible.
 2. Allow the vial to fill and reach equilibrium with its surrounding reservoir for several seconds.
 3. Place the cap over the mouth of the vial so that the septum is properly oriented and screw down the cap firmly.
 4. Invert the vial to discover any entrapped air bubbles. If such is the case, the sample will be discarded and another 40 ml vial selected and filled.
 5. Collect a replicate sample per instructions above.
 - Label the sample vials with the appropriate designated sample tag.
 - Place the properly labeled sample vials in an appropriate carrying container maintained at 4°C throughout the sampling and transportation period.
- Analyze samples within 14 days.

Quality Control:

- Standard quality assurance practices should be used with this method. Field replicates should be collected to validate the precision of the sampling techniques.
- Samples can be contaminated by diffusion of volatile organics (particularly methylene chloride) through the septum seal into the sample during shipment and storage. A field blank* prepared from organic-free water and carried through the sampling and handling protocol can serve as a check on such contamination.

*Field Blank. The field blank is defined as an appropriate volume of "organic-free" water which has been sent to the sampling site and back to the analytical laboratory in a container and bottle identical to the type used to collect the samples. Field blanks and samples must be shipped in separate containers. When received in the lab, the field blank is analyzed, as if it were an actual sample.

Method 625 Extractables (Base/Neutrals, Acids and Pesticides)

Apparatus and Materials:

- Sampling equipment, for discrete or composite sampling.

Grab sample bottle - Amber glass, 1 liter to 1 gallon volume. French or Boston Round design is recommended. The container must be washed and solvent rinsed before use to minimize interferences.

Bottle caps - Threaded to fit sample bottles. Caps must be lined with Teflon. Aluminum foil may be substituted if sample is not corrosive.

Compositing equipment - Automatic or manual compositing system. Must incorporate glass sample containers for the collection of a minimum of 1000 ml. Sample containers must be kept refrigerated during sampling. No plastic or rubber tubing other than Teflon may be used in the system.

Sample Collection, Preservation, and Handling:

- Grab samples must be collected in glass containers. Conventional sampling practices should be followed, except that the bottle must not be prerinsed with sample before collection. Composite samples should be collected in refrigerated glass containers. Automatic sampling equipment must be free of Tygon and other potential sources of contamination.
- The sample must be iced or refrigerated from the time of collection until extraction.
- All samples must be extracted within 7 days and completely analyzed within 30 days of collection.

Quality Control:

- Standard quality assurance practices should be used with this method.

- Glassware must be scrupulously clean. Clean all glassware as soon as possible after use by rinsing with the last solvent used. This should be followed by detergent washing in hot water. Rinse with tap water, distilled water, acetone and finally pesticide quality hexane. Heavily contaminated glassware may require treatment in a muffle furnace at 400°C for 15 to 30 minutes. Some high boiling materials, such as PCB's, may not be eliminated by this treatment. Glassware should be sealed/stored in a clean environment immediately after drying or cooling to prevent any accumulation of dust or other contaminants. Store inverted or capped with aluminum foil.

Method 608 Organochlorine Pesticides and PCBs

Apparatus and Materials:

- Sampling equipment, for discrete or composite sampling.

Grab sample bottle - Amber glass, 1 liter or 1 quart volume. French or Boston Round design is recommended. The container must be washed and solvent rinsed before use to minimize interferences.

Bottle caps - Threaded to screw on to the sample bottles. Caps must be lined with Teflon. Foil may be substituted if sample is not corrosive.

Compositing equipment - Automatic or manual compositing system. Must incorporate glass sample containers for the collection of a minimum of 250 ml. Sample containers must be kept refrigerated during sampling. No Tygon or rubber tubing may be used in the system.

Sample Collection, Preservation, and Handling:

- Grab samples must be collected in glass containers. Conventional sampling practices should be followed, except that the bottle must not be prewashed with sample before collection. Composite samples should be collected in refrigerated glass containers. Automatic sampling equipment must be free of Tygon and other potential sources of contamination.
- The samples must be iced or refrigerated from the time of collection until extraction.
- All samples must be extracted within 7 days and completely analyzed within 30 days of collection.

Quality Control:

- Standard quality assurance practices should be used with this method.

Hydrogen Ion (pH)

Apparatus and Materials:

- Polyethylene or glass bottles.
- Electronic pH meter with temperature compensation adjustment. Glass electrode: Glass electrodes are available for measurement over the entire pH range. Use minimum-sodium-ion-error type electrodes for high-pH high-sodium samples. Reference electrode: Use calomel, silver-silver chloride, or other constant-potential electrode.
- Standard buffer solutions of known pH.

Sample Collection, Preservation, and Handling:

- The electrometric measurement of pH is the only method approved by EPA. The determination should be made onsite. The maximum holding time for any sample is 2 hours.
- Because of the difference between the many makes and models of commercially available pH meters, it is impossible to provide detailed instructions for the proper operation of every instrument. In each case, follow the manufacturer's instructions. Thoroughly wet the glass and reference electrodes by immersing the tips in water overnight or in accordance with instructions. Thereafter, when the meter is not in use for pH measurement, keep the tips of the electrodes immersed in water.

Before use, remove the electrodes from the water and rinse with distilled or demineralized water. Dry the electrodes by gently blotting with a soft tissue. Standardize the instrument with the electrodes immersed in a buffer solution with a pH approaching that of the sample and note the temperature of the buffer and the pH at the measured temperature. Remove the electrodes from the buffer, rinse thoroughly, and blot dry. Immerse in a second buffer approximately 3 pH units different from the first and note the temperature of the buffer and the pH at the measured temperature; the reading should be within 0.1 unit of the pH for the second buffer. Rinse electrodes thoroughly, blot dry, and immerse in the sample. Agitate the sample sufficiently to provide homogeneity and keep solids in suspension. If the sample temperature is different from that of the buffers, let the electrodes equilibrate with the sample. Measure the sample temperature and set the temperature compensator on the pH meter to the measured temperature. Note and record the pH and temperature. Rinse electrodes and immerse in water until the next measurement.

When only occasional pH measurements are made, standardize the instrument before each measurement. Where frequent measurements are made, less frequent standardization (every 1 or 2 hours) is satisfactory. However, if sample pH values vary widely, standardize more frequently with a buffer having a pH within 1 to 2 pH units of that sample. Measure with two or more buffers of different pH at least once daily and more frequently if samples contain abrasive solids or dissolved fluorides, in order to check the linearity of response. When electrode response to two buffers 3 pH units apart show differences greater than 0.1 pH unit, replace the glass electrode.

Measurements of pH in high purity waters, such as condensate or demineralizer effluents, are subject to atmospheric contamination and require special procedures for accurate pH measurement.

Quality Control:

- The glass electrode is relatively free from interference from color, turbidity, colloidal matter, oxidants, reductants, or high salinity, except for a sodium error at high pH. This error at a pH above 10 may be reduced by using "low sodium error" electrodes. When using ordinary glass electrodes, make approximate corrections for the sodium error in accordance with information supplied by the manufacturer. Temperature exerts two significant effects on pH measurement. The pH potential, i.e., the change in potential per pH unit, varies with temperature, and ionization in the sample also varies. The first effect can be overcome by a temperature compensation adjustment provided on the better commercial instruments. The second effect is inherent in the sample and is taken into consideration by recording both the temperature and pH of each sample.

Phenols

Apparatus and Materials:

- Glass bottles.
- Concentrated Sulfuric Acid, H_2SO_4 (ACS).

Sample Collection, Preservation, and Handling:

- Acidify sample with concentrated H_2SO_4 acid to a pH of 2.0 or less.
- Oxidizing agents, such as chlorine, should be removed immediately after sampling by the addition of an excess of ferrous ammonium sulfate.
- Store samples at 4°C.
- All samples should be analyzed within 28 days of collection.

Quality Control:

- Phenols in concentrations usually encountered in wastewaters are subject to biological and chemical oxidation. It is recommended that preserved and stored samples be analyzed as soon as possible.

Orthophosphate

Apparatus and Materials:

- Polyethylene or glass bottles.

Sample Collection, Preservation, and Handling:

- Store samples at 4°C.
- All samples must be analyzed within 48 hours of collection.

Quality Control:

- Do not store samples containing low concentrations of phosphorus in plastic bottles because phosphate may be adsorbed onto the walls of the bottles. Rinse all glass containers with hot dilute HCl, then rinse several times in distilled water. Never use commercial detergents containing phosphate for cleansing glassware used in phosphate analyses.

Phosphorus, Total

Apparatus and Materials:

- Polyethylene or glass bottles.
- conc. sulfuric acid (H_2SO_4) (ACS).

Sample Collection, Preservation, and Handling:

- Acidify sample with conc. sulfuric acid to a pH of 2.0 or less.
- Store samples at 4°C.
- All samples must be analyzed within 28 days of collection.

Quality Control:

- Do not store samples containing low concentrations of phosphorus in plastic bottles because phosphate may be adsorbed onto the walls of the bottles. Rinse all glass containers with hot dilute HCl, then rinse several times in distilled water. Never use commercial detergents containing phosphate for cleansing glassware used in phosphate analyses.

Radioactivity

Apparatus and Materials:

- Polyethylene or glass bottles.
- Conc. nitric acid (HNO_3) (ACS).

Sample Collection, Preservation, and Handling:

- Acidify samples with conc. nitric acid to a pH of 2.0 or less.
- All samples must be analyzed within 6 months of collection.

Quality Control:

- The principles of representative sampling of water and wastewater apply to sampling for radioactivity examinations. When radioactive industrial wastes or comparable materials are sampled, consideration should be given to the deposition of radioactivity on the walls and surfaces of glassware, plastic containers, and equipment. Because a radioactive element is often present in submicrogram quantities, a significant fraction of it may be readily lost by adsorption on the surface of containers or glassware used in the examination. This may cause a loss of radioactivity and possible contamination of subsequent samples due to reuse of inadequately cleansed containers.

Silica

Apparatus and Materials:

- Polyethylene bottles.

Sample Collection, Preservation, and Handling:

- Collect samples in bottles of polyethylene plastic only; do not use glassware for any sample handling.
- Store samples at 4°C.
- All samples must be analyzed within 28 days of collection.

Quality Control:

- If samples are stored in glass, silica may leach into the sample and raise concentrations, therefore glassware cannot be used.

Solids

Apparatus and Materials:

- Polyethylene or glass bottles.

Sample Collection, Preservation, and Handling:

- Store samples at 4°C.
- Samples must be analyzed within the following times, according to the analysis to be performed:

Dissolved	7 days
Volatile Dissolved	7 days
Suspended	7 days
Volatile Suspended	7 days
Total	7 days
Volatile Total	7 days
Settleable	48 hours

Quality Control:

- Sample should be analyzed as soon as possible after collection for best results.
- Exclude unrepresentative particles such as leaves, sticks, or large solids.
- Glass bottles are desirable. Plastic bottles are satisfactory provided that the material in suspension in the sample does not adhere to the walls of the container. Store samples that are likely to contain iron or manganese so that oxygen will not come into contact with the water. Analyze these samples promptly to minimize the possibility of chemical or physical change during storage.

Sulfate

Apparatus and Materials:

- Polyethylene or glass bottles.

Sample Collection, Preservation, and Handling:

- In the presence of organic matter, certain bacteria may reduce sulfate to sulfide. To avoid this, samples are stored at 4°C.
- All samples must be analyzed within 28 days of collection.

Quality Control:

- No special precautions.

Sulfide

Apparatus and Materials:

- Polyethylene or glass bottles.
- Zinc Acetate [$\text{Zn}(\text{C}_2\text{H}_3\text{O}_2)_2$], 2 N.
- Sodium hydroxide (NaOH), 6 N.

Sample Collection, Preservation, and Handling:

- Take sample with a minimum of aeration. Preserve sample by addition of 2 ml of 2N zinc acetate; raise pH to 9 using NaOH. Fill sample bottle completely allowing no headspace.
- Store sample at 4°C.
- All samples must be analyzed within 7 days of collection.

Quality Control:

- It is important that all sample bottles are sealed airtight, with no entrapped air.

Sulfite

Apparatus and Materials:

- Polyethylene or glass bottles.

Sample Collection, Preservation, and Handling:

- Collect a fresh water sample, allow as little contact with air as possible, as air will oxidize the sulfite to sulfate.
- All samples should be analyzed onsite.

Quality Control:

- It is important that all sample bottles be sealed airtight, with no entrapped air.

Surfactants

Apparatus and Materials:

- Polyethylene or glass bottles.

Sample Collection, Preservation, and Handling:

- Store samples at 4°C.
- All samples must be analyzed within 48 hours of collection.

Quality Control:

- No special precautions.

Total Organic Carbon (TOC)

Apparatus and Materials:

- Glass bottles, with Teflon lined caps.
- conc. hydrochloric acid (**H₂SO₄**) (ACS).

Sample Collection, Preservation, and Handling:

- Acidify samples with conc. hydrochloric acid to a pH of 2.0 or less.
- Store samples at 4°C.
- All samples should be analyzed within 28 days of collection.

Quality Control:

- Avoid exposure of the sample to light and atmosphere, minimize storage time.

Total Organic Halide (TOX)

Apparatus and Materials:

- Glass bottles, amber, with Teflon lined caps.
- Sodium sulfite, **Na₂SO₃**, 0.1 M

Sample Collection, Preservation, and Handling:

- If amber glass bottles are not available, samples should be protected from light.
- Samples should be stored at 4°C without headspace.
- Reduce residual chlorine by the addition of 1 ml of 0.1 M sodium sulfite per liter of sample.
- TOX may increase with storage, therefore, samples should be analyzed as soon as possible after collection; maximum holding time should not exceed 7 days.

Quality Control:

- Glassware used in TOX sampling and analysis must be thoroughly cleaned. All glassware should be washed using detergent and hot water, rinsed with tap water and, as a final rinse, deionized water. Drain dry and heat at 105°C for 1 hour. Glassware should be sealed and stored in a clean area after drying and cooling.

Turbidity

Apparatus and Materials:

- Polyethylene or glass bottles.

Sample Collection, Preservation, and Handling:

- Store samples at 4°C.
- All samples must be analyzed within 48 hours of collection.

Quality Control:

- Turbidity analysis should be performed on the day the sample is taken. If longer storage is unavoidable, store samples in the dark for up to 48 hours. Prolonged storage before measurement is not recommended because irreversible changes in turbidity may occur.

TABLE A-1. RECOMMENDED SAMPLING AND PRESERVATION PROCEDURES FOR WATER AND WASTEWATER

Parameter	Collection technique	Container ^a	Preservation	Holding time ^b	Minimum required volume (mL)
Acidity	Grab or composite	P,G	Cool, 4°C	14 days	100
Alkalinity	Grab or composite	P,G	Cool, 4°C	14 days	100
Asbestos	Grab or composite	P	Cool, 4°C	48 hours	1000
Bacteria	Grab only	Pro, G	Cool, 4°C, 10% Na ₂ S ₂ O ₃ , EDTA	6 hours	200
Bicarbonate	Grab only	P,G	Determine onsite	No holding	100
BOD	Grab only	P,G	Cool, 4°C	48 hours	1000
Bromide	Grab or composite	P,G	None required	28 days	100
Carbonate	Grab only	P,G	Determine onsite	No holding	100
Chloride	Grab or composite	P,G	None required	28 days	50
Chlorine demand	Grab only	P,G	Determine onsite	No holding	200
Chromium VI	Grab or composite	P,G	Cool, 4°C	24 hours	100
COD	Grab only	P,G	H ₂ SO ₄ to pH <2; Cool, 4°C	28 days	50
Color	Grab or composite	P,G	Cool, 4°C	48 hours	50
Conductance	Grab or composite	P,G	Cool, 4°C	28 days	100
Cyanide	Grab or composite	P,G	NaOH to pH >12, 0.6g Ascorbic acid ^d	14 days	500
Fluoride	Grab or composite	P	None required	28 days	300
Hardness	Grab or composite	P,G	HNO ₃ to pH <2	6 months	100
Hydrazine	Grab or composite	P,G	If not analyzed immediately, collect under acid. Add 90 ml of sample to 10 ml (1 + 9) HCl	7 days	100

(continued)

TABLE A-1 (continued)

Parameter	Collection technique	Container ^a	Preservation	Holding time ^b	Minimum required volume (mL)
Iodide	Grab or composite	P,G	Cool 4°C	24 hours	100
Iodine	Grab only	P,G	Determine onsite	No holding	500
<u>Metals (Except Cr VI)</u>					
Dissolved	Grab or composite	P,G	Filter onsite, HNO ₃ to pH <2	6 months, except Hg--28 days	200
Suspended	Grab or composite	P,G	Filter onsite	6 months, except Hg--28 days	200
Total	Grab or composite	P,G	HNO ₃ to pH <2	6 months, except Hg--28 days	100
<u>Nitrogen</u>					
Ammonia	Grab or composite	P,G	Cool, 4°C, H ₂ SO ₄ to pH <2	28 days	400
Kjeldahl (total)	Grab or composite	P,G	Cool, 4°C, H ₂ SO ₄ to pH <2	28 days	500
Nitrate plus Nitrite	Grab or composite	P,G	Cool, 4°C, H ₂ SO ₄ to pH <2	28 days	100
Nitrate	Grab or composite	P,G	Cool, 4°C, H ₂ SO ₄ to pH <2	48 hours	100
Nitrite	Grab or composite	P,G	Cool 4°C, H ₂ SO ₄ to pH <2	48 hours	50
Oil and Grease	Grab only	G	Cool 4°C, H ₂ SO ₄ to pH <2	28 days	1000
<u>Organics</u>					
Extractables (base/neutral and acids)	Grab or composite	G, Teflon-lined cap	Cool, 4°C	7 days until extraction, 30 days after extraction	1000

(continued)

TABLE A-1 (continued)

Parameter	Collection technique	Container ^a	Preservation	Holding time ^b	Minimum required volume (mL)
Organics (cont.)					
Purgeables (halocarbons-aromatics)	Grab only	G, Teflon-lined cap	Cool, 4°C	14 days	40
Purgeables (acrolein and acrylonitrile)	Grab only	G, Teflon-lined cap	Cool, 4°C	14 days	40
Pesticides and PCBs	Grab or composite	G, Teflon-lined cap	Cool, 4°C	7 days until extraction, 30 days after extraction	250
pH	Grab only	P,G	Determine onsite	2 hours	25
Phenol	Grab or composite	G	Cool, 4°C, H ₂ SO ₄ to pH <2	24 hours	500
Phosphorus					
Ortho phosphate	Grab or composite	P,G	Filter onsite, cool, 4°C	48 hours	50
Phosphorus, Total	Grab or composite	P,G	Cool, 4°C, H ₂ SO ₄ to pH <2	28 days	50
Radioactivity	Grab or composite	P,G	HNO ₃ to pH <2	6 months	1 gal
Silica					
Dissolved	Grab or composite	P	Cool, 4°C	28 days	50
Total	Grab or composite	P	Cool, 4°C	28 days	50
Solids					
Dissolved	Grab or composite	P,G	Cool, 4°C	7 days	100
Volatile Dissolved	Grab or composite	P,G	Cool, 4°C	7 days	100
Suspended	Grab or composite	P,G	Cool, 4°C	7 days	100

TABLE A-1 (continued)

Parameter	Collection technique	Container ^a	Preservation	Holding time ^b	Minimum required volume (mL)
Solids (cont.)					
Volatile Suspended	Grab or composite	P,G	Cool, 4°C	7 days	100
Total	Grab or composite	P,G	Cool, 4°C	7 days	100
Volatile Total	Grab or composite	P,G	Cool, 4°C	7 days	100
Settleable	Grab or composite	P,G	Cool, 4°C	48 hours	100
Sulfate	Grab or composite	P,G	Cool, 4°C	28 days	50
Sulfide	Grab or composite	P,G	Cool, 4°C, 2 ml zinc acetate plus NaOH to pH >9	7 days	500
Sulfite	Grab or composite	P,G	Determine onsite	No holding	50
Surfactants	Grab or composite	P,G	Cool, 4°C	48 hours	250
TOC	Grab or composite	G, Teflon-lined cap	Cool, 4°C, HCl to pH <2	28 days	25
TOX	Grab or composite	G, Amber, Teflon-lined cap	Cool, 4°C, add 1 ml 0.1 M sodium sulfite	7 days	100
Turbidity	Grab or composite	P,G	Cool, 4°C	48 hours	100

^aP = Polyethylene, G = Glass, Pro = Polypropylene

^bThe holding times are those listed in Technical Additions to Methods for Chemical Analysis of Water and Wastes, EPA-600/4-82-055 and Methods for Organic Chemical Analysis of Municipal and Industrial Wastewater, EPA-600/4-82-057.

^cIf samples cannot be filtered within 48 hours, add 1 ml of a 2.71% solution of mercuric chloride to inhibit bacterial growth.

^dShould only be used in the presence of residual chlorine.

APPENDIX B
EQUIPMENT AVAILABILITY AND FABRICATION

EQUIPMENT AVAILABILITY

Apparatus

- Stainless Steel Scoops, Trays, Beakers, Ladles
8, 9, 15
- Thin Wall Tube Samplers, Soil Augers, Hand Corers
45, 50
- Gravity Corers, Dredges and Grabs
40, 45
- Thieftakers and Triers
9, 34
- Water Level Indicators
38, 45
- Down Hole Submersible Probes
23, 25, 43, 51
- Bailers, Coliwasa
26, 34, 48
- Peristaltic Pumps
8, 9, 15, 29
- Gas Displacement Pumps
5, 48
- Combustible Gas Detectors
3, 13, 16, 17, 33, 36, 41
- Oxygen Monitors
6, 13, 16, 17, 33, 36, 41
- Portable Flame Ionization Detectors
1, 2
- Portable Photoionization Detectors
22, 37
- Stain Detector Tubes
7, 17, 31, 33, 35
- Personal Sampling Pumps
7, 11, 14, 19, 28, 32, 33, 39, 46
- High Volume Air Samplers
18, 39, 44

- Radiation Dosimeters
4, 10, 20, 47, 49
- Radiation Film Badges
12, 20, 24, 42
- Radiation Survey Instruments
4, 10, 12, 20, 21, 27, 30, 47, 49

Vendors

1. Analabs, Inc.
80 Republic Drive
North Haven, CT 06473
(203) 288-8463
2. Analytical Instrument Development, Inc.
Rt. 41 and Newark Rd.
Avondale, PA 19311
(215) 268-3181
3. Bacharach Instrument Company
301 Alpha Drive
Pittsburgh, PA 15238
(412) 782-3500
4. Baird Atomic
125 Middlesex Turnpike
Bedford, MA 01730
(617) 276-6000
5. BarCad System, Inc.
P.O. Box 424
Concord, MA 01742
(617) 969-0050
6. Beckman Instruments, Inc.
Process Instrument Division
2500 Harbor Boulevard
Fullerton, CA 92634
7. Bendix Corporation
Environmental and Process Instruments Division
P.O. Drawer 831
Ronceverte, WV 24970
(304) 647-4358
8. Cole Palmer
7425 North Oak Park Ave.
Chicago, Illinois 60648
(800) 323-4340

9. Curtin Matheson Scientific
Major Metropolitan Areas
10. Dosimeter Corporation of America
P.O. Box 42377
Cincinnati, OH 45242
(513) 489-8100
11. DuPont Company
Applied Technology Division
Concord Plaza - Clayton Bldg.
Wilmington, DE 19898
(302) 772-5989
12. Eberline Instruments
P.O. Box 2108
Santa Fe, NM 87501
(505) 471-3232
13. Energetic Science
Six Skyline Drive
Hawthorne, NY 10532
14. Environmental Measurements, Inc.
215 Leidesdorff Street
San Francisco, CA 94111
(415) 398-7664
15. Fisher Scientific
Major Metropolitan Areas
16. Gas Measurement Instruments Ltd.
Inchinnan Estate
RenfrewPA49RG
(041) 812-3211
17. GasTech Inc.
Johnson Instrument Division
331 Fairchild Drive
Mountain View, CA 94043
(415) 967-6794
18. General Metal Works Inc.
8368 Bridgetown Road
Village of Cleves, OH 45002
(513) 941-2229
19. Gilian Instrument Corp.
1275 Route 23
Wayne, NJ 07470
(201) 696-9244

20. Gulf Nuclear
202 Medical Center Boulevard
Webster, TX 77598
(713) 332-3581
21. Health Physics Instruments
124 San Felicia Drive
Goleta, CA 93117
(805) 685-2612
22. HNU Systems, Inc.
30 Ossipee Road
Newton Upper Falls, MA 02164
(617) 964-6690
23. Hydrolab Corporation
P.O. Box 9406
Austin, TX 78766
(512) 255-8841
24. ICN Dosimetry Service
26201 Niles Road
Cleveland, OH 44128
(216) 831-3000
25. Industrial and Environmental Analysts Inc.
P.O. Box 626
Essex Junction, VT 05452
(802) 878-5138
26. Johnson Division
UOP, Inc.
St. Paul, MN 55164
(612) 636-3900
27. Johnston Laboratories
P.O. Box 20086
383 Hillen Road
Towson, MD 21204
(301) 337-8700
28. Kurz Instruments Inc.
P.O. Box 849
Carmel Valley, CA 93924
(408) 659-3421
29. Leonard Mold and Die
960 West 48th Avenue
Denver, CO 80221
(303) 433-7101

30. Ludlum Measurements
P.O. Box 248
Sweetwater, TX 79556
(915) 235-5494
31. Matheson Safety Products
P.O. Box 85
932 Paterson Plank Road
East Rutherford, NJ 07073
(201) 933-2400
32. MDA Scientific, Inc.
1815 Elmdale Ave.
Glenview, IL 60025
33. Mine Safety Appliance Co.
600 Penn Center Boulevard
Pittsburgh, PA 15235
34. Nasco
901 Janesville Ave.
Fort Atkinson, WI 53538
(414) 563-2446
35. National Draeger, Inc.
101 Technology Drive
Pittsburgh, PA 15275
(412) 787-8383
36. National Mine Service Company
Industrial Safety Division
355 N. Old Steubenville Pike
Oakdale, PA 15071
(412) 788-4353
37. Photovac, Incorp.
134 Doncaster Ave.
Unit 2
Thornhill
Ontario, Canada L3T1L3
38. Powers Electric Products Company
P.O. Box 11591
Fresno, CA 93774
39. Research Appliance Company
Moose Lodge Road
Cambridge, MD 21613
(301) 228-9505

40. Research Instrument Manufacturing Co. Ltd.
RR No. 2 Guelph
Ontario, Canada N1H6H8
(519) 822-1547
41. Rexnord Safety Products/Biomarine Ind.
45 Great Valley Parkway
Malvern, PA 19355
(215) 647-7200
42. R. S. Landauer Jr. Company
Division of Technical Operations, Inc.
Science Road
Glenwood, IL 60425
(312) 755-7000
43. Sensorex
9713 Bolsa Ave.
Westminster, CA 92683
(714) 554-7090
44. Sierra Instruments Inc.
P.O. Box 909
Carmel Valley, CA 93924
(408) 659-3177
45. Soiltest, Inc
2205 Lee Street
Evanston, IL 60202
(312) 869-5500
46. Spectrex Corporation
3594 Haven Ave.
Redwood City, CA 94063
(415) 365-6567
47. Technical Associates
7051 Eton Avenue
Canoy Park, CA 91303
(213) 883-7043
48. Timco Manufacturing Company, Inc.
P.O. Box 35
Prairie Du Sac, WI 53578
(608)-643-8534
49. Victoreen, Inc.
10101 Woodland Ave.
Cleveland, OH 44104
(216) 795-8200

50. Wildco
301 Cass Street
Saginaw, MI 48602
(517) 799-8100
51. Yellow Springs Instrument Co.
Yellow Springs, OH 45387
(513) 767-7241

EQUIPMENT FABRICATION

Many of the instruments and devices listed previously can also be readily fabricated in-house. This usually affords considerable cost savings as well as allows for custom designs and alterations.

Bailers, coliwasas and hand corers can be constructed from available stainless steel and teflon stock. The diagrams and drawings which accompany their description in the text show nominal dimensions and construction materials. Sizes can however be altered to fit particular needs. The sources cited with these drawings as well as the references at the end of the method comment further on their construction and use.

The device used in Method IV-13: Sampling of Headspace Gases in Sealed Vessels, is not currently available through commercial sources. The fabrication details are therefore included in this Appendix.

Sealed Vessel Tapping Device Assembly

1. Fabricate mounting plate.
2. Position Portalign on mounting plate, drill 6.4 mm holes through Portalign base and mounting plate. Tap holes for 7.14 mm thread in mounting plate. Secure Portalign to mounting plate with 7.14 mm SAE bolts.
3. Thread ball valve into mounting plate.
4. Thread Swagelok cross assembly onto ball valve.
5. Insert drill bit into chuck of drill.
6. Insert drill into Portalign assembly per manufacturer's instruction. Pass drill bit through Teflon ferrule.
7. Place part 101-6 so that it stops drill bit travel approximately 10 mm below bottom of gasket material on mounting plate.
8. Mount entire assembly onto container using standard steel strap packaging equipment.
9. Place springs over Portalign guide rods.
10. Push springs down until good tension is obtained. Secure with extra 101-6 and 101-8 parts.
11. Finger tighten compression nut containing Teflon ferrules.

DRUM DRILLING MECHANISM

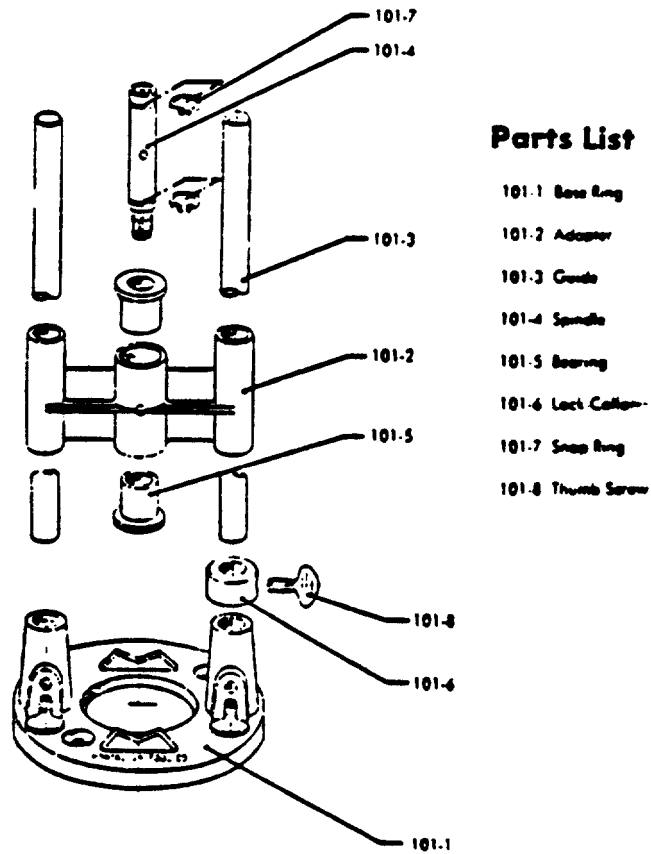
Parts

1. Mounting Plate-- 12.7 mm thick x 76 mm wide x 127 mm long, mild steel.

19.1 mm x 3.2 mm deep channel on top of each side. A 6.4 mm NPT hole in center of plate.

Bottom of mounting plate covered with 4.8 mm thick closed cell Neoprene gasket.
2. Ball Valve-- 316 stainless steel, 6.4 mm male NPT thread one end, 6.4 mm female NPT other end.
3. Swagelok Cross-- 316 stainless steel, three sides 6.4 mm male NPT, 6.4 mm Swagelok side.

Assemble as Follows:
 - A. 0-50 psig pressure gauge, 6.4 mm female NPT to one side of cross.
 - B. 316 stainless steel, 6.4 mm male NPT to 6.4 Swagelok needle valve, mount opposite pressure gauge.
 - C. 6.4 mm Teflon ferrules into 6.4 mm Swagelok fitting.
4. Drill Bit-- 4 mm drill bit, 140 mm long, flutes approximately 12 mm long.
5. Drill-- Skill Model No. 2002 hand drill, cordless. Wired to operate remotely at 300 rpm. Interlocked with microswitch attached to depth stop.
6. Portalign Drill Assembly-- Portalign, Portalign Tool Company, San Diego, California, as shown below.



Parts List

- 101-1 Base Ring
- 101-2 Adapter
- 101-3 Guide
- 101-4 Spindle
- 101-5 Bearing
- 101-6 Lock Collar
- 101-7 Snap Ring
- 101-8 Thumb Screw

Portalign drill guide

Additional Parts Required Per Assembly

2 each 101-8 }
2 each 101-6 } From Portalign

2 each Springs to fit over guide rods of Portalign approximately
30 kg force each spring when compressed.

APPENDIX C

PACKING, MARKING, LABELING, AND SHIPPING OF HAZARDOUS MATERIAL SAMPLES

Portions of this Appendix have been taken,
by permission, from EPA/ERT
"Hazardous Materials Incident Response Operations"
Training Course Manual (165.1)

INTRODUCTION

Samples collected during a response to a hazardous material incident may have to be transported elsewhere for analysis. The Environmental Protection Agency (EPA) encourages compliance with Department of Transportation (DOT) regulations governing the shipment of hazardous materials. These regulations (49 CFR parts 171 through 179) describe proper marking, labeling, packaging and shipment of hazardous materials, substances and wastes. In-particular, part 172.402(h) of 49 CFR is intended to cover shipment of samples of unknown materials destined for laboratory analysis.

ENVIRONMENTAL SAMPLES VERSUS HAZARDOUS MATERIAL SAMPLES

Samples collected at an incident should be classified as either environmental or hazardous material (or waste) samples. In general, environmental samples are collected offsite (for example from streams, ponds, or wells) and are not expected to be grossly contaminated with high levels of hazardous materials. Onsite samples (for example, soil, water, and materials from drums or bulk storage tanks, obviously contaminated ponds, lagoons, pools, and leachates from hazardous waste sites) are considered hazardous. A distinction must be made between the two types of samples in order to:

- Determine appropriate procedures for transportation of samples. If there is any doubt, a sample should be considered hazardous and shipped accordingly.
- Protect the health and safety of laboratory personnel receiving the samples. Special precautions are used at laboratories when samples other than environmental samples are received.

The following section describes the packaging, labeling and shipping requirements for these two sample types. Specific DOT regulations for hazardous materials shipping papers and general marking requirements are presented as Attachments C-1 and C-2.

ENVIRONMENTAL SAMPLES

Environmental samples must be packaged and shipped according to the following procedures.

Packaging

Environmental samples may be packaged following the procedures outlined later for samples classified as "flammable liquids" or "flammable solids," but the requirements for marking, labeling, and shipping papers do not apply.

Environmental samples may also be packaged without being placed inside metal cans as required for flammable liquids or solids.

- Place sample container, properly identified and with a sealed lid, in a polyethylene bag, and seal bag.

- Place sample in a fiberboard container or metal picnic cooler which has been lined with a large polyethylene bag.
- Pack with enough noncombustible, absorbent, cushioning material to minimize the possibility of the container breaking.
- Seal large bag.
- Seal or close outside container.

Marking/Labeling

Sample containers must have a completed sample identification tag and the outside container must be marked "Environmental Sample." The appropriate side of the container must be marked "This End Up" and arrows placed accordingly. No DOT marking or labeling are required.

Shipping Papers

No DOT shipping papers are required.

Transportation

There are no DOT restrictions on mode of transportation.

RATIONALE: HAZARDOUS MATERIAL SAMPLES

Samples not determined to be environmental samples or samples known or expected to contain hazardous materials must be considered hazardous substance samples and transported according to the following requirements:

- If the substance in the sample is known or can be identified, package, mark, label, and ship according to the specific instructions for that material (if it is listed) in the DOT Hazardous Materials Table, 49 CFR 172.101.
- For samples of hazardous materials of unknown content, part 172.402 of 49 CFR allows the designation of hazard class based on the shipper's knowledge of the material and selection of the appropriate hazard class from part 173.2 (see Table C-1).

The correct shipping classification for an unknown sample is selected through a process of elimination, utilizing the DOT classification system (Table C-1). Unless known or demonstrated otherwise (through the use of radiation survey instruments), the sample is considered radioactive and appropriate shipping regulations for "radioactive material" followed. If radioactive material is eliminated, the sample is considered to contain "Poison A" materials (Table C-2), the next classification on the list. DOT defines "Poison A" as extremely dangerous poisonous gases or liquids of such a nature that a very small amount of gas, or vapor of the liquid, mixed with air is dangerous to life.

TABLE C-1. DOT PRIORITY RANKING OF HAZARDOUS MATERIALS

=====			
	Category	Definition	Application regulations
	General		49 CFR 173.1-173.34, 177
1	Radioactive Material	49 CFR 173.389	49 CFR 173.390-173.398
2	Poison A	49 CFR 173.326	49 CFR 173.327-173.337
3	Flammable Gas	49 CFR 173.300	49 CFR 173.300-173.316
4	Nonflammable Gas	49 CFR 173.300	49 CFR 173.300-173.316
5	Flammable Liquid	49 CFR 173.115	49 CFR 173.116-173.119, 173.121-173.149a
6	Oxidizer	49 CFR 173.151	49 CFR 173.152-173.239a
7	Flammable Solid	49 CFR 173.150	49 CFR 173.152-173.239a
8	Corrosive Material (Liquid)	49 CFR 173.240	49 CFR 173.241-173.299a
9	Poison B	49 CFR 173.343	49 CFR 173.344-173.379
10	Corrosive Material (Solid)	49 CFR 173.240	49 CFR 173.241-173.299a
11	Irritating Materials	49 CFR 173.381	49 CFR 173.381-173.385
12	Combustible Liquid (in containers exceeding 100 gal capacity)	49 CFR 173.115	49 CFR 173.116-173.118a, 173.121-173.149a
13	ORM-B	49 CFR 173.800	49 CFR 173.510, 173.800-173.862
14	ORM-A	49 CFR 163.605	49 CFR 173.510, 173.605-173.655
15	Combustible Liquid (in containers having capacities of 110 gal or less)	49 CFR 173.115	49 CFR 173.116-173.118a, 173.121-173.149a
16	ORM-E	49 CFR 173.1300	49 CFR 173.510
=====			

TABLE C-2. DOT LIST OF CLASS "A" POISONS (49 CFR 172.101)

Material	Physical state at standard temperature
Arsine	Gas
Bromoacetone	Liquid
Chloropicrin and methyl chloride mixture	Gas
Chloropicrin and nonflammable, nonliquefied compressed gas mixture	Gas
Cyanogen chloride	Gas (13.1 °C)
Cyanogen gas	Gas
Gas identification set	Gas
Germane	-
Grenade (with Poison "A" gas charge)	-
Hexaethyl tetraphosphate/compressed gas mixture	Gas
Hydrocyanic acid (prussic) solution	Liquid
Hydrocyanic acid, liquefied	Gas
Insecticide liquefied gas containing Poison "A" or Poison "B" material	Gas
Methyldichloroarsine	Liquid
Nitric oxide	Gas
Nitrogen peroxide	Gas
Nitrogen tetroxide	Gas
Nitrogen dioxide, liquid	Gas
Parathion/compressed gas mixture	Gas
Phosgene (diphosgene)	Liquid

Most poison A materials are gases or compressed gases and would not be found in drum-type containers. Liquid poison A's would be found only in closed containers. All samples taken from closed drums do not have to be shipped as poison A's, which provides for a "worst case" situation. Based upon information available, a judgment must be made whether a sample from a closed container is a poison A.

If poison A is eliminated as a shipment category, the next two classifications are "flammable" or "nonflammable" gases. Since few gas samples are collected, "flammable liquid" would be the next applicable category. With the elimination of radioactive material, poison A, flammable gas, and nonflammable gas, the sample can be classified as flammable liquid (or solid) and shipped accordingly. These procedures would also suffice for shipping any other samples classified below flammable liquids in the DOT classification table.

For samples containing unknown material, other categories listed below flammable liquids/solids on the table are generally not considered because eliminating other substances as flammable liquids requires flashpoint testing, which may be impractical and possibly dangerous at a site. Thus, unless the sample is known to consist of material listed below flammable liquid on the table, it is considered a flammable liquid (or solid) and shipped as such.

PROCEDURES: SAMPLES CLASSIFIED AS FLAMMABLE LIQUID (OR SOLID)

The following procedure is designed to meet the requirements for a "limited quantity" exclusion for shipment of flammable liquids and solids, as set forth in parts 173.118 and 173.153 of 49 CFR. By meeting these requirements, the DOT constraints on packaging are greatly reduced. Packaging according to the limited quantity exclusion requires notification on the shipping papers.

Packaging

1. Collect sample in a glass container (16 ounces or less) with a nonmetallic, teflon-lined screw cap. To prevent leakage, fill container no more than 90 percent full at 130°F. If an air space in the sample container would affect sample integrity, place that container within a second container to meet 90 percent requirement.
2. Complete sample identification tag and attach securely to sample container.
3. Seal container and place in 2-mil thick (or thicker) polyethylene bag, one sample per bag. Position identification tag so it can be read through bag. Seal bag.
4. Place sealed bag inside metal can and cushion it with enough noncombustible, absorbent material (for example, vermiculite or diatomaceous earth) between the bottom and sides of the can and bag

to prevent breakage and to absorb leakage. Pack one bag per can. Use clips, tape, or other positive means to hold can lid securely, tightly, and permanently.

5. Place one or more metal cans into a strong outside container, such as a metal picnic cooler or a DOT approved fiberboard box. Surround cans with noncombustible, absorbent, cushioning material for stability during transport.
6. Limited quantities of flammable liquids, for the purpose of the exclusion, are defined as one pint or less (49 CFR part 173.118(a)(2)).
7. Limited quantities of flammable solids, for the purpose of this exclusion, are defined as one pound net weight in inner containers and no greater than 25 pounds net weight in the outer container (49 CFR part 173.153(a)(1)).

Marking/Labeling

1. Use abbreviations only where specified.
2. Place following information, either hand printed or in label form, on the metal card.
 - Laboratory Name and Address
 - "Flammable Liquid, n.o.s. UN1993" or "Flammable Solid, n.o.s. UN1325."

Not otherwise specified (n.o.s.) is not used if the flammable liquid (or solid) is identified. Then the name of the specific material is listed before the category (for example, Acetone, Flammable Liquid) followed by its appropriate UN number found in the DOT hazardous materials table (172.101).
3. Place the following DOT labels (if applicable) on outside of can (or bottle).
 - "Flammable Liquid" or "Flammable Solid."
 - "Dangerous When Wet." Must be used with "Flammable Solid" label if material meets the definition of a water-reactive material.
 - "Cargo Aircraft Only." Must be used if net quantity of sample in each outer container is greater than 1 quart (for "Flammable Liquid, n.o.s.") or 25 pounds (for "Flammable Solid, n.o.s.)."
4. Place all information on outside shipping container as on can (or bottle), specifically,

- Proper shipping name.
- UN or NA number.
- Proper label(s).
- Addressee and addressor.

(Note that the previous two steps (2 and 3) are EPA recommendations. Step 4 is a DOT requirement.

5. Print "Laboratory Samples and "This End Up" or "This Side Up" clearly on top of shipping container. Put upward pointing arrows on all four sides of container.

Shipping Papers

1. Use abbreviations only where specified.

Complete carrier-provided bill of lading and sign certification statement (if carrier does not provide, use standard industry form). Provide the following information in the order listed. (One form may be used for more than one exterior container.)

- "Flammable Liquid, n.o.s. UN1993" or "Flammable Solid, n.o.s. UN1325."
 - "Limited Quantity" (or "Ltd. Qty.").
 - Net weight or net volume (weight or volume may be abbreviated) just before or just after "Flammable Liquid, n.o.s. UN1325" or "Flammable Solid, n.o.s. UN1325"
 - Further descriptions such as "Laboratory Samples" or "Cargo Aircraft Only" (if applicable) are allowed if they do not contradict required information.
3. Include chain-of-custody record, properly executed, in outside container if legal use of samples is required or anticipated.

Transportation

1. Transport unknown hazardous substance samples classified as flammable liquids by rented or common carrier truck, railroad, or express overnight package services.
2. Do not transport by any passenger-carrying air transport system, even if they have cargo only aircraft. DOT regulations permit regular airline cargo only aircraft, but difficulties with most suggest avoiding them. Instead, ship by airlines that only carry cargo.

3. Transport by government-owned vehicle, including aircraft. DOT regulations do not apply, but EPA personnel will still use procedures described except for execution of the bill of lading with certification.

Other Considerations

1. Check with analytical laboratory for size of sample to be collected and if sample should be preserved or packed in ice.
2. For EPA employees, accompany shipping containers to carrier and, if required, open outside container(s) for inspection.
3. For overnight package services, determine weight restrictions--at least one service limits weight to 70 pounds per package.

PROCEDURES: SAMPLES CLASSIFIED AS POISON "A"

Packaging

1. Collect samples in a polyethylene or glass container with an outer diameter narrower than the valve hole on a DOT specification No. 3A1800 or No. 3AA1800 metal cylinder. To prevent leakage, fill container no more than 90 percent full (at 130°F).
2. Seal sample container.
3. Complete sample identification tag and attach securely to sample container.
4. Attach string or flexible wire to neck of the sample container; lower it into metal cylinder partially filled with noncombustible, absorbent cushioning material (for example, diatomaceous earth or vermiculite). Place only one container in a metal cylinder. Pack with enough absorbing material between the bottom and sides of the sample container and the metal cylinder to prevent breakage and absorb leakage. After the cushioning material is in place, drop the end of the string or wire into the cylinder valve hole.
5. Replace valve, torque to 250 ft/lb (for 1-inch opening), and replace valve protector on metal cylinder, using Teflon tape.
6. Place one or more cylinders in a sturdy outside container.

Marking/Labeling

1. Use abbreviations only where specified.
2. Place following information, either hand printed or in label form, on the side of the cylinder or on a tag wired to the cylinder valve protector.

- "Poisonous Liquid, n.o.s. NA1955" or "Poisonous Gas, n.o.s. NA1955."
 - Laboratory name and address.
 - DOT label "Poisonous Gas" (even if sample is liquid) on cylinder.
3. Put all information on metal cylinder on outside container.
 4. Print "Laboratory Sample" and "Inside Packages Comply With Prescribed Specifications" on top and/or front of outside container. Mark "This Side Up" on top of container and upward-pointing arrows on all four sides.

Shipping Papers

1. Use abbreviations only as specified.
2. Complete carrier-provided bill of lading and sign certification statement (if carrier does not provide, use standard industry form). Provide following information in order listed. (One form may be used for more than one exterior container.)
 - "Poisonous Liquid, n.o.s. NA1955."
 - Net weight or net volume (weight or volume may be abbreviated), just before or just after "Poisonous Liquid, n.o.s. NA1955."
3. Include a chain-of-custody record, properly executed, in container or with cylinder if legal use of samples is required or anticipated.
4. For EPA employees, accompany shipping container to carrier and, if required, open outside container(s) for inspection.

Transportation

1. Transport unknown hazardous substance samples classified as poison A only by ground transport or Government-owned aircraft. Do not use air cargo, other common carrier aircraft, or rented aircraft.

SAMPLE IDENTIFICATION

The sample tag is the means for identifying and recording the sample and the pertinent information about it. The sample tag should be legibly written and completed with an indelible pencil or waterproof ink. The information should also be recorded in a logbook. The tag should be firmly affixed to the sample container. As a minimum, it should include:

- Exact location of sample.
- Time and date sample was collected.
- Name of sampler and witnesses (if necessary).
- Project codes, sample station, number, and identifying code (if applicable).
- Type of sample (if known).
- Hazardous substance or environmental sample.
- Tag number (if sequential tag system is used).
- Laboratory number (if applicable).
- Any other pertinent information.



U.S. Department
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Research and
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Administration

ATTACHMENT C-1

GUIDE FOR

Appendix C

Revision 0

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HAZARDOUS MATERIALS SHIPPING PAPERS

The following information has been abstracted from the Code of Federal Regulations, Title 49, Parts 100-177

1. DEFINITIONS

- A. SHIPPING PAPER (Sec. 171.8) A shipping paper may be a shipping order, bill of lading, manifest, or other shipping document serving a similar purpose containing the information required by Sec. 172.202, 172.203 and 172.204.
- B. HAZARDOUS WASTE MANIFEST (CFR, Title 40, Sec. 262.20) A hazardous waste manifest is a document (shipping paper) on which all hazardous waste is identified. A copy of the manifest must accompany each shipment of waste from the point of pick-up to the destination. (CFR, Title 49, Sec. 172.205)

2. SHIPPER'S RESPONSIBILITY [Sec. 172.200(a)] The shipper has the responsibility to properly prepare the shipping paper when offering a hazardous material for transport.
NOTE: For shipments of hazardous waste, the hazardous waste manifest is the only authorized documentation. (CFR, Title 40, Sec. 262.23)

3. HAZARDOUS MATERIALS DESCRIPTION (Sec. 172.202) The shipping description of a hazardous material on a shipping paper must include the following information:
 - A. Proper shipping name- Sec. 172.101 or Sec. 172.102 (when authorized);
 - B. The hazard class prescribed for the material in the same section; [See exceptions Sec. 172.202(a)(2)].
 - C. The identification number for the material (preceded by "UN" or "NA" as appropriate); and
 - D. Except for empty packagings, the total quantity (by weight, volume, or as otherwise appropriate) of the hazardous materials covered by the description.
 - E. Except as otherwise provided in the regulations, the basic description in 3A, B and C above must be shown in sequence. For example "Acetone, Flammable Liquid, UN1090."
 - F. The total quantity of the material covered by one description must appear before or after (or both before and after) the basic description as indicated in 3A, B and C above.
 - (1) Abbreviations may be used to specify the type of packaging, weight or volume.
Example: "40 Cyl. Nitrogen Nonflammable gas UN 1066, 800 pounds"; "1 box Cement liquid, n.o.s., Flammable liquid, NAI133, 25 lbs."
 - (2) Type of packaging and destination marks may be entered in any appropriate manner before or after the basic description.
 - G. Technical and chemical group names may be entered in parentheses between the proper shipping name and hazard class. Example: Corrosive liquid, n.o.s. (capryl chloride), corrosive material.

4. GENERAL ENTRIES ON SHIPPING PAPERS (Sec. 172.201)

- A. CONTENTS When describing a hazardous material on the shipping paper(s), that description must conform to the following requirements:
 - (1) When a hazardous material, including materials not subject to the regulations, is described on the same shipping paper, the hazardous material description entries required by Sec. 172.202 and those additional entries that may be required by Sec. 172.203.
 - a. Must be entered first (See Figure 1), or
 - b. Must be entered in a contrasting color, except that a description on a reproduction of a shipping paper may be highlighted, rather than printed, in a contrasting color (these requirements apply only to the basic description required by Sec. 172.202(a)(1), (2) and (3), (See Figure 1); or
 - c. Must be identified by the entry "X" placed before the proper shipping name in a column captioned "HM" [the "X" may be replaced by "RQ" (Reportable Quantity), if appropriate] See Figure 1.
 - (2) The required shipping description on a shipping paper and all copies that are used for transportation purposes must be legible and printed (manually or mechanically) in English.
 - (3) Unless it is specifically authorized or required, the required shipping description may not contain any code or abbreviation.

BASIC DESCRIPTION

Type of Packagings May Be Abbreviated If Used.

Proper Shipping Name

Hazard Class

Total Quantity

HAZARDOUS MATERIALS LISTED FIRST

No. of Packages & Description	HAZARDOUS MATERIALS LISTED FIRST	HAZARDOUS MATERIALS LISTED FIRST	HAZARDOUS MATERIALS LISTED FIRST	HAZARDOUS MATERIALS LISTED FIRST	HAZARDOUS MATERIALS LISTED FIRST
10	Drum, Gasoline, Flammable Liquid, UN1203	4500 lbs.			
40	Cyl. Nitrogen, Non-Flammable Gas, UN1066	800 lbs.			
1	Drum, Flammable Solid, n.s.s., UN1325	452 lbs.			
4	Boxes, Advertising Material, Paper, R01	60 lbs.			
1	Roll, Paper Printing, Newspaper	690 lbs.			
12	Sets, Carbon Paper	22 lbs.			

CERTIFICATION

COO

BASIC DESCRIPTION

Type of Packagings May Be Abbreviated If Used.

Proper Shipping Name

Hazard Class

Total Quantity

HAZARDOUS MATERIALS ENTRIES-CONTRASTING COLOR

No. of Packages & Description	HAZARDOUS MATERIALS ENTRIES-CONTRASTING COLOR	HAZARDOUS MATERIALS ENTRIES-CONTRASTING COLOR	HAZARDOUS MATERIALS ENTRIES-CONTRASTING COLOR	HAZARDOUS MATERIALS ENTRIES-CONTRASTING COLOR	HAZARDOUS MATERIALS ENTRIES-CONTRASTING COLOR
10	Drums, Gasoline, Flammable Liquid, UN1203	4500 lbs.			
4	Boxes, Advertising Material, Paper, R01	60 lbs.			
1	Drum, Flammable Solid, n.s.s., UN1325	452 lbs.			
12	Sets, Carbon Paper	22 lbs.			
40	Cyl. Nitrogen, Non-Flammable Gas, UN1066	800 lbs.			

CERTIFICATION

COO

HAZARDOUS MATERIALS ENTRIES-CONTRASTING COLOR

BASIC DESCRIPTION

Type of Packagings May Be Abbreviated If Used.

Proper Shipping Name

Hazard Class

Total Quantity

HAZARDOUS MATERIALS LISTED ON SHIPPING PAPERS

No. of Packages & Description	HAZARDOUS MATERIALS LISTED ON SHIPPING PAPERS	HAZARDOUS MATERIALS LISTED ON SHIPPING PAPERS	HAZARDOUS MATERIALS LISTED ON SHIPPING PAPERS	HAZARDOUS MATERIALS LISTED ON SHIPPING PAPERS	HAZARDOUS MATERIALS LISTED ON SHIPPING PAPERS
4	Boxes, Advertising Material, Paper, R01	60 lbs.			
10	Drums; Gasoline, Flammable Liquid, UN1203	4500 lbs.			
1	Drum, Flammable Solid, n.s.s., UN1325	452 lbs.			
12	Sets, Carbon paper	22 lbs.			
40	Cyl. Nitrogen, Nonflammable Gas, UN1066	800 lbs.			
1	Roll, Paper, Printing, Newspaper	690 lbs.			

CERTIFICATION

COO

HAZARDOUS MATERIALS LISTED ON SHIPPING PAPERS

- (4) A shipping paper may contain additional information concerning the material provided the information is not inconsistent with the required description. Unless otherwise permitted or required, additional information must be placed after the basic description required by Sec. 172.202(a).
 - a. When appropriate, the entries "IMCO" or "IMCO Class" may be entered immediately before or immediately following the class entry in the basic description.
 - b. If a material meets the definition of more than one hazard class, the additional hazard class or classes may be entered after the hazard class in the basic description.
 - B. NAME OF SHIPPER A shipping paper for a shipment by water must contain the name of the shipper.
- 5. ADDITIONAL DESCRIPTION REQUIREMENTS (Sec. 172.203) (ALL MODES)
 - A. Exemptions - Each shipping paper issued in connection with a shipment made under an exemption must bear the notation "DOT-E" followed by the exemption number assigned (Example: DOT-E 4648) and so located that the exemption number is clearly associated with the description to which the exemption applies.
 - B. Limited Quantities - Descriptions for materials defined as "Limited Quantities" ... must include the words "Limited Quantities" or "Ltd. Qty." following the basic description.
 - C. Hazardous Substances
 - (1) If the proper shipping name for a mixture or solution that is a hazardous substance does not identify the constituents, making it a hazardous substance, the name or names of such constituents shall be entered in association with the basic description.
 - (2) The letters "RQ" (Reportable Quantity) shall be entered on the shipping paper either before or after the basic description required by Sec. 172.202 for each hazardous substance. (See definition Sec. 171.8) Example: RQ, Cresol, Corrosive Material, NA2076; or Adipic Acid, ORM-E, NA9077, RQ.
 - D. Radioactive Materials - For additional description for radioactive materials, refer to Sec. 172.203(d).
 - E. Empty Packagings
 - (1) Except for a tank car, or any packaging that still contains a hazardous substance, the description on the shipping paper for an empty packaging containing the residue of a hazardous material may include the word(s) "EMPTY" or "EMPTY: Last Contained (Name of Substance)" as appropriate in association with the basic description of the hazardous material last contained in the packaging.
 - (2) For empty tank cars, see Sec. 174.25(c).
 - (3) If a packaging, including a tank car, contains a residue that is a hazardous substance the description on the shipping paper shall be prefaced with the phrase "EMPTY: Last Contained (Name of Substance)" and shall have "RQ" entered before or after the basic description.
 - F. Dangerous When Wet - The words "Dangerous When Wet" shall be entered on the shipping paper in association with the basic description when a package covered by the basic description is required to be labeled with a "DANGEROUS WHEN WET" label.
 - G. Poisonous Materials - Notwithstanding the class to which a material is assigned:
 - (1) If the name of the compound or principal constituent that causes the material to meet the definition of a poison is not included in the proper shipping name for the material, the name of that compound or constituent shall be entered on the shipping paper in association with the shipping description for the material.
 - (2) The name of the compound or principal constituent may be either a technical name or any name for the material that is listed in the NIOSH Registry. (Registry of Toxic Effects of Chemical Substances, 1978 Edition) [Sec. 172.203(k)]
NOTE: For additional details, see Sec. 172.203(k)
 - H. Exceptions: OTHER REGULATED MATERIAL (ORM-A, B, C, AND D)
 - (1) Shipping paper requirements do not apply to any material other than a hazardous waste or a hazardous substance that is:
 - a. An ORM-A, B or C unless it is offered or intended for transportation by air or water when it is subject to the regulations pertaining to transportation by air or water as specified in Sec. 172.101 (Hazardous Materials Table); or
 - b. An ORM-D unless it is offered or intended for transportation by air.

MODAL REQUIREMENTS
(ADDITIONAL INFORMATION)

NOTE: In addition to the basic requirements for shipping papers, additional information is listed for each mode.

6. TRANSPORTATION BY RAIL

A. SHIPPING PAPERS (Sec. 176.24)

- (1) Except as provided in paragraph (b) of this section, no person may accept for transportation by rail any hazardous material which is subject to this subchapter unless he has received a shipping paper prepared in a manner specified in Sec. 172.200. In addition, the shipping paper must include a certificate, if required by Sec. 172.204. However, no member of the train crew of a train transporting the hazardous material is required to have a shippers certificate on the shipping paper in his possession if the original shipping paper containing the certificate is in the originating carriers possession.

- (2) This subpart does not apply to materials classed as ORM-A, B, C or D.

B. ADDITIONAL DESCRIPTION FOR SHIPPING PAPERS [Sec. 172.203(g)]

- (1) The shipping paper for a rail car containing a hazardous material must contain the notation "Placarded" followed by the name of the placard required for the rail car.
- (2) The shipping paper for each specification DOT 112A or 114A tank car (without head shields) containing a flammable compressed gas must contain the notation "DOT 112A" or "DOT 114A", as appropriate, and either "Must be handled in accordance with FRA E.O. No. 5" or "Shove to rest per E.O. No. 5."

NOTE: For additional details, refer to Part 174.

7. TRANSPORTATION BY AIR

A. SHIPPING PAPERS ABOARD AIRCRAFT (Sec. 175.35) A copy of the shipping papers required by Sec. 175.30(a)(2) must accompany the shipment it covers during transportation aboard an aircraft.

NOTE: The documents required (shipping papers and notification of pilot-in-command) may be combined into one document if it is given to the pilot-in-command before departure of the aircraft. [Sec. 175.35(b)].

B. NOTIFICATION OF PILOT-IN-COMMAND (Sec. 175.33) The operator of the aircraft shall give the pilot-in-command the following information in writing before takeoff (Sec. 175.35):

- (1) Description of hazardous material on shipping papers (Sec. 172.202 and 172.203);
- (2) Location of the hazardous material in the aircraft; and
- (3) The results of the inspection requirements by Sec. 175.30(b).

NOTE: For additional details, refer to Part 175.

8. TRANSPORTATION BY WATER

A. SHIPPING PAPERS (Sec. 176.24) A carrier may not transport a hazardous material by vessel unless the material is properly described on the shipping paper in the manner prescribed in Part 172.

B. CERTIFICATE (Sec. 176.27)

- (1) A carrier may not transport a hazardous material by vessel unless he has received a certificate prepared in accordance with Sec. 172.204.
- (2) In the case of an import or export shipment of hazardous materials which will not be transported by rail, highway, or air, the shipper may certify on the bill of lading or other shipping paper that the hazardous material is properly classed, described, marked, packaged and labeled according to Part 172 or in accordance with the requirements of the IMCO Code. (See Sec. 171.12)

C. DANGEROUS CARGO MANIFEST (Sec. 176.30) The master of a vessel transporting hazardous materials or his authorized representative shall prepare a dangerous cargo manifest, list, or stowage plan. This document may not include a material which is not subject to the requirements of CFR, Title 49, or the IMCO Code. This document must be kept in a designated holder on or near the vessel's bridge. (See Sec. 176.30 for details)

D. EXEMPTIONS (Sec. 176.31) If a hazardous material is being transported by vessel under the authority of an exemption and a copy of the exemption is required to be on board the vessel, it must be kept with the dangerous cargo manifest.

NOTE: For additional details, refer to Part 176.

E. ADDITIONAL DESCRIPTION FOR SHIPPING PAPERS [Sec. 172.203(i)]

- (1) Each shipment by water must have the following additional shipping paper entries:
 - a. Identification of the type of packages such as barrels, drums, cylinders, and boxes,
 - b. The number of each type of packages including those in freight container or on a pallet, and
 - c. The gross weight of each type of package or the individual gross weight of each package.
- (2) The shipping papers for a hazardous material offered for transportation by water to any country outside the United States must have in parenthesis the technical name of the material following the proper shipping name when the material is described by a "n.o.s." entry in Sec. 172.101 (Hazardous Materials Table). For example: Corrosive liquid, n.o.s. (acrylyl chloride), Corrosive material. However, for a mixture, only the technical name of any hazardous material giving the mixture its hazardous properties must be identified.

9. TRANSPORTATION BY HIGHWAY

A. SHIPPING PAPERS (Sec. 177.817)

- (1) General - A carrier may not transport a hazardous material unless it is accompanied by a shipping paper that is prepared in accordance with Sec. 172.201, 172.202 and 172.203.
- (2) Shipper's certification - An initial carrier may not accept hazardous materials offered for transportation unless the shipping paper describing the material includes a shipper's certification which meets the requirements in Sec. 172.204 of this subchapter. The certification is not required for shipments to be transported entirely by private carriage and for bulk shipments to be transported in a cargo tank supplied by the carrier. [Sec. 177.817(c)]
- (3) Interlining with carriers by rail - A motor carrier shall mark on the shipping paper required by this section, if it offers or delivers a freight container or transport vehicle to a rail carrier for further transportation: [Sec. 177.817(c)]
 - a. A description of the freight container or transport vehicle; and
 - b. The kind of placard affixed to the freight container or transport vehicle.
- (4) This subpart does not apply to materials classed as an ORM-A, B, C or D.
- (5) Accessibility of shipping papers: The driver and each carrier using the vehicle shall ensure that the shipping paper is readily available and recognizable by authorities in the case of an accident or inspection. [See Sec. 177.817(e) for details]

B. ADDITIONAL DESCRIPTION FOR SHIPPING PAPERS [Sec. 172.203(h)] For additional descriptions for Anhydrous ammonia see Sec. 172.203(h)(1); Liquefied petroleum gas see Sec. 172.203(h)(2) and Exemptions see Sec. 172.203(a).

10. SHIPPER'S CERTIFICATION (Sec. 172.204)

A. GENERAL (Except B and D below)

- (1) Except as provided in paragraphs (b) and (c) of Sec. 172.204, each person who offers a hazardous material for transportation shall certify that the material offered for transportation is in accordance with the regulations by printing (manually or mechanically) the following statement on the shipping paper containing the required description:

This is to certify that the above-named materials are properly classified, described, packaged, marked and labeled, and are in proper condition for transportation according to the applicable regulations of the Department of Transportation.*

NOTE: The words "herein-named" may be substituted for the words "above named".

*NOTE: For hazardous waste shipments, the words "and the EPA" must be added to the end of the certification. [See CFR, Title 40, Sec. 262.21(b)]

B. AIR TRANSPORTATION

- (1) General - Certification containing the following language may be used in place of the certification required by paragraph A(1) above:
I hereby certify that the contents of this consignment are fully and accurately described above by proper shipping name and are classified, packed, marked and labeled, and in proper condition for carriage by air according to applicable national governmental regulations.
- (2) Duplicate Certificate - Each person who offers a hazardous material to an aircraft operator for transportation by air shall provide two (2) copies of the certificate. (Sec. 175.30)
- (3) Passenger and Cargo Aircraft - If hazardous materials are offered for transportation by air, add to the certificate the following statement:
This shipment is within the limitations prescribed for passenger/cargo-only aircraft. (delete non-applicable)
- (4) Radioactive Material - Each person who offers any radioactive material for transportation aboard a passenger-carrying aircraft shall sign (mechanically or manually) a printed certificate stating that the shipment contains radioactive material intended for use in, or incident to, research, medical diagnosis or treatment.
NOTE: See Sec. 175.10 for exceptions.

C. SIGNATURE - The certifications required above must be legibly signed (mechanically or manually) by a principal, officer, partner or employee of the shipper or his agent. [Sec. 172.204(d)]

D. EXCEPTIONS - Except for a hazardous waste, no certification is required for hazardous material offered for transportation by motor vehicle and transported:

- (1) In a cargo tank supplied by the carrier, or
- (2) By the shipper as a private carrier except for hazardous material that is to be reshipped or transferred from one carrier to another.
- (3) No certification is required for the return of an empty tank car which previously contained a hazardous material and which has not been cleaned or purged.

HAZARDOUS WASTE MANIFEST INFORMATION

The following information has been abstracted from the Code of Federal Regulations (CFR), Title 49, Parts 100-177 and CFR, Title 40, Part 262.

1. DEFINITIONS

- A. HAZARDOUS WASTE MANIFEST (CFR Title 40, §262.20)
A hazardous waste manifest is a shipping document on which all hazardous wastes are identified.
- B. SHIPPING PAPER - A shipping order, bill of lading, manifest, or other shipping document serving a similar purpose and containing the information required by §172.202, §172.203 and §172.204.

2. DOT HAZARDOUS MATERIALS MANIFEST REQUIREMENTS (§172.205)

- A. No person may offer, transport, transfer or deliver a hazardous waste unless a hazardous waste manifest is prepared, signed, carried and given as required of that person by §172.205.
- B. The shipper (generator) must prepare the manifest in accordance with the EPA Regulations, CFR Title 40, Part 262.
- C. The original copy of the manifest must be dated by, and bear the handwritten signature of the person representing the:
 - (1) Shipper (generator) of waste at the time it is offered for transportation, and
 - (2) Initial carrier accepting the waste for transportation.
- D. A copy of the manifest must be dated by, and bear the handwritten signature of the person representing:
 - (1) Each subsequent carrier accepting the waste for transportation, at the time of acceptance, and
 - (2) The designated facility receiving the waste, upon receipt.

- E. A copy of the manifest bearing all required dates and signatures must be:
 - (1) Given to a person representing each carrier accepting the waste for transportation,
 - (2) Carried during transportation in the same manner as required for shipping papers,
 - (3) Given to a person representing the designated facility receiving the waste,
 - (4) Returned to the shipper (generator) by the carrier that transported the waste from the United States to a foreign destination with a notation of the date of departure from the United States, and
 - (5) Retained by the shipper (generator) and by the initial and each subsequent carrier for three (3) years from the date the waste was accepted by the initial carrier. Each retained copy must bear all required signatures and dates up to and including those entered by the next person who received the waste.
 - F. The requirements of §172.205(d) and (3) do not apply to a rail carrier when waste is delivered to a designated facility by railroad if:
 - (1) All of the information required to be entered on the manifest (except generator and carrier identification numbers and the generator's certification) is entered on the shipping paper carried in accordance with §174.26(c);
 - (2) The delivering rail carrier obtains and retains a receipt for the waste that is dated by and bears the handwritten signature of the person representing the designated facility; and
 - (3) A copy of the shipping paper is retained for three (3) years by each railroad transporting the waste.
 - G. The person delivering a hazardous waste to an initial rail carrier shall send a copy of the manifest, dated and signed by a representative of the rail carrier, to the person representing the designated facility.
 - H. A hazardous waste manifest required by CFR, Title 40, Part 262 containing all the information required by CFR, Title 49, Subpart C, may be used as the shipping paper.
3. THE MANIFEST-GENERAL REQUIREMENTS (§262.20)
- A. A generator (shipper) who transports, or offers for transportation, hazardous waste for off-site treatment, storage, or disposal must prepare a manifest before transporting the waste off-site.
 - B. A generator (shipper) must designate on the manifest one facility which is permitted to handle the waste described on the manifest.
 - C. A generator (shipper) may also designate on the manifest one alternate facility which is permitted to handle his waste in the event an emergency prevents delivery of the waste to the primary designated facility.
 - D. If the transporter (carrier) is unable to deliver the waste to the designated facility, the generator must either designate another facility or instruct the transporter to return the waste.
4. MANIFEST INFORMATION (§262.21)
- A. The manifest must contain:
 - (1) Manifest document number;
 - (2) Generator's (Shipper's) name, mailing address, telephone number, and the EPA identification number;
 - (3) Name and EPA identification number of each transporter (carrier);
 - (4) Name, address and EPA identification number of the designated facility and an alternate facility, if any;
 - (5) Description of the waste(s) (e.g. proper shipping name required by the Department of Transportation Hazardous Materials Regulations CFR, Title 49, §172.101, §172.202 and §172.203); and
 - (6) Total quantity of each hazardous waste by units of weight or volume, and the type and size of containers loaded into or onto the transport vehicle.
 - B. Certification [§262.21(b)] The following certification must appear on the manifest:
"This is to certify that the above named materials are properly classified, described, packaged, marked, labeled and are in proper condition for transportation according to the applicable regulations of the Department of Transportation and the EPA"

5. COPIES OF MANIFEST REQUIRED (§262.22)

The manifest must consist of at least the number of copies to provide the generator, each transporter, and the owner or operator of the designated facility with one copy each for their records and another copy to be returned to the generator.

6. USE OF THE MANIFEST (§262.23)

A. The generator must:

- (1) Sign the manifest certification by hand;
- (2) obtain the handwritten signature of the initial transporter and date of acceptance of manifest; and
- (3) Retain one copy in accordance with §262.40(a).

B. The generator must give the transporter the remaining copies of the manifest.

C. Shipment of hazardous waste within the United States solely by railroad or water (bulk shipments only); the generator must send three (3) copies of the manifest dated and signed in accordance with §262.20 to the owner or operator of the designated facility.

NOTE: Copies of the manifest are not required for each transporter. For special provisions for rail or water (bulk shipment) transporters see §263.20(e).

7. PREPARATION OF HAZARDOUS WASTE FOR SHIPMENT (§262.30)

A. Packaging Hazardous Waste - The generator (shipper) has the responsibility for the classification and packaging of hazardous waste prior to offering for transportation. The requirements for packaging will be found in the Department of Transportation Regulations CFR, Title 49, Parts 172, 173, 178 and 179.

B. Labeling Requirements (§262.31) - Prior to offering a hazardous waste for transportation off-site, the generator (shipper) must label each package in accordance with CFR Title 49, Part 172, Subpart E.

C. Marking Requirements (§262.32) - Prior to offering hazardous waste for transportation off-site, the generator must:

- (1) Mark each package of the hazardous waste; and
- (2) Mark each container 110 gallons or less offered for transportation with the following words and information displayed in accordance with the requirements of CFR, Title 49, Sec. 172.304.

"HAZARDOUS WASTE-Federal Law Prohibits Improper Disposal. If found, contact the nearest police or public safety authority or the United States Environmental Protection Agency"

Generator's Name and Address _____

Manifest Document Number _____

D. Placarding Requirements (§262.33) - Prior to offering a hazardous waste for transportation off-site, the generator must:

- (1) Placard the shipment; or
- (2) Offer the initial transporter (carrier) the appropriate placards. (CFR Title 49, Part 172, Subpart F)

NOTE: This handout is designed as a training aid only. It does not relieve persons from complying with the Department of Transportation's Hazardous Materials Regulations. Final authority for use of shipping papers is found in the Code of Federal Regulation, Title 49, Part 100-177.

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DEPARTMENT OF TRANSPORTATION
RESEARCH AND SPECIAL PROGRAMS ADMINISTRATION
MATERIALS TRANSPORTATION BUREAU
OFFICE OF OPERATIONS AND ENFORCEMENT
INFORMATION SERVICES DIVISION, DMT-11
WASHINGTON, D.C. 20590

REVISED MAY 1981



U.S. Department
of Transportation
**Research and
Special Programs
Administration**

ATTACHMENT C-2

Appendix C
Revision 0
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GUIDE FOR MARKINGS

The following information has been abstracted from the Code of Federal Regulations (CFR), Title 49 Transportation, Parts 100-199. Refer to the appropriate Sections for details.

NOTE: Rulemaking proposals are outstanding or are contemplated concerning the regulations. Keep up to date with the changes.

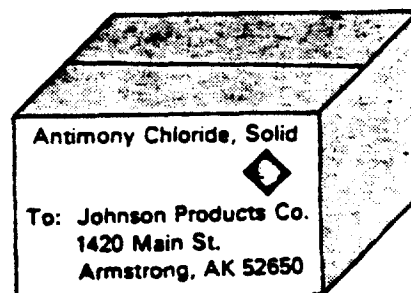
MARKING - means the application of the descriptive name, proper shipping name, hazard class, identification number (when authorized), instructions, cautions, weight or a combination thereof on the outside shipping container. Marking also includes the specification mark for both the inside and outside shipping containers required by the Hazardous Materials Regulation.

DESCRIPTIVE INFORMATION

GENERAL REQUIREMENTS (§172.300-172.304)

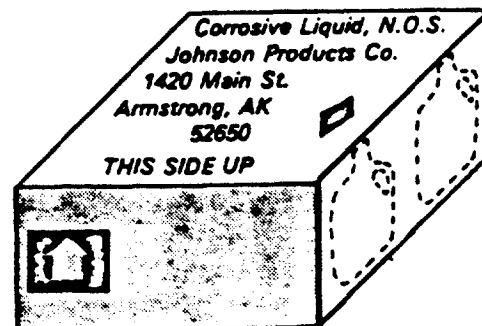
All containers of hazardous materials, i.e. packages, freight containers, or transport vehicles, must, unless specifically exempted, be marked with the proper shipping name(s) of the contents and the name and address or either the consignee or consignor. All markings must be:

1. Durable, in English, and printed on or affixed to the surface of the package or on a label, tag or sign.
2. On a background of a sharply contrasting color and unobscured by labels or attachments.
3. Away from other markings that could reduce its effectiveness.



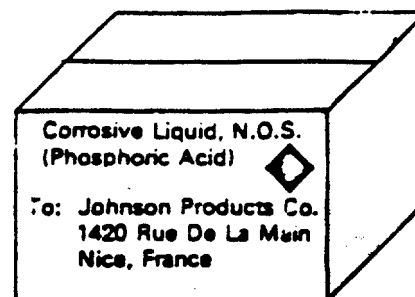
LIQUIDS - INSIDE CONTAINERS (§172.312)

1. Inside containers must be packed with closures in the upright position.
2. Must be marked on the outside with "THIS END UP" or "THIS SIDE UP"
3. Arrows must be used only to show orientation of package. An arrow symbol indicated by ANSI Standard MH6.11968 "THIS WAY UP". Pictorial (arrows) of goods is recommended.



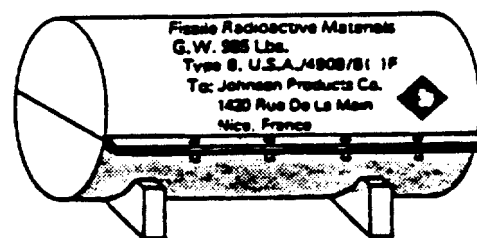
EXPORT BY WATER (§172.302)

All n.o.s. entries, when authorized in §172.101 or §172.102, must have the technical name(s) of the material immediately following the proper shipping name for export by water. For mixtures (two or more) hazardous materials, the technical name of at least two components must be identified.



RADIOACTIVE MATERIALS (§172.310)

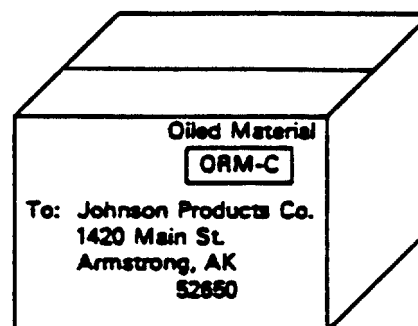
1. Containers weighing over 110 pounds (gross weight) must be marked on the container.
2. Must be marked "TYPE A" or "TYPE B" as required in letters at least 1/2" high.
3. For export, the letters "USA" must follow the specification markings or package certification.



OTHER REGULATED MATERIALS (ORM'S) (§172.316)

ORM materials must be designated immediately following or below the proper shipping name marking within a rectangular border approximately 1/4 inch larger on each side of the designation. The appropriate designation must be one of the following:

- | | |
|--------------------|--------------|
| 1. ORM-A | 5. ORM-D |
| 2. ORM-B -KEEP DRY | 6. ORM-D-AIR |
| 3. ORM-B | 7. ORM-E |
| 4. ORM-C | |



NOTE: These markings serve as the certification by the shipper that the material is properly described, classed, packaged, marked and labeled (when appropriate) and in proper condition for transportation. Use of this type of certification does not preclude the requirement for a certificate on the shipping paper [§172.316(c)].



EXAMPLE

AUTHORIZED CONTAINERS IN OUTSIDE CONTAINERS

When a DOT specification container is required for a hazardous material and that container is overpacked in another container meeting the requirements of §173.21 and §173.24, the outside container must be marked in accordance with §173.25.

EXAMPLES: "THIS SIDE UP" or "THIS END UP" or "INSIDE PACKAGES COMPLY WITH PRESCRIBED SPECIFICATIONS"

CYLINDERS - All cylinders must be marked in accordance with §173.34 and §§173.301 through 173.306. Cylinders passing reinspection and retesting must be marked in accordance with §173.34(e)(6).

PORTABLE TANKS (§172.326 and §172.332) - Portable tanks must display the proper shipping name in letters at least 2 inches high and placed on two opposite sides. Identification numbers [§§171.101 and 171.102 (when authorized)] are required on each side and each end for capacities of 1,000 gallons or more and on two opposing sides in association with the proper shipping name for capacities of less than 1,000 gallons. The name of the owner or lessee must be displayed. Tanks carrying compressed gases (DOT-51) must have all inlets and outlets, except safety relief valves, marked to designate whether or not they communicate with vapor or liquid. [§178.245-6(b)].

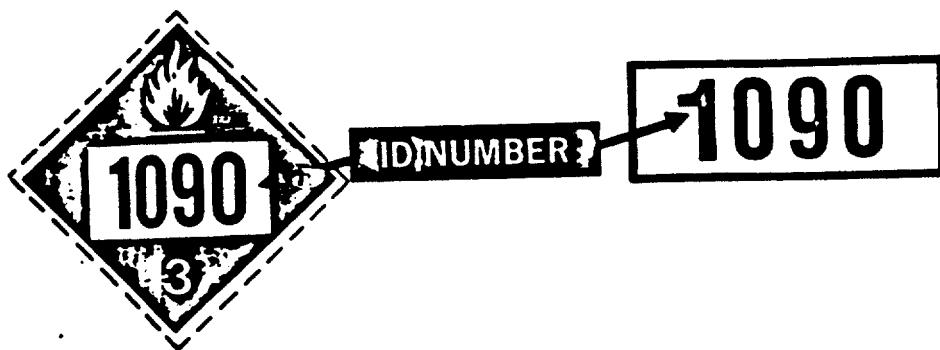
NOTE: When different hazardous materials are transported in marked portable tanks, the shipping name and the identification number displayed must identify the material.

CARGO TANKS - HIGHWAY (COMPRESSED GASES) (§172.328) - Cargo tanks must be marked, in letters no less than 2 inches high, with either the proper shipping name of the gas or an appropriate common name, such as "Refrigerant Gas". Cargo tanks must only be marked, i.e. proper shipping name and identification number [when authorized (§§171.101 and 171.102)] for the material contained therein. DOT MC 331 tanks must have inlets and outlets, except safety relief valves, marked to designate whether they communicate with liquid or vapor when the tank is filled to its maximum permitted silling density. [§178.337-9(c)].

TANK CARS - RAIL (§172.330) - Tank cars, when required to be marked with the proper shipping name by Parts 173 and 179, must be marked in letters at least 4 inches high with at least 5/8 inch stroke with the proper shipping name or the appropriate common name. Identification number markings (when authorized) must be displayed on each side and each end [§§171.101 and 171.102 (when authorized)]. Tank cars must only be marked for the material contained therein.

NOTE: See referenced Sections for requirements for DOT-106 and DOT 110 tank car tanks.

EXAMPLE OF PLACARD AND PANEL WITH IDENTIFICATION NUMBER



NOTE: The Identification Number (ID No.) may be displayed on placards or on orange panels on tanks. Check the sides of the transport vehicle if the ID number is not displayed on the ends of the vehicle.

OTHER MARKING REQUIREMENTS

REQUALIFIED CONTAINERS - Reusable cylinders, portable tanks, cargo tanks and tank cars are required to be either visually inspected or retested at periodic intervals. When this is accomplished, the date of the requalification must be shown on the container as required in §§173.24, 173.31, 173.32, 173.33 and 173.34.

REUSE OF CONTAINERS - Some steel containers in the DOT Series (DOT-17C, 17E and 17H) may be qualified for reuse by a reconditioner of drums who is registered with the Department of Transportation. These drums must meet the requirements of §173.28(m) i.e. old labels removed, exemption number (if any) and descriptive markings removed and the drum reconditioned. Other containers may be reused under varying conditions. See §173.28 for details.

CARGO HEATERS - Cargo heaters authorized for use with flammable liquid or gas must be marked in accordance with §177.834(l)(2)(e) and (f).

MOTOR VEHICLES - Marking of motor vehicles and special requirements are found in §§177.823 and 177.824.

SPECIFICATION CONTAINERS

Markings on specification containers must generally identify: (1) the DOT specification number to which the container is made (Parts 178 and 179); (2) the manufacturers name and address or symbol (registered with the Associate Director for the Office of Hazardous Material Regulation). Duplicate symbols are not authorized. All containers must comply with the marking requirements of §173.24 and the appropriate Section(s) of Parts 178 and 179. Exceptions for Canadian and other import/export situations may be found in §§171.12 and 173.8.

NOTE: For certain containers, specific detailed information such as original test date information and type of material which may be required can be found in Parts 178 and 179.

This publication does not contain all the marking requirements. It is designed as a guide only. For details for all markings, consult Code of Federal Regulations, Title 49, Parts 100-199.

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Department of Transportation
Research and Special Programs Administration
Materials Transportation Bureau
Office of Operation and Enforcement
Information Services Division, DMT-11
Washington, D.C. 20590

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APPENDIX D

DOCUMENT CONTROL/CHAIN-OF-CUSTODY PROCEDURES

GENERAL

Adherence to strict document control and chain-of-custody procedures is extremely important especially in relation to surveys at hazardous waste sites. The legal implications alone demand that accountability be given an utmost priority. The basic aspects of document control and chain-of-custody have therefore been included in this section. For additional information, the following publication, from which this section was developed, should be consulted.

- NEIC Policies and Procedures Manual, EPA-330-78-001R, May 1978 (revised December 1981), Section II

DOCUMENT CONTROL

The purpose of document control is to assure all project documents will be accounted for when the project is complete. Document control should include the use of serialized documents, a document inventory procedure and an adequate document filing system, all issued by, under the control of, and maintained by an appointed Document Control Officer (DCO). Table D-1 lists the principal items subject to document control during a specific project.

Serialized Documents

Sample collection and analytical tags, and chain-of-custody records should have preprinted serial numbers. It is not necessary that a sample tag number match a custody record number, however, it is necessary that all issued numbers be appropriately accounted for by the DCO. It is also necessary that in the event a tag or custody record is damaged, lost or destroyed prior to its use, its serial number and disposition are recorded.

Other Documents

Other documents used during the conduct of a project (e.g., field logbooks, laboratory notebooks, data sheets, etc.) should be appropriately coded with a unique identifier to ensure accountability. The project DCO will be responsible for development of the document identification system, paying particular attention to its utility and consistency for the specified program. An example of a document identification system is as follows:

<u>Project Code</u>	<u>Subcontractor Code (if necessary)</u>	<u>Document Code</u>	<u>Serial Number</u>
00-000-000-	-00-	-A-	-00001

In addition, a listing of suggested codes is shown in Table D-2.

TABLE D-1. DOCUMENTS SUBJECT TO CONTROL

Project Work Plan
Project Logbooks
Field Logbooks
Sample Data Sheets
Sample Tags
Chain-of-Custody Records
Laboratory Logbooks
Laboratory Data, Calculations, Graphs, etc.
Sample Checkout
Sample Inventory
Internal Memos
External Written Communication
Confidential Information
Photographs, Drawings, Maps
Quality Assurance Plan
Litigation Document
Final Report

TABLE D-2. SUGGESTED DOCUMENT CODES

Document	Code letter
Project Work Plans	A
Project Logbooks	B
Sampling Logbooks	C
Sampling Data Sheets	D1, D2 etc.
Sampling Coding Form	E
Laboratory Notebooks	G
Laboratory Data Sheets	H1, H2 etc.
Sample Logs	L1, L2 etc.
Internal Memos	M
External Written Communication	N
Confidential Information	O
Photos, Maps, Drawings	P
QA Plan	Q
Reports	R
Final Report	FR
Miscellaneous	X

CHAIN-OF-CUSTODY

The primary need for the implementation of chain-of-custody procedures stems from the possibility that a sample or a piece of data derived from the collection of a sample will be used as physical evidence in an enforcement action. The purpose of chain-of-custody in these instances is to trace the possession of a sample from the time of collection, until it or the derived data is introduced as evidence in legal proceedings. Custody records should, therefore, trace a sample from its collection, through all transfers of custody, until it is delivered to the analytical laboratory. At this point, internal laboratory records should document sample custody until its final disposition.

In order to establish that a sample is valid, it is also necessary to document the measures taken to prevent and/or detect tampering--either to the sample itself, the sampling equipment used or the environment sampled. This is done by the use of evidence tape, locks and custody seals, and documented entries noting their condition in field and laboratory log books. The custody record must document any tampering that may have occurred; the absence of any such comments indicates no tampering observed or noticed during the period of custody.

Since it may not always be possible to know ahead of time if a sample will be used as evidence in future legal actions, it is a good common sense practice to institute a proper chain-of-custody in all instances. Use of such practices as standard operating procedures on a project to project basis will contribute to the consistency and quality of the generated data.

Sample Identification

Preprinted, preserialized sample collection tags are recommended to identify samples collected for shipment to the analytical laboratory. Specific analysis tags may also be issued by the analytical laboratory after the sample has arrived. All collected samples, including duplicates and field blanks should be completely filled in with a minimum of the following information:

- Project Code
- Location Number Assigned by the
Document Control Officer (DCO)
- Date of Collection
- Time of Collection
- Location Description
- Signature of Sampler

- Lab Sample Number--Assigned by the Analytical Laboratory
- Remarks Section

An example of an appropriate sample collection tag and analysis tag is shown in Figure D-1, respectively.

After sample analysis and appropriate quality assurance checks have been made, original sample collection tags are to be stored in a document file maintained by the DCO and the tag serial number is recorded in a master log for future reference. Maintaining such files and records is an important aspect of sample traceability and provides a needed cross referencing tool that can be used to correlate any one of the identifying numbers and sources (e. g., collection tag, laboratory number, master log, etc.) with a specific sample.

Chain-of-Custody Forms

There are many transfers of custody during the course of a sampling program, from time of collection through final sample disposition, and all samples should be accompanied by a Chain-of-Custody Record to document these transfers. In some instances, such as in the collection of air samples on solid sorbents, it becomes necessary to initiate custody procedures from collection media preparation on as the sorbent itself becomes part of the sample after collection is complete. Laboratories providing QC samples must also initiate a custody record. The use of a customized record sheet, such as the one shown in Figure D-2 fulfills these requirements by providing a convenient format for recording pertinent information.

The custody records are used for a packaged lot of samples; more than one sample will usually be recorded on one form. More than one custody record sheet may be used for one package, if necessary. Their purpose is to document the transfer of a group of samples traveling together; when the group of samples changes, a new custody record is initiated. The original of the custody record always travels with the samples; the initiator of the record keeps the copy. When custody of the same group of samples changes hands several times, some people will not have a copy of the custody record. This is acceptable as long as the original custody record shows that each person who had received custody has properly relinquished it.

In general, the following procedures should be followed when using the custody record sheets.

- The originator fills in all requested information from the sample tags (except in the case of air collection media and external QC samples which will be accompanied by custody forms from the originating facility).
- The person receiving custody checks the sample tag information against the custody record. He also checks sample condition and notes anything unusual under "Remarks" on the custody form.

Figure D-1. Sample Collection Tag.

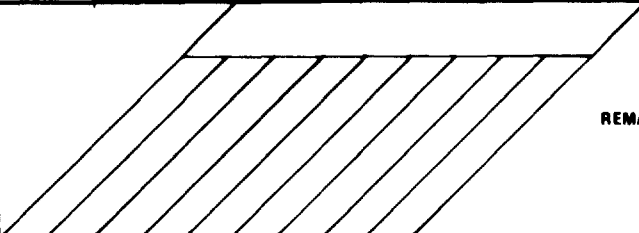
Project Code	Project Name						 REMARKS									
SAMPLERS (Signature)																
Relinquished by: (Signature)		Date/Time		Received by (Signature)		Relinquished by (Signature)		Date/Time		Received by (Signature)						
Relinquished by: (Signature)		Date/Time		Received by (Signature)		Relinquished by (Signature)		Date/Time		Received by (Signature)						
Relinquished by: (Signature)		Date/Time		Received for Laboratory by: (Signature)		Date/Time		Remarks								

Figure D-2. Chain-of-Custody Form.

N 1000

- The originator signs in the top left "Relinquished by" box and keeps the copy.
- The person receiving custody signs in the adjacent "Received by" box and keeps the original.
- The Date/Time will be the same for both signatures since custody must be transferred to another person.
- When custody is transferred to the Sample Bank or an analytical laboratory, blank signature spaces may be left and the last "Received by" signature box used. Another approach is to run a line through the unused signature boxes.
- In all cases, it must be readily seen that the same person receiving custody has relinquished it to the next custodian.
- If samples are left unattended or a person refuses to sign, this must be documented and explained on the custody record.

Receipt for Samples Form

When it becomes necessary to split samples with another source, a separate receipt for samples from (Figure D-3) is prepared and marked to indicate with whom the samples have been split. The signature of the person receiving the samples is required and if this person refuses to sign, it should be noted in the "Received by" space.

This form also complies with requirements of both Section 3007(a)(2) of RCRA and Section 104 of the Comprehensive Environmental Response Compensation and Liability Act. These sections both state that "...If the officer, employee or representative obtains any samples prior to leaving the premises, he shall give to the owner, operator, or agent-in-charge a receipt describing the samples obtained and, if requested, a portion of such sample equal in volume or weight to the portion retained." A copy of the completed form must be given to one of the above described individuals, even if the offer for split samples is declined.

Custody Seals

Custody seals are narrow strips of adhesive paper used to demonstrate that no tampering has occurred. They may be used on sampling equipment or a house door, but they are intended for use on a sample transport container which is not secured by a padlock. They are not intended for use on individual sample containers.

[illegible]

Distribution Original to Coordinator Field Files. Copy to Facility

Figure D-3. Receipt for Sample form.

Laboratory Custody Procedures

An onsite sample bank, the sampling laboratory area and any analytical laboratory used for analyses are considered to be working "laboratories" subject to laboratory custody procedures. Each laboratory should have a designated sample custodian who implements a system to maintain control of the samples.

This includes accepting custody of arriving samples, verifying that information on the sample tags match the Chain-of-Custody Record, assigning unique laboratory numbers and laboratory sample tags and distributing the samples to the analyst.

The designated custodian is also responsible for retaining all original identifying tags, data sheets and laboratory records as part of the permanent project file.

Questions/Problems Concerning Custody Records

If a discrepancy between sample tag numbers and custody record listings is found, the person receiving custody should document this and properly store the samples. The samples should not be analyzed until the problem is resolved.

The responsible person receiving custody should attempt to resolve the problem by checking all available information (other markings on sample container, type of sample, etc.). He should then document the situation on the custody record and in his project logbook and notify the project QA Manager by the fastest available means, followed by written notification.

Changes may be written in the "Remarks" section of the Custody record and should be initialed and dated. A copy of this record should accompany the written notification to the QA Manager.

APPENDIX E
DECONTAMINATION PROCEDURES

Source: Interim Standard Operating Safety Guides
Revised September, 1982
Office of Emergency and Remedial Response

INTRODUCTION

Personnel responding to hazardous substance incidents may become contaminated in a number of ways, including:

- Contacting vapors, gases, mists, or particulate in the air.
- Being splashed by materials while sampling or opening containers.
- Walking through puddles of liquids or on contaminated soil.
- Using contaminated instruments or equipment.

Protective clothing and respirators help prevent the wearer from becoming contaminated or inhaling contaminants, while good work practices help reduce contamination on protective clothing, instruments, and equipment.

Even with these safeguards, contamination may occur. Harmful materials can be transferred into clean areas, exposing unprotected personnel. Or in removing contaminated clothing, personnel may contact contaminants on the clothing and/or inhale them. To prevent such occurrences, methods to reduce contamination and decontamination procedures must be developed and implemented before anyone enters a site and must continue (modified when necessary) throughout site operations.

Decontamination consists of physically removing contaminants and/or changing their chemical nature to innocuous substances. How extensive decontamination must be depends on a number of factors, the most important being the type of contaminants involved. The more harmful the contaminant, the more extensive and thorough decontamination must be. Less harmful contaminants may require less decontamination. Combining decontamination, the correct method of doffing personnel protective equipment, and the use of site work zones minimizes cross-contamination from protective clothing to wearer, equipment to personnel, and one area to another. Only general guidance can be given on methods and techniques for decontamination. The exact procedure to use must be determined after evaluating a number of factors specific to the incident. In addition, the decontamination procedures for sample equipment should be developed in conjunction with the analytical lab(s).

PRELIMINARY CONCERNS

Initial Planning

The initial decontamination plan assumes all personnel and equipment leaving the Exclusion Zone (area of potential contamination) are grossly contaminated. A system is then set up to wash and rinse, at least once, all the personnel protective equipment worn. This is done in combination with a sequential doffing of equipment, starting at the first station with the most heavily contaminated item and progressing to the last station with the least contaminated article. Each piece of clothing or operation requires a separate station. Figure E-1 diagrams a contamination control program showing the layout of the contamination reduction zone.

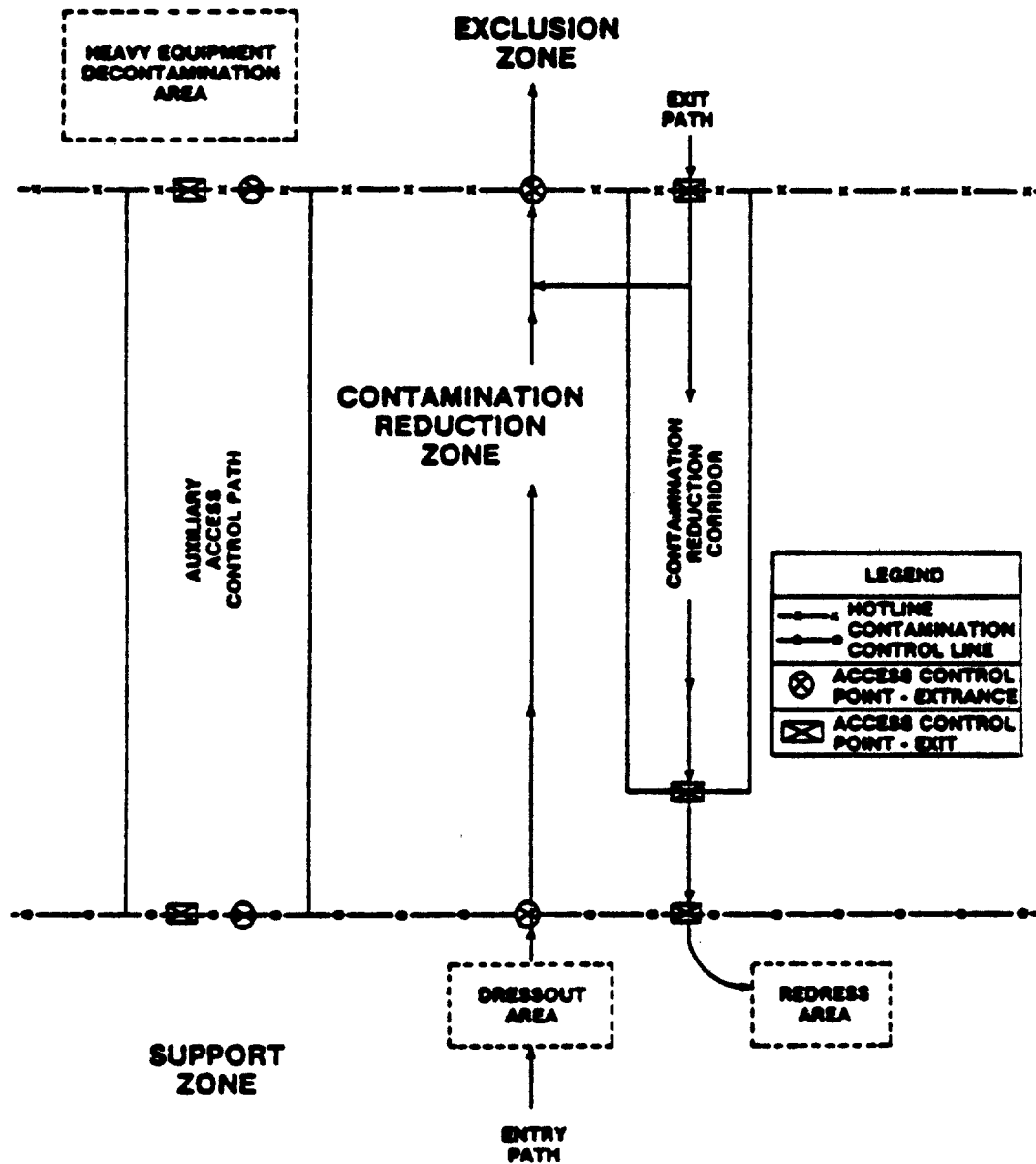


Figure E-1. Contamination reduction zone layout.

The spread of contaminants during the washing/doffing process is further reduced by separating each decontamination station by a minimum of 3 feet. Ideally, contamination should decrease as a person moves from one station to another farther along the line.

While planning site operations, methods should be developed to prevent the contamination of people and equipment. For example, using remote sampling techniques, not opening containers by hand, bagging monitoring instruments, using drum grapplers, watering down dusty areas, and not walking through areas of obvious contamination will reduce the probability of becoming contaminated which would require a less elaborate decontamination procedure.

The initial decontamination plan is based on a worst-case situation (if no information is available about the incident). Specific conditions at the site are then evaluated, including:

- Type of contaminant.
- The amount of contamination.
- Levels of protection required.
- Type of protective clothing worn.

The initial system is modified, eliminating unnecessary stations or otherwise adapting it to site conditions. For instance, the initial plan might require a complete wash and rinse of chemical protective garments. If disposable garments are worn, the wash/rinse step could be omitted. Wearing disposable boot covers and gloves could eliminate washing and rinsing both gloves and disposable boots and reduce the number of stations needed.

Contamination Reduction Corridor

An area within the Contamination Reduction Zone is designated the Contamination Reduction Corridor (CRC). The CRC controls access into and out of the Exclusion Zone and confines personnel decontamination activities to a limited area. The size of the corridor depends on the number of stations in the decontamination procedure, overall dimensions of work control zones, and amount of space available at the site. A corridor of 75 feet by 15 feet should be adequate for full decontamination. Whenever possible, it should be a straight path.

The CRC boundaries should be conspicuously marked, with entry and exit restricted. The far end is the hotline--the boundary between the Exclusion Zone and the Contamination Reduction Zone. Personnel exiting the Exclusion Zone must go through the CRC. Anyone in the CRC should be wearing the Level of Protection designated for the decontamination crew. Another corridor may be required for the entrance and exit of heavy equipment needing decontamination. Within the CRC, distinct areas are set aside for decontamination of personnel, portable field equipment, removed clothing, etc. All activities within the corridor are confined to decontamination.

Personnel protective clothing, respirators, monitoring equipment, sampling supplies, etc. are all maintained outside of the CRC. Personnel don their protective equipment away from the CRC and enter the Exclusion Zone through a separate access control point at the hotline.

EXTENT OF DECONTAMINATION REQUIRED

Modifications of Initial Plan

The original decontamination plan must be adapted to specific conditions found at incidents. These conditions may require more or less personnel decontamination than planned, depending on a number of factors.

Type of Contaminant--

The extent of personnel decontamination depends on the effects the contaminants have on the body. Contaminants do not all exhibit the same degree of toxicity (or other hazard). The more toxic a substance is the more extensive or thorough decontamination must be. Whenever it is known or suspected that personnel can become contaminated with highly toxic or skin-destructive substances, a full decontamination procedure should be followed. If less hazardous materials are involved, the procedure can be downgraded.

Amount of Contamination--

The amount of contamination on protective clothing is usually determined visually. If it is badly contaminated, a thorough decontamination is generally required. Gross material remaining on the protective clothing for any extended period of time may degrade or permeate it. This likelihood increases with higher air concentrations and greater amounts of liquid contamination. Gross contamination also increases the probability of personnel contact. Swipe tests may help determine the type and quantity of surface contaminants.

Level of Protection--

The Level of Protection and specific pieces of clothing worn determine on a preliminary basis the layout of the decontamination line. Each Level of Protection incorporates different problems in decontamination and doffing of the equipment. For example, decontamination of the harness straps and backpack assembly of the self-contained breathing apparatus is difficult. A butyl rubber apron worn over the harness makes decontamination easier. Clothing variations and different Levels of Protection may require adding or deleting stations in the original decontamination procedure.

Work Function--

The work each person does determines the potential for contact with hazardous materials. In turn, this dictates the layout of the decontamination line. Observers, photographers, operators of air samplers, or others in the Exclusion Zone performing tasks that will not bring them in contact with contaminants may not need, for example, to have their garments washed or rinsed. Others in the Exclusion Zone with a potential for direct contact with the hazardous material will require more thorough decontamination. Different decontamination lines could be set up for different job functions, or certain stations in a line could be omitted for personnel performing certain tasks.

Location of Contamination--

Contamination on the upper areas of protective clothing poses a greater risk to the worker because volatile compounds may generate a hazardous breathing concentration both for the worker and for the decontamination personnel. There is also an increased probability of contact with skin when doffing the upper part of clothing.

Reason for Leaving Site--

The reason for leaving the Exclusion Zone also determines the need and extent of decontamination. A worker leaving the Exclusion Zone to pick up or drop off tools or instruments and immediately return may not require decontamination. However, a worker leaving to get a new air cylinder or to change a respirator or canister may require some degree of decontamination. Individuals departing the CRC for a break, lunch, end of day, etc., must be thoroughly decontaminated.

Effectiveness of Decontamination

There is no method to immediately determine how effective decontamination is in removing contaminants. Discolorations, stains, corrosive effects, and substances adhering to objects may indicate contaminants have not been removed. However, observable effects only indicate surface contamination and not permeation (absorption) into clothing. Also many contaminants are not easily observed.

A method for determining effectiveness of surface decontamination is swipe testing. Cloth or paper patches--swipes--are wiped over predetermined surfaces of the suspect object and analyzed in a laboratory. Both the inner and outer surfaces of protective clothing should be swipe tested. Positive indications of both sets of swipes would indicate surface contamination has not been removed and substances have penetrated or permeated through the garment. Swipe tests can also be done on skin or inside clothing. Permeation of protective garments requires laboratory analysis of a piece of the material. Both swipe and permeation testing provide after-the-fact information. Along with visual observations, results of these tests can help evaluate the effectiveness of decontamination.

Equipment

Decontamination equipment, materials, and supplies are generally selected based on availability. Other considerations are ease of equipment decontamination or disposability. Most equipment and supplies can be easily procured. For example, soft-bristle scrub brushes or long-handle brushes are used to remove contaminants. Water in buckets or garden sprayers is used for rinsing. Large galvanized wash tubs or stock tanks can hold wash and rinse solutions. Children's wading pools can also be used. Large plastic garbage cans or other similar containers lined with plastic bags store contaminated clothing and equipment. Contaminated liquids can be stored temporarily in metal or plastic cans or drums. Other gear includes paper or cloth towels for drying protective clothing and equipment.

Decontamination Solution

Personnel protective equipment, sampling tools, and other equipment are usually decontaminated by scrubbing with detergent-water using a soft-bristle brush followed by rinsing with copious amounts of water. While this process may not be fully effective in removing some contaminants (or in a few cases, contaminants may react with water), it is a relatively safe option compared with using a chemical decontaminating solution. Using chemicals requires that the contaminant be identified. A decon chemical is then needed that will change the contaminant into a less harmful substance. Especially troublesome are unknown substances or mixtures from a variety of known or unknown substances. The appropriate decontamination solution must be selected in consultation with an experienced chemist.

Establishment of Procedures

Once decontamination procedures have been established, all personnel requiring decontamination must be given precise instructions (and practice, if necessary). Compliance must be frequently checked. The time it takes for decontamination must be ascertained. Personnel wearing SCBAs must leave their work area with sufficient air to walk to CRC and go through decontamination.

CONTAMINATION DURING MEDICAL EMERGENCIES

Basic Considerations

Part of overall planning for incident response is managing medical emergencies. The plan should provide for:

- Some response team members fully trained in first aid and CPR.
- Arrangements with the nearest medical facility for transportation and treatment of injured, and for treatment of personnel suffering from exposure to chemicals.
- Consultation services with a toxicologist.
- Emergency eye washes, showers, and/or wash stations.
- First aid kits, blankets, stretcher, and resuscitator.

In addition, the plan should have established methods for decontaminating personnel with medical problems and injuries. There is the possibility that the decontamination may aggravate or cause more serious health effects. If prompt life-saving first aid and/or medical treatment is required, decontamination procedures should be omitted. Whenever possible, response personnel should accompany contaminated victims to the medical facility to advise on matters involving decontamination.

Physical Injury

Physical injuries can range from a sprained ankle to a compound fracture, from a minor cut to massive bleeding. Depending on the seriousness of the injury, treatment may be given at the site by trained response personnel. For more serious injuries, additional assistance may be required at the site or the victim may have to be treated at a medical facility.

Life-saving care should be instituted immediately without considering decontamination. The outside garments can be removed (depending on the weather) if they do not cause delays, interfere with treatment, or aggravate the problem. Respiratory masks and backpack assemblies must always be removed. Fully encapsulating suits or chemical-resistant clothing can be cut away. If the outer contaminated garments cannot be safely removed, the individual should be wrapped in plastic, rubber, or blankets to help prevent contaminating the inside of ambulances and/or medical personnel. Outside garments are then removed at the medical facility. No attempt should be made to wash or rinse the victim. One exception would be if it is known that the individual has been contaminated with an extremely toxic or corrosive material which could also cause severe injury or loss of life. For minor medical problems or injuries, the normal decontamination procedure should be followed.

Heat Stress

Heat-related illnesses range from heat fatigue to heat stroke, the most serious. Heat stroke requires prompt treatment to prevent irreversible damage or death. Protective clothing may have to be cut off. Less serious forms of heat stress require prompt attention or they may lead to a heat stroke. Unless the victim is obviously contaminated, decontamination should be omitted or minimized and treatment begun immediately.

Chemical Exposure

Exposure to chemicals can be divided into two categories:

- Injuries from direct contact, such as acid burns or inhalation of toxic chemicals.
- Potential injury due to gross contamination on clothing or equipment

For the contaminant inhaled treatment can only be by qualified physicians. If the contaminant is on the skin or in the eyes, immediate measures must be taken to counteract the substance's effect. First aid treatment usually is flooding the affected area with water; however, for a few chemicals, water may cause more severe problems.

When protective clothing is grossly contaminated, contaminants may be transferred to treatment personnel or the wearer and cause injuries. Unless severe medical problems have occurred simultaneously with splashes, the protective clothing should be washed off as rapidly as possible and carefully removed.

PROTECTION FOR DECONTAMINATION WORKERS

The Level of Protection worn by decontamination workers is determined by:

- Expected or visible contamination on workers.
- Type of contaminant and associated respiratory and skin hazards.
- Total vapor/gas concentrations in the CRC.
- Particulate and specific inorganic or organic vapors in the CRC.
- Results of swipe tests.
- The presence (or suspected presence) of highly toxic or skin-destructive materials.

Level C Use

Level C includes a full-face, canister-type air-purifying respirator, hard hat with face shield (if splash is a problem), chemical-resistant boots and gloves, and protective clothing. The body covering recommended is chemical-resistant overalls with an apron, or chemical-resistant overalls and jacket.

A face shield is recommended to protect against splashes because respirators alone may not provide this protection. The respirator should have a canister approved for filtering any specific known contaminants such as ammonia, organic vapors, acid gases, and particulate.

Level B Use

In situations where site workers may be contaminated with unknowns, highly volatile liquids, or highly toxic materials, decontamination workers should wear Level B protection.

Level B protection includes SCBA, hard hat with face shield, chemical-resistant gloves, and protective covering. The clothing suggested is chemical-resistant overalls, jacket, and a rubber apron. The rubber apron protects the SCBA harness assembly and regulatory from becoming contaminated.

DECONTAMINATION OF EQUIPMENT

Insofar as possible, measures should be taken to prevent contamination of sampling and monitoring equipment. Sampling devices become contaminated, but monitoring instruments, unless they are splashed, usually do not. Once contaminated, instruments are difficult to clean without damaging them. Any delicate instrument which cannot be decontaminated easily should be protected while it is being used. It should be bagged, and the bag taped and secured around the instrument. Openings are made in the bag for sampling intake.

Decontamination Procedures

Sampling Devices--

Sampling devices required special cleaning. Decontamination procedures including solution and solvent selection must be developed in conjunction with the designated analytical laboratory.

Tools

Wooden tools are difficult to decontaminate because they absorb chemicals. They should be kept on site and handled only by protected workers. At the end of the response, wooden tools should be discarded. For decontaminating other tools, consult with the analytical laboratory and the site safety officer.

Heavy Equipment

Bulldozers, trucks, backhoes, bulking chambers, and other heavy equipment are difficult to decontaminate. The method generally used is to wash them with water under high pressure and/or to scrub accessible parts with detergent/water solution under pressure, if possible. In some cases, shovels, scoops, and lifts have been sandblasted or steam cleaned. Particular care must be given to those components in direct contact with contaminants such as tires and scoops. Swipe tests should be utilized to measure effectiveness.

Sanitizing of Personnel Protective Equipment

Respirators, reusable protective clothing, and other personal articles not only must be decontaminated before being reused, but also sanitized. The inside of masks and clothing becomes soiled due to exhalation, body oils, and perspiration. The manufacturer's instructions should be used to sanitize the respirator mask. If practical, protective clothing should be machine washed after a thorough decontamination; otherwise it must be cleaned by hand.

Persistent Contamination

In some instances, clothing and equipment will become contaminated with substances that cannot be removed by normal decontamination procedures. A solvent may be used to remove such contamination from equipment if it does not destroy or degrade the protective material. If persistent contamination is expected, disposable garments should be used. Testing for persistent contamination of protective clothing and appropriate decontamination must be done by qualified laboratory personnel.

Disposal of Contaminated Materials

All materials and equipment used for decontamination must be disposed of properly. Clothing, tools, buckets, brushes, and all other equipment that is contaminated must be secured in drums or other containers and labeled. Clothing not completely decontaminated onsite should be secured in plastic bags before being removed from the site.

Contaminated wash and rinse solutions should be contained by using step-in-containers (for example, child's wading pool) to hold spent solutions. Another containment method is to dig a trench about 4 inches deep and line it with plastic. In both cases the spent solutions are transferred to drums, which are labeled and disposed of with other substances onsite.

APPENDIX F
INSTRUMENT CERTIFICATION

Source: "Hazardous Materials Incident Response Operations"
Training Course Manual (165.1)

INHERENT SAFETY

The portable instrumentation used to evaluate hazardous material spills or waste sites must be demonstrated as being safe to use in those hostile environments. Electrical devices, such as the monitoring instruments, must be constructed in such a fashion as to eliminate the possibility of igniting a combustible atmosphere. The sources of this ignition could be an arc generated by the power source itself or the associated electronics, and/or a flame or heat source inherent in the instrument and necessary for its proper functioning.

Several engineering, insurance, and safety industries have standardized test methods, established inclusive definitions, and developed codes for testing electrical devices used in hazardous locations. The National Fire Protection Association (NFPA), a forerunner in this endeavor, has created minimum standards in its National Electrical Code (NEC), which is published every 3 years.

This code spells out, among other things, the following:

- Types of controls acceptable for use in hazardous atmospheres.
- Types of areas in which hazardous atmospheres can be generated and the types of materials that generate these atmospheres.

HAZARDOUS ATMOSPHERES

Depending upon the response worker's background, the term "hazardous atmosphere" conjures up situations ranging from toxic air contaminants to flammable atmospheres. For our purposes, an atmosphere is hazardous if it meets the following criteria:

- It is a mixture of any flammable material in air (see Class and Group below) whose composition is within this material's flammable range (LEL-LFL).
- A critical volume of the mixture is sufficiently heated by an outside ignition source.
- The resulting exothermic reaction propagates the flame beyond where it started.

Hazardous atmospheres can be produced by one of three general types of materials:

- Flammable gases/vapors
- Combustible dusts
- Ignitable fibers

Whereas the flammable material may define the hazard associated with a given product, the occurrence of release (how often the material generates a hazardous atmosphere) dictates the risk. Two types of releases are associated with hazardous atmospheres:

- Continuous: Those existing continuously in an open unconfined area during normal operating conditions.
- Confined: Those existing in closed containers, systems, or piping, where only ruptures, leaks, or other failures result in a hazardous atmosphere outside the closed system.

There are six possible environments in which a hazardous atmosphere can be generated. However, not every type of control will prevent an ignition in every environment. To adequately describe the characteristics of those environments and what controls can be used, the National Electrical Code defines each characteristic:

Class is a category describing the type of flammable material that produces the hazardous atmosphere:

- Class I is flammable vapors and gases, such as gasoline, and hydrogen. Class I is further divided into groups A, B, C, and D on the basis of similar flammability characteristics (Table F-1).
- Class II consists of combustible dusts like coal or grain and is divided into groups E, F, and G.
- Class III is ignitable fibers such as produced by cotton milling.

Division is the term describing the "location" of generation and release of the flammable material.

- Division 1 is a location where the generation and release are continuous, intermittent, or periodic into an open, unconfined area under normal conditions.
- Division 2 is a location where the generation and release are in closed systems or containers and only from ruptures, leaks, or other failures.

Using this system, a hazardous atmosphere can be routinely and adequately defined. As an example, a spray-painting operation using acetone carrier would be classified as a Class I, Division 1, Group D environment. Additionally, an abandoned waste site containing intact closed drums of methyl ethyl ketone, toluene, and xylene would be considered a Class I, Division 2, Group D environment. Once the containers begin to leak and produce a hazardous atmosphere, the environment changes to Class I, Division 1, Group D.

TABLE F-1. CLASS I CHEMICALS BY GROUP

<u>Group A Atmospheres</u>	<u>Group D Atmospheres</u>
Acetylene	Acetic Acid (glacial)
	Acetone
<u>Group B Atmospheres</u>	Acrylonitrile
Acrolein (inhibited)	Ammonia
Arsine	Benzene
Ethylene oxide	Butane
Hydrogen	1-Butanol (butyl alcohol)
Manufactured gases containing more	2-Butanol (secondary butyl alcohol)
than 30% hydrogen (by volume)	n-Butyl acetate
Propylene oxide	Isobutyl acetate
Propyl nitrate	di-Isobutylene
	Ethane
<u>Group C Atmospheres</u>	Ethanol (ethyl alcohol)
Acetaldehyde	Ethyl acetate
Allyl alcohol	Ethyl acrylate (inhibited)
n-Butyraldehyde	Ethyl diamine
Carbon monoxide	Ethylene dichloride
Cyclopropane	Ethylene glycol monomethyl ether
Diethyl ether	Gasoline
Diethylamine	Heptanes
Epichlorohydrin	Hexanes
Ethylene	Isoprene
Ethylamine	Isopropyl ether
Ethyl mercaptan	Mesityl oxide
Ethyl sulfide	Methane (natural gas)
Hydrogen cyanide	Methanol (methyl alcohol)
Hydrogen sulfide	3-Methyl-1-butanol (isoamyl alcohol)
Morpholine	Methyl ethyl ketone
2-Nitropropane	Methyl isobutyl ketone
Tetrahydrofuran	2-Methyl-1-propanol (isobutyl alcohol)
Unsymmetrical dimethyl hydrazine	2-Methyl-2-propanol (tertiary butyl alcohol)
(UDMH, 1-, 1-dimethyl hydrazine)	Octanes
	Petroleum naphtha ^a
	Pentanes
	1-Pentanol (amyl alcohol)
	Propane
	1-Propanol (propyl alcohol)
	2-Propanol (isopropyl alcohol)
	Propylene
	Pyridine
	Styrene
	Toluene
	Vinyl acetate
	Vinyl chloride
	Xylenes

^a A saturated hydrocarbon mixture boiling in the range 20° - 135°C (68° - 275°F). Also known by the synonyms benzine, ligroin, petroleum ether, or naphtha.

Source: National Electrical Code, Vol. 70, Table 500-2. National Fire Protection Association, 470 Atlantic Avenue, Boston, MA 02210 (1981).

CONTROLS

Three methods exist to prevent a potential ignition source from igniting a flammable atmosphere:

- Explosion-proof: Encase the ignition source in a rigidly built container. "Explosion-proof" instruments allow the flammable atmosphere to enter. If and when an arc is generated, the ensuing explosion is contained within the specially designed and built enclosure. Within it, any flames or hot gases are cooled prior to exiting into the ambient flammable atmosphere so that the explosion does not spread into the environment.
- Intrinsically Safe: Reduce the potential for arcing among components by encasing them in a solid insulating material. Also, reducing the instrument's operational current and voltage below the energy level necessary for ignition of the flammable atmosphere provides equal protection. An "intrinsically safe" device, as defined by the National Electrical Code, is incapable "of releasing sufficient electrical or thermal energy under normal or abnormal conditions to cause ignition of a specific hazardous atmospheric mixture in its most easily ignited concentration. Abnormal conditions shall include accidental damage to any ... wiring, failure of electrical components, application of over-voltage, adjustment and maintenance operations and other similar conditions."
- Purged: Buffer the arcing or flame-producing device from the flammable atmosphere with an inert gas. In a pressurized or "purged" system, a steady stream of, for example, nitrogen or helium is passed by the potential arcing device, keeping the flammable atmosphere from the ignition source. This type of control, however, does not satisfactorily control analytical devices that use a flame or heat for analysis such as a combustible gas indicator (CGI) or gas chromatography (GC).

CERTIFICATION

National groups such as Underwriters Laboratories (UL), Factory Mutual (FM), and the American National Standards Institute (ANSI), together with NFPA, have developed test protocols for certifying explosion-proof, intrinsically safe, or purged devices to meet minimum standards of acceptance.

An electrical device certified under one of these test methods carries a permanently affixed plate showing the logo of the laboratory granting certification and the Class(es), Division(s), and Group(s) it was tested against. See Figure F-1.

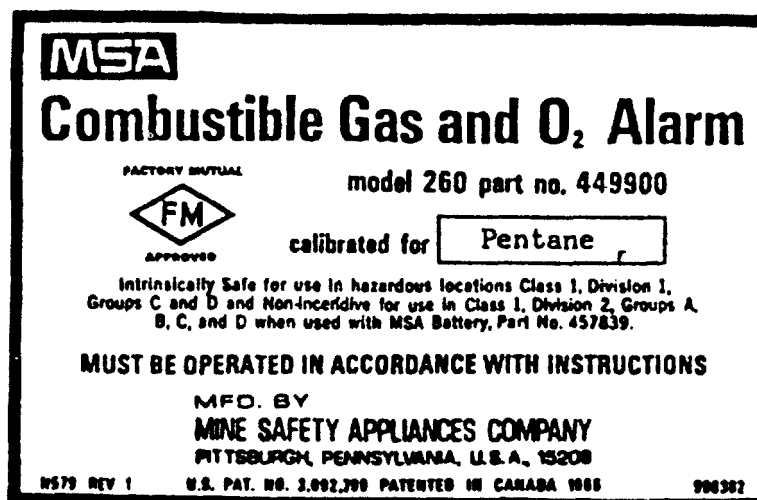


Figure F-1. Example Device Certification Plate

Certification means that if a device is certified as explosion-proof, intrinsically safe, or purged for a given Class, Division, and Group, and is used, maintained, and serviced according to the manufacturer's instructions, it will not contribute to ignition. The device is not, however, certified for use in atmospheres other than those indicated.

Any manufacturer wishing to have an electrical device certified by FM or UL must submit a prototype for testing. If the unit passes, it is certified as submitted. However, the manufacturer agrees to allow the testing laboratory to randomly check the manufacturing plant at any time, as well as any marketed units. Furthermore, any change in the unit requires the manufacturer to notify the test laboratory, which can continue the certification or withdraw it until the modified unit can be retested.

A unit may be certified either by UL, FM, or both. Both Laboratories follow test protocols established by NFPA and ANSI. Therefore one certification is no better or worse than the other. The important consideration is that the device is approved for the Class(es), Division(s), and Group(s) it will be used in.

The mention of FM or UL in the manufacturer's equipment literature does not guarantee certification. All certified devices that are used in hazardous (flammable) locations must be marked to show Class, Division, and Group, per NEC Table 500-2(b).

Other organizations such as the Mine Safety and Health Administration (MSHA), Canadian Standards Association (CSA), National Electrical Manufacturers Association (NEMA), and the U.S. Coast Guard (USCG) have developed their own testing and certification schemes for electrical devices in hazardous locations common to their jurisdiction.

MSHA tests and certifies electrical equipment to be used in hazardous atmospheres associated with underground mining. These atmospheres usually contain methane gas and coal dust; hence, the tests and certification are specific to those two contaminants.

Often the same monitoring equipment is used both in mines and above ground and would carry more than one certification, such as FM and MSHA.

To ensure personnel safety, it is recommended that only approved (FM or UL) instruments be used onsite and only in atmospheres for which they have been certified. When investigating incidents involving unknown hazards, the monitoring instruments should be rated for use in the most hazardous locations. The following points will assist in selection of equipment that will not contribute to ignition of a hazardous atmosphere:

- In an area designated Division 1, there is a greater probability of generating a hazardous atmosphere than in Division 2. Therefore, the test protocols for Division 1 certification are more stringent than those for Division 2. Thus, a device approved for Division 1 is also permitted for use in Division 2, but not vice versa. For most response work this means that devices approved for Class I (vapors, gases), Division 1 (areas of ignitable concentrations), Groups A, B, C, D should be chosen whenever possible. At a minimum, an instrument should be approved for use in Division 2 locations.
- All instruments to be used in a methane environment should be approved by the Mine Safety and Health Administration (MSHA) as being safe in such atmospheres.
- There are so many Groups, Classes, and Divisions that it is impossible to certify an all-inclusive instrument. Therefore, select a certified device based on the chemicals and conditions most likely to be encountered. For example, a device certified for a Class II, Division 1, Group E (combustible metal dust) would offer little protection around a flammable vapor or gas.

APPENDIX G
APPLICABLE TABLES

TABLE G-1. RANDOM NUMBERS

=====														
03	47	43	73	86	36	96	47	36	61	46	98	63	71	62
97	74	24	67	62	42	81	14	57	20	42	53	32	37	32
16	76	62	27	66	56	50	26	71	07	32	90	79	78	53
12	56	85	99	26	96	96	68	27	31	05	03	72	93	15
55	59	56	35	64	38	54	82	46	22	31	62	43	09	90
16	22	77	94	39	49	54	43	54	82	17	37	93	23	78
84	42	17	53	31	57	24	55	06	88	77	04	74	47	67
63	01	63	78	59	16	95	55	67	19	98	10	50	71	75
33	21	12	34	29	78	64	56	07	82	52	42	07	44	38
57	60	86	32	44	09	47	27	96	54	49	17	46	09	62
18	18	07	92	46	44	17	16	58	09	79	83	86	19	62
26	62	38	97	75	84	16	07	44	99	83	11	46	32	24
23	42	40	64	74	82	97	77	77	81	07	45	32	14	08
52	36	28	19	95	50	92	26	11	97	00	56	76	31	38
37	85	94	35	12	83	39	50	08	30	42	34	07	96	88
70	29	17	12	13	40	33	20	38	26	13	89	51	03	74
56	62	18	37	35	96	83	50	87	75	97	12	25	93	47
99	49	57	22	77	88	42	95	45	72	16	64	36	16	00
16	08	15	04	72	33	27	14	34	09	45	59	34	68	49
31	16	93	32	43	50	27	89	87	19	20	15	37	00	49
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INSTRUCTIONS FOR THE USE OF THE RANDOM NUMBER TABLE

1. Number the members of the lot (i.e., the drums onsite, the sections within a grid) in a numerical order.
2. Decide on the number of samples necessary. This should be a number sufficient to give statistical significant data. Ten to 20 percent is usually adequate. This number should be predetermined in the test plan or should coincide with the time and equipment available.
3. Using the set of random numbers above; choose any number as a starting point, then proceed to select the next number in any predetermined direction (i.e., down the column, across the rows) until the selection process has yielded the desired number of samples.

TABLE G-2. CONVERSION FACTORS/TABLES OF MEASUREMENT

MISCELLANEOUS

<p> Units kilo — means one thousand centi — means one-hundredth milli — means one-thousandth micro — means one-millionth </p> <p> Length and area 1 statute mile (mi) = 5280 feet = 1.609 kilometers 1 foot (ft) = 12 inches = 30.48 centimeters 1 inch (in.) = 25.40 millimeters 100 ft per min = 0.508 meter per sec 1 square foot = 144 sq inches = 0.0929 sq meter 1 square inch = 6.45 sq centimeters 1 kilometer (km) = 1000 meters = 0.621 statute mile 1 meter (m) = 100 centimeters (cm) = 1000 millimeters (mm) = 1.094 yards = 3.281 feet = 39.37 inches 1 centimeter = 1 x 10⁸ angstroms (Å) 1 micron = 0.001 millimeter = 0.000039 inch 1 meter per sec = 166.9 ft per min </p> <p> Weight 1 U.S. long ton = 2240 pounds = 1016 kilograms 1 U.S. short ton = 2000 pounds = 907 kilograms 1 pound (lb) = 16 ounces = 7000 grains = 0.454 kilogram 1 ounce (oz) = 0.0625 pound = 28.35 grams 1 kg per cu m = 1 gram per liter = 1 part per thousand 1 g per cu m = 1 mg per liter = 1 part per million (ppm) 1 ppm = 8.33 lb per million gal 1 grain per gal = 143 lb per million gal 1 lb per million gal = 0.12 ppm 1 lb per million gal = 0.007 grain per gallon 1 lb per thousand gal = 120 ppm </p> <p> Pressure 1 atmosphere = 760 mm (29.92 in.) mercury with density 13.595 grams per cc </p>	<p> 1 grain = 64.8 milligrams = 0.0023 ounce 1 pound per foot = 1.488 kg per meter 1 metric ton (tonne) = 1000 kilograms = 0.984 long ton = 1.102 U.S. short tons = 2205 pounds 1 kilogram (kg) = 1000 grams = 2.205 pounds 1 gram (g) = 1000 milligrams (mg) = 0.03527 ounce = 15.43 grains 1 kg per meter = 0.672 pound per ft </p> <p> Volume 1 cubic yard = 27 cu ft = 0.765 cu m 1 cubic foot = 1728 cu in. = 28.32 liters = 7.48 U.S. gallons 1 cubic inch = 16.39 cu centimeters (cu cm) 1 imperial gallon = 277.4 cu in. = 4.55 liters 1 U.S. gallon = 0.833 imperial gallon = 3.785 liters = 231 cu in. = 0.1337 cu ft 1 U.S. barrel (petroleum) = 42 U.S. gallons = 35 imperial gallons 1 cubic meter (cu m) = 1000 liters = 35.31 cu ft 1 liter = 1000 cc = 0.2200 imperial gallon = 0.2642 U.S. gallon = 61.0 cu in. = 14.696 lb per sq in. = 1.033 kg per sq cm 1 atmosphere (metric) = 1 kg per sq cm = 10,000 kg per sq m = 10 m head of water = 14.22 lb per sq in. 1 ft. per square foot = 0.1924 in. of water = 4.88 kg per sq m 1 lb per square inch = 2.036 in. head of mercury = 2.309 ft head of water = 0.0703 kg per sq cm = 0.0690 bar 1 ton per square inch = 1.406 kg per sq mm 1 inch head of water = 5.20 lb per sq ft </p>	<p> 1 board foot = 12 in. x 12 in. x 1 in. thick = 144 cu in. 1 cu ft per min = 1.680 cu m per hour 1 cu m per hour = 0.588 cu ft per min 1 cu ft per sec = 646.316 gallons per day = 448.83 gallons per min 1 gallon per min = 0.00144 million gallons per day </p> <p> Density (weight/volume) 1 cu ft per lb = 0.0824 cu m per kg 1 lb per cu ft = 16.02 kg per cu m 1 grain per cu ft = 2.208 grams per cu m 1 grain per U.S. gallon = 17.11 grams per cu m = 17.11 mg/liter 1 cu m per kg = 16.02 cu ft per lb 1 kg per cu m = 0.0824 lb per cu ft 1 gram per cu m = 0.437 grain per cu ft = 0.0584 grain per U.S. gallon 1 gram per cc = 62.4 lb per cu ft 1 gram per liter = 58.4 grains per U.S. gallon </p> <p> Water at 62 F (16.7 C) 1 cubic foot = 62.3 lb 1 pound = 0.01604 cu ft 1 U.S. gallon = 8.34 lb </p> <p> Water at 39.2 F (4 C), maximum density 1 cubic foot = 62.4 lb 1 cubic meter = 1000 kg 1 pound = 0.01602 cu ft 1 liter = 1.0 kg 1 foot head of water = 0.433 lb per sq in 1 m head of water = 0.1 kg per sq cm 1 in. head of mercury = 0.491 lb per sq in 1 m head of mercury = 1.360 kg per sq cm 1 kilogram per sq m = 1-mm head of water = 0.2048 lb per sq ft 1 kilogram per sq cm = 735.5 mm of mercury = 14.22 lb per sq in 1 kg per sq mm = 0.711 ton per sq m </p> <p> In these conversions, inches and feet of water are measured at 62 F (16.7 C), millimeters and meters of water at 39.2 F (4 C), and inches, millimeters and meters of mercury at 32 F (0 C). </p>
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Source : Betz Handbook of Industrial Water Conditioning, 1976 seventh edition, Betz Laboratories, Inc., Trevose, PA

Example:

Soil samples are to be collected from a field 10 meters by 15 meters in area. Equipment and laboratory arrangements have been made to handle eight samples.

- A. The area is divided into an imaginary 1 meter grid.
- B. Each quadrant in the grid is assigned a number in a numerical order; West to East, North to South (or left to right, top to bottom).
- C. Referring to the Random Number table it is arbitrarily decided to start at the first number in the third row, then proceed down the column.

This would result in the selection of 43 as the first number followed by 24, 62, 85, 56, 77, 17 and 63 as the eighth and final selection.

The grids corresponding to these numbers would then be sampled.