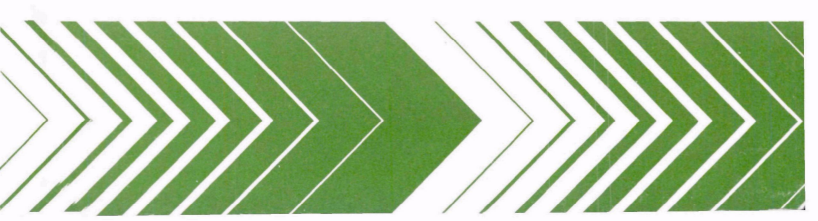
Research and Development



# Procedure for the Evaluation of Environmental Monitoring Laboratories

**Environmental Monitoring Series** 



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## PROCEDURE FOR THE EVALUATION OF ENVIRONMENTAL MONITORING LABORATORIES

bу

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Contract No. 68-03-2171

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CINCINNATI, OHIO 45268

#### DISCLAIMER

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#### FOREWORD

Environmental measurements are required to determine the quality of ambient waters and the character of waste effluents. The Environmental Monitoring and Support Laboratory-Cincinnati conducts research to:

- Develop and evaluate techniques to measure the presence and concentration of physical, chemical and radiological pollutants in water, wastewater, bottom sediments and solid waste.
- Investigate methods for the concentration, recovery and identification of viruses, bacteria and other microbiological organisms in water. Conduct studies to determine the responses of aquatic organisms to water quality.
- Conduct an Agency-wide quality assurance program to assure standardization and quality control of systems for monitoring water and wastewater.

The latest quality assurance report on procedures for evaluation of environmental monitoring laboratories was prepared by Tracor Jitco, Inc. The report, in detail, contains registration and preliminary questionnaire forms, on-site visit checklist, evaluator's guide, and a scoring system for assessment of the laboratory's management, personnel, facilities, analytical methodology and instruments, and its quality control procedures.

This research report is not an official EPA Manual. Rather, it is a report which is but one of a series being used as input to develop EPA Manuals and Guidelines for Certification Programs.

Dwight G. Ballinger Director, EMSL-Cincinnati

#### ABSTRACT

Tracor Jitco, Inc., examined in depth existing evaluation procedures of EPA, Federal and State Agencies with the aim of incorporating their best features in a procedure for general use in evaluating laboratories engaged in measuring environmental pollution.

The procedures developed are suitable for the media of air, water, radiation, and pesticides. They are intended for use by EPA Regions in evaluating state laboratories and by the states in evaluating local or private laboratories. They are useful as a management tool to control or upgrade laboratory performance or they could be used as part of a laboratory accreditation or certification system. The inclusion of a scoring plan makes it possible, with suitable training of evaluators in uniform application of the procedures, to make comparisons with standards of performance.

The laboratories are required to provide information on physical plant, equipment, personnel, quality control and other general aspects of laboratory performance on check-off types of forms provided. This is followed by an on-site inspection during which information on less quantifiable aspects are obtained. This phase of the evaluation is oriented to the specific methodology for which the laboratory is to be qualified.

The scoring system includes inherent weighing of criteria. The procedure is designed to be compatible with programs of proficiency testing and taken as a part of a total quality assurance program will contribute to the objectivity of the determination of laboratory capability.

This research report is not an official EPA manual. Rather, it is a report which is but one of a series being used as input to develop EPA Manuals and Guidelines for Certification Programs.

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#### ACKNOWLEDGMENTS

Thanks are due to the Project Officer, Mr. Edward Berg, and to the staff of the Environmental Monitoring and Support Laboratory, Cincinnati, for their support of this project. The many EPA Research Laboratories and Regional Laboratories visited at various stages of development of the procedure were all most generous in providing information and in giving advise. We also found very helpful the review of the draft of the procedure by the Wisconsin Department of Natural Resources and the Illinois Environmental Protection Agency. We encountered many points of view and have done our best to reconcile differences and to come up with a procedure that is standardized, widely useful, and fair in its application.

# SECTION 1 INTRODUCTION

An evaluation procedure has been developed based on EPA experience in evaluating its Regional Laboratories, the experience of other governmental and private evaluating agencies, and the combined experience of the contractor's senior staff.

Nevertheless, the result has been arrived at independently, specifically without reference to EPA's conclusions about its own evaluation efforts. This was done consciously so that the result of the contractor's efforts will stand on its own merits. Moreover, it has the advantage that a completely disinterested point of view has been brought to bear on the problem.

Two objectives of the project have had a strong bearing on the nature of the procedure that has been developed:

- A major objective was to produce combined forms containing sections with general application and sections with applications to specific media, in recognition of the fact that the areas of uniformity in an evaluation protocol outweigh the differences required by the media covered, namely air, water, radiation, and pesticides.
- 2. A plan of scoring was required, using rating criteria based on standards of acceptability in operation in EPA and elsewhere and based on the contractor's own experienced judgment.

The resulting procedure has several unique aspects.

- It collects information about areas of management, personnel, facilities, methodology, instrumentation, and quality control oriented toward the requirements of environmental monitoring laboratories.
- It presents criteria against which the individual laboratory may be judged in each area.

- It contains an Evaluator's Guide which explains the intent of inspection in each area and suggests specific questions to be asked to enable the evaluator to arrive at the necessary judgments.
- It is sectionalized as to methodology and equipment so that only the parts applicable to even a small laboratory or to a laboratory devoted to a single medium need be used, thus avoiding unnecessary burden on the laboratory.
- The scoring system is adjustable to the size and scope of the laboratory yet provides a final score which is comparable under any circumstances of use of the procedure.

# SECTION 2 CONCLUSIONS

The procedure which has been presented in this manual is directly applicable to the evaluation of laboratories of all sizes. In its entirety it will apply to large laboratories. In this application it is lengthy, but its length is justified by the necessity for a thorough inspection of all aspects of laboratory personnel, facilities, equipment, and operations. To do less would be to slight some important aspect and make difficult a balanced, meaningful scoring system.

For smaller laboratories or specialized laboratories, only the applicable portions of the procedure need to be used. The Registration Form is intended to provide information that will make it necessary to send out only the pertinent parts of the Preliminary Questionnaire. The information provided by the Preliminary Questionnaire, assimilated by the evaluator, or evaluation team, before the onsite visit should limit the first hand inspection to the aspects of the laboratory's operations that appear to deviate from standard.

When used conscientiously by evaluators with pertinent scientific back-ground the procedure and its scoring system should result in the ability to discern those laboratories that are acceptable participants in the environmental monitoring programs.

# SECTION 3 BACKGROUND AND SCOPE

The pollution of the atmosphere, the contamination of the waters, and the littering of the land have become problems international in scope. The continued violence done by man to the total environment must be checked if this planet is to remain a fit place in which to live. One of the first steps that can be taken is the qualitative and quantitative monitoring of the environment.

Successful monitoring of the environment requires the identification of the contaminants, an accurate measurement of the amounts present, and pin-pointing of the sources of the pollution. Because of the increase in number of contaminating substances, many of which require sophisticated analysis, and because of the reduction in levels of pollution that can be tolerated, the involvement of an increasingly large number of people and of laboratories is required.

In the United States, the U.S. Environmental Protection Agency has the responsibility for enforcement of national laws and regulations designed to restore and protect the environment. Its work is assisted and supplemented by environmental programs carried out by the states. The private sector is also depended upon to carry a part of the laboratory analytical workload. The wide diffusion of monitoring and analysis leads to a need for standards of performance. Extensive laboratory inspection and evaluation must be done to ascertain the capabilities of the participating laboratories. In order to avoid arbitrary inspections and to protect both the evaluating agency and the laboratories from capricious judgements, this procedure which standardizes requirements has been prepared.

The procedure provides a basis for inspection and evaluation of environmental monitoring laboratories at national, state, and private levels. It is applicable to laboratories concerned with the various media, particularly air, water, pesticides, and radiation. The experience of EPA and of other standardizing institutions has been used as a basis for this procedure. It is thorough, yet as concise as the intended wide range of applicability permits.

This procedure employs standards which will make evaluation as objective as it can be made. It includes a scoring system for assessment of the laboratory's management, personnel, facilities, analytical methodology and instruments, and its quality control procedures. An acceptable score will signify that there are no serious deficiencies in the organization, physical plant, or technical operations of the laboratory.

#### PURPOSE OF EVALUATION

Enhancement of the performance of environmental monitoring laboratories is the primary goal of the laboratory evaluation. Its purpose is to ascertain that the laboratory follows sound scientific procedures in its analytical work; that it operates under the auspices of good management and professional supervisors; that it utilizes proper equipment; and that it maintains and uses accurate records. The evaluation procedure provides the laboratory an opportunity for improvement by identifying weaknesses in its organization or performance and to obtain information and assistance for overall improvement. In this sense, the evaluation may serve not only to assure laboratory competence; but also to promote professionalism in the laboratory by facilitating the establishment of standards of excellence.

Any system designed for the evaluation of laboratories will inevitably identify certain laboratories which fail to meet the established standards. This procedure for evaluation of laboratories does not necessitate a definitive rejection of unqualified laboratories. It provides an opportunity for the laboratory to correct existing deficiencies. If the laboratory complies with recommended modifications, it may receive an acceptable rating.

The uniform scoring system employed in the procedure considers a large number of characteristics which are given preassigned weights. This contributes to the objectiveness and comparability of the evaluation which are among its principal purposes.

#### INTENDED APPLICATIONS OF THE PROCEDURE

The procedure for laboratory evaluation is a versatile instrument. The preliminary questionnaire coupled with the onsite checklists is suitable to a number of situations. It may serve as a self-evaluation for Environmental Protection Agency laboratories. It may be used by EPA for the evaluation of state laboratories. It could be used by state laboratory personnel to evaluate commercial laboratories. The procedure was not designed for use in a formal certification program, however, it could readily be adapted for that purpose.

Although it is recognized that the different media of air, water, pesticides, and radiation have some unique methodologies, the areas of uniformity in all laboratories outweigh the differences and a generally applicable procedure has been developed. Sections with application to specific media can be used for inspection to the extent necessary.

The procedure does not purport to be a panacea. For example, although recognition is given to the necessity for participation in interlaboratory proficiency testing programs, the scores obtained in such programs do not enter directly into the scoring recommended in this procedure. The procedure simply provides a methodology necessary for environmentally concerned scientists to ensure that a laboratory has the capability for valid analyses. The combination of the score from applying this procedure with scores from inter-laboratory testing programs should be the object of further consideration.

The extremely large number of data points collected may have to be collated by computer. Although this is not one of the requirements of this procedure, most of the data will have been recorded in such a way that it can readily be computerized. Most of the answers to the questionnaire require only a checkmark and not involved descriptions of the laboratory.

The media covered include the broad application of environmental monitoring to air, water, pesticides, and radiation. This involves chemical methodology appropriate to potable water, wastewater, ambient water, ambient air, stack emissions and other source emissions into the atmosphere, sediments, pesticides and other organic chemicals, both natural and industrial. It involves biology, including aquatic biology and virology. It also includes bacteriology as applied to potable water, waste water and ambient water. Finally, it includes radiation measurement.

The analytical methodology required is in a state of flux. Some methods are EPA approved, some are used as interim methods and others are in various states of development and are in more or less wide use. Although this procedure lends itself to the evaluation of performance of all methodology required in the various areas, the material actually presented on methodology is limited to those methods referenced in the Federal Register. These are the presently EPA approved methods. For water and radiation test methods, see Federal Register, Vol. 35, No. 199, October 16, 1973. Interim methods for algicides, chlorinated organic compounds, and pesticides can be obtained from the Environmental Monitoring and Support Laboratory, USEPA, 1014 Broadway, Cincinnati, Ohio 45268. Air test methods are referenced in Federal Register, Vol. 36, No. 228, November 25, 1971 and Vol. 38, No. 110, June 8, 1973.

Methods for measurement of emissions from stationary sources differ in important aspects from methods for measurement in ambient air. These source methods are to be found in Federal Register, Vol. 36, No. 247, Part II, December 1971; Vol. 38, No. 111, June 11, 1973; Vol. 39, No. 47, March 8, 1974; Vol. 40, No. 152, August 6, 1975; and Vol. 40, No. 194, October 6, 1975.

Biology is an important area not covered by referenced methods. However, see Bibliography items 6-7-8-9 for methods in use that may be consulted for methodological requirements of satisfactory laboratory performance in this area.

Modified or alternate methods ("equivalent" methods) may be used if specifically approved under published regulations. A laboratory under evaluation is required to provide information on any such methods in use. The evaluator must refer to this information in order to judge whether the laboratory's use of the methods produces satisfactory results.

Although some state environmental monitoring laboratories are a part of, or are closely associated with, Health Laboratories, this procedure is not intended for use in any health oriented analyses. Procedures exist for evaluation of health laboratories where this is required for certification or licensing.

Although the word "laboratory" is used, it is emphasized that the field aspects as well as the laboratory aspects of environmental monitoring must be a part of any complete evaluation. The procedure developed herein is compatible with the "total system" concept. The evaluator should go into the field to look at monitoring equipment including flow measurement instrumentation and automatic sample compositing equipment.

#### USE OF PROCEDURE

Experts, such as those found in the larger environmental protection agencies, who are experienced in all of the media may not always be available for inspection and evaluation duties. It may be necessary to employ evaluators who have not had long years of experience in all the details of methodology of environmental monitoring. Therefore, this procedure has been designed for use by individuals who are skilled in science, but who will find guidelines useful for the evaluation of specialized laboratories. An elaborate "Guide for Evaluators" is an essential part of the Manual.

As a part of this "Guide" there is included, where available, very detailed background material in some of the specialized methodologies. For example, the EPA Check List for Bacteriological Examination of Water is to be found in Part 4. Also, recommended laboratory performance standards are included, such as "A Schedule of Suggested Instrument Calibrations" and a "Table of Recommendations for Sampling and Sample Preservation" from the EPA Manual of Methods for Chemical Analysis of Water and Wastes. Other such helps could be added, if so desired, as they became available.

The ideal evaluator should possess a broad understanding of scientific methods and an appreciation of the complexity of analytic procedures. A strong background in an applied science, preferably chemistry, coupled with some experience in laboratory management should equip the inspector with the insight required to thoroughly assess a laboratory's operation.

The qualifications required for the position of evaluator should be strictly observed. Failure to do so would be a disservice to both the EPA and the laboratory undergoing evaluation. For even the most detailed and efficient guidelines cannot guarantee a quality evaluation if administered by an unqualified individual.

The evaluation procedure, though not extremely complex, is lengthy and time-consuming. It will run most smoothly if the evaluators have had some training in its use. The introductory material, the various instruction sheets, and particularly the "Evaluator's Guide" may be used as a text in training sessions for evaluators. If such training sessions are not arranged, at least the evaluator should study the entire procedure thoroughly before embarking on an evaluation.

# SECTION 4 REGISTRATION AND PRELIMINARY EVALUATION

The U.S. Environmental Protection Agency is engaged in the monumental task of pollution abatement and control on a national scale. The workload grows with the increase of substances which require sophisticated analyses, with the growing technical complexity of analytical procedures and with the reduction of tolerated contamination levels. The cooperation of many laboratories, state and commercial, must be enlisted to further EPA's efforts to maintain the integrity of the environment.

Laboratories which participate in environmental monitoring must meet rigid standards of excellence. The data gleaned from their analyses must be defendable for it may serve as evidence in a court of law. To ensure the analytic capabilities of collaboratoring laboratories, EPA has instituted a systematic evaluation procedure.

The evaluation procedure is a standardized instrument designed to produce an objective appraisal of a laboratory's performance. It strives to utilize the insights of a qualified evaluator without falling prey to the caprices of a subjective appraisal. It employs a numerical scoring system to organize the myriad details and to produce a manageable result. The scoring framework supplies a strong influence toward uniformity in the application of criteria from laboratory to laboratory.

A laboratory evaluation is a time consuming endeavor. To minimize this time factor, the EPA procedure consists of a three step process: Registration, Completion of a Preliminary Questionnaire by the laboratory, and an Onsite survey by personnel of the evaluating agency.

A laboratory interested in participating in an evaluation may identify itself by completion of a brief registration form. This form will indicate to the evaluating agency the extent of the evaluation required, i.e., whether it is to cover all media or a few tests for one medium.

Parts 1, 2, 3, and 6 will go to all laboratories. Those parts of Part 4 (Chart C - Analytical Methodology) and of Part 5 (Chart D - Analytical Instruments) applicable to the media with which the laboratory is involved will be selected and sent to the laboratory for completion.

Return of the completed questionnaire triggers the final phase of the evaluation.

The evaluator carefully studies the information provided by the laboratory and notes any items which require special attention. The onsite visit is then scheduled.

During the onsite visit, the evaluator implements the numerical scoring system to assess the laboratory operation. Any deficiencies which require improvement prior to scoring are identified and discussed with the laboratory.

When the evaluation has been completed, a written report of deficiencies and recommendations will be sent to the laboratory director. Upon return of satisfactory evidence that all reported deficiencies have been taken care of, a final score will be issued.

An acceptable score will signify that the laboratory is fully qualified to participate in the vital work of preserving a safe, liveable environment.

### **REGISTRATION FORM**

The evaluation of Environmental Monitoring Laboratories is designed to assist the participating laboratories to upgrade their overall performance in order to safeguard the scientific and legal validity of their data. Submission of this registration form is the first step in the evaluation process. A preliminary questionnaire which requests background information about the laboratory's staff, facilities, and operating procedures is the second step. Upon completion of the preliminary questionnaire, an onsite visit to assess the performance capability of the laboratory will be scheduled at the convenience of the laboratory.

٦.	Name of Laboratory		
2.	Address	<del></del>	
3.	Telephone Number		<del></del>
4.	Name of Laboratory Director		
5.	If Private, Name of Owner		<del></del>
6.	Type of Laboratory		
	Commercial (privately owned,	works on fee or contract ba	asis)
	☐ Noncommercial (publicly cont	trolled; usually does not wor	k on a fee basis)
7.	Provide a brief functional description	on of the activities of the lab	oratory
		<del> </del>	
8.	Media to be covered in evaluation		
	∐ Water		
	Chemistry		
	Bacteriology		
	☐ Biology		
	Air		
	Pesticides		
	☐ Radiation		
_	Other (specify)		
9.	If evaluation is not desired for com tests for which you wish to be eval		
	are availiable is given on overleaf.*		• •
	evaluation.)	(Do not list for any intention	Troi which you desire complete
10.	Total Number of employees	Technical	Administrative
	* EPA approved water and radiation test meth- interim methods for algicides, chlorinated orga Monitoring and Support Laboratory, U. S. Env EPA approved air test methods are referenced in No. 110, June 8, 1973.	nic compounds, and pesticides can be ironmental Protection Agency, 1014	obtained from Environmental Broadway, Cincinnati, Ohio 45268.
Sign	ature of Director		Date

#### PRELIMINARY QUESTIONNAIRE

This questionnaire is designed to elicit all the information required prior to an onsite survey. Please make a concerted effort to furnish the information as accurately and concisely as possible.

For convenience, the questionnaire has been divided into six parts:

- 1) General Laboratory Information
- 2) Personnel
- 3) Laboratory Space and Facilities
- 4) Technical Services
- 5) Analytical Instruments and Special Apparatus
- 6) Quality Control

In each section, the questions are styled for the ease of the laboratory's response. In many cases only a check ( $\checkmark$ ) is required. Other questions call for a short answer; clarity and brevity should hallmark your response. If you need more space, please continue on blank sheets and attach them to the questionnaire.

Each section is independent, so that the different sections may be distributed to the most knowledgeable persons in the laboratory who can complete their parts independently. Finally, management can assemble and check all responses before returning the completed forms.

Upon return of the completed questionnaire, the onsite visit will be scheduled at your convenience. The time involved in the onsite evaluation can be minimized by a thorough presentation of the information sought in the preliminary questionnaire. Therefore, it is advantageous to both your laboratory and the evaluating agency if these questions are answered precisely and completely.

Thank you for your cooperation.

## PART 1. GENERAL INFORMATION ABOUT THE LABORATORY

1.	Name	e of Laboratory				<del></del>
2.	Addr	ess				·
3.	Telep	phone Number		<del></del>		
4.	Name	e of Laboratory Director	<del></del>		<del></del>	
5.	affilia	de an organization chart of th ations to show how the labora e check.	e laboi tory fi	ratory, including any field its into the general organiza	operati Itional	ions or other internal structure. If attached,
6.	List r	names and addresses of externa	al orga	inizations used for significa	nt supp	porting technical services
						<del></del>
7.		names of principal users of serv		•		
8.	Has the by wh	he laboratory been evaluated p	oreviou	usly? Yes 🗌 No 🔲 If	yes, w	hen
9.	Do yo	ou perform monitoring activiti toring activity:	es?	Yes 🗆 No 🗀 II	f yes, p	lease check nature of
		Water Quality		Air-Ambient		Radiation
		Estuaries		Air-Source		Other (specify)
		Oceans		Pesticides		
		NPDES				

		Lab Name
10.	Do you participate in specify.	enforcement actions, emergency episodes, or special studies? Please
11.	Provide a copy of the	latest annual report of the laboratory.
	☐ Attached	□ Not Available

TITLE

Completed by \_\_\_\_

NAME

\_\_\_\_\_ Date \_\_\_

Lab Name	

## PART 2. PERSONNEL

1.	Laboratory staff. Complete Chart A for all technical personnel, including the laboratory director.
2.	Provide brief summary job description for each supervisory, professional, and technical position. If attached, please check.
3.	What is the total number of laboratory employees? Has this number increased over the past five years? Check if yes
4.	What portion of your staff participated in a formal training program related to improving work performance during the past year? Number %
5.	What was your turnover rate during the last 12 months?  a) Administrative Staff Number
6.	What portion of your staff was formally evaluated for performance during the past year?  Number %
7.	What portion of your staff received merit increases in grade or salary during the past year?  Number %
8.	What portion of your staff received service increases in grade or salary during the past year?  Number %
Con	npleted by Date
	NAME TITLE

Lab Name			
Lad Name		 	

## **CHART A**

Complete Chart A for all technical personnel, including the laboratory director. Use a separate block for each employee and arrange the presentation to reflect the lines of organizational responsibility.

					,	Date	No of	_ pages
	Training			Years of Experience			lentify Analyses Performed by	
Name	Degree (Circle One)	Major	Position	Present Job	Previous Jobs		umbers From Attached Index	
	Ph.D. MS BS Assoc. HS					, .	·	
	Ph. D. MS BS Assoc. HS		,			·		
	Ph. D. MS BS Assoc. HS		,					

Lab	Name	
Lab	iname	

## PART 3. LABORATORY SPACE AND FACILITIES

#### **CHART B**

Complete Chart B. Please indicate both the availability and the adequacy of laboratory equipment and facilities.

			Adeq	Additional	
ltem	Desc	ription	Yes	No	Information
Buildings in Use Total m <sup>2</sup> (Sq. Ft.)					<u> </u>
Office Space Total m <sup>2</sup> (Sq. Ft.)					· — — — — — — — — — — — — — — — — — — —
Lab Space Total m <sup>2</sup> (Sq. Ft.)					-
Bench-top Space Total m <sup>2</sup> (Sq. Ft.)					
Bench Hoods No Capacity (m/sec.) (lin. ft./min.)					
	Ava	ilable	Adeq	uate	Additional
	Yes	No	Yes	. No	Information
Storage Space Chemicals					•
Sample Storage - General					
Secured Space					
Refrigerated Space					
Hazardous Samples					
Controlled Space - Temperature					
Humidity					
Noise Insulation					•
Shielded .					
Clean Rooms	٠.				
Heat					
Air-Conditioning					
Electrical Services					· · · · · · · · · · · · · · · · · · ·
Gas			·		
Compressed Air					

ltem .	Avai	ilable	Adec	quate	Additional		
	Yes	No	Yes	No	Information		
Vacuum							
Safety Equipment - Fire Alarm							
Fire Extinquishing Equipment							
Emergency Showers							
Eye Fountains							
Personal Equipment: glasses, gloves							
Hazardous Area Escape							
Flammable Material Storage							
Safety Cans							
Ventilation							
Smoking Areas							
Handling Equipment for Acids,							
Caustic							
OSHA Signs							
Water Supply - Distilled							
Deionized							
Ammonia - free							
CO <sub>2</sub> - free							
Bacteriologically Suitable							
Glassware Supply							
Glassware Washing Equipment							
Disposal Equipment - Broken Glass							
Contaminated Material, Solvents							
Library							
Conference Room							
Employee Lounge							
Employee Lockers							
Drinking Fountains							
Lunch Room							
Data Processing Equipment							

Logistic Services - Telephone
Intercom
Emergency Line
Motor Vehicle
Facilities as a Whole

Avai	lable	Adeq	Juate	Additional		
Yes	No	No Yes		Informatio		
			<del></del>			
		i 		<del> </del>		
			<u> </u>	<del> </del>		
		L		<u> </u>		

Completed by	****	<del></del>	Date	
	MAME	TITLE		

#### PART 4. TECHNICAL SERVICES OFFERED

#### Instructions

In Chart C, Table of Analytical Methods, you are asked to indicate the tests which are performed by this laboratory and the specific method(s) which you use for each test. This may be done simply by circling the appropriate references under Method Used in This Laboratory. In cases where you follow an EPA method which refers to ASTM or Standard Methods for the detailed procedure, you may circle the EPA reference only.

The Standard Methods, ASTM, and EPA references are given for your convenience. Standard Methods refers to Standard Methods for the Examination of Water and Wastewater, 13th Edition, 1971, published jointly by the American Public Health Association, the American Water Works Association, and the Water Pollution Control Federation. ASTM refers to the Annual Book of ASTM Standards, Part 31, Water, 1974, published by the American Society for Testing and Materials. EPA refers to Methods for Chemical Analysis of Water and Wastes, 1974, published by the Environmental Monitoring and Support Laboratory (National Environmental Research Center, Cincinnati, Ohio) and the Office of Technology Transfer, U.S. Environmental Protection Agency or to the Federal Register (for air tests). References in Standard Methods and ASTM are to method numbers, whereas references in the EPA Manual are to page numbers in the 1974 edition.

If this laboratory uses an alternate method or a modification of a referenced method, write "Other" under "Method Used in This Laboratory" and provide the requested information for each such case on a copy of the form "Alternate Analytical Method", page 40.

Under "Sample Frequency," please enter, in the #/Month column, the average number of samples per month tested by the specified method over the last 12 months. In the Peak Load column, give the maximum number of samples analyzed in a one-month period during the last 12 months. Your best estimates of these numbers will be satisfactory.

The tests listed in Chart C are limited to those referenced in the Federal Register. Referenced in the Federal Register but not included in Chart C are the variations in air methods suitable for measurement of emissions from stationary sources. Refer to Federal Register Vol. 36, No. 247, Part II, December 23, 1971; Vol. 38, No. 111, June 11, 1973; Vol. 39, No. 47, March 8, 1974; Vol. 40, No. 152, August 6, 1975; and Vol. 80, No. 194, October 6, 1975. There are, however, important areas not yet covered by such references. Biology is one such area. The bibliography appended to this procedure lists some of the sources of information on missing tests. Method 406, Standard Plate Count, is found in Standard Methods (Ref. 4). Refer also to the so-called "Equivalency Document," Federal Register, February 18, 1975.

At the end of Chart C a blank chart is included, page 20, on which information may be supplied on important tests performed by the laboratory which are not included in the check list.

1.	Complete Chart C indicating analytical methodology which the laboratory wishes to have
2.	evaluated.  Provide a brief description of any special or unusual technical capability provided by the laboratory.
3.	Provide a brief description of methods that you use for pretreatment of samples before analysis for trace metals, Tests No. 16-43
_	

Lab Name

TITLE

Completed by

NAME

\_\_\_ Date \_

	1			Method Used in This Lab			1	nple
Test and Unit		Method	Circle Appropriate Reference <sup>1</sup> Check next col. if copies available in lab.			Copy Avail-	Frequency	
			Standard Method	ASTM	EPA	able	# / Month	Peak Load
General Analytical Tests:								
1. Alkalinity as CaCO <sub>3</sub>	(a)	Electrometric Titration, Manual	201	D1067-70B	p. 3			
(mg CaCO3/liter)	(b)	Electrometric Titration, Automated	201		p. 3	1		
<del></del>	(c)	Automated, Methyl Orange			p. 5			
2. Biochemical Oxygen	<u> </u>	Modified Winkler with Full-Bottle	219		p. 11, 51			
Demand (B.O.D.) 5- 20° C (mg/liter)	day (b)	Probe Method			p. 11, 56			
3. Chemical Oxygen De (C.O.D.) (mg/liter)	emand (a)	Dichromate Reflux (organic C > 15 mg/liter)	220	D1252-67	p. 20			
	(b)	Low Level Modification			p. 21			
	(c)	Saline Water Modification (C1 > 2000 mg/liter)			p. 25			
4. Total Solids (Total Residue) (mg/liter)	(a)	Gravimetric, Dried at 103-105° C	224A		p. 270			
5. Total Dissolved Solid (Total Filterable Res (mg/liter)	(-,	Glass Fiber Filtration, Dried at 180° C	224E		p. 266			

			Method Used in This Lab  Circle Appropriate Reference 1 Check next col. if copies available in lab.				Sam	•
	Test and Unit	Method				Copy Avail-	Frequency	
			Standard Method	ASTM	EPA	able	# / Month	Peak Load
6.	Total Suspended Solids (Total Nonfilterable Residue) (mg/liter)	(a) Glass Fiber Filtration, Dried at 103-105° C	224C		p. 268			
7.	Total Volatile Solids (Volatile Residue) (mg/liter)	(a) Gravimetric, Dried at 550° C	224B		p. 272			
8.	Ammonia (as N) (mg/liter)	<ul> <li>(a) Distillation and Titration</li> <li>(b) Distillation and Nesslerization</li> <li>(c) Distillation and Ammonia Electrode</li> <li>(d) Automated Colorimetric Phenate</li> <li>Method</li> </ul>			p. 159 p. 159 p. 159, 165 p. 168			
9.	Total Kjeldahl Nitrogen (as N) (mg/liter)	<ul> <li>(a) Digestion, Distillation &amp; Titration</li> <li>(b) Digestion, Distillation &amp; Nesslerization</li> <li>(c) Digestion, Distillation &amp; Ammonia Electrode</li> <li>(d) Automated Phenate Method</li> </ul>	216		p. 175-181 p. 175-181 p. 165, 175- 181 p. 182			
10	. Nitrate (as N) (mg/liter)	(a) Cadmium Reduction Method (Nitrate- Nitrite)	213B	D992-71	p. 201			

	Met		Sample Frequency			
Method	Circle Appropriate Reference <sup>1</sup> Check next col. if copies available in lab.				Avail-	
	Standard Method	Standard ASTA		able	# / Month	Peak Load
(b) Automated Cadmium Reduction  Method (Nitrate-Nitrite)			p. 207			
(c) Brucine Method			р. 197			
(d) Automated Hydrazine Reduction  Method			p. 185 <sup>2</sup>			
(a) Single Reagent (Ascorbic Acid Reduction Method)	223CIII 223F		p. 249			
(b) Automated Colorimetric Ascorbic Acid Reduction Method	-		p. 256			
(c) Automated SnCl <sub>2</sub> Method	223E					
(a) Hydrogen Peroxide Digestion & Electrometric Titration		D1067-70E	p. 1			
(b) Hydrogen Peroxide Digestion & Phenolphthalein End-Point Titration		D1067-70E				
(a) Combustion and Infrared Method	138A	D2579-74	p. 236			
(b) Combustion & Flame Ionization Method (CH <sub>4</sub> )			p. 236			
	(b) Automated Cadmium Reduction Method (Nitrate-Nitrite) (c) Brucine Method (d) Automated Hydrazine Reduction Method  (a) Single Reagent (Ascorbic Acid Reduction Method) (b) Automated Colorimetric Ascorbic Acid Reduction Method (c) Automated SnCl <sub>2</sub> Method  (a) Hydrogen Peroxide Digestion & Electrometric Titration (b) Hydrogen Peroxide Digestion & Phenolphthalein End-Point Titration  (a) Combustion and Infrared Method CO <sub>2</sub> (b) Combustion & Flame Ionization	Check next Standard Method  (b) Automated Cadmium Reduction Method (Nitrate-Nitrite)  (c) Brucine Method  (d) Automated Hydrazine Reduction Method  (a) Single Reagent (Ascorbic Acid Reduction Method)  (b) Automated Colorimetric Ascorbic Acid Reduction Method  (c) Automated SnCl <sub>2</sub> Method  223E  (a) Hydrogen Peroxide Digestion & Electrometric Titration  (b) Hydrogen Peroxide Digestion & Phenolphthalein End-Point Titration  (a) Combustion and Infrared Method CO <sub>2</sub> (b) Combustion & Flame Ionization	Check next col. if copies avail.  Standard Method  ASTM  (b) Automated Cadmium Reduction Method (Nitrate-Nitrite)  (c) Brucine Method  (d) Automated Hydrazine Reduction Method  (a) Single Reagent (Ascorbic Acid Reduction Method)  (b) Automated Colorimetric Ascorbic Acid Reduction Method  (c) Automated SnCl <sub>2</sub> Method  (d) Automated SnCl <sub>2</sub> Method  (e) Automated SnCl <sub>2</sub> Method  (f) Hydrogen Peroxide Digestion & D1067-70E Phenolphthalein End-Point Titration  (a) Combustion and Infrared Method CO <sub>2</sub> (b) Combustion & Flame Ionization	Check next col. if copies available in lab.  Standard Method  ASTM  EPA  (b) Automated Cadmium Reduction Method (Nitrate-Nitrite)  (c) Brucine Method  p. 197  (d) Automated Hydrazine Reduction Method  (a) Single Reagent (Ascorbic Acid Reduction Method)  (b) Automated Colorimetric Ascorbic Acid Reduction Method  (c) Automated SnCl <sub>2</sub> Method  223E  (a) Hydrogen Peroxide Digestion & Electrometric Titration  (b) Hydrogen Peroxide Digestion & Phenolphthalein End-Point Titration  (a) Combustion and Infrared Method CO <sub>2</sub> (b) Combustion & Flame Ionization  Check next col. if copies available in lab.  Standard Method  p. 207  223CIII p. 249  223F  p. 256  D1067-70E  D1067-70E  P. 1	Check next col. if copies available in lab.  Standard Method ASTM EPA  (b) Automated Cadmium Reduction Method (Nitrate-Nitrite) (c) Brucine Method p. 197 (d) Automated Hydrazine Reduction Method  (a) Single Reagent (Ascorbic Acid Reduction Method) (b) Automated Colorimetric Ascorbic Acid Reduction Method (c) Automated SnCl <sub>2</sub> Method  (a) Hydrogen Peroxide Digestion & Electrometric Titration  (b) Hydrogen Peroxide Digestion & Phenolphthalein End-Point Titration  (a) Combustion and Infrared Method CO <sub>2</sub> (b) Combustion & Flame Ionization  (b) Combustion & Flame Ionization  (c) Combustion and Infrared Method P. 236	Circle Appropriate Reference   Available in lab.   Available   Astandard   Method   Astandard   Asta

		Me	Сору	Sample			
Test and Unit	Method	Circle App Check nex	Circle Appropriate Reference 1 Check next col. if copies available in lab.				iency
		Standard Method	ASTM	EPA	able	# / Month	Peak Load
14. Total Hardness	(a) EDTA Titration	122	D1126-67B	p. 68			
(mg CaCO <sub>3</sub> /liter)	(b) Automated Colorimetric (c) Atomic Absorption (Ca + Mg)			p. 70 p. 78-91, 103, 114			
15. Nitrite (as N) (mg/liter)	(a) Manual Colorimetric Diazotization (b) Automated Colorimetric Diazotization			p. 215 p. 207*			
	-						

Lab Name: \_\_\_\_\_

		Me	Lab		Sample		
Test and Unit	Method	Circle App Check nex	ropriate Reference t col. if copies avai	1 lable in lab.	Copy Avail- able	Frequ	
		Standard Method	ASTM	EPA		# / Month	Peak Load
Tests for Trace Metals:							
16. Aluminum (mg/liter)	(a) Atomic Absorption	103A		р. 92 Ψ			
17. Antimony (mg/liter)	(a) Atomic Absorption			р. 94 Ψ			
18. Arsenic (mg/liter)	(a) Atomic Absorption (Gaseous Hydride Method)			р. 95 <b>Ψ</b>			
	(b) Gaseous Hydride - Silver Diethyl- dithiocarbamate Colorimetric	104A		p. 9			
19. Barium (mg/liter)	(a) Atomic Absorption	129A		р. 97 <b>Ψ</b>	<u> </u>		
20. Beryllium (mg/liter)	(a) Atomic Absorption (b) Aluminon Method	129A 106B		р. 99 Ψ			
21. Boron (mg/liter)	(a) Curcumin Method	107A		p. 13			
22. Cadmium (mg/liter)	(a) Atomic Absorption (b) Dithizone Colorimetric Method	129A 211(II)B	D2576-70	р. 101 Ф	_	-	
23. Calcium (mg/liter)	(a) Atomic Absorption		D2576-70	р. 103 <b>Ψ</b>			
	(b) EDTA Titration	110C		p. 19			
				1	1	1	ļ

		Met	ab		Sam	•	
Test and Unit	Method	Circle Appr Check next	ropriate Reference <sup>1</sup> col. if copies avail	able in lab.	Copy Avail-	Frequ	
		Standard ASTM		EPA able		# / Month	Peak Load
24. Chromium VI (mg/liter)	(a) Extraction and Atomic Absorption			р. 78-91, 105 <del>V</del>			
	(b) Diphenylcarbazide Colorimetric	211(II)D					
25. Chromium, Total (mg/liter)	(a) Atomic Absorption	129A	D2576-70	р. 78-91, 105 <del>V</del>			
	(b) Oxidation & Diphenylcarbazide Colorimetric	211(II)C	D1687-67	p. 105			
26. Cobalt (mg/liter)	(a) Atomic Absorption		D2576-70	р. 107 Ψ			
27. Copper (mg/liter)	(a) Atomic Absorption	129A	D2576-70	р. 108 Ψ	ļ		
	(b) Neocuproine Colorimetric	211(II)E	D1688-68		<u> </u>	ļ	<u> </u>
28. Iron (mg/liter)	(a) Atomic Absorption (b) O-Phenanthroline Colorimetric	129A 211(II)F	D2576-70F D1068-68A	р. 110 Ψ			
29. Lead (mg/liter)	(a) Atomic Absorption (b) Dithizone Colorimetric	129A 211(II)G	D2576-70G	р. 112 Ѱ			
30. Magnesium (mg/liter)	(a) Atomic Absorption	129A	D2567-70	р. 114 Ф			
	(b) Gravimetric	127A	D511-52		+	-	<b>├</b>
31. Manganese (mg/liter)	(a) Atomic Absorption	129A	D2567-70	р. 116 Ф	<u> </u>		

		Method Used in This Lab				Sample Frequency	
Test and Unit	Method	Circle Ap Check ne	propriate Reference kt col. if copies ava	e1 iilable in lab.	Copy Avail- able		
		Standard Method	ASTM	EPA		# / Month	Peak Load
32. Mercury (mg/liter)	(a) Flameless Atomic Absorption: Man- ual Cold Vapor Technique (Hg in Water)		D3223-73	р. 118 <b>Ψ</b>			
	(b) Flameless Atomic Absorption: Automated Cold Vapor Technique (Hg in Water) (not approved generally)			р. 127 <b>У</b>			
	(c) Flameless Atomic Absorption: Man- ual Cold Vapor Technique (Hg in Sediment)		D3223-73	р. 134 Ψ		!	
33. Molybdenum (mg/liter)	(a) Atomic Absorption			р. 139 <b>Ψ</b>			
34. Nickel (mg/liter)	(a) Atomic Absorption (b) Heptoxime Colorimetric	211(II)I	D2576-70	р. 141 <b>Ψ</b>			
35. Potassium (mg/liter)	(a) Atomic Absorption (b) Colorimetric (c) Flame Photometric	147B 147A	D1428-64	р. 143 Ψ			
36. Selenium (mg/liter)	(a) Atomic Absorption (Gaseous Hydride Method)			р. 145 <b>Ψ</b>			
37. Silver (mg/liter)	(a) Atomic Absorption	129A		р. 146 Ψ			

		Method Used in This Lab				Sam	•
Test and Unit	Method	Circle App Check nex	ropriate Referenc t col. if copies ava	e1 ilable in lab.	Copy Avail-		iency
		Standard Method	ASTM	EPA	able	# / Month	Peak Load
38. Sodium (mg/liter)	(a) Atomic Absorption			р. 147 <b>Ψ</b>			
	(b) Flame Photometric	153A	D1428-64				
39. Thallium (mg/liter)	(a) Atomic Absorption			р. 149 Ψ			
40. Tin (mg/liter)	(a) Atomic Absorption		<u> </u>	р. 150 Ψ			
41. Titanium (mg/liter)	(a) Atomic Absorption		ļ	р. 151 <b>Ψ</b>			
42. Vanadium (mg/liter)	(a) Atomic Absorption	1644		р. 153 <b>Ψ</b>			
	(b) Colorimetric (Catalysis of Gallic Acid Oxidation)	164A				ļ	<u> </u>
43. Zinc (mg/liter)	(a) Atomic Absorption	129A		р. 155 <b>Ψ</b>			
	(b) Dithizone Colorimetric Method	165B	D1691-67		<del> </del>	<b></b>	<u> </u>
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		Me	Method Used in This Lab  Circle Appropriate Reference 1  Check next col. if copies available in lab.			Sam	•
Test and Unit	Method	Circle App Check nex				Frequ	,
		Standard Method	ASTM	EPA	able	# / Month	Peak Load
Tests for Nutrients, Anions, and Organics							
44. Organic Nitrogen (as N) (mg/liter)	(a) Kjeldahl Nitrogen minus Ammonia Nitrogen	215		See (8) and (9) above			
45. Orthophosphate (as P) (mg/liter)	(a) Single Reagent Ascorbic Acid Reduction Method	223F	D515-72A	p. 249			
	(b) Automated Colorimetric Ascorbic Acid Reduction Method			p. 256			
46. Sulphate (as SO <sub>4</sub> )	(a) Gravimetric	156A	D516-68A	p. 283			
(mg/liter)	(b) Turbidimetric	156C	D516-68B	p. 277			
	(c) Automated Colorimetric Barium Chloranilate			p. 279			
47. Sulfide (as S) (mg/liter)	(a) Titrimetric Iodine	228A		p. 284			
48. Sulfite (as SO <sub>3</sub> ) (mg/liter)	(a) Titrimetric lodide-lodate	158	D1339-72C	p. 285			
49. Bromide (mg/liter)	(a) Titrimetric Iodide-Iodate		D1246-68C	р. 14			
50. Chloride (mg/liter)	(a) Silver Nitrate	112A	D512-67B				
	(b) Mercuric Nitrate	112B	D512-67A	p. 29			
	(c) Automated Colorimetric Ferricyanide	2		p. 31			

		Me	.ab		Sample		
Test and Unit	Method	Circle App Check nex	ropriate Reference t col. if copies avail	l able in lab.	Copy Avail- able	Frequ	
		Standard Method	ASTM	EPA		# / Month	Peak Load
51. Cyanide, Total (mg/liter)	(a) Distillation & Silver Nitrate Titration  (b) Distillation & Pyridine-Pyrazolone (or Pyridine - Barbituric Acid) Colorimetric	207A, 207B 207A, 207C	D2036-74A D2036-74A	p. 40 p. 40			
52. Fluoride (mg/liter)	(a) Distillation-SPADNS	121A, 121C	D1179-72A	p. 59			
	(b) Automated Complexone Method (c) Fluoride Electrode			p. 61 p. 65	Ī		
53. Chlorine, Total Residual (mg/liter)	(a) Starch-lodide Titration (b) Amperometric Titration	204A 204A	D1427-68A D1427-68B	p. 35			
54. Oil and Grease (mg/liter)	(a) Gravimetric (Separatory Funnel Extraction)	137		p. 229			
	(b) Infrared (Separatory Funnel Extraction)			p. 232			
55. Phenols (mg/liter)	(a) Colorimetric (4-AAP Method with Distillation)	222E	D1783-70	p. 241			

		Me	thod Used in This	Lab		Sam	
Test and Unit	Method	Circle App Check nex	ropriate Reference t col. if copies ava	e1 ilable in lab.	Copy Avail-	Frequ	· · · - · ·
		Standard Method	ASTM	EPA	able	# / Month	Peak Load
56. Surfactants (mg/liter)	(a) Methylene Blue Colorimetric	159A	D2330-68	p. 157			
57. Algicides (mg/liter)	(a) Gas Chromatography			§			
58. Benzidine (mg/liter)	(a) Diazotization & Colorimetric			+			
59. Chlorinated Organic Compounds (Except Pesticides) (mg/liter)	(a) Gas Chromatography			ş			
60. Pesticides (mg/liter)	(a) Gas Chromatography			5			
	(b) Thin Layer Chromatography			Ş			
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		Method Used in This Lab					nple Jency
Test and Unit	Method	Circle App Check nex	ropriate Reference t col. if copies avai	1 lable in lab.	Copy Avail-	<u> </u>	Г
		Standard Method	ASTM	EPA	able	# / Month	Peak Load
Physical and Biological Tests							
61. Color	(a) Platinum-cobalt Colorimetric (b) Spectrophotometric (Dominant wavelength, hue, luminance, purity)	118 206A		p. 36 p. 39			
62. Specific Conductance (mho/cm @ 25° C)	(a) Wheatstone Bridge	154	D1125-64	p. 275			
63. Turbidity (Jackson Units)	(a) Turbidimeter Method	163A	D1889-71 (Sect. 10- 16)	p. 295			
64. Streptococci Bacteria,	(a) MPN	409A					_
Fecal (number/100 ml)	(b) Membrane Filter	409B					
	(c) Plate Count	409C	<b> </b>			<b> </b>	ļ
65. Coliform Bacteria,	(a) MPN	407C					
Fecal (number/100 ml)	(b) Membrane Filter	408B					
66. Coliform Bacteria,	(a) MPN	407A					<u> </u>
Total (number/100 ml)	(b) Membrane Filter	408A	ļ <u>.                                    </u>			ļ	ļ
							<u> </u> 

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		Me	thod Used in This La	ab		Sam Frequ	•
Test and Unit	Method	Circle App Check nex	ropriate Reference <sup>1</sup> t col. if copies availa	ble in lab.	Copy Avail-	<del></del>	
		Standard Method	ASTM	EPA	able	# / Month	Peak Load
Radiological Tests							
67. Alpha, Total (pCi/liter)	(a) Proportional Counter	302	D1943-66				
	(b) Scintillation Counter		D1943-66		<del> </del>		
68. Alpha, Counting Error (pCi/liter)	(a) Proportional Counter (b) Scintillation Counter	302	D3085-72T		-		
69. Beta, Total (pCi/liter)	(a) Proportional Counter	302	D1890-66				
70. Beta Counting Error (pCi/liter)	(a) Proportional Counter	302	D3085-72T				
71. Radium, Total (pCi/liter)	(a) Proportional Counter	304	D2460-70				
	(b) Scintillation Counter	304,305	D2460-70				
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Lab Name:	 
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	Method Used in This Lab		Сору	Sample Frequency			
Test and Unit	Method	Circle App Check nex	Circle Appropriate Reference <sup>2</sup> Check next col. if copies available in lab.				
		Standard Method	ASTM	EPA	able	# / Month	Peak Load
Tests for Other Characteristics							
72. Temperature	(a) Thermometer or Thermistor	162		p. 286			
73. pH	(a) Electrometric	144A	D1293-65	p. 239			
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		Method Used in This Lab				Sample	
Test and Unit	Method	Circle Appropriate Reference Check next col. if copies available in lab.			Copy Avail-	Frequency # / Pea	
· ·		Standard ASTM		ЕРА	EPA able		Peak Load
Tests for Air Characteristics							
74. Sulphur Dioxide	(a) Pararosaniline Method { Manual Automated			b.22385-7 φ			
75. Suspended Particulates	(a) High Volume Method			b. 22388-90 φ	ļ		
76. Carbon Monoxide	(a) Nondispersive Infrared Spectrometry			b.22391 φ			
77. Photochemical Oxidants (Ozone)	(a) Chemiuminescence, Continuous			b.22392 φ			
78. Hydrocarbons (minus Methane)	(a) GC-FID			b.22394 φ			
79. Nitrogen Dioxide	(a) Arsenite 24-Hr Sampling Method  Manual Automated			b.15175 ▽ 22396 φ			
	(b) Chemluminescence, Continuous			b. 15177▽			

		Method Used in This Lab				Sam	iple
Test and Unit  Non-referenced Tests in Use	Method	Give Metho Check next	Copy Avail-	Frequency			
		Standard Method	ASTM	EPA	able	# / Month	Peak Load
Non-referenced Tests in Use							
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					<del> </del>	<del>                                     </del>	<del></del>
					<del> </del>	<u> </u>	
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#### REFERENCE MARKS IN CHART C

- 1 Federal Register, Vol. 35, No. 199, October 16, 1973.
- 2 EPA\_Methods\_Manual, 1971.
- 4 An introduction to atomic absorption spectrophotometry and a general procedure for trace metal analysis by atomic absorption is given in EPA Manual, pp. 78-91.
- 5 . Interim procedures for algicides, chlorinated organic compounds, and pesticides obtained from the Environmental Monitoring and Support Laboratory, USEPA, Cincinnati, Ohio 45268.
- Estimated by the method of M.A. El-Dib, "Colorimetric Determination of Aniline Derivatives in Natural Waters," <u>Journal of the Association of Official Analytical Chemists</u>, Vol. 54, No. 6, November, 1971, pp. 1383-1387.
- Federal Register, Vol. 38, No. 110, June 8, 1973.
- φ <u>Federal Register</u>, Vol. 36, No. 228, November 25, 1971.
- . Without Cd reduction.

# **ALTERNATE ANALYTICAL METHOD**

Nan	e of Laboratory	
(a)	# Test:	
	If this is a modification of a referenced method,	
	(1) Which referenced method (give manual name and pages)?	
	(2) Purpose of modification:	
	(3) Brief description of modification:	
	(4) Literature reference, if any:	
(c)	If this is an alternate method,	
	(1) Purpose of use of alternate method:	
	(2) Brief description of method:	
	(3) Literature reference, if any:	
(d)	Have you applied to EPA for approval of this procedure? (c.f., Federal Register, Vol. 38, No. 199, October 16, 1973, p. 28760)	

Lab Name	

## PART 5. ANALYTICAL INSTRUMENTS AND SPECIAL APPARATUS

(1)	Complete Chart D indicating analytical instruments and special apparatus available in the
	laboratory. See complete list of equipment, by analytical method, in the Appendix.

Completed by \_\_\_\_\_\_ Date \_\_\_\_\_

# INDEX OF ANALYTICAL INSTRUMENTS AND SPECIAL APPARATUS

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39.	Autoclave	58
40.	Light Microscope	58
41.	Misc. Microbiological Containers	59
42.	Membrane Filters	59
43.	Colony Counters	59
44.	Other Microbiological Instruments	60

### Special Radiological Equipment

45. 46. 47.	Alpha/Beta Particle Counters Spectrometer Systems Other Radiological Instruments and Apparatus	61 61 62
• • • • • • • • • • • • • • • • • • • •	Special Air Equipment	
48.	Sulfur Dioxide Monitoring	63
49.	High Volume Sampler (Particulates)	63
50.	Carbon Monoxide Monitor	63
51.	Total Hydrocarbons Monitor	64
52.	Photochemical Oxidants Monitor	64
53.	Nitrogen Dioxide Monitor	64
54.	Other Air Monitoring Equipment Including Calibration Equipment	65

### CHART D. ANALYTICAL INSTRUMENTS AND SPECIAL APPARATUS

Identify the instruments and apparatus in use and in good working condition in your laboratory.

	Instrument		Manu	facturer	Model	Year Purchased	Operating Manual Avail. in Lab.	
1.	Technicon Autoanalyzer							
	AAII Units							
	Samplers				<u> </u>			
	Manifolds for:			_				
	Alkalinity			Mercury	y (Cold '	Vapor Tech	nique)	
	Ammonia Nitrogen (Colorimetric Phenate)			Sulphur (Chloranilate)				
	Kjeldahl Nitrogen (Colorimetric Phenate)			Chloride (Ferricyanide)				
	Kjeldahl Nitrogen (Selenium Metho	od)		Fluoride (Complexone)				
	Nitrate-Nitrite (Cd Reduction)			Phenois (4-AAP)				
	Total Phosphorus or Orthophospha	ate		Others	(Specify	·):		
	Total Hardness							
	Analytical Cartridges for:							
	Ammonia Nitrogen (Colorimetric F	Phenate)		Nitrate	-Nitrite	(Cd Reduct	ion)	
	Total Phosphorus or Orthophospha	ate		Others	(Specify	/):		
				•				

Accessory		Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
Colorimeters					
Cells, tubular flow (Give number of	each ty	pe.)			
15 mm	,	, ,	Γ		50 mm
Filters:					
Wavelength of Max. Transmittance			ngth of ransmitta	ince	
			-		
			_		
		<u></u>	•		
Accessory	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
Recorders					
<del></del> 1					
Range Expansion			Digital P	rinter	
Associated Apparatus:					
Continuous Filter		Heati	ng Bath		
Proportioning Pump		45° -	80° C R	ange	
Planetary Pump		With	Distillati	on Coil & F	Head
Vapor-liquid Separator (for Hg Co	ld Vapo	r) With	Double	Delay Coil	
Continuous Digester			Tempera lation Co	iture with 2	?
Others (Specify):		<del></del>			
<del></del>					

Instrumen	t	# of Units	Manufacturer	Model	Year Purchased	Operating Manua Avail. in Lab.
2. Colorimeters/Filte Photometers	r					
Range:				ļ		
Range:						
Filters:						
Wavelength of Max. Transmittance	Bandwic	lth		ngth of ransmitta	nce	Bandwidth
			-			
				<del></del>		
pecial Associated Appar	atus (Specif	y):	<del></del>			
	<del></del>					
<del></del>					<u> </u>	
Instrumen	:	# of Units	Manufacturer	Model	Year Purchased	Operating Manua Avail. in Lab.
3. Spectrophotomete (UV - visible)	rs					
Recording (Range:	)			-		
Manual (Range:	)					
mandar (reange:						
pecial Attachments (Spe	cify):					<del></del>

Lamps			Fuels					
Metal	Hollow Cathode	Electric Discharge	Other (Specify)	Acetylene	Air	Nitrous Oxide	Argon	Hydrogen
Magnesium								
Manganese								,
Mercury (Cold Vapor)								
Molybdenum								
Nickel								
Potassium								
Selenium (Gaseous Hydride)					·			
Silver	, -							
Sodium								
Thallium	_							
Tin								
Titanium								
Vanadium								
Zinc								
Associated Equipment:								
Spectrophotometric gas cells, 10 cm, quartz windows (for Hg Cold Vapor)  Others (Specify):								

	Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
4.	Atomic Absorption Spectrophotometers					

Recorders (Specify):	<del></del>
	•
	<del></del>

Indicate lamps and fuels used for each metal:

		Lamps		Fuels				
Metal	Hollow Cathode	Electric Discharge	Other (Specify)	Acetylene	Air	Nitrous Oxide	Argon	Hydrogen
Aluminum								
Antimony								
Arsenic (Gaseous Hydride)								
Barium								
Beryllium								
Cadmium								
Calcium								
Chromium VI								
Chromium, Total								
Cobalt								
Copper								
Iron								
Lead								

	Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
5.	Mercury Analyzers			ĺ		
	Technique:	<u> </u>		<u> </u>		
	Range:	-		<del></del>		
	Sensitivity:		<del></del>	<u>.</u>		
		<u> </u>	<del></del>	<del>r -</del>	<del></del>	<del></del>
	Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
6.	Flame Photometers	Ġ				
}	Direct Reading:	<u> </u>				
	Internal Standard:					
	(Air blower & filter, Oxy-hydroger flame, polyethylene, or Teflon app		Others (Sp			
	Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
7.	Infrared Spectrophotometers					
	Single Beam (Range: )	ļ				
	Double Beam (Range: )				<u> </u>	
<u> </u>	Special Features:					
	IR Cells (Specify):					
ĺ						

	Instrument		# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
8.	Conductivity Meters						
	Field (Cell Type: )	<del></del> _	<del> </del>				
	Laboratory (Cell Type:	)	L	<u></u>	<u> </u>	<u></u>	<u> </u>
	Associated Apparatus (Spe	ecify): _					····
		.,				_	

	Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
9.	Electrometric Apparatus.					
	Electrometers: Field - ASTM Type I					
	ASTM Type II					
	Laboratory - ASTM I					

Electrodes:	Manufacturer	Туре
рН		
Dissolved Oxygen		
Ammonia		
Fluoride		
Cyanide		
Other (Specify):		

	Instr	ument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
10.	Automatic Tit	rimeters					
	Recorders (Spe	ecify):					
ļ		ed Electrodes					
	(Specify):						
					_		
	Instru	ument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
11.	Amperometric Apparatus	Titration					
	Instru	ıment	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
12.	Analytic Balan	ces					
	Capacity	Sensitivity					
	Certified	l Weights		Certification			
•							
	Instru	iment	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
13.	Carbon Analyz						
	Infrared (as CC	O <sub>2</sub> )	<del> </del>		<del> </del>		
	Flame Ionizati (as CH4)						

	Instrument	# Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
14.	Nephelometers/ Turbidimeters  Range:					
	Sensitivity Below 1 NTU:				<del></del> -	·
	Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
15.	Blenders					
	Instruments	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
16.	Vacuum Pumps					
	Type:					
l	Type:	<b>!</b>	<u></u>	l	<del></del>	
	Apparatus	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
17.	Magnetic Stirrers  With Heater					
	With Timer					
						·
18.	Drying Ovens					
	98° C		103°-105° C			] 180° C

		·
19.	Muffle Furnance 550° C	
20.	Hot Plate (persulphate digestion)  Autoclave (persulphate digestion)	
21.	Water Baths/Incubators  10°-15° C  100° C, well stirred, with Neoprene coated wire rack for 40-50 ml sample tubes (for Brucine Nitrate Method)	25° C with rack (for conductance measurements)  20° C incubator (for B.O.D.) with circulator
22.	B.O.D. Incubation Bottles	Number
23.	Gravimetric Evaporating/Weighing Dishes  Porcelain  Vycor  Platinum	Number

	Apparatus	Number
24.	Dessicators	
	Type:	
{	Type:	
	<u>.</u>	
	Apparatus	Number
25.	Kjeldahl Distillation Apparatus	
	Macro	
}	Micro	
	Apparatus	Number
26.	Arsine Generator & Absorption Apparatus	
		19,
	Apparatus	Number
27	Cyanide Distillation Apparatus	
. 41.	Cyamor Distillation Apparatus	
	Apparatus	Number
28.	Soxhlet Extraction Apparatus	
	Thimble Size:	
	Apparatus	Number
29	Phenol Distillation Setups	

	Apparatus	<del></del>	Number							
	···									
30.	Nessler Tubes, matched sets, APHA standard									
	50 ml, tall									
	100 ml, tall									
	Apparatus	Model	Temperature	Cubic Feet						
31.	Refrigerators									
				<del>-                                   </del>						
· · · · · · · · · · · · · · · · · · ·										
	Apparatus		Certification	Range						
32.	Special Thermometers									
	Apparatus									
33.	Thin-Layer Chromatography Apparatus (Describe chambers; plates, commercial or homemade; spray reagents and apparatus; spotting apparatus; special equipment)									
				<del></del>						
				<del></del>						
		<del></del>								

4.	Column Chromatography Apparatus (Describe columns; adsorbents - type, source, grade, special handling; solvent evaporation apparatus; special equipment)
	<del></del>
<u> </u>	Gas Chromatographs (Describe for each instrument: make and model; column type-capillary, 1/8 in., 1/4 in., etc., temperature programming; detector type and model; recorder; most commonly used columns; special equipment)
5.	capillary, 1/8 in., 1/4 in., etc., temperature programming; detector type and model; recorder; most commonly used columns; special equipment)
5.	capillary, 1/8 in., 1/4 in., etc., temperature programming; detector type and model; re-
5.	capillary, 1/8 in., 1/4 in., etc., temperature programming; detector type and model; recorder; most commonly used columns; special equipment)
35.	capillary, 1/8 in., 1/4 in., etc., temperature programming; detector type and model; recorder; most commonly used columns; special equipment)
35.	capillary, 1/8 in., 1/4 in., etc., temperature programming; detector type and model; recorder; most commonly used columns; special equipment)
35.	capillary, 1/8 in., 1/4 in., etc., temperature programming; detector type and model; recorder; most commonly used columns; special equipment)
35.	capillary, 1/8 in., 1/4 in., etc., temperature programming; detector type and model; recorder; most commonly used columns; special equipment)

	Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
36.	Other Special Instruments (Mass Spec., NMR, Flowmeters, Electrom Microscope, etc.)					
	<u> </u>		···		ļ	
		-	<u> </u>		<u> </u>	
		<del>                                     </del>				
					-	
	·		l		,	
		-				
•						

### CHART D. ANALYTICAL INSTRUMENTS AND SPECIAL APPARATUS

	Lab Name:		
· · · · · · · · · · · · · · · · · · ·			·
tive humidity	)		· · · · · · · · · · · · · · · · · · ·
44	4.5 ± 0.2° C		
# of Units	Manufacturer	Magnification	Light Source
	# of	tive humidity)  44.5 ± 0.2° C	# of Manufacturer Magnification

				Plastic		Gl	ass	Other (Specify)
41.	Miscellaneous Microbiological Con							
•	Inoculation Tubes							
	Dilution Bottles							
	Containers for Media		į					
	Petri Dishes							-
	Other (Specify)							-
•				Mani	ufacturer		-	Туре
42.	Membrane Filters							<u>V</u>
								<del></del>
<del></del>	Instrument	# of Units	Mar	nufacturer	Model	Year Purc	r chased	Operating Manual Avail. in Lab.
43.	Colony Counters							
					•			

	Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
44.	Other Microbiological/Biological Instrumentation	,				
			·			
				·		

### CHART D. ANALYTICAL INSTRUMENTS AND SPECIAL APPARATUS

				Lab Nai	me:	
Spec	ial Radiological Equipment				-, ·	
		# of			Year	Operating Manual
	Instrument	Units	Manufacturer	Model	Purchased	Avail. in Lab.
45.	Alpha & Beta Particle Counters Windowless Gas-Flow Proportional Counter					
	Thin Window Gas-Flow Proportional Counter					
	Alpha Scintillation Counter					
	Beta Scintillation Counter					
	Liquid Scintillation Counter					
			<u> </u>	<u></u>		<u> </u>
	Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
46.	Spectrometer Systems					
	Alpha Spectrometer (Surface Barrier Type)					
	Detector			ļ		
	Analyzer					
	Other Pertinent Information	n				

Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
46. Spectrometer Systems (Con't.)					
Gamma Spectrometer  Detector					
Analyzer					
Other Pertinent Information					

Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
47. Other Radiological Instrumentation (Radon Gas Counters, Survey Instruments, etc.)					
				<u> </u>	

## **CHART D. ANALYTICAL INSTRUMENTS AND SPECIAL APPARATUS**

	Lab Name:							
	Special Air Monitoring Equipment							
·								
	Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.		
48.	(Field Sampling/Lab Analysis)					·		
	Field Sampler	.44_			L	<u> </u>		
	Lab Analytical Method:	T				J		
<b>.</b>	(Field Sampling/Field Analysis)	, ,						
	Field Sampler/Analyzer	<u> </u>		<u> </u>	<u></u>	<u> </u> 		
	Analytical Method:							
					,			
	Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.		
40	Sugar and ad Dantisulator				-			
49.	Suspended Particulates (High Volume Sampler)	1		ļ.		İ		
	Filter Type				<u> </u>			
				•				
	Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.		
50.	Carbon Monoxide Monitor							

Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
51. Total Hydrocarbons (corrected for CH <sub>4</sub> ) Monitor					

Instrument		Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
52. Photochemical Oxidants (O <sub>3</sub> ) Monitor					

Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail. in Lab.
53. Nitrogen Dioxide Monitor (Field Sampling/ Lab Analysis)  Field Sampler					
Lab Analytical Method:			T	· · · · · · · · · · · · · · · · · · ·	
(Field Sampling/ Field Analysis)					
Field Sampler/Analyzer		<u></u>	<u> </u>		
Analytical Method:		<del>,.</del> .			, —- ····.

Instrument	# of Units	Manufacturer	Model	Year Purchased	Operating Manual Avail, in Lab.
54. Other Air Monitoring Equipment Including Calibration Equipment					
Permeation Tubes					
Standard Cylinders					
Gas Phase Titration-Commercial				. 1	
-Home Made					
Air Dilution Systems					
Variable Temperature Bath: 25° C ± 0.1° C					

#### INTERNAL AND EXTERNAL CONTROLS

The first three sections of this chart contain lists on which the availability of written operating procedures is to be checked. These parts cover instrument maintenance and calibration, all aspects of sampling, and the quality control program of the laboratory. You may be asked to show these documents to the evaluator during the onsite inspection and to discuss them with him.

This check list should not be looked upon as a demand for written procedures (for example, a Quality Control Program) in a particular standard format. The important thing is that the principal laboratory controls should be documented in a permanent way. Some procedures may be brief or may not include all of the items to be checked. In the list please check those items which you believe to be adequately documented. The onsite visit will provide an opportunity to discuss the completeness of the documentation with the evaluator.

Part 4 of this chart asks for information on participation in interlaboratory proficiency testing programs. Information is required on the test methods covered in any plan in which you have participated, the organization conducting the program and the date of the last check sample reported upon.

You will be rated on the extent of your participation in such programs. However, as of the present, the actual standing you have achieved in proficiency tests is not a part of the scoring system for this evaluation.


Available

# PART 6. INTERNAL AND EXTERNAL CONTROLS

#### CHART E

			Yes	No
1.	Con	trol of Analytical Methods and Instruments		-
	(1)	Written Instrument Maintenance and		
		Calibration Procedures and Log Books		
	(2)	Written Bench Operating Procedures		
2.	Cor	ntrol of Sampling and Sample Preservation		
		Written Sampling Procedures Covering:		
	, .	Sampling Plans and Sampling Equipment		
		Sample Collection and Preservation		· · · · · · · · · · · · · · · · · · ·
		Identification and Storage of Samples	<del></del>	
		Laboratory Handling of Samples		•
		(Request for analysis, sample		
		preparation, timely performance, etc.)		<del> </del>
	(2)	Written Description of the Chain of Custody		
		of Samples		
	(3)	Written Procedures for Field Measurement		
(		(Flow, critical tests: D.O., Residual C1,		
		etc.)	<del></del>	<u> </u>
	(4)			
		(Water supply, effluents, ambient air,		
		stacks, mobile vehicles, pesticides,		l
		radiation, etc. )	<del></del>	<del></del>
3.	Qua	lity Control		
	(1)	Written Quality Control Program Covering:		
		Quality Policy		
		Assignment of Responsibility		
		Training in Quality Control Methods		
		Control of Purchased Chemicals/Reagents	<del></del>	<del></del>
		Internal Field and Laboratory Checks:		ı
		Precision/Accuracy	<del></del>	
		Routine Duplicates, Spiked, and		i
		Standard Samples	<del></del>	<del></del>
		Statistical Methods, Including Control Charts and/or Computer Methods		
	(2)	Written Description of Lab Record System		
	(2)	(Data handling/calculations, data review, validation		
		and audit)		•
	(3)	Written Description of Lab Report Systems		
	(4)	If you have a Quality Control Manual,		
	· · /	please provide a copy. Attached		
		· · · · · · · · · · · · · · · · · · ·		

# 4. Inter-laboratory Proficiency Testing Programs

Test Method		Participated in Program of:			Date of Last Check	Within	
	EPA	CDC	State	USGS	Other (Specify)	Sample	Acceptabl Limits
		ļ					
	İ						
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		]			!		
	]						

Completed by: _		 	Date	
. , -	NAME	 TITLE		

# SECTION 5 EVALUATOR'S GUIDE

#### PART 1. GENERAL INFORMATION ABOUT THE LABORATORY

#### Appropriateness of Organization

<u>Intent</u>. To determine appropriateness of the organization to render the services offered by the laboratory. This protocol is primarily concerned with the laboratory's monitoring activities: analyses performed to determine compliance with laws and regulations. The organization should be suited to the media that the evaluation covers: air, water, pesticides, or radiation.

Request a short discussion of the organization as seen from management's viewpoint.

- Is the organization chart supplied with the preliminary information up-to-date? Does it agree with the actual organization?
- Do functions performed in the laboratory follow the organization chart exactly?
- Are problems handled strictly through chain of command or do sections of the laboratory interact to get timely solutions?
- Does the laboratory experience difficulty in meeting performance requirements?

#### Impairment of Functions

<u>Intent</u>. To determine whether management perceives problems that might lead to impairment of laboratory functions.

Request a brief oral description of any problems encountered in operating the laboratory. Ask specifically about the following:

• Does the laboratory have difficulties in obtaining a sufficient number of well qualified staff in all disciplines?

- Are the facilities, equipment and services adequate to perform the services offered in the media covered by the laboratory?
   Water? Air? Pesticides? Radiation?
- Does the laboratory have difficulties in getting adequate services from outside supporting organizations? Specifically, is it satisfied with the validity of sampling, performed for it by others? With testing? With calibration? Are reports from outside signed?
- Does the laboratory have any problems in budgeting for next year? Does it have separate budgets for routine operations and for equipment and apparatus? Who is responsible for preparing the different parts of the budget? Is there input from all levels of the organization?
- Does the laboratory have any problems in satisfying those who use its services?

#### Strength of Management

Intent. To discover something about the strength of management.

Request discussion on the following items.

- Does the laboratory experience difficulties in maintaining cooperation between different laboratory groups? Between supervisors and analysts?
- Does the laboratory have specific plans and procedures for rapid to-the-point internal communications?
- Does the laboratory prepare an annual plan for operation of the laboratory? A long-range plan? What is management's experience with performance according to plan?
- Does the laboratory have a policy manual? Does sufficient informal control exist to ensure that things that need to be done quickly get done, for example cross over lines of authority in the lab or change-orders for sample analyses?
- Refer to Preliminary Questionnaire, Part 1, Items 9 and 10, dealing with involvement in activities outside the laboratory. Is the level of involvement in these activities consistent with the expectations the user should have of the laboratory?

#### Objectivity of the Laboratory

<u>Intent</u>. To determine whether there are reasons for questioning or discounting the objectivity of the laboratory.

Request a brief discussion of the relationship of the laboratory with its own organization and with its customers. If laboratory is privately owned the enquiry should be deeper than for publicly controlled laboratories.

#### Inquire more about:

- Ownership
- Managerial structure and individuals in sensitive and controlling positions
- Any other affiliations of principal officers and directors and those in supervisory positions in laboratory
- Any chance of conflict of interest of individuals in management in laboratory work
- · Basis for funding other than fees for direct services performed.

#### Cooperation Obtained

<u>Intent</u>. To determine the degree of cooperation of the entire laboratory in the total evaluation procedure. If such cooperation is not evident, capability of management is questionable.

Request a brief oral description of how the preliminary questionnaire was handled. How were various sections distributed for completion? Who decided who would answer the different sections? How were personnel advised of the importance of cooperation in the evaluation? Try to discover:

- Reasons for not providing complete information
- Any plans for using results of evaluation for benefit of the laboratory.

#### PART 2. PERSONNEL

Before visit is made some scores can be assigned from data already received on Chart A, Preliminary Questionnaire, Section 4.

#### Supervisor Training

All personnel in supervisory positions should have a university degree. Under unusual circumstances experience in specific methodology used in environmental monitoring laboratories and experience in such a laboratory may be substituted; however, a non-degree person should have had 5-10 years of experience.

#### Supervisor Experience

Determine from questioning of each supervisor the pertinence of his experience to environmental monitoring problems, e.g.

- Laboratory and field experience
- Involvement in investigation of emergency episodes and enforcement actions
- Leadership in special studies
- Length of experience in operation of laboratory functions now engaged in.

#### Job Descriptions

Study job descriptions carefully to determine if indeed the jobs are carried out according to the description. Make notes about jobs beside each name on Chart A, Section 4, so that you can ascertain through conversation whether what he actually does compares closely with the position as described.

#### Training Program

Check information received on training programs. If no formal training program exists, more time will be required on the visit to determine what is done for training. Question individuals to determine if they have received any training since joining the laboratory. Ask how much time is devoted to training on starting employment at the laboratory and how it is continued.

#### Turnover Rate

The rate of turnover may serve as an unobtrusive measure of effective personnel management. A consistently high turnover rate may indicate operating problems which the management is not successfully

handling. However, this is not the whole story because turnover rate may be due to causes beyond the laboratory management's control. Discreet questions may be asked about employment problems both of supervision and of analysts. Do personnel ceilings or funding problems interfere with administration of a sound salary policy? Are sound hiring procedures used for obtaining new personnel? Is hiring according to the crony system? Do Civil Service regulations apply? Is there a functioning affirmative action program?

#### General Morale

This question is related to the general morale and well being of the people employed at the laboratory. Determine by questions if any positive stpes are taken by the laboratory management to indicate some concern for individuals.

- Does management urge that training programs are taken either on site or at a nearby school?
- Is there a definite program for advancement?
- Are careful records kept on advancement?
- What is done regarding health programs?

The morale question is wider than whether policies exist and records are kept. Are employee organizations or unions in existence? What percentage of employees belong to unions? Ask analysts as well as supervisors about the state of relations between workers and management. Is there an opportunity for input by laboratory personnel into technical and management concerns of the laboratory?

#### PART 3. LABORATORY SPACE AND FACILITIES

Refer to Chart B, Preliminary Questionnaire, Section 4.

#### General Characteristics

The location of the laboratory, proximity to public transportation, its outside appearance, and a walk through of the building should help the evaluator to determine whether it is generally acceptable as an environmental monitoring laboratory.

Although many general features of the laboratory may have been checked in Chart B of the Preliminary Questionnaire, some discreet questions of laboratory personnel may be helpful.

- Is the location such that housing is available to the staff without excessive travel?
- Are there public eating facilities nearby, available for the entire staff?
- Is the neighborhood one that would cause no worry to any staff member who worked late?

Consider whether general support facilities are appropriate to the size and nature of the laboratory, i.e., secretarial and technician support, duplication facilities, photographic facilities, machine shop, electrical/electronics shop, glass blowing, etc.

Observe the adequacy of the visitor reception area, conference room, employee lounge or lunch room, locker space, drinking fountains, heating and air conditioning, service for electricity (Voltage stable?), gas, compressed air, and vacuum, etc. (filters installed?) Use Chart B as a guide, if desired.

#### Office Space

How does the square foot of office space per person compare to the adopted standard of  $16.7 \text{ m}^2$  (180 sq. ft)?

#### Laboratory Space

How does the square foot of laboratory space per person compare to the adopted standard of  $18.5 \text{ m}^2$  (200 sq. ft)?

#### Bench Top Space

How does the length of bench top per person compare to the adopted standard of 1.2m (4 ft)?

#### Hood Space and Operation

Examine hoods to make certain they operate properly. Ask, if in the opinion of the lab staff they are sufficient both in space and exhaust capability. Are records kept showing hood monitoring with velometer, last cleaning of ducts, general condition of glass, services, etc.? Filters last changed? Adopted standard 0.5 m/s (100 ft./min.).

#### Storage Space for Chemicals, Reagents, Glassware, and Supplies

The laboratory should have separate storage spaces for general chemicals, volatile chemicals and solvents, reagents, glassware and general supplies.

Closed cabinets should be used to keep bottles, glassware etc., free of dust and contamination from fumes.

Storage of volatile chemicals should meet OSHA standards: closed metal cabinet, under negative pressure, and away from flame/heat or sparks. (This may be storage under hood if under constant negative pressure).

No more than one liter each of volatile chemicals and solvents should be stored in the laboratory area. Larger amounts should be held in a separate storage facility away from the laboratory.

Use of carcinogenic/mutagenic chemicals should be kept to a minimum. If used these should be stored, handled and weighed in a glove box under constant negative pressure. Wherever possible, substitute chemicals or procedures should be used.

There should be sufficient in-lab storage available to permit the clearing of bench tops between test series. This is important for assurance of good control over procedures and for safety of the worker.

Storage areas should be inspected and corrected for overcrowding, breakage, outdated chemicals and general condition as a part of the routine lab clean-up.

#### Sample Storage

It is necessary not only that there be sufficient, accessible, well arranged storage space for general samples but also that provisions be made for special requirements of some samples, such as secured areas, refrigerated areas, and facilities for isolated storage of contaminated samples.

#### Controlled Space

The need for temperature and humidity control, for noise or electrical shielding and for clean rooms will depend on the media

handled by the laboratory. Using answers given in preliminary questionnaire, Chart B, question staff about requirement if space is not available.

#### Safety Equipment/Procedures

An opportunity was given in the preliminary questionnaire to check availability of specific items of equipment. Check the condition of this equipment.

In addition, observe, or ask questions about safety related matters. Examples:

- Eye protections, respiratory protection, floors not slippery, trash cans adequate and emptied regularly, first aid kit available?
- Does lighting in the laboratory meet standard of 100 ft. candles at bench top? If possible, carry a light meter to place on benches and desks to actually measure amount of light.
- Are fire prevention regulations posted? Smoking rules?
- Is area use clearly marked?
- Is the fire alarm clearly audible?
- Are exits marked and illuminated?
- Are fire extinguishers conspicuously located and in working order? Inspected last?
- Are emergency telephone numbers posted?

Fire	Medical
LILE	rieutcat

- Are regular fire drills conducted? When? Has local fire department ever visited laboratory? When?
- Does the laboratory give the appearance of having a constant awareness of the importance of safety?

#### Distilled Water/Deionized Water

Determine who is responsible for the stills which supply distilled water. Is there a central supply of deionized water? Are checking procedures written and a record maintained?

#### Glassware Supply and Washing

Is a sufficient supply of all the necessary types of glassware available? Is the glassware washing area convenient to work areas served? Is sufficient space provided for washing and drying? Are water supply, drains, drying ovens (165°C) and racks adequate? Are there written procedures for handling special glassware? Are contaminated containers sterilized or disinfected prior to washing? Are water spots present on recently washed ware? Are items tested for detergent removal (by appropriate indicator)? Is rinse water supply adequate? Are chipped or scratched items discarded? Are pipettes stored in aluminum or stainless steel (not copper) cans?

#### Housekeeping

Are passageways kept clear? Are broken glass and contaminated materials properly collected and disposed of? Are floors clean and well maintained? Are rooms and benches clean and uncluttered?

#### Data Processing Equipment and Logistic Services

Does laboratory have its own data processing facilities or access to a shared system? Is telephone service adequate? Is there an intercom system? Is there an emergency outside line? Is there a motor vehicle pool of any sort?

#### PART, 4. ANALYTICAL METHODS

Refer to Chart C, Preliminary Questionnaire, Section 4.

<u>Intent</u>. The intent of the part is to determine actual laboratory practices in the conduct of tests. Discuss each test for which the laboratory is being evaluated with individual "bench analysts", or their immediate supervisors.

- Are copies of the correct methods readily available to analysts?
- Do the analysts follow the methods exactly?
- Does the laboratory require adherence to a specific control program for each sampling procedure and analytical test? (Note: specific questions about sampling, calibration, and laboratory quality control procedures are asked in Part 6, Section 6.)

#### Reference Methods or Approved Alternates

All methods must be Federal Register referenced methods or specific EPA approval must have been obtained for modifications or alternates.

#### Reagent and Media Preparation

<u>Intent</u>. To assess the care taken in preparation, use, and storage of reagents and microbiological culture media.

#### Suggested Approach

The evaluator should inspect reagent bottles and media containers for clear and complete labelling, including date of preparation (or reference to a log containing dates of preparation). Containers should be appropriate for the particular reagent or medium and should be stored under appropriate conditions (temperature, light, etc.). Questions might include:

- Is there a written schedule for preparation of fresh reagents and media? (Some must be prepared on the day of the analysis, while others may be kept for extended periods under proper conditions.)
- Are new reagent batches always checked immediately against reference standards?
- Is a record kept of reagent batches and dates used?

- Is responsibility clearly assigned for preparing and maintaining fresh supplies of reagents and media?
- Are reagents rechecked at intervals against standards for possible contamination or degradation?

#### PART 5. FORMS FOR ONSITE EVALUATION

The following pages contain questions that may be asked about performance of specific tests, arranged by media.

In putting this material together, we have drawn on many sources, some of which, such as the EPA form for the bacteriological survey for water laboratories, have been used successfully in practice for some time. Other parts are drawn from recent USEPA or State EPA experience wherever the material appears to have been assembled in form most closely fitting the purpose of this procedure. Although, in the following outline of this material, we have indicated the primary source from which we obtained the material, we realize that many people may have been involved and we acknowledge our debt to the many individuals and groups with whom we have held discussions during the course of this task.

Part A. Medium- Water (Chemistry). Illinois EPA, Springfield Laboratory.

Part B. Medium- Water (Bacteriology). USEPA, Water Quality Office, Water Hygiene Division, Cincinnati.

Part C. Medium- Water (Biology). USEPA Environmental Monitoring Laboratory, Cincinnati.

Part D. Medium- Air. USEPA Environmental Monitoring Laboratory, Research Triangle Park.

Part E. Medium- Pesticides. USEPA Environmental Monitoring Laboratory, Research Triangle Park

Part F. Medium- Radiation. USEPA Environmental Monitoring Laboratory, Las Vegas.

Some of the material available to us was in rough draft form and, as updated versions become available, it may be desirable to include them in this Evaluator's Guide.

# PART A. MEDIUM - WATER

# (CHEMISTRY)

Type of samples	
Surface or ground water	
Industrial waste	· ·
Domestic mixed sewage	
Marine or estuary water	
Sediment, sludge, or semi-solid	<del></del>
Equipment-Analytical balance-	
Annual service, documented	
Certified weights available	. ——
Monthly check with certified weights, documented	
Autoclave	
Checked yearly by manufacturer	•
with maximum registering thermo- meter	
Safety valve works	
Operating instructions posted or available	
Deionizer	r
Million ohm water checked daily and documented	
KMn0 <sub>4</sub> 60 min. color retention check- ed daily and documented	
Still	
Checked daily and quality documented	
Operating instructions posted or available	<del></del>

Distilled water	
Checked for copper, ammonia,	
and chlorine documented	
Conductivity bridge	
Checked daily, documented	
Double deionized water	
Available for trace anal-	
ysis	<del></del>
pH meter	
Standardized for each use with buffer,	
documented	
Checked daily against second buffer	
for linearity, documented	
Flouride electrode	
standardized with each use,	
documented	<del></del>
Colorimeter	
Calibration curves checked with at	
least one standard each time used	· · · · · · · · · · · · · · · · · · ·
Drying ovens	
Temperature checked daily and re-	
corded	
record indicates satisfactory oper-	
ation and temperature controller	
functioning correctly	<del></del>
Muffle furnace pyrometer	
Pan balances	
Clean and in servicable condition	

Checked each month with two anal-	
ytical balance weights	
Automated analyzers	
Standard and blanks run	
each time	
Test frequency allows instru-	
ment to return to baseline	
between tests	
Record maintained of readings	
of standards for each test	
each time instrument is oper-	
ated	
Maintenance schedule followed	
for pump tube replacement,	
colorimeter cell cleaning,	
etc.	
Incubator BOD	
Thermometer calibrated, Documented	<del></del>
Daily record	
Uniformity of temperature check,	
documented	
Certified thermometer	
Certification on file	
Record of thermometer	
checks	
Pipette containers-	
Alumimum or stainless steel,	
no copper	

Dry heat sterilizer	
Temperature documented with	
recorder, charts filed	
accuracy of recorder checked	
Microscope	
Binocular wide field	
Fluorescent light source	<del>-</del>

#### **CHEMISTRY METHODS**

BOD

	Dilution water depletion on 5 days not more than 0.2 mg/l	
	If D.O. probe is used, calibration documented	
	If D.O. probe is used, correlation with Winkler method documented	
	Water seals on bottles protected	
	Dilutions for calculation are in the range which shows depletion of at least 2 mg/l and residual D.O. of l mg/l	
	Supersaturated samples deaerated before setting up	
	Chlorinated effluents checked for residual chlorine	
	Incubator temperature 20 ± 1° C documented	
	Method checked periodically by running glucose-glutamic acid standard	
	Seeding used when required On chlorinated effluents	
	On other sterile samples	_
	Sodium thiosulfate stock preserved, standardized, refrigerated, and documented	
	Samples refrigerated at 4°C immediately at point of collection and delivered to lab within 12 hours	
	Analyst does not pipet samples by mouth  Samples pipetted by	
COD - dichr	omate reflux method	
	Samples preserved by acidification, refrigeration, or both	_
	Silver sulfate catalyst used	
	Mercuric sulfate used to depress chloride interference	_
	Standardization of dichromate documented	
	Use boiling chips for smooth boiling	_

	Daily standardization of ferrous ammonium sulfate documented	
	Sample, reagents, and sulfuric acid mixed thoroughly before heat is applied	
	Analyst uses safety glasses or eye protection	
	Analyst does not pipet sample or reagents by mouth	
	Wastes properly disposed of	<del></del>
pH - elec	trometric method	
	Instrument manufacturer's instructions available and followed	
	Instrument checked for linearity with two buffers, documented	
	Instrument standardized daily, documented	
	Calomel electrode - liquid junction functioning	
	Calomel electrode - contains at least a crystal of KC1 but not solid with KC1	
	Electrodes rinsed between samples with distilled water and/or sample to be measured	
	Measurements made on successive portions of sample until two successive portions give equal readings	
	Sample temperature compensation applied	
	Solution pressure inside the calomel liquid junction in excess of that outside the junction	
	Immersible tips of electrodes stored in reagent water between periods of use	
	Sample agitated while making measurement	

# PART B. MEDIUM - WATER

(Bacteriological Examination)

#### **ENVIRONMENTAL PROTECTION AGENCY**

Water Quality Office Water Hygiene Division

Bacteriological Survey for Water Laboratories

Indicating conformity with the 13th edition of "Standard Methods for the Examination of Water and Wastewater," 1971.

Surv	Survey By $X = Deviation$ $U = Under$		U = Undetermin			
O = Not Used		sed				
Labo	oratory	Location	Date			
	Sampling a	nd Monitoring Response	<u>.</u>			
1.	Location and Frequency					
•	Representative points on system					
	Frequency of sampling adequate					
2.	Collection Procedure		,			
	Faucets with aerators should not be u	ısed	<u></u>			
	Flush tap 1 min. prior to sampling.					
	Pump well 1 min. to waste prior to sa					
	River, stream, lake, or reservior samp	. •				
	6 inches below surface and toward					
	Minimum sample not less than 100 m					
	Ample air space in bottle for mixing					
	Promptly identify sample legibly and					
3.	Sample Bottles	•				
	Wide mouth, glass or plastic bottles o	f capacity				
	Sample bottles capable of sterilization					
	Closure:					
	a. Glass stoppered bottles protect	cted with metal foil,				
	rubberized cloth or kraft type	paper				
	b. Metal or plastic screw cap wit	h leakproof liner				
	Sodium thiosulfate added for dechlor	rination				
	Concentration of 100 mg/l added	before sterilization	· · · · <u></u>			
	Chelation agent for stream samples (c	optional)				
	Concentration 372 mg/l added bef	ore sterilization				
4.	Transportation and Storage					
	Complete and accurate data accompa	nies sample	· · · · <u></u>			
	Transit time for potable water sample					
	48 hrs, preferably within 30 hrs .		· · · · · ———			
	Transit time for source waters, reserve					
	bathing waters should not exceed (					
	All samples examined within 2 hrs of					
	Sample refrigeration mandatory on st					
	optional on potable water samples					
<b>5</b> .	Record of Laboratory Examination					
	Results assembled and available for in					
	Number of tests per year		· · · · <del></del>			

Labo	pratory	Location		Date
 5.	Record of Laboratory Examination (	 Continued)		L
-				
	Confirmed (+) (-)	(Total)		
	Completed (+) (-)	(Total)		
	MF Test - Type of sample	(''ota')		
	Direct Count (+) (-)	(Total)		
	Verified Count (+) (-)	(Total)		
			ions	
	• •	•		•
	·			•
	Prompt resampling for unsatisfactory	samples		•
6.	Laboratory Evaluation Service			•
٠.		ories which examine		
				_
	Total labs known to ex	zamine water		·
				•
	provisionar labor	atories	· .	
	Labo	oratory Apparatus		
7.	Incubator			
	Sufficient size for daily work load .			•
				•
		•		
		_		
		ype of sample  (+)		
	•	•		
_				
8.	Incubator Room (Optional) Manufac			
	Daily record of temperature at selected	ed areas or use		
		0.5° C changes		·
9.	Water Bath			
	Manufacturer	Model		<del>_</del>
	Sufficient size for fecal coliform tests			
	Maintain uniform temperature 44.5°	$C \pm 0.2^{\circ} C$		
	Daily record of temperature or use of			
	thermometer sensitive to 0.2° C ch	ianges		

Labo	ratory	Location		Date
10.	Hot Air Sterilizing Oven Manufacturer	· Madal		
	Size sufficient to prevent crowding			
	Constructed to insure a stable steril			
	Equipped with accurate thermomet			•
	or with recording thermometer			
11.	Autoclave			<u> </u>
	Manufacturer	Model		_
	Size sufficient to prevent crowding	of interior		
	Constructed to provide uniform ten including 121° C			·
	Equipped with accurate thermomet	er with bulb properly located	t	
	to register minimal temperature			
	Pressure gage and operational safety			•
	Steam source from saturated steam			
	electrically heated steam generate Reach sterilization temperature in 3			
	Pressure cooker may be used only i			•
	gage and thermometer with bulb	•		
12.	Thermometers	inch above water level .		•
	Accuracy checked with thermomet	er certified by National		
	Bureau of Standards or one of ed			
	Liquid column free of discontinuou			
	marks legible		,	
13.	pH Meter		•	
	Manufactuer	Model		<del>_</del>
	Electronic pH meter accurate to 0.1	l pH units		
14.	Balance			
	Balance with 2 g sensitivity at 150 g			
	media preparations, Type	initial at 10 plant and		•
	Analytical balance with 1 mg sensit for weighing quantities less than			
	Appropriate weights of good qualit			•
15.	Microscope and Lamp	y for cacif balance		•
13.	Preferably binocular wide field, 10	to 15 diameters magnifi-		
	cation for MF colony counts, Ty			
	Fluorescent light source for sheen of			
16.	Colony Count			
	Quebec colony counter, dark-field	model preferred for		
	standard plate counts			
17.	Inoculating Equipment			
	Wire loop of 22 or 24 gauge chrome iridium, sterilized by flame			
	Single-service transfer loops of alun			
	sterilized by dry heat or steam.			
	Disposable single service hardwood			
4.0	sterilized by dry heat only			
18.	Membrane Filtration Units	Toma		
	Manufacturer			
	Leakproof during filtration			
	micial plating not worn to expose of	43C IIICLAI		

Labo	ratory	Location	Date
<del></del> 19.	Membrane Filters	<u> </u>	
	Manufacturer	Type	
	Full bacterial retention, satisfactory fi		
	Stable in use, glycerin free		· ·
	Grid marked with non-toxic ink		
	Presterilized or autoclaved 121° C for	10 min	· ·
20.	Absorbent Pads		
	Manufacturer	Type	
	Filter paper free from growth inhibito		
	Thickness uniform to permit 1.8 - 2.2	ml medium absorption	<u> </u>
21.	Presterilized or autoclaved with memb Forceps	rane filters	• •
	Preferably round tip without corrugat	ions	
	Forceps are alcohol flamed for use in l		
	. c. cops are allegated that the assetting	procedure	• •
	Glassware, Meta	Utensils and Plastic Items	
22.	Media Preparation Utensils		
	Borosilicate glass		· ·
	Stainless steel		
	Utensils clean and free from foreign re	sidues or	
	dried medium		
<b>23</b> .	Pipettes		
	Brand	Туре	<del></del>
	Calibration error not exceeding 2.5%		
	Tips unbroken, graduation distinctly r		
	Deliver accurately and quickly		
	Mouth end plugged with cotton (option	onal)	· · ·
24.	Pipette Containers		
	Box, aluminum or stainless steel		<del></del>
05	Paper wrapping of good quality sulfite	paper (optional)	• •
25.	Petri Dishes	_	
	Brand	Type	<del></del>
	Use 100 mm x 15 mm dishes for pour	plates . · , , , ,	• •
	Use 60 mm x 15 mm dishes for MF cu		
	Clear, flat bottom, free from bubbles a Plastic dishes may be reused if sterilize		• •
	30 min. or by ultraviolet radiation		
26.	Petri Dish Containers		• •
20.	Aluminum or stainless steel cans with	sovers coarsely woven	
	wire baskets, char-resistant paper sa		
<b>27</b> .	Culture Tubes	cks of wrappings	• •
21.	Size sufficient for total volume of med	fium and sample portions	
	Borosilicate glass or other corrosive res		
28.	Dilution Bottles or Tubes	515tant 81455	• •
<b>_</b> U.	Borosilicate or other corrosive resistan	t glass	
	Screw cap with leakproof liner free from	-	• • —
	on sterilization		
	Graduation level indelibly marked on		

Labo	ratory	Location	Date
	Materials ar	nd Media Preparation	I
29.	Cleaning Glassware		
	Dishwasher manufacturer	Model	
	Thoroughly washed in detergent at 16	0° F, cycle time	· ·
	Rinse in clean water at 180° F, cycle t	ime	· ·
	Final rinse in distilled water, cycle tim		
	Detergent brand	<u> </u>	· ·
	Washing procedure leaves no toxic resi		
30.	Glassware free from acidity or alkalini Sterilization of Materials	ty	
50.	Dry heat sterilization (1 hr at 170° C)		
	Glassware not in metal containers		
	Dry heat sterilization (2 hrs at 170° C)		· ·
	Glassware in metal containers		
	Glass sample bottles		· · · · · · · · · · · · · · · · · · ·
	Autoclaving at 121° C for 15 min		
	Plastic sample bottles		
	Dilution water blanks		
31.	Laboratory Water Quality		-
	Still manufacturer	Construction material	
	Demineralizer with		
	Protected storage tank		
	Supply adequate for all laboratory nee	ds	
	Free from traces of dissolved metals or	chlorine	· ·
	Free from bactericidal compounds as r	neasured	
	by bacteriological suitability test.		
	Bacteriological quality of water measu		
	by suitability test or sooner if neces	sary	
<b>32</b> .	Buffered Dilution Water		
	Stock phosphate buffer solution pH 7.		
	Prepare fresh stock buffer when turbid	lity appears	
	Stock buffer autoclaved and stored at	5 - 10° C	· ·
	1.25 ml stock buffer per 1 liter distille		
	Dispense to give $99 \pm 2$ ml or $9 \pm 0.2$	ml after autoclaving	· · ·
33.	pH Measurements		
	Calibrate pH meter against appropriate		
	Standard buffer brand		<del></del>
	Check the pH of each sterile medium to from each new medium lot number		
	Maintain a pH record of each sterile me		· · · · · ·
	the date and lot number		
34.	Sterilization of Media		
-	Carbohydrate medium sterilized 121°	C for 12 min	
	All other media autoclaved 121° C for	15 min	
	Tubes packed loosely in baskets for un		
	Timing starts when autoclave reaches 1		
	Total exposure of carbohydrate media		
	Media removed and cooled as soon as a		

Labo	ratory	Location	Date
35.	Storage	· · · · · · · · · · · · · · · · · · ·	
	Dehydrated media bottles kept tightly	clósed and stored	
	Dehydrated media not used if discolor	ed or caked	· · <del></del>
	Sterile culture media stored in clean ar		· · <del></del>
	contamination and excessive evapor	ation	· · ·
	Sterile batches used in less than 1 weel	<	· · <u> </u>
	All media protected from sunlight .		
	It media is stored at low temperatures,	, it must be incubated	
	overnight and any tubes with air bu	bbles discarded	• •
	Culture M	ledia - Specifications	•
36.	Lactose Broth		
	Manufacturer		
	Single strength composition 13 g per li Single strength pH 6.9 ± 0.1, double st		
	Not less than 10 ml medium per tube		
	Composition of medium after 10 ml sa		
37.	Lauryl Tryptose Broth		•
	Manufacturer	Lot No	<u></u>
	Single strength composition 35.6 g per		
	Single strength pH 6.8 ± 0.1, double st		
	Not less than 10 ml medium per tube		· ·
	Composition of medium after 10 ml sa	ample is added must	
	contain 0.0356 g per ml of dry ingre	edients	• •
38.	Brilliant Green Lactose Bile Broth	Lat Na	
	Manufacturer		
	Not less than 10 ml medium per tube		
39.	Eosin Methylene Blue Agar		
	Manufacturer	Lot No	
	Medium contains no sucrose, Cat. No.		
	Correct composition, sterility and pH		· · · <del></del>
40.	Plate Count Agar (Tryptose Glucose Y		
	Manufacturer		
	Correct composition, sterility and pH		
	Free from precipitate		
41.	EC Medium	time arter stermzation	· · ·
• • • •	Manufacturer	Lot No.	
	Correct composition, sterility and pH	6.9	
	Not less than 10 ml medium per tube		
<b>42</b> .	M-Endo Medium		
	Manufacturer		
	Correct composition and pH 7.1 - 7.3		
	Reconstituted in distilled water contai		
	Heat to boiling point, promptly remove Store in dark at 2 - 10° C		
	Unused medium discarded after 96 hrs		
	Chasea mediam discarded after 50 ms	· • • • • • • • • • • • • • • • • • • •	

Labo	ratory	Location	Date
—— 43.	M-FC Broth		
		Lot No	
	Reconstituted in 100 ml distilled was		· <u>———</u>
	Stock solution of rosolic acid discard		
		brown	
		ove and cool	
		rs	
44.	<u> </u>		
		Lot. No	_
	Correct composition and pH		
<b>45</b> .	<del></del>	Agar	
	Manufacturer	Lot No	
	Correct composition and pH		
	Multipl	e Tube Coliform Test	
46.	Presumptive Procedure		
	Lactose broth	lauryl tryptose broth	
	Shake sample vigorously		
		either 10 or 100 ml	
		ns	
		4 ± 2 hrs	
		bble positive	
	Examine for gas at 48 ± 3 hr from or	riginal incubation	
<b>47</b> .	Confirmed Test	-	
	Promptly submit all presumptive tub		
	before or at 24 hr and 48 hr period	ds to confirmed test	
	a. Brilliant green lactose broth		
		be or mix by rotating	
		ive broth or one dip of applicator	
		brilliant green lactose broth	
		check at 24 hrs for gas production	
	Reincubate negative tubes fo		
	and check for gas producti		
	, ,	itive tube results	
		e agar plates adequate streaking	
		separated by 0.5 cm	
		24 ± 2 hrs	
	* *	ith or without sheen are coliforms	
	If atypical unnucleated pink		
	· ·	est must be applied	
	If no colonies or only colorle		
	confirmed test is negative		
48.	Completed Test		
	Applied to all potable water samples		
	months to establish the validity of		
	determining their sanitary quality		

Labo	ratory	Location	Date			
54.	Standard Plate Count (Continued)	<u> </u>	<del> </del>			
J-1.	Melted medium stored for no more than 3 hrs at 43 - 45° C					
	Liquid agar and sample portion thoroughly mixed by gently					
	Count only plates with between 30					
	• •	30 colonies	· · ·			
	Record only two significant figures					
		sample"	<u></u>			
<b>55</b> .	Fecal Coliform Test	•	,			
	a. Multiple Tube Procedure					
	Applied as an EC broth conf	irmation of all positive				
		· · · · · · · · · · · · · · · · · · ·	, , , <u></u>			
	Place EC tubes in water bath	within 30 min. of transfers	· · ·			
	Incubate at 44.5° C ± 0.2° C	C for 24 hrs				
	Gas production is positive te	st for fecal coliforms				
	Calculate MPN based on con	nbination of positive EC tubes	<u></u>			
•	b. Membrane Filter Procedure					
	Following filtration place M					
	M-FC broth					
		roof plastic bag and submerge				
		nin				
		for 24 hrs				
		iforms				
		nsity per 100 ml	• • • ———			
<b>56</b> .	Delayed-Icubation Coliform Test					
	After filtration, place MF over pad of	<del>-</del>				
	of a 12% sodium benzoate solution		· · ·			
	Addition of 50 mg cycloheximide p	•				
		optional				
	Transport culture by mail service to	•	• • •			
	Transfer MF cultures to standard M-Endo medium					
	at laboratory		· · · · · <del></del>			
			· · · · <del></del>			
	If at time of transfer growth is visible					
	till end of work day then incubat					
	Count sheen colonies, verify if neces		· · · · ———			
		per 100 ml				
<b>57</b> .	•	er 100 mi				
57.	Additional Test Capabilities	Mathad				
	Fecal streptococci Pseudomonas aeruginosa	Mathod				
	Staphylococcus					
	SalmonellaeBiochemical tests					
	Serological tests					
	Other					
	Otilici	rui pose	<del></del>			

Laboratory		ocation	Date
	Laboratory S	Staff and Facilities	
58.	Personnel  Adequately trained or supervised for bacce examination of water  Laboratory staff (Total) Prep		
59.	Reference Material Copy of the current edition of Standard in the laboratory State or federal manuals on bacteriologic water available for staff use	Methods available	
60.	Physical Facilities  Bench-top area adequate for periods of processing samples	eak work in	
61.	Laboratory Safety Proper receptacles for contaminated glass Adequately functioning autoclaves with pand maintenance Accessible facilities for hand washing Proper maintenance of electrical equipment gas and electrical shock	sware and pipettes	

First aid supplies available and not out-dated . . . . . . . . . . . . . . . . .

**62**.

Remarks

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# PART C. MEDIUM-WATER BIOLOGY

# **BIOLOGICAL FIELD EQUIPMENT**

	Α.	Mobile L	.abs								
		1. Nu	mber available ()								
(Check)			uipped for:								
		a. '	Static bioassay								
		b.	Flow-through bioassay								
		c.					•				
		d.	Macroinvertebrate bioassa	ıvs							
		e.	Algal assay								
	В.	Boat and									
			Туре	Age	Condition	Hull	Length	Beam	HP	Float	ation
				<del>-   - ° -</del>			<u>-</u> -			Integral	Other
		1.			<u> </u>						
		2.									
		3.			ļ		<b></b>	ļ		ļ	
		4.			<b>_</b>		ļ	<b></b>			
		5.			<u>L</u>	<u> </u>	<u> </u>	<u> </u>	L	<b></b>	
	C.	Scuba G	ear								
	•										
			description of available gear	•							
	D.		g Equipment								
		1. Plai	nkton								
		a.	Water Bottles							-	
					Manalli	<u>. T</u>	Dia salis	7 7/-1		7	
					Metalli	IC	Plastic		ume ers)	1	
			(1) Kemmerer	•	1	- 1		} '''		1	
			(a)		<del></del> -			╅───		-	
			(b)		<del></del>			+		1	
			(c)		<del>-  </del>			<del>                                     </del>		┪	
			(2) Van Dorn				-			1	
			(a)						-	┪	
			(b)							1	
			(c)							1	
		b.	Pump sampler							_	
		о. с.	Integrated (Tubular) samp	aler							
		d.	Plankton Net	J.C.							
		ч.	Wisconsin type (net mesh	cize	1						
			Clark-Bumbus type (net r	nesh size	<del></del> / ,						
			Cambus type theth	・・・ショ・・ コールシー							

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# **BIOLOGICAL FIELD EQUIPMENT**

	D.	San	Sampling Equipment (Continued)							
		2.	Peripa.  b. c. d.	Substrate type (1) Glass (2) Plexiglass (3) Other (specify) Substrate dimensions cm X cm Substrate exposure depth, cm Substrate Orientation (1) Vertical (2) Horizontal						
		3.	Macı	rophyton:						
			Spec	cify:						
		4.	Mac	croinvertebrates:						
			a.	Grabs						
				Туре		Area of bite (m <sup>2</sup> )				
					.0232	.0523	.0929	Other		
				Ponar						
				Petersen						
				Ekman	<u> </u>					
				Tall Ekman				i		
				Other (specify)						
			b.	Corers						
				Specify:						

#### **BIOLOGICAL FIELD EQUIPMENT**

D.	Sampling Equipment (Continued)					
	4. Macroinvertebrates (Continued) c. Artificial Substrates (1) Multiplate (2) Masonite (3) Other (Specify)	·				
	d. Basket (1) Limestone (2) Other (Specify)	<del></del>				
	e. Surber sampler f. Other samplers g. Sieves (1) #30 Standard (2) #40 Standard (3)(Other)					
5.	. Fish					
	a. Shocker (1) AC (2) DC (3) Operating voltage (4) Manufacturer					
	b. Gill nets					
	(1) (2) (3) (4)	Mesh size (cm)	Length (m)			

#### **BIOLOGICAL FIELD EQUIPMENT**

#### c. Trammel nets

	Mesh size (cm)	Length (m)		
(1)				
(2)		1		
(3)				
(4)				

#### d. Seines

	Mesh size (cm)	Length (m)
(1)		
(2)		1
(3)		
(4)		

e.	Trawls	
	(1) Specify	

#### E. Miscellaneous

	Instrument	Manufacturer	Type/Model	Age	Condition
1. 5	Submarine Photometer	·		<u> </u> 	
2. (	Current meter				
3. 9	Secchi measurement	,			4.51
4.	Benthic respirometer				ž v
5.	Secchi disk				

**BIOLOGICAL LABORATORY EQUIPMENT** 

Counting and Identification Age/ Ocul				Γ.	Objectives			Phase	Nomarski	Nomarski Equipped for photomicrogr.	Whipple
1. Microscopes	Age/ Cond.	Manuf.	Ocular(s) Magnif.(X)	x x x x		هٔ ا	ž	Equip photo	֓֞֞֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓֓		
a. Compound, monocular											Γ
(1)	·	<u> </u>		<u> </u>							
(2)											
(3)										}	
b. Compound, binocular											Γ
(1)											
(2)											
(3)											Ī
c. Portable field microscope											Ī
d. Stereo (dissection) microscope						nifica range					Ī
(1) Rotating nosepiece											Ì
(a)				<u>.</u>		_					
(b)											
(c)											Ī
(2) Zoom											Ī
(a)	į										
(b)											T
(c)											T

#### **BIOLOGICAL LABORATORY EQUIPMENT**

#### B. Biomass Determination

1.	Balance	No. Avail.	Make	Type/Model Size, etc.	Age	Condition
	a.					
	b.					
	с.					
2.	Drying oven					
3.	Vacuum oven					
4.	Muffle furnace					
	Temp. control (?)					
5.	Desiccators					
6.	ATP measuring instruments					
	a					
	b. ·					
7.	Centrifuge					
L	Refrigerated (?)					
8.	Freeze drier				<u> </u>	

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#### **BIOLOGICAL LABORATORY EQUIPMENT**

#### C. Chlorophyll Measurements

	Instrument	No. Avail.	Make	Type/Model Size, etc.	Age	Condition
1.	Spectrophotometer.		 			
	a. ·					
	b. ·				<u></u>	
	с.					
2.	Fluorometer					
	a.	[ 				
	b.					
1	c.					
	(List excitation and emission filters for fluorometers)					
3.	Tissue grinder					
4.	Sonifier					

#### D. Culturing and Rearing Equipment

- 1. Algal culture chamber
- 2. Macroinvertebrate
- 3. Fish

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#### **BIOLOGICAL LABORATORY EQUIPMENT**

- E. Bioassay Facilities
  - 1. Algal Assay (Describe briefly)

2. Macroinvertebrate bioassay - (Describe briefly)

- 3. ¡Fish bioassay
  - a. Static bioassay (Briefly describe size and number of dilution water supply chambers, number of replicates, number of tests that can be run simultaneously, temperature control, and other supporting equipment.)
  - b. Flow-through (Briefly describe size and number of chambers, dilution water supply, temperature control, diluters, etc. )

#### BIOLOGICAL SAMPLE ANALYSIS

(Work load and methodology)

		No.		Methodology	,	
	Type of Analysis	Samples/ yr.	EPA	Stand. Methods	Other	Comments
Α.	Plankton				}	·
	<ol> <li>Phytoplankton count &amp; identification</li> <li>Diatom species proportional count</li> <li>Zooplankton count and identification</li> <li>Ash-free weight</li> <li>Chlorophyll determination</li> <li>ATP determination</li> <li>Primary productivity, oxygen method</li> </ol>					
	<ul><li>8. Primary productivity, carbon-14 method</li><li>9. Algal assay</li></ul>					
В.	Periphyton  1. Cell counts and identification  2. Diatom species proportional counts  3. Ash-free weight					
	<ul><li>4. Chlorophyll determinations</li><li>5. ATP determination</li></ul>					<del></del>

		No.		Metholology	,	
	Type of Analysis	Samples/ yr.	EPA	Stand. Methods	Other	Comments
C.	<ul><li>Macrophyton</li><li>1. Identification</li><li>2. Ash-free weight</li><li>3. Chlorophyll</li></ul>					
D.	<ol> <li>Macroinvertebrates</li> <li>Counts and Identification</li> <li>Ash-free weight</li> <li>Flesh tainting</li> <li>Tissue analysis</li> <li>Bioassay, static</li> <li>Bioassay, flow-through</li> </ol>					
E.	Fish 1. Counts/ID/wgt/lgth 2. Flesh tainting 3. Tissue analysis 4. Bioassay, static 5. Bioassay, flow-through					

#### LIST OF BIOLOGICAL PROCEDURES

	A.	Phy	toplankton
(Check)		1.	Sample Volume(liters)
		2.	Preservative
			a. Formalin
			b. Merthiolate
			c. Other (specify)
		3.	Counting Techniques used
			a. Sedgwick-Rafter Cell
			b. Palmer-Maloney Cell
			c. Membrane Filter Counts
			d. Inverted Microscope Method
			e. Other (specify)
		4.	Counting units used
			a. Natural Unit (clump count)
			b. Areal Unit
			c. Cell count
			d. Cell volume
		5.	Identification Level
			a. Total phytoplankton count
			(1) Identify to genus
			(2) Identify to species
			(3) Identify to major groups only
			b. Diatom species proportional count
		6.	Biomass measurements
			a. Dry weight
			b. Ash-free weight
			c. ATP
			d. DNA
			e. Chlorophyll
			(1) Solvent used
			(2) Fluorometric, in vivo method
			(3) Fluorometric, in vitro method
			(4) Fluorometric, in pheophytin correction
			(5) Spectrophotometric, Trichromatic, Strickland/Parsons
			(6) Spectrophotometric, Trichromatic, SCQR/UNESCO
			(7) Spectrophotometric, pheophytin correction
		7.	Metabolic Rates
			a. Productivity, oxygen method
			b. Productivity, Carbon-14 method
			c. Nitrogen fixation, acetylene reduction
		8.	Algal Assay
			a. Trophic level (biostimulation test)
			b. Limiting nutrient test
			c. Toxicity test
			d. Bottle method
			e. Flow-through method

### LIST OF BIOLOGICAL PROCEDURES (2)

	B.	Peri	phyton
(Check)		1.	Sample areamm <sup>2</sup>
		2.	Preservative
			a. Formalin
			b. Merthiolate
			c. Other (specify)
		3.	Counting Techniques used
			a. Sedgwick-Rafter Cell
			b. Palmer-Maloney Cell
			c. Membrane Filter Counts
			d. Inverted Microscope Method
		4	e. Other (specify)
		4.	Counting units used
			a. Natural Unit (clump count)
			b. Areal Unit
			c. Cell count d. Cell volume
		5.	Identification Level
,		٥.	
			a. Total phytoplankton count (1) Identify to genus
			(2) Identify to species
			(3) Identify to species (3) Identify to major groups only
			b. Diatom species proportional count
		6.	Biomass measurements
		٥.	a. Dry weight
			b. Ash-free weight
			c. ATP
			d. DNA
			e. Chlorophyll
			(1) Solvent used
			(2) Fluorometric, in vivo method
			(3) Fluorometric, in vitro method
			(4) Fluorometric, pheophytin correction
			(5) Spectrophotometric, Trichromatic, Strickland/Parsons
			(6) Spectrophotometric, Trichromatic, SCQR/UNESCO
			(7) Spectrophotometric, pheophytin correction
		7.	Metabolic Rates
<del></del>			a. Productivity, oxygen method
			b. Productivity, Carbon-14 method
			c. Nitrogen fixation, acetylene reduction
		8.	Bioassay
			a. Trophic level (biostimulation test)
			b. Limiting nutrient test
			c. Toxicity test
			d. Bottle method
			e. Flow-through method

#### **LIST OF BIOLOGICAL PROCEDURES (3)**

	C.	Mad	croin	vertebrates				
(Check)		1.	San	nple preservation				
			a.	Formalin%				
			b.	Ethanol%				
			C.	Other (specify)				
		2.	Sie	ve employed				
			a.	Standard #30				
			b.	Standard #40				
			c.	Other (specify)				
		3.	Sor	ting techniques				
			a.	Stain with Rose bengal				
			b.	Fluorescent stain				
			C.	Other stain (specify)				
			d.	Sugar floatation				
			e.	Other separation method	(specify)_			
		4.	Ide	ntification		(Cho	eck)	
						Level of Identification		
				Group	Order	Family	Genus	Species
			 a.	Diptera (excl. midges)				
			b.	Midges		:		<del>                                     </del>
			c.	Trichoptera	<u> </u>			
			d.	Plecoptera	<b>†</b>			1
			e.	Ephemeroptera				
			f.	Odonata				<b>†</b>
			g.	Neuroptera				1
			ĥ.	Hemiptera		<u> </u>		†
			i.	Crustacea				
			j.	Hirudinea	Ī			
			k.	Nematoda				
			١.	Bivalvia				
			m.	Gastropoda		-		
		5.	— Mai	intain reference collection	of organism	s for identif	ication	
		6.		"outside" consultants for				
		7.		ar larvae to adult stage to a				
		8.		sue analysis for toxic subst				
		9.		assay				
			a.	Static				
			ъ. b.	Flow-through				

#### LIST OF BIOLOGICAL PROCEDURES (4)

	D.	risn	
(Check)		1.	Preservative
			a. Formalin
			b. Other (specify)
		2.	Age determinations
			a. Scales
			b. Other (specify)
		3.	Condition factor (length-weight relationship)
		4.	Flesh tainting
		5.	Histopathological studies (describe):
		6.	Bioassays
			a. Laboratory
			(1) Static tests (describe):
			(2) Flow-through tests (describe):
			b. In-situ tests (describe):

## PART D. MEDIUM AIR

## $\mathsf{MANUAL} - \mathsf{SO}_2 \text{ or } \mathsf{NO}_2$

Sampling			
Volume (No. of Samples)			·
Method of Delivery(Field to Lab)			
Sampler Used			
Frits-Impingers	· · · · · · · · · · · · · · · · · · ·		
Time Between Sampling and Analysis			
Storage Method			
Analysis			
Method		<u> </u>	`
Copy Available			
Calculations			
Equipment(Automated-Manual)		•	
	<u>Items</u>	· ·	<u>Schedule</u>
(Automated-Manual)		or	
(Automated-Manual)  Preventive Maintenance	<u>Items</u> Reagents	or	Schedule Gases
(Automated-Manual)  Preventive Maintenance  Chemical Purity of Reagents	<u>Items</u> Reagents	or	Schedule Gases
(Automated-Manual)  Preventive Maintenance  Chemical Purity of Reagents  Reagent Makeup Procedure	<u>Items</u> Reagents	or	Schedule Gases
(Automated-Manual)  Preventive Maintenance  Chemical Purity of Reagents  Reagent Makeup Procedure  Reagent Standardization Procedure	<u>Items</u> <u>Reagents</u>	or	Schedule Gases
(Automated-Manual)  Preventive Maintenance  Chemical Purity of Reagents  Reagent Makeup Procedure  Reagent Standardization Procedure  Calibration  Procedure (Samplers)	<u>Items</u> Reagents	or	Schedule Gases
(Automated-Manual)  Preventive Maintenance  Chemical Purity of Reagents  Reagent Makeup Procedure  Reagent Standardization Procedure  Calibration	<u>Items</u> <u>Reagents</u>	or	Schedule Gases
(Automated-Manual)  Preventive Maintenance  Chemical Purity of Reagents  Reagent Makeup Procedure  Reagent Standardization Procedure  Calibration  Procedure (Samplers)  Procedure (Analysis)	<u>Items</u> Reagents	or	Schedule Gases
(Automated-Manual)  Preventive Maintenance  Chemical Purity of Reagents  Reagent Makeup Procedure  Reagent Standardization Procedure  Calibration  Procedure (Samplers)  Procedure (Analysis)  Copies Available	<u>Items</u> Reagents	or	Schedule Gases

# $\mathsf{MANUAL} - \mathsf{SO}_2 \text{ or } \mathsf{NO}_2 \text{ (Continued)}$

Data Processing			
Mode Utilized(Strip Chart, Mag. Tape)		·	
Discrepancies	 		
SAROAD Format			
Reduction Procedure			·· <u>·</u> ··
Reporting of Data	 . ,		

# Continuous $SO_2$ , $NO_2$ , CO, or $O_3$

Type of Analyzers	proper seather a single
No. of Analyzers	
No. of Field Stations	
Containing Analyzers	
Frequency of Sampling	
Manpower (Attended-Unattended)	
Frequency of Calibration	· · · · · · · · · · · · · · · · · · ·
Method of Calibration	
Traceability of Calibration	
Curves Available	
Documentation	
Frequency of Zero and Span	
Corrective Action Plan  If Out of Specs	······································
Maintenance Log	
Data Collection Device(Strip Chart, Mag. Tape)	
Data Reduction	· · · · · · · · · · · · · · · · · · ·
Reports	<del>,</del>

#### MANUAL - HI VOL

Туре		<u>.</u>			 
No. of Analyzers					
No. of Sites					 
Frequency	<del></del>		<del></del>		· ——
Type of Filter			·		 
Pre-exposure Checks and Procedures	· · · · · · · · · · · · · · · · · · ·				 
Collection Procedures				<del> </del>	 
Calibrating Procedures			<del></del>		 
Weighing Procedures				<u>-</u>	 
Frequency			_ <del>.</del>	<del>_</del>	
Data Handling					 

#### PART E. MEDIUM - PESTICIDES

The following questions may be asked in toto if the personnel do not seem to know much about gas chromatography. If personnel seem versed in GLC, it may be necessary only to pick out some questions in each subsection.

I. A.	GLC Calibration & Maintenance Detector (EC)
1.	Frequency of preparation of linearity curves for pesticides of interest - weekly  monthly never
2.	Frequency of determination of standing current profile - weekly  monthly never other (describe)
3.	Frequency of construction of voltage/response curve - weekly
4.	Comments on method of selection of optimum polarizing voltage.
<b>B.</b> 1. 2. 3.	Detector (FPD)  Date unit purchased  Manufacturer of power supply unit volt. Awareness of operator
4.	Has a determination been made of the signal to noise ratio as a criterion for optimal selection?  Yes  No Comments:
5.	Have heat shields and filters been checked on a spectrophotometer for light transmission at specified wavelengths? Yes \( \subseteq \text{No} \subseteq \text{If yes, at what wavelengths?} \)  P (526 mu) Actual %T  S (394 mu) Actual %T
6.	Have velocity of gases been adjusted to give optimum signal to noise ratio? Yes \( \sqrt{No} \sqrt{\sqrt{No}
7.	Date detector last cleaned? O-rings changed?
8.	Sensitivity in terms of % F.S.D. for 2.5 ng of ethylparathion.
9. 10. 11.	Baseline noise % F.S.D.  Is flame extinguished overnight? Yes  No  No  If no, how is flame-out avoided?  Does instrument have vent valve? Yes  No  If no, how is flame-out avoided?
<b>C</b> . 1.	Alkaline Flame Detector Which salt is used?
2. 3.	Is flame extinguished overnight? Yes \( \subseteq \text{No } \subseteq \)  Give frequency of cleaning of loop collector to detector \( \subseteq \text{Loop} \)
4.	If electrical current to collector loop is supplied by batteries, give frequency of battery changing or recharging
5. 6.	Give operating baseline current amps.  Baseline noise % F.S.D.
7.	Give sensitivity in terms of % F.S.D. from 2.5 ng ethylparathion.

<b>D.</b> 1.	Flame Ionization Detector Give frequency of cleaning of collector loop
2.	If electrical current to collector loop is supplied by batteries, give frequency of battery changing or recharging
3.	Give operating baseline current amps.
E.	Coulson Electrolytic Conductivity Detector
1.	Date of purchase
2.	Mode of operation
3.	Sensitivity in terms of % F.S.D. resulting from 1 ng of aldrin
4.	Normal baseline noise % F.S.D.
5.	Pyrolysis furnace temperature °C
6.	Pyrolysis furnace temperature °C.  Block temperature °C.
7.	Flow rates in m1/min. of purge and carrier gas
8.	Flow rates in m1/min. of Q <sub>2</sub> or H <sub>2</sub>
9.	Pretreatment of water used in cell
10.	, <u> </u>
11.	, ———
12.	Identity of GLC column(s) used
F.	Electrometers
1.	Frequency of zeroing daily \( \square\) weekly \( \square\) monthly \( \square\) never \( \square\) other \( \square\) (describe)
2.	Frequency of determination of attenuator linearity daily  weekly monthly never other (describe)
3.	Frequency of repair
G.	Strip Chart Recorders
1.	Frequency of zeroing baseline daily  weekly  monthly  never  other  (describe)
_	
2.	Describe method of determining optimum gain control setting
3.	Frequency of cleaning of slide wire
J. Н.	
1.	Is column efficiency determined before routine use? Yes \( \simeq \) No \( \simeq \) If yes, describe method
2.	Are response characteristics determined before use? Yes \( \subseteq \text{No} \subseteq \text{If yes, describe method} \)
3.	Frequency of changing demister tube, if used
4.	Frequency of changing glass wool plug at column inlet
5.	Is any determination made of compound degradation characteristics of column-endrin, p.p'-DDT?
6.	If column used for FPD, are response characteristics determined prior to use? Yes \( \subseteq \text{No} \subseteq \text{If} \) answer is yes, describe method.
7.	In using the column for tentative identification of peaks, are RRTA or EP data utilized?  Yes  No  If answer is no, describe the alternative used.

l.	GLC Operation - General
1.	Is any method used to monitor accuracy of instrument pyrometer? Yes  No  If answer is yes, describe.
2.	Is carrier flow velocity monitored by bubble meter? Yes \( \sum \) No \( \subseteq \) If answer is no, request operator to set what he thinks is 70 ml/min. and make actual check with bubble meter.
3.	Assessment of flow system plumbing - molecular sieve filter, neatness of layout, knowledge of operator pertinent to checking for leaks, etc.
4.	Is a log maintained for each instrument showing chronological data such as change of detector, etc.
5.	General assessment of GLC operation capability for pesticide work:
6.	Is any check made in the early A.M. with a working standard solution to relate response characteristics to those of the pervious day's operation? Yes \(\Boxed{D}\) No \(\Boxed{D}\) Comment \(\Boxed{D}\)
	actorisates to those of the pervious day's operation. Test in two in comments

#### PART F. RADIATION

#### Counting Room Facilities and Equipment

A.

#### **Counting Room Facilities** 1. Are counting instruments located and operated in a separate counting room facility? Yes No 🗆 2. Number and size of counting rooms: Number Size 3. Are instruments operated from regulated power? No 🗆 Yes $\square$ Is there an adequate ground available to all counting 4. instruments? No $\square$ Yes 🗌 Can the light in the counting room that houses the liquid scintillation systems be readily controlled (for sample loading, etc. )? Yes 🗌 No 🗆 6. Are counting room facilities adequately protected (by location or shielding) from higher radiation areas and sources? Yes $\square$ No 🔲 7. Is there adequate temperature control in the counting room(s)?

Yes \( \Bar{\cap} \) No \( \Bar{\cap} \)

<b>B.</b> ·	Spec	ecial Questions							
	1.	What beta emitter is used for gross beta calibration?							
	2.	What alpha emitter is used for gross alpha calibration?							
	3.	Are individual analyses logged in permanent type laboratory notebooks and initialed and dated by the analyst?							
Comme	•	Yes No No							
	4.	Are working copies of all methods used readily available to the laboratory analyst?							
		Yes No No							
Comme	nts : ˈ								
	5.	Are standard solutions prepared and stored in an area separate from areas where analysis of samples and blanks is being performed?							
		Yes No No							
Comme	nts:								
	6.	How often are standards preparation areas and sample working areas being swiped and checked for radioactivity contamination?							
Comme	nts:								

#### Equipment

Refer to Chart D, Section on Special Radiological Equipment, in the Preliminary Questionnaire.

For Alpha and Beta-particle Counters

	Sample C	hanging				
		anual				
		utomatic				
		apacity				
	O.	ipuoity				
	Instrumer	nt Background	1	A 1 ln		Data
		_		Alpha	l	Beta
	•	perating volta	ge		<del>-</del> ·	
	cp	om				
Back	ground Co					
		equency				
	Lo	og kept				<del></del>
Fora	all Instrum	ients				
	Fr	equency of c	alibration			
	Fr	equency of se	ervice mainten	ance		
Alpha and	d Beta Part	icle Counters				
				1		
Wind	dowless Ga	s-flow Propor	tional Counté	r		
	Co	ounting gas				
		0.0				
	Sa	ample dish dia	meter			
Thin	Window C	Sas-flow Prop	ortional Cour	nter		
	Co	ounting gas				
		,				
	W	indow density	/ (g/cm <sup>2</sup> )			
			(6)			<del></del>
Δinh	a Scintilla	tion Counter				
Діріі	ia Jennina	tion Counter				
	ΔΙ	pha Phospho	r location			
	Λ.	Photo tube				
			e			
		Samples				
_	<b>.</b>	_				
Beta	Scintillati	on Counter				
	,					
	Be	eta Phosphor				
		Type				
		Thickness				
		Diamatar				
		Diameter				

Liquid Scintillation Counter	
Discrimination channels 1 2 3	
Data readout	
Visual	
Printout, Channel 1 2 3	
Spectrometer Systems	
Alpha Spectrometer (Surface Barrier Type)	
Detector	
Active diameter	
Detectors/chamber	
Analyzer	
Channels	
Gamma Spectrometer	
Detector- size	
Analyzer-Channels	
Radon Gas Counter	
Gas counting cells/system	
Manufacturer of gas counting cells	<del></del>

#### METHODS USED IN THE CALCULATION OF RADIATION DATA

"Hand" or Computer	Matrix or Least Squares	"Spectrum Stripping"	"Compton Subtraction"	Precision/Accuracy Reported	Opportunity for Final Recheck
		, 			
:					
i	;				
	"Hand" or Computer	"Hand" or Computer Matrix or Least Squares	"Hand" or Least Squares "Spectrum Stripping"	"Hand" or Computer	"Hand" or Computer Least Squares "Spectrum Stripping" "Compton Subtraction" Reported

#### CALIBRATED RADIOACTIVE SOURCES

Radionuclide	Supplier	Where Stored	Comments
			·
	,		:
			·
		ı	

#### SAMPLING GUIDELINES

Radionuclide ·	Media	Media Site	Site Selection Criteria	Sampling Procedures			
Radionuciide				Grab	Continuous	Other	Custody
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				<b> </b> 			
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# SECTION 6 INSTRUCTIONS AND RATING SYSTEM

#### GENERAL INSTRUCTIONS

This section of the procedure provides instructions for the evaluators who will conduct the onsite survey. The laboratory visit is the most important part of the evaluation procedure. Thorough preparation is the key to its success. Unless ample time is devoted to prior preparation, an accurate laboratory evaluation will be impossible. "Ample" cannot be rigidly defined, for the time involved will vary according to: the talents and experience of individual evaluators; the information provided in the preliminary questionnaire; the number and variety of analyses that a laboratory performs; and the number of inspectors performing the site evaluation. A minimum of several hours preparation should be allotted for even the most straightforward situation.

Initially, the evaluators must completely familiarize themselves with the format and questions of the onsite checklist. This familiarity will facilitate the flow of the interviews. It will help the evaluators to anticipate laboratory reactions and to know when further probing of a response is necessary. When an evaluator is required to exercise judgment in rating a laboratory, close adherence to the procedural guidelines will enhance the objectivity of the judgement.

Upon receipt of the preliminary questionnaire, the evaluator must carefully study all the information provided. Although responsibility for different aspects of the laboratory evaluation may be divided among several members of an evaluation team, each evaluator should be familiar with the information contained in the entire report. A broad understanding of the background information provides a valuable resource for the evaluator who must assess a particular function such as quality control or analytical procedures.

For convenience, the onsite survey has been divided into three areas:

- (1) Management and Organization, (2) Technical Services, and
- (3) Internal/External Controls. Detailed instructions, suggested questions and scoring procedures are provided for each of these areas.

Prior to the onsite visit, some scores may be calculated from the data recorded in the answers to the preliminary questionnaire. Careful study of the detailed instructions will indicate those items which can be pregraded. In addition to the preliminary assignment of scores much can be done in preparation for the visit. For example, in the technical services section, the apparatus list can be compared with the requirements for each analysis allegedly performed to determine that the necessary equipment is on hand. If a laboratory states that it performs atomic absorption to determine cadmium but lists no cadmium hollow cathode, the analysis could not be done. Awareness of such discrepancies before the visit can highlight areas in need of the evaluator's scrutiny.

All questions suggested in the Evaluator's Guide will not require numerical scores. Some demand only the evaluator's positive or negative judgement in support of the overall laboratory score. However, there are a sufficient number to be rated on the score sheets to allow an objective laboratory evaluation which can be used for comparative purposes.

Numerical scores should not be computed onsite. The evaluator should gather information and check appropriate entries for possible scoring levels. At the end of the laboratory visit, the data should be reviewed, all confusion dispelled, and discrepancies resolved. The score can then be tabulated and the laboratory informed in writing of the outcome of the evaluation process. Items which require correction or improvement prior to final scoring should also be highlighted to afford the laboratory the opportunity to make the adjustments necessary for subsequent acceptance.

Some items are more crucial to laboratory operation and security than others. The onsite check list specifies certain conditions which <u>must</u> be present and satisfactory for a laboratory to be deemed qualified. These items are marked with an asterisk. For example, regardless of the excellence of facilities and analytical competency of an establishment, if it lacks adequate sample custody and control, it cannot be found acceptable. If an item in any part of the check list is marked with an asterisk, the problem must be resolved before the final score is calculated.

#### Suggested Sequence of Onsite Interviews

Part 1. General Information about the Laboratory

Part 2. Personnel (in part)

Part 3. Laboratory Space and Facilities (in part)

Interview #2 (or Series of Interviews) - Supervisors with their laboratory personnel in the laboratories. (If number of supervised personnel is large, subdivide the group for convenience.)

Part to be covered:

Part 2. Personnel (in part)

Part 3. Laboratory Space and Facilities (in part)

Part 4. Technical Service

Part 5. Equipment List

Part 6. Quality Control (in part)

Interview #3 Internal and External Controls - This may involve a designated quality control officer, a section with responsibility for review of operations, or an individual or individuals with parttime responsibility for quality control in the lab.

SPECIFIC INSTRUCTIONS AND RATING SYSTEM

#### Management and Organization Area - Parts 1, 2 and 3

This area does not readily lend itself to an objective evaluation. The questions frequently cannot be designed to elicit a "yes" or "no" response. Therefore, the judgements made by the evaluator are of great importance.

Guides are provided to help standardize the scores of the individual evaluators. The scoring system is designed so that the values assigned to any individual characteristic of the lab will affect the total score by only a small increment. Thus, although many laboratories with many different evaluators may be involved, scores should be comparable.

The experience gleaned from the onsite visit, from witnessing the attitude and manner of responses by laboratory personnel to questions, from watching the interplay between individuals when more than one is present during the onsite evaluation is essential to the assignment of scores to answers. The evaluator is responsible for the integration of all these factors to arrive at the decision to score each question with 5, 3, or 1.

The rationale for particular questions and the approach to their formulation may not always be apparent. Therefore, a statement of intent for each series of questions is provided in the "Evaluator's Guide", Section 5. This will help the evaluator to ask suggested questions and to develop a personal line of questioning.

If the laboratory is privately owned some determination of its financial stability should be made. Some information can be gained from the annual operating budget, fees charged for services and the number of analyses performed per year. Also the age and condition of the real estate and laboratory apparatus could indicate, in general terms, the health of the organization. Laboratory apparatus is expensive and a large investment is required to start an analytical laboratory. Much can be learned without demanding an audit or inspection of the books. A study of the annual report requested in the preliminary questionnaire should provide a good deal of information about the financial condition of the organization. If no fiscal information is contained in the annual report, some inquiries concerning the financial condition of the laboratory should be made.

# PART 1. GENERAL INFORMATION ABOUT THE LABORATORY

(1)	Appro	Appropriateness of Organization				
		Best Description of Laboratory	Score			
		Responses to questions indicate the organization is as reported, and that its functioning is not so rigid as to interfere with operational requirements.	5			
		Some doubt that organization as described is really followed. Chain-of-command is followed without deviation to the detriment of good performance.	3			
٠		Serious doubts concerning organization and control of people.	1*			
(2)	Impair	ment of Functions				
		Best Description of Laboratory	Score			
		Responses generally satisfactory, no real problems in any of these areas. Certainly nothing said that would indicate impairment of laboratory functions.	5			
		Some problems evident in one or two places. These may make it difficult to operate effectively.	3			
		Management problems are obvious in several areas, hard to get help, customers complain, etc., and performance is likely to suffer.	1*			
(3)	Streng	th of Management				
		Best Description of Laboratory	Score			
		Firm stand taken concerning internal communications and cooperation between groups. Both annual and long-range plans are made and followed. Firm authority demonstrated without the feeling of an "absolute monarch." Impression given of "wide awake" management.	5			
		Some weaknesses indicated in a few of the items.	3			
		It appears that management is weak - no plans made for future - little cooperation or internal communications.	1*			

(4)	Obje	Objectivity of the Laboratory							
		Best Description of Laboratory	Score						
		Responses open and direct - no reason to doubt objectivity.	5						
		Some aspects of relationships unclear.  Objectivity not seriously questioned but some doubts.	3						
		A conflict of interest exists, is clearly apparent in any part of the organization.	1*						
(5)	Coop	Cooperation Obtained							
		Best Description of Laboratory	Score						
		All information provided promptly.  Cooperative attitude displayed by all personnel. Preliminary questionnaire distributed to proper persons for completion.							
		Most information supplied readily. Satisfactory candid responses.	3						
		Important information not provided and difficult to draw out answers. Cooperation of all not evident. Evaluation presented and no plans to make constructive use of the results.	1*						
	•	A score of 1, when marked with an asterisk (1*) in any of the check list must be resolved to the satisfaction of the later before a final score is calculated.							

#### **Evaluator's Notes**

## PART 2. PERSONNEL

(1)	Super	visor Training	
		Best Description of Laboratory	Score
		Supervisors have degrees. Sufficient experience in place of degree.	5
		No degree and less than 5 years experience.	3
		No degree and insufficient experience.	1
(2)	Super	visor Experience	
		Best Description of Laboratory	Score
		The supervisors as individuals and as a group are highly trained and experienced.	5
		The supervisors meet general requirements but are weak in environmental monitoring work.	3
		Some of the group appear deficient in training and experience.	1*
(3)	Job D	escriptions	
		Best Description of Laboratory	Score
		There is good agreement between description and what is done.	5
		There is some agreement between description and what is done but considerable deviation.	3
		General agreement only.	1
(4)	Traini	ng Program	
		Best Description of Laboratory	Score
		Formal training program exists and is followed.	5
		No formal training program but obviously some training is continued.	3
		Little evidence that training is done.	1
(5)	Turno	ver Rate	
		Best Description of Laboratory	Score
		Rate is less than 25%.	5
		Rate is 25-50%.	3
		Rate is greater than 50%.	1*

#### (6) General Morale

Best Description of Laboratory	Score
Management exhibits real concern for individuals, evidenced by central records kept for each individual showing advancement dates, promotions, training programs taken, participation in health programs. General morale of personnel is high.	5
No formal central records are maintained but some effort made to encourage people.  No evidence of serious morale problems.	3
Little concern demonstrated for individuals.  Morale problems are evident.	1

#### **Evaluator's Notes**

## PART 3. LABORATORY SPACE AND FACILITIES

(1)	General Characteristics				
		Best Descript	tion of Laboratory	Score	
		General characteristics of laborator satisfactory.	у	5	
		Impression of laboratory is only av	erage.	3	
		General features of the laboratory	are poor.	1	
(2)	Office	Space			
		Best Descrip	tion of Laboratory	Score	
		16.7m <sup>2</sup> (180 sq. ft.) or greater/pers	son.	5	
		12.5-16.7m <sup>2</sup> (135-180 sq. ft.)/pers	on.	3	
		Less than 12.5m <sup>2</sup> (135 sq. ft.)/pers	son.	1	
(3)	Labora	atory Space Rest Descrip	tion of Laboratory	Score	
		18.6m <sup>2</sup> (200 sq. ft.) or greater/pers	•	5	
		13.9-18.6m <sup>2</sup> (150-200 sq. ft.)/perso	on.	3	
		Less than 13.9m <sup>2</sup> (150 sq. ft.)/perso	on.	1*	
(4)	Bench-	-top Space Best Descript	tion of Laboratory	Score	
		1.2m (4 lin. ft.) or greater/person.		5	
		0.9-1.2m (3-4 lin. ft.)/person.		3	
		Less than 0.9m (3 lin. ft.)/person.		1	
(5)	Hood :	Space and Operation  Best Descrip	tion of Laboratory	Score	
		Hoods sufficient in number and cap	•	5	
		Some additional hoods and/or capa	city needed.	3	
		Hoods inadequate for purpose inter	nded.	1	

(6)	Storage Space Chemicals, Reagents, Glassware, Supplies †				
		Best Description of Laboratory	Score		
		Storage space adequate, accessible, and kept orderly.	5		
		Storage space available but overtaxed.	3		
		Storage space insufficient.	1		
		† NOTE			
		Further questions about inventory policy and materials identification are asked in Part 6.			
(7)	Samp	ole Storage †			
		Best Description of Laboratory	Score		
		Sample storage space is adequate and necessary provisions are made for samples requiring special attention.	. 5		
		Sample storage satisfactory in general but some special requirements are not fully met.	3		
		Sample storage space is inadequate and inefficiently arranged.	1*		
		† NOTE			
		Further questions about control of samples appear in Part 6.			
(8)	Cont	rolled Space			
		Best Description of Laboratory	Score		
		Controlled space necessary for performance of services offered by the laboratory is available. Responsibility for operation of these rooms is assigned and continuous check of conditions is maintained.	5		
		Necessary rooms are available but control is slack and checks are made only daily.	3		
		There are unsatisfied needs for controlled space and/or responsibility is not well defined and checking is less frequent than daily.	1*		

(9)	Librar	y .	
		Best Description of Library	Score
		A library is available; it is easily accessible, orderly, and well looked after.	5
		There is a library but it is disorganized and difficult to use.	3
		No organized library exists and each section or staff member keeps own references, periodicals, etc.	1
(10)	Safety	Equipment and Procedures	
		Best Description of Laboratory	Score
		Safety equipment is available, regulations are posted, and regular drills are held.	5
		Safety equipment is good but improvements in the lab safety program are needed.	3
		Safety equipment is not complete and an effective program does not exist.	1*
(11)	Distill	ed Water/Deionized Water	
		Best Description of Laboratory	Score
		Apparatus and water checked every day and kept in proper condition by one designated individual who keeps a record.	5
		Greater interval than one day between checks by designated individual.	3
		No one person responsible and no written procedure or records maintained.	1*
(12)	Glassy	ware Supply and Washing	
		Best Description of Laboratory	Score
		Glassware supply and washing are satisfactory in all respects.	5
		More attention needs to be given to washing equipment or procedures.	3

(13) House	ekeeping	
	Best Description of Laboratory	Score
	Laboratory has clean, neat appearance; movement and work are not impeded by clutter.	5
	Laboratory is clean but not as neat as should be expected.	3
	Poor housekeeping evident by dirt or clutter.	1
(14) Data l	Processing Equipment and Logistic Services	
	Best Description of Laboratory	Score
	Communication facilities within and outside the laboratory good. Computing capability present.	5
	Communications within and outside laboratory are limited. Data processing facilities inconvenient to use.	3
	Laboratory and sections thereof are isolated and computing capability sufficient for timely results not available.	. 1

## **Evaluator's Notes**

#### Technical Service Area - Parts 4 and 5

The technical services area encompasses analytical methods and instrumentation. The questions are straight-forward. They seek to determine the nature of analytical methods employed in the laboratory and the adequacy of the instruments used in these procedures.

All analytical methods must be Federal Register referenced or alternates which have been specifically approved by the Environmental Protection Agency. EPA approved water and radiation test methods are referenced in the Federal Register, Vol. 35, No. 199, October 16, 1973. Air test methods with EOA approval are referenced in the Federal Register Vol. 36, No. 228, November 25, 1971 and Vol. 38, No. 110, June 8, 1973. Air methods for stationary sources are referenced in Federal Register, Vol. 36, No. 247, Part II, December 23, 1971; Vol. 38, No. III, June 11, 1973; Vol. 39, No. 47, March 8, 1974; Vol. 40, No. 152, August 6, 1975; and Vol. 40, No. 194, October 6, 1975. The interim methods for algicides, chlorinated organic compounds and pesticides were issued by EPA's Environmental Monitoring and Support Laboratory. For non-referenced Biological Tests see Bibliography entries No. 6, 7, 8, and 9. See Ref. 4, Standard Methods, for Test No. 406, Standard Plate Count.

Prior to the onsite visit, the evaluator should compare the apparatus list in the preliminary questionnaire with the requirements for each analysis that the laboratory performs. The absence of essential equipment should be thoroughly investigated. If all necessary apparatus is available, the evaluator should carefully assess the condition of the instruments. To function properly, the analytical equipment should be inspected and serviced regularly.

In laboratories concerned with more than one medium, it may be desirable to score the different sections of the laboratory individually for Part 4 Analytical Methods and Part 5 Instruments. The separation of Charts C and D in the Preliminary Questionnaire, Section 4, by media will facilitate this.

The evaluator must complete a set of score sheets for Parts 4 and 5 for each section of the laboratory, if separate scores are desired.

Depending on the circumstance, the evaluators report might contain one over-all score for the laboratory or two or more scores, one for each media with which the laboratory is concerned.

A list of major equipment requirements for each analytical method is given in the Appendix. Prior to the onsite visit this list should be checked against the Analytical Methods circled in Chart C and the Instruments checked in Chart D of the Preliminary Questionnaire, Section 4, to verify that equipment is on hand to perform all the tests for which evaluation is being made. Ask questions about any observed discrepancies. Check the condition of the equipment and ascertain its capabilities in every instance for:

- Required Instruments
- Function Tests and Standardization of Instruments It is important that calibration curves be available for all major instruments and that they have been checked recently and updated if necessary.
- Calibration Equipment The availability of suitable calibration equipment is important. Standard weights and special thermometers should be traceable back to a standardizing agency such as the National Bureau of Standards. In air monitoring, especially, the calibration equipment available should be checked carefully. Is equipment available for basic calibration of flow measurement devices? Is the laboratory able to produce satisfactory standard atmospheres? Note mention of required calibration equipment in the Appendix, Major Equipment Requirements for Each Analytical Method.

# PART 4. ANALYTICAL METHODS

(1)	Refere	ence Methods or Approved Alternates	
		Best Description of Laboratory	Score
		All methods used are Federal Register referenced methods or specific EPA approval has been obtained.	5
		Some easily correctable minor deviations from referenced methods exist and steps are being taken to conform to standards.	3
		Some nonstandard methods which do not have specific EPA approval are in use.	1*
(2)	Reage	nt and Media Preparation	
		Best Description of Laboratory	Score
		Laboratory personnel are clearly aware of the importance of proper preparation, use, and storage of reagents and media, and laboratory procedures and practices are adequate to ensure same.	5
		Although personnel awareness and laboratory procedures regarding preparation, use, and storage of reagents and media are generally satisfactory, one or two examples of improper preparation, careless use, improper storage (time, temperature, container, etc.), inadequate records, or other unacceptable procedures or attitudes were noted.	3
		Personnel attitudes and/or laboratory procedures for ensuring proper preparation, use, and storage of reagents and media are not adequate.	1
(3)	Perfor	mance According to Standard	
		Best Description of Laboratory	Score
		Performance of analysts is closely supervised and all testing conforms to standards.	5
		The laboratory does not have a specific control program for each sampling procedure and analytical test and performance is uneven.	3
		Supervision of the analysts is lax and confusion exists about specific details of some control procedures.	1

## Evaluator's Notes

# PART 5. ANALYTICAL INSTRUMENTS

(1)	Required Instrumentation						
		Best Description of Laboratory	Score				
		All required instrumentation is in good working condition.	5				
		Some instrumentation is of doubtful quality or is in some degree of disrepair.	3				
		Needed items of equipment are missing, are not adequate for satisfactory work, or are improperly maintained.	1*				
(2)	Functi Instru	·					
		Best Description of Laboratory	Score				
		Instruments are maintained operative, accurate, and precise by regular functioning checks and by use of standard before unknown samples. Standard curves are available wherever indicated.	5				
	_		_				
		Instruments are periodically checked against zero point or other reference and examined for evidence of physical wear or inadequate maintenance.	3				
		Instruments are checked only when they stop working or when excessive difficulties are experienced.	1				
(3)	Calibra	ation Equipment					
(-)		Best Description of Laboratory	Score				
		Necessary calibration equipment is available and in good working condition.	5				
		Calibration equipment is of doubtful quality or is in some degree of disrepair.	3				
		Needed items for calibration are missing, are not adequate for precise work, or are improperly maintained.	1				

## Evaluator's Notes

### Internal and External Controls - Part 6

Quality control is an indispensable aspect of laboratory performance. Initiated by management's interest and concern and embodied in distinct operating procedures, commitment to quality performance should pervade all levels of the laboratory.

#### Concern for quality has many manifestations:

- Responsibility for quality control is clearly assigned.
- Analytical apparatus is adequately maintained and calibrations are performed frequently.
- Samples are carefully collected and identified, and promptly processed.
- Tests for precision and accuracy are employed to ascertain the validity of data.
- Laboratory uses quality control check (reference) samples on a scheduled basis.
- Laboratory records are assiduously kept and reports are completed regularly.
- The laboratory participates periodically in inter-laboratory proficiency tests.
- A training program exists for new employees; trainees' performance is monitored and evaluated.
- Corrective action procedures are available and can be quickly implemented when necessary

With the guidance of Internal and External Controls Part 6, the evaluator should explore the laboratory's provisions for quality control.

In addition to the operational components of a quality control plan, the evaluator must assess a number of intangibles. An atmosphere conducive to quality performance requires interest and enthusiasm, a cooperative working relationship between supervisor, analyst and technician, dedication, and a free flow of communication. Through insight and discussion, the evaluator must determine whether or not a sincere concern for quality control exists. The following guidance should assist the evaluator to make this judgment.

#### Control of Analytical Methods and Instruments

An effective environmental monitoring program must include a quality assurance plan to protect the validity of its data. Quality assurance has many components: calibration standards, standard reference material, careful maintenance of records, sample taking, sample processing and control, interlaboratory comparison studies and data validation.

Maintenance and calibration of analytical apparatus are critical to the generation of good data. The evaluator must determine whether instruments and apparatus are maintained and how well, whether calibrations are performed in an appropriate manner and with sufficient frequency, and whether records and documentation of maintenance and calibration are adequate. If maintenance is done on an outside contract, determine for what instruments such contracts exist. The following items should be considered to assess the laboratory's quality assurance measurements.

- Assignment of Responsibility The evaluator's first task will be to determine who has the responsibility to see that each of the instruments in Chart D in the preliminary forms is properly maintained and calibrated on schedule. This may or may not be the same person who actually does the maintenance and calibration. Here the intent is to evaluate whether the responsibility is clearly assigned or not. It may be useful to question several people, bench analysts and supervisors alike, on this point to see if the assignment of responsibilities is clearly and uniformly understood by all.
- Maintenance and Calibration Logs For legal and scientific reasons, it is important to keep careful records of maintenance and calibration of instruments and apparatus. Generally, these records should be kept in permanent (bound) notebooks in ink with each entry signed and dated. A separate log (or a separate section of a log) should be assigned to each instrument or piece of apparatus that requires any sort of periodic calibration or maintenance, whether that activity is performed by laboratory personnel or by an outside agency under contract. It is convenient to include all calibration, maintenance, and repair actions on an instrument in the log, as a complete and accessible record of the conditions of that instrument. This includes evidence of traceability of standards to the National Bureau of Standards or other recognized source.

Each entry must specify clearly what action was taken when and by whom. For example, if a new calibration curve was established which will be the basis for future analyses, either the curve or a reference to a notebook containing the curve should be included, along with an explanation of how the curve was established (identification of reference standards, methodology) and when the analyst began using the curve in "real sample" analysis.

Adequacy of Calibration and Maintenance Practices - The evaluator now must assess the laboratory's actual procedures and practices for calibrating and maintaining its instruments and apparatus. The critical factors for purposes of this evaluation are the procedure itself. What maintenance checks are routinely performed? How is calibration done and the frequency and regularity with which it is carried out? This information should appear in the instrument calibration and maintenance logs and the laboratory quality control manual. If not, it will have to be obtained directly in conversations with the analysts and their supervisors. In either case, it will be important to discuss laboratory calibration and maintenance practices in the onsite visit and how to ascertain insofar as possible what is actually done and how frequently for each instrument. The evaluator should look for calibration tags on major pieces of instrumentation.

Ideally, the evaluation would involve simply comparing this laboratory's practices to generally accepted standards, summarized in some Table or reference test. Unfortunately, there is no such Table or text that covers all instruments and apparatus employed in environmental monitoring.

However, calibration recommendations for some of the major instruments are included in Table I. These "recommendations" are not to be considered rigid rules but rather guidelines for the evaluator in estimating laboratory performance. It is recognized that optimum procedures may vary somewhat as a function of instrument manufacturer and model. Additional materials that could be useful to the evaluator are operation and maintenance manuals for the various instruments and references in the Bibliography.

TABLE 1. INSTRUMENT CALIBRATIONS\*

Instrument	Procedure	Frequency
1) Analytical Balances	<ul><li>(a) Zero</li><li>(b) Standard weights</li><li>(c) Full calibration and adjustment</li></ul>	Before each use Monthly Annually
2) pH Meters	At pH 4,7, and 10	Daily
3) Conductivity Meters	(a) Obtain cell constant with potassium chloride reference solutions	Daily
	(b) Construct temperature curve if measurements are to be made other than at $25 \pm 0.5^{\circ}$ C	Monthly
4) Nephelometer/ Turbidimeters	(a) Check instrument scales or develop calibration curve with formazin stds. (<40NTU)	Monthly
	(b) If manufacturer's stds. are not formazine, check against formazine stds. (< 40NTU)	Annually
5) Colorimeters/Filter Photometers	Curves determined with 5-6 laboratory-prepared std. solutions for each parameter in conc. range of samples	Daily
6) UV/Visible Spectrophotometers	(a) Wavelength calibration with holimum oxide glass or solution, low-pressure mercury arc, benzene vapor (UV), or hydrogen arc (visible)	Quarterly
·	(b) Absorbance vs. concentra- tion curves with 5-6 std. Solutions for each param- eter at analytical wave- length in conc. range of samples	Daily
*Continued	(c) Full servicing and adjust- ment	Annually

TABLE 1. (Continued)\*

	Instrument	Procedure	Frequency
7)	Infrared Spectro- photometers	(a) Wavelength calibration with polystyrene or indene	Daily
		(b) Absorbance vs. concentration curves with 5-6 std. solutions for each parameter at analytical wavelength in conc. range of samples	Daily
		(c) Full servicing and adjust- ment	Semi-annually
8)	Atomic Absorption Spectrophotometers	(a) Response vs. concentration curves with 6-8 std. solutions for each metal (std. mixtures are acceptable, but with same acid as samples to be run) in conc. range of samples	Daily
		(b) Full servicing and adjust- ment	Annually
9)	Carbon Analyzers	Curves determined with 5-6 std. solutions in conc. range of samples	Daily
10)	DO Meters	Calibrated against modified Winkler method on aerated distilled or tap water	Daily
11)	Other Selective Ion Electrodes and Electrometers	Curves determined with 5-6 std. solutions in conc. range of samples	Daily
12)	Thermometers	Calibrate in constant temperature baths at two temperatures against precision thermometers certified by NBS.	Quarterly
13)	Technicon Auto Analyzers	(a) Curves determined with std. solutions for each parameter.	Each set of samples
		(b) Full service and adjust- ment (esp. colorimeter)	Annually

<sup>\*</sup>Continued

## TABLE 1. (Continued)\*

Instrument		Procedure	Frequency
14)	Gas Chromatographs	<ul> <li>(a) Retention times and detector response checked with std. solutions</li> <li>(b) Response curves for each parameter determined with std. solutions</li> </ul>	Daily Monthly
15)	Radiological Equipment	(See Standard Methods, Sect. 300	)
16)	Sulfur Dioxide in Air Sampler/Analyzers (Pararosaniline Method)	<ul> <li>(a) Calibrate flowmeters and hypodermic needle against a wet test meter</li> <li>(b) Spectrophotometric calibration curve with 5-6 std. sulfite-TCM solutions at controlled temperature (+ 1°C)</li> </ul>	Quarterly (Nee- dles before and after each run) Monthly
		(c) Sampling calibration curve with 5-6 std. atomospheres from permeation tubes or cylinders	Monthly
		(d) Calibrate associated thermometers, barometers, and spectrophotometer (wave- length)	- Quarterly
17)	Suspended Particulates (High-volume Sampler Method)	(a) Calibrate sampler (curve of true airflow rate vs. rota- meter or recorder reading) with orifice calibration unit and differential manom- eter at 6 air flow rates.	Monthly
		(b) Calibrate orifice calibration unit with positive displacement primary standard and differential manometers	Annually
		(c) Calibrate relative humidity indicator in the condition- ing environment against wet- bulb/dry-bulb psychrometer	Semi-annually
		<ul><li>(d) Check elapsed time indicator</li><li>(e) Calibrate associated analytical balances, thermometers, barometers</li></ul>	Semi-annually As needed

# TABLE 1. (Continued)\*

Instrument	Procedure	Frequency
18) Carbon monoxide (Non-dispersive IR)	(a) Determine linearity of detector response (cali- bration curve) with cali- bration gases (0, 10, 20, 40, and 80% of full scale, certified to ±2% and checked against auditing gases certified to +1%)	Monthly
	<ul> <li>(b) Perform zero and span calibrations</li> <li>(c) Calibrate rotameter and sample cell pressure gauge</li> </ul>	Daily or every three days Semi-annually
19) Photochemical Oxidants (Ozone)	(a) Calibrate standard KI/I2 solutions in terms of calculated O3 equivalents at 352 nm	At same time as ozone generator
	(b) Calibrate instrument response with 6-8 test atmospheres from ozone generator, spanning expected ranges of sample concentrations (usually 0.05-0.5 ppm 03)	Monthly
	(c) Calibrate flowmeters, barom- eter, thermometer	Semi-annually
	(d) Calibrate and service spec- trophotometer	As specified
	(e) Calibrate ozone generator	Monthly .
20) Hydrocarbons (corrected for Methane)	(a) Determine linearity of detector response (calibration curve) with calibration gases (0, 10, 20, 40, and 80% of fuscale, certified to ±2% and checked against auditing gages certified to +1%)	
	(b) Perform zero and span cali- brations	Before and after each sampling
	(c) Calibrate flowmeters and other associated apparatus	period Semi-annually

TABLE 1. (Continued)

Instrument			Procedure	Frequency
21)	Nitrogen Dioxide (Arsenite 24 hr.	(a)	Calibrate flowmeter with wet test meter	Monthly
	Sampling Method)	(b)	Calibrate Hypodermic needle (flow restrictor) with flowmeter	Each new needle and before and after each run
		(c)	Obtain colorimetric calibra- tion curves with 5-6 std. nitrite solutions	Weekly
22)	Nitrogen Dioxide (Chemiluminescence, Continuous)	(a)	Determine linearity of detector response (calibration curve) with calibration gases (0, 10, 20, 40, and 80% of full scale, certified to ±2% and checked against auditing gases certified to ±1%)	Monthly
		(b)	Perform zero and span cali- brations	Daily or every three days
		(c)	Calibrate rotameter and sample cell pressure gauge	Semi-annually
23)	Autoclaves and Sterilizers	(a)	Sterilization effectiveness checked (e.g., B. stearo-thermophilus, color-indicator tape for ethylene oxide)	Daily
		(b)	Temperature-recording device calibrated	Semi-annually

#### Control of Sampling

Sampling Plans and Sampling Equipment - The intent of this
item is to determine whether adequate attention has been
given to planning for sampling, whether appropriate sampling
instruments are available, and whether they are used properly.

Sampling is the operation of removing a part which is of convenient size for testing from a much larger whole substance in such a way that the measure of the characteristic of interest (such as pH or chemical analysis) in the sample is identical, within measureable limits of error, to that characteristic's presence in the whole substance. It is necessary that sampling be planned carefully in order to measure and control sampling errors and minimize the cost of sampling and testing.

If the substance to be sampled consists of discrete, constant, identifiable units (as do agricultural commodities tested for pesticide residues) standard sampling tables may be used to determine sample size. However, in environmental sampling the media are of a bulk nature (air, water, etc.) and the sampling units must be created by means of a sampling device, such as a bottle or sampling tube. The quantity and often the form of the sample units depends on the particular device, how it was used, and on the location and condition of the substance being sampled.

Sampling may be instantaneous at a given station (grab sampling) or continuous and automatic. Validity of sampling depends on randomness of selection of the samples. Where stratification exists, random samples must be taken from each stratum in proportion to its size. When the statistical criteria have been met, the required sample size may be calculated.

The design of sampling is seen to require some special skills and the person responsible for it must have considerable sophistication in handling the statistical aspects.

Generally, sampling instruments and their use are described in the analytical methods and questions related to sampling should be asked for each test or group of test methods.

• Sample Collection and Preservation - The evaluator will want to determine sample taking and preservation practices for at least some of the tests performed by the laboratory. For evaluation purposes, these practices can be compared with the recommendations incorporated in Table 2 of the EPA Manual - Methods for the Chemical Analysis of Water and Wastes,

for most water parameters; the Federal Register for air parameters (Sect. 4, Precision, Accuracy and Stability, for each method); Standard Methods: Sections 405 (microbiological), 200 and 300A (radiological); and the specified references in the Analytical Methods Table C, Section 4, for the remaining parameters (57-60).

The holding time given in Table 2 of the EPA Manual is interpreted as the recommended maximum period between sampling and analysis. Preservatives, where specified, are required to ensure stability for the holding time. Look at records, at sample bottles, etc., to assure yourself that good procedures are actually followed. If holding times are exceeded, a notation of that fact should be made on data sheets before they are transmitted.

For some tests, to exceed the maximum holding time would very seriously compromise the accuracy of the measurement. If the laboratory is exceeding the maximum holding time for these tests, the laboratory must be given a score of 1\* and the problem must be resolved before a final score is calculated. The parameters to which this applies include the following:

Biochemical Oxygen Demand (Dissolved Oxygen)

Cyanide, Total

Chlorine, Total Residual

Pheno1s

Turbidity

Streptococci

Coliform Bacteria

Temperature

pН

• Identification and Storage of Samples - All samples should be clearly marked with a code number at the time of sampling. Labels should be securely attached to the sample container. In the field, information about the sample should be entered immediately in a field notebook. In handling and storing the samples precautions should be taken against mix-up in identification.

Storage space should permit storage of samples in a separate area, refrigerated if necessary for preservation, and secured against tampering.

- Laboratory Handling of Sampling The flow of samples through the laboratory should be organized. Forms should be available for requests for analysis and for reporting of results. Sample handling procedures should be formalized so that samples arriving at the lab are accepted, prepared and analyzed promptly. Holding times given in Table 2 should be adhered to. For air, requirements given in the referenced methods should be followed.
- Chain of Custody Assignment of responsibility for custody of samples should be clear and the importance of a tight system of control should be understood by all. The procedures to be followed should be written. Samples should be logged in and their progress through the labs should be recorded and the samples themselves should be in a secure location when not signed out to an analyst.
- Control of Field Sampling/Measurements The requirements of sampling in the field are as demanding as those of sampling in the laboratory. Most sample taking is a field operation. Sometimes measurement also must be done in the field. Certain special analytical methods or modifications of standard methods apply. Also, other measurements, such as flow rates, not made in the laboratory must be done in the field.

Questions should be directed toward an understanding of how well the field aspects of sampling and testing are attended to when done by laboratory personnel or when done by a service agency.

• Control of Monitoring - The important thing to be checked for in this item is whether written procedures cover all monitoring activities in which the laboratory is engaged and whether they are being followed exactly.

TABLE 2. RECOMMENDATION FOR SAMPLING AND PRESERVATION OF SAMPLES ACCORDING TO MEASUREMENT (1)\*

Measurement	Vol. Req. (ml)	Container	Preservative	Holding Time(6)
Acidity	100	P, G <sup>(2)</sup>	Cool, 4°C	24 Hrs.
Alkalinity	100	P, G	Cool, 4°C	24 Hrs.
Arsenic	100	P, G	$HNO_3$ to $pH < 2$	6 Mos.
BOD	1000	P, G	Cool, 4°C	6 Hrs. (3)
Bromide	100	P, G	Cool, 4°C	24 Hrs.
COD	50	P, G	$H_2SO_4$ to pH < 2	7 Days
Chloride	50	P, G	None Req.	7 Days
Chlorine Req.	50	P, G	Cool, 4°C	24 Hrs.
Color	50	P, G	Cool, 4°C	24 Hrs.
Cyanides	500	P, G	Cool, 4 <sup>O</sup> C NaOH to pH 12	24 Hrs.
Dissolved Oxygen Probe	300	G only	Det. on site	No Holding
Winkler	300	G only	Fix on site	No Holding
Fluoride	300	P, G	Cool, 4°C	7 Days
Hardness	100	P, G	Cool, 4°C	7 Days
Iodide	100	P, G	Cool, 4 <sup>o</sup> C	24 Hrs.
MBAS	250	P, G	Cool, 4°C	24 Hrs.
Metals Dissolved	200	P, G	Filter on site HNO <sub>3</sub> to pH < 2	6 Mos.
Suspended			Filter on site	6 Mos.
Total	100		$HNO_3$ to $pH < 2$	6 Mos.
*Continued				

TABLE 2. (Continued)\*

Measurement	Vol. Req. (ml)	Container	Preservative	Holding Time(6)
Mercury	· · · · · · · · · · · · · · · · · · ·			•
Dissolved	100	P, G	Filter	38 Days
•			$HNO_3$ to $pH < 2$	(Glass)
•			to the second second	13 Days
×.	•			(Hard Plastic)
Total	100	P, G	HNO <sub>3</sub> to pH < 2	38 Days
			, and the second	(Glass)
				13 Days
			(	(Hard Plastic)
Nitrogen				
Ammonia	400	P, G	Cool, 4°C	24 Hrs. (4)
			$H_2SO_4$ to pH < 2	
Kjeldahl	500	P, G	Cool, 4°C	24 Hrs. (4)
			$H_2SO_4$ to pH < 2	
Nitrate	100	P, G	Cool, 4°C	24 Hrs. (4)
			$H_2SO_4$ to pH < 2	
Nitrite	50	P, G	Cool, 4°C	24 Hrs. (4)
NTA	50	P, 'G	Cool, 4°C	24 Hrs.
Oil & Grease	1000	G only	Cool, 4°C	24 Hrs.
			$H_2SO_4$ to pH < 2	
Organic Carbon	25	P, G	Cool, $4^{\circ}$ C $H_2$ SO <sub>4</sub> to pH < 2	24 Hrs.

TABLE 2. (Continued)\*

<del></del>	Vol.	<del></del>	<del></del>	
Measurement	Req. (m1)	Container	Preservative	Holding Time(6)
рН	25	P, G	Cool, 4°C	6 Hrs. (3)
•			Det. on site	
Phenolics	500	G only	Cool, 4°C H <sub>3</sub> Po <sub>4</sub> to pH < 4 1.0 g CuSO <sub>4</sub> /1	24 Hrs.
Phosphorus				
Ortho- phosphate	50	P, G	Filter on site	24 Hrs. (4)
Dissolved			Cool, 4 <sup>o</sup> C	
Hydrolyzable	50	P, G	Cool, $4^{\circ}$ C $H_2$ SO <sub>4</sub> to pH < 2	24 Hrs. (4)
Total	50	P, G	Cool, 4 <sup>0</sup> C	24 Hrs. (4
Total, Dissolved	<b>50</b>	P, G	Filter on site	24 Hrs. (4
Residue				•
Filterable	100	P, G	Cool, 4°C	7 Days
Non- Filterable	100	P, G	Cool, 4°C	7 Days
Total	100	P, G	Cool, 4°C	7 Days
Volatile	100	P, G	Cool, 4°C	7 Days
Settleable Matter	1000	P, G	None Req.	24 Hrs.
Selenium	50	P, G	HNO <sub>3</sub> to pH < 2	6 Mos.
Silica	50	P only	Cool, 4°C	7 Days
Specific Conductance	100	P, G	Cool, 4°C	24 Hrs. (5
*Continued		· 158		

TABLE 2. (Continued)

Measurement	Vol. Req. (ml)	Container	Preservative	Holding Time(6)
Sulfate	50	P, G	Cool, 4°C	7 Days
Sulfide	50	P, G	2 ml zinc acetate	24 Hrs.
Sulfite	50	P, G	Cool, 4°C	24 Hrs.
Temperature	1000	P, G	Det. on site	No Holding
Threshold Odor	200	G only	Cool, 4°C	24 Hrs.
Turbidity	100	P, G	Cool, 4°C	7 Days

<sup>1.</sup> More specific instructions for preservation and sampling are found with each procedure as detailed in this manual. A general discussion on sampling water and industrial wastewater may be found in ASTM, Part 23, p. 72-91 (1973).

#### 2. Plastic or Glass

- 3. If samples cannot be returned to the laboratory in less than 6 hours and holding time exceeds this limit, the final reported data should indicate the actual holding time.
- 4. Mercuric chloride may be used as an alternate preservative at a concentration of 40 mg/l, especially if a longer holding time is required. However, the use of mercuric chloride is discouraged whenever possible.
- 5. If the sample is stabilized by cooling, it should be warmed to 25°C for reading, or temperature correction made and results reported at 25°C.
- 6. It has been shown that samples properly preserved may be held for extended periods beyond the recommended holding time.

#### Quality Control

- Quality Policy To ascertain that quality control is a pervasive concern; one that merits attention not only at critical points, but daily in the routine performance of analyses. There should be a clear statement of policy by management.
- Quality Program Manual To identify the means by which quality control procedures are disseminated in the laboratory.
- Responsibility for Quality To determine which person or group of people assumes responsibility for quality control.
- Training in Quality Control To determine what measures are used to prepare employees to meet quality control standards.
- Control of Chemicals and Reagents To assess the laboratory's methods for monitoring the flow of chemicals and reagents. Procurement control includes equipment and other materials as well as chemicals and reagents.
- Intra-Laboratory Checks; Precision and Accuracy An analytical laboratory must have a well-organized and clearly defined program to check the validity of the data it produces. Validity is usually expressed in terms of precision and accuracy. According to the EPA Handbook for Analytical Quality Control in Water and Wastewater Laboratories, "precision refers to the reproducibility among replicate observations", and "accuracy refers to a degree of difference between observed and known, or actual values".

An analyst initially may establish the precision of a particular method by 5-10 replicate determinations on a single "real sample". Generally, it will be necessary to repeat this procedure on each of the various types of samples that will be analyzed by this method (e.g., surface water, industrial effluent, sea water, etc) and preferably on several samples of each type from a variety of sources. Comparison of the precision obtained with reference standards and that obtained with actual samples will reveal any interferences from contaminants in the complex samples.

The accuracy of a method may be determined initially by 5-10 replicate analyses of samples to which known amounts of reference standards have been added (spiked samples). The EPA AQC Handbook mentioned above suggests reporting the results as "percent recovery at the final concentration of the spiked sample". The spiking of actual samples for these determinations allows for a more realistic measurement of accuracy than the exclusive use of pure reference standards, although again comparison of the accuracy obtained with spiked samples and that obtained with reference standards may be of interest in

identifying the source of errors. Analysis of blanks also will be important for many parameters where the apparent background level may be non-zero and where a blank correction may be necessary.

Routine Checks of Testing Performance - After the precision and accuracy of the method are established, the analyst will need to incorporate replicates, spikes, standards, and blanks, as appropriate, into the sequence of routine analyses to ensure that valid data is being generated. The frequency and procedures required for adequate monitoring of the quality of the data will depend on the method itself. The evaluator will find some guidance as to what is adequate in the references in the Bibliography, particularly in the EPA AQC Handbook mentioned above, the EPA Guidelines for the Development of a Quality Assurance Program (for various air parameters), and the Methods Manuals (EPA, ASTM, Standard Methods). The experience of conscientious analysts and statisticians in the field of environmental monitoring is an invaluable source in this matter. For example, one group of water chemists experienced on the Technicon Auto Analyzer usually runs a duplicate, a spiked sample, and a reference standard every 8 samples in a large series of similar samples, or once in each set of samples, whichever is more frequent. A chemist experienced in the analysis of Phenols and Cyanide suggests verifying the standard curves each day that these parameters are analyzed with a low and a high reference standard and a blank and running a duplicate and a spike with each small set of samples. Gas chromatography often requires multiple injections of the sample with and without an internal standard, in addition to spiked samples and a blank, for each sample analyzed. These examples are given only to demonstrate how quality control protocols will vary considerably with the method and the experience of the analyst. The nature of the samples (simple or complex mixtures), the condition of the instrument, the importance of the sample (e.g., for enforcement action), the breadth of the precision and accuracy control limits, and many other factors may also affect the quality control requirements.

Because there are no universal guidelines for the frequency and procedures required in the use of quality control samples, it is very important that each laboratory develop its own internal guidelines based on sound statistical methods and experience. These should be in the form of written, explicit protocols for each test or group of tests. Statistical methods for the development of such protocols are discussed in the quality control references in the Bibliography and in standard quality control texts.

For purposes of this evaluation it will be of primary importance to determine if the analyst and the laboratory have a proper appreciation of the importance of replicates, spikes, standards, and blanks in assuring the validity. of their analytical data. Since the evaluator is not expected to be an expert with long experience in the performance of every method, this evaluation does not place heavy emphasis on the content of the detailed protocols for replicates, spikes, standards, and blanks used by the laboratory. Rather, emphasis is laid on an assessment of the concern for and awareness of quality control evidenced and practiced by the analyst and the laboratory as a whole, as discussed below. The evaluator is asked to make a judgment as to whether quality control samples are run with sufficient frequency, but it is recognized that the evaluator may have little experience in many methods and may wish to place proportionately little weight on this judgment. The evaluator, nonetheless, should carefully record and document laboratory practices, so that patterns of quality control procedures can be developed.

The evaluator will want to discuss in the onsite visit the actual laboratory protocol for the use of replicates, spiked samples, reference standards, and blanks for each test. Some tests, of course, can be considered in groups with similar requirements (e.g., metals determined by atomic absorption or many of the tests determined on the Technicon Auto Analyzer). Questions to be asked by the evaluator for each parameter (method) include the following:

Is there a formal protocol in this lab for the control of analytical performance of this method, including specifications of the <u>frequency</u> of and <u>procedures</u> for replicate sample, spiked sample, reference standard, and blank analyses, where applicable?

Are the analysts familiar with the protocol? Does the protocol appear to vary from analyst to analyst?

Have the precision and accuracy of the method been determined in this laboratory? By each analyst using the method? How frequently?

Are replicates, spiked samples, reference standards, and blanks, if applicable, run with sufficient frequency to assure that precision and accuracy are remaining within the control limits?

Is there a well-defined and clearly understood procedure for evaluating the data and for handling "out-of-control" data?

Have you developed acceptance criteria for data (could be three-sigma limits)? Is corrective action taken on lack of control? One of the basic procedures of statistical quality control is to associate troubles with specific causes. Does the laboratory try to do this?

The answers to these and other questions the evaluator may develop should offer a clear impression of the effort devoted by laboratory and analyst to assuring that valid data is produced for each parameter.

The score given is to be based on the laboratory's quality control procedures, particularly as they relate to replicates, spiked samples, reference standards, and blanks, if applicable, and the analyst's familiarity and understanding of the procedures.

- Statistical Methods A popular method of monitoring daily performance is the use of Quality Control Charts. Basically, these charts, constructed separately for each test, display control limits for precision and accuracy. The precision and accuracy measured from day to day are plotted on these charts which provide a continuous visual picture of the control of data quality. Details will be found in textbooks on Quality Control and in the two EPA publications, Handbook for Analytical Quality Control in Water and Wastewater Analysis and Quality Control Practices in Processing Air Pollution Samples. The control chart method is particularly helpful is assisting in identifying causes of trouble in the measurement process: both special causes within the power of the analyst to correct and general causes, such as fluctuations in the laboratory environment, which are the duty of management to correct.
- Inter-Laboratory Proficiency Tests Refer to Chart E of the Preliminary Questionnaire. Question the lab about results of participation in formal programs. Ask questions about cooperation with peer laboratories in the exchange of split samples, as another sort of inter-laboratory control.
- Laboratory Records Accurate records provide a means for the laboratory to monitor its workload, locate errors, and evaluate its own progress. All three functions contribute to quality control and, therefore, should be assessed from this perspective. How does management decide whether data are satisfactory? Can data be rejected in this laboratory? (i.e., Are new samples collected and analyzed if results are suspect?) Are results recorded in an acceptable manner (in a notebook, on bench cards, or on NCR data forms)?

• Laboratory Reports - Regularly scheduled laboratory reports may function as a catalyst to continuous awareness of the importance of quality control. They are evidence both of managements' demand and analysts' effort to achieve excellence in quality control.

## **Evaluator's Notes**

# PART 6. INTERNAL AND EXTERNAL CONTROLS

## 1. Control of Analytical Methods and Instruments

# (1) Assignment of Responsibility for Maintenance and Calibration

•	and Ca	alibration	
		Best Description of Laboratory	Score
		Responsibility is clearly assigned in this laboratory and understood by all personnel.	5
-		Responsibility is assigned but not clearly recognized or understood by assignee(s) or other personnel.	3
		Responsibility is not clearly assigned or recognized in this laboratory.	. 1
(2)	Mainte	enance and Calibration Logs	
		Best Description of Laboratory	Score
		The instrument logs are properly executed, complete, and up-to-date.	5
		An instrument log exists but is faulty.	3
		An instrument log does not exist.	1
(3)	Adequ	acy of Calibration and Maintenance Practices  Best Description of Laboratory	Score
		Calibration and maintenance of instruments is adequate.	5
		Marginal.	3
		Inadequate.	1*

# 2. Control of Sampling

(4)	Sampling Plans and Sampling Equipment			
		Best Description of Laboratory	Score	
		Samples are carefully designed, suitable sampling equipment is on hand and is used properly.	5	
		Sampling is taken for granted and no particular efforts are made to assure validity of samples.	3	
		Sampling is not organized, equipment is poor, or insufficient care is taken in obtaining the samples.	. 1	
(5)	Sample	e Collection and Preservation		
		Best Description of Laboratory	Score	
		Samples are kept in proper containers using the recommended preservative for no longer than the recommended maximum holding time.	. 5	
		When possible, the recommended procedures for collection and preservation are followed, although circumstances (laboratory manpower, lack of control over sample taking, variability of workload, etc.) do not always allow strict adherence.	3	
		The laboratory often does not follow EPA recommendations for maximum holding time, preservation technique, and/or container type.	1	

(6)	Identi'ication and Storage of Samples			
		Best Description of Laboratory	Score	
		Samples are carefully and clearly identified by code number and stored so as to protect their identity and security.	5	
		Sample identification system and storage of samples not well organized.	3	
		There are serious defects in sample identification and storage practices that could lead to serious mix-ups.	1*	
(7)	Labora	atory Handling of Samples		
		Best Description of Laboratory	Score	
		Activities of the laboratory are well organized so that samples are given the attention required and work proceeds smoothly from sample receipt to report of results.	5	
		Procedures for assuring smooth flow of samples through the laboratory are not complete.	3	
		The system and load are not well matched so that there is a backlog of work and time requirements are sometimes missed.	1	
(8)	Chain	of Custody		
-		Best Description of Laboratory	Score	
		A chain of custody procedure is followed precisely, with clearly assigned responsibility, complete recording of activities, and careful security of samples.	5	
		A chain of custody procedure exists but it is lax and not strictly followed.	3	
		Chain of custody is not formally organized.	1*	

(9)	Contr	ol of Field Sampling/Measurements  Best Description of Laboratory	Score
·		Written procedures for field sampling/ measurements are complete and are followed meticulously under surveillance by the laboratory.	5
		Field sampling/measurement are subject to standard methods but surveillance by the laboratory is lax.	3
		Field sampling/measurement is not treated as a major concern of the laboratory.	, 1
		† NOTE	
		If the laboratory does not participate in this activity, do not score it and subtract 5 from the denominator of the fraction in the formula for calculating its score for internal and external controls.	
(10)	Contr	ol of Monitoring †	
		Best Description of Laboratory	Score
		Written procedures which are followed exactly are available for all monitoring activities in which this laboratory is engaged.	5
		Written procedures exist but they are incomplete and not followed exactly.	3
		No written procedures exist.	1
		† <sub>NOTE</sub>	
		If the laboratory does not participate in this activity, do not score it and subtract 5 from the denominator of the fraction in the formula for calculating its score for internal and external controls.	

# 3. Quality Control

(11) Qual	ity Policy	•
	Best Description of Laboratory	Score
	A clear statement of quality objectives by the top executive exists with continuing visible evidence of its sincerity to all levels of the organization.	5
	Periodic meetings among the section heads of service, research and development, and quality assurance are held to discuss quality objectives and progress toward their achievement.	3
	There was a "one-shot" statement of the desire for product quality by the top executive after which the quality control staff is on its own.	1
(12) Qual	ity Program Manual	
	Best Description of Laboratory	Score
	Formalized and documented by a set of procedures which clearly describe the activities necessary and sufficient to achieve desired quality objectives.  This may be in the form of a Quality Control Manual.	5
	The Quality Program is contained in methods procedures or is implicit in those procedures. Experience with the materials, product and equipment is needed for continuity of control.	3
	The Quality Program is undefined in any procedures and is left to the current managers or supervisors to determine as the situation dictates.	1*

(13) Respo	onsibility for Quality  Best Description of Laboratory	Score
	Responsibility for quality is a full-time assignment of a quality control department with well-defined authority or in smaller laboratories is clearly defined for all	
. 🗆	Responsibility for quality is assigned to a part-time quality control coordinator who must use whatever means possible to achieve quality goals.	3
	Responsibility for quality is not defined.	1*
(14) Train	ing for Quality Control	
	Best Description of Laboratory	Score
	The people who have an impact on quality (bench chemists, supervisors, etc.) are trained in the reasons for and the benefits of standards of quality and the methods by which high quality can be achieved.	5
	Personnel are told about quality only when their work falls below acceptable levels.	3
	Personnel are reprimanded when quality deficiencies are directly traceable to the chemists' analytical work.	1
(15) Contr	rol of Chemicals and Reagents	
	Best Description of Laboratory	Score
	Reagents and chemicals are inspected upon receipt and accepted only if they conform to all specifications. In inventory they are identified as to type and age and issued on a first in/first out plan.	5
	Reagents and chemicals are only spot checked for quantity and shipping damage; in storage they are identified as to material only and are issued randomly.	3
	Reagents and chemicals are not checked on receipt, are not clearly identified, and are issued on a last in/first out basis.	1

(16) Intral	aboratory Checks - Precision and Accuracy	
	Best Description of Laboratory	Score
	Laboratory has a well-organized program to check the validity of data it produces.	5
	Incomplete information is available on precision and accuracy of the tests in use.	3
	Laboratory has no plan to check on validity of its data.	. 1
(17) Rout	ine Checks of Testing Performance	
	Best Description of Laboratory	Score
	Procedures are excellent and should provide adequate assurance that the data is valid.	5
	Procedures are fair and should provide some indication of the validity of the data.	3
	Procedures are poor or poorly defined and do not provide adequate assurance that the data is valid.	1*
(18) Statis	stical Methods	
` '	Best Description of Laboratory	Score
	Use is made of statistical methods, such as control charts to insure continuing validity of tests.	5
	Some statistical checks of measure- ments are made but level of assurance of quality is uncertain	3
	No efforts are made to use statistical methods of quality control.	1

(19) Inter	laboratory Proficiency Tests	
	Best Description of Laboratory	Score
	The laboratory has a good record of participation in formal proficiency tseting and has a good record of performance.	5
	Laboratory participates only sporadically and not recently. Performance in programs not outstanding.	3
	Laboratory does not participate in proficiency testing programs.	1*
(20) Labo	oratory Records	
	Best Description of Laboratory	Score
	Analytical results are entered in a lab notebook or in a card system which is signed and witnessed. Results are summarized and entered in appropriate data system promptly.	5
	Analytical results are complete but they are not routinely signed and witnessed. Data processing is not always prompt.	3
	Data keeping is not organized, i.e., results kept on loose sheets of paper and incompletely reviewed and analyzed.	1
(21) Lab	oratory Reports	
	Best Description of Laboratory	Score
	Lab activities are reported regularly and periodic quality reports are made to feed forward to management and to feed back to bench analysts quality of the work reported.	5
	Laboratory reports are sporadic and quality reports do not result in bringing necessary information for action on quality to all levels of the organization.	3
	Reports are very irregular and no system for quality reporting exists.	1

#### FOLLOW-UP ON DEFICIENCIES

The goal of laboratory evaluation is the improvement of laboratory performance. Identification of deficiencies is not intended to bar a laboratory from participation in environmental monitoring. Rather, it indicates that improvements are necessary to enable the laboratory to fulfill its role optimally.

Certain aspects of laboratory activity are more crucial to successful environmental monitoring than are others. It is the evaluator's responsibility to insist that rigid standards are met in these critical areas before the laboratory receives a final score. In the Onsite Check List, problems which must be resolved to the evaluator's satisfaction prior to approval are marked by an asterisk next to the lowest possible score (1\*).

Unacceptable deficiencies may be indicated in each area of laboratory evaluation: Consistently high turnover rates, customer complaints, lack of cooperation among laboratory employees, and obstacles to internal communication are symptoms of poor organization and management which could seriously impair laboratory operation. Supervisors who have neither degrees nor sufficient experience may jeopardize the laboratory's analytical capabilities. Inadequate space, whether it be laboratory space, storage space or controlled space, impedes orderly laboratory functioning. Incomplete safety equipment may endanger both successful analyses and laboratory personnel. The use of nonstandard methods, the absence of essential instruments, or the malfunction of instruments as a result of improper maintenance, may compromise all analytical results. Failure to employ rigid quality control procedures may also raise serious doubts concerning the validity of laboratory data. If the quality assurance plan is not clearly defined, and responsibility for its execution is not assigned; if a chain of custody of samples is not established and followed; if sample storage exceeds the recommended maximum holding time; or if calibration is inadequate; the reliability of the laboratory's work may be impugned.

To protect the scientific and legal defensibility of the data, the evaluator must ensure that environmental monitoring laboratories are free of these deficiencies. Any inadequacies discovered by the evaluator should be brought to the attention of the laboratory management immediately, before completion of the onsite visit. The evaluator may offer recommendations for remedial action or stipulate essential adjustments which must be made before the laboratory may be scored.

After discussion with laboratory management, the evaluator should make note of the exchange and then compute a <u>tentative</u> score for the laboratory. The final score cannot be computed, nor approval given, until the laboratory has submitted evidence that all deficiencies have been corrected.

## SECTION 7 CALCULATION OF SCORE

#### ACCEPTABILITY OF A LABORATORY

The Procedure for the Evaluation of Environmental Monitoring Laboratories strives to construct a standardized system for the objective appraisal of laboratory management, personnel, equipment, analytical capabilities and quality control procedures. The numerical scoring system plays an integral role in achieving this end. It provides a means to organize the multiplicity of data and to produce a manageable result. The values assigned to individual characteristics of the laboratory affect the total score by very small increments. This affords a measure of uniformity to laboratory assessment which is essential for the comparison of results compiled by a variety of evaluators in diverse situations.

The numberical scoring system is based upon 100 points. Each item may be rated with 5 points, 3 points, or 1 point. While onsite, the evaluator should check on the score sheets the scoring level for each item. If the level checked is scored one followed by an asterisk (1\*) the laboratory fails to meet required specifications. The laboratory must resolve the deficiency before a final score can be computed.

After the onsite survey has been completed, the evaluator should use the summary sheets to calculate the numerical scores. On these forms, each item's score is weighted according to its importance for successful laboratory operation. After summing the weighted scores, performing the calculation at the bottom of the page produces the final score for each section.

Addition of the scores for each section provides the laboratory's final evaluation score. The highest possible score is 100 points. The minimum acceptable score is 60 points. Laboratories which score below this minimum require major improvements to be capable of participation in environmental monitoring programs.

If separate scores are desired for each section of a laboratory which deals with different media, the evaluator must have completed during the onsite visit a set of score sheets for Part 4 Analytical Methods and Part 5 Instruments for each section. In the event that this has been done, a total score is obtained for each section of the laboratory by adding to the separate scores on Parts 4 and 5, the general scores given the Laboratory on Management and Organization (Parts 1, 2, 3) plus Part 6, Internal and External Controls. Thus, a laboratory may obtain an overall score or two or more scores covering individual media with which it is concerned.

When the evaluator has computed the score for a laboratory, this score coupled with the evaluator's recommendations and comments should be sent to the participating lab. Laboratories which fail to meet required standards may later submit proof of adjustments made in compliance with the evaluator's recommendations to receive an upward revision.

In special circumstances, such as in evaluating very small laboratories, it may be desirable to drop one or more questions from the onsite score sheets. This should be done only after due deliberation by the evaluating agency. In no instance should the evaluation team arbitrarily eliminate or "forget" any question. If, for valid reasons, a question is dropped from a Part, the prorating of the score on the Scoring Forms may be accomplished as follows:

Multiply by five (5) the assigned weight in Column (2) of the question dropped and subtract the product from the denominator in the calculation of score for that part. Make such an adjustment for each question dropped. For example, if Question 4, Bench-Top Space, is dropped from Part 3, Laboratory Space and Facilities, the weight (Col. 2) is 1, the denominator 100 is reduced by 5x1 to 95 and calculation of score proceeds as indicated.

A report to the laboratory management might contain the following sections:

- 1. Recommendations to improve overall performance
- 2. Amplification of recommendations for any equipment or instrument purchases.

# PART 1. SCORE FOR GENERAL INFORMATION ABOUT THE LABORATORY

Name of Labo	oratory	· · · · · · · · · · · · · · · · · · ·	<del> </del>	:-	• ,_		
			(1) Score	X	(2) Weight	=	(3) Extension
Question 1. Question 2. Question 3. Question 4. Question 5.	Appropriateness of Organization Impairment of Functions* Strength of Management* Objectivity of Laboratory* Cooperation Obtained*	1*			2 2 4 1	• • • •	;
		TOTAL		,** ·	٠.	•	
Calculation:						•	••
	$\frac{\text{Total Col (3)}}{50}  X  20  = $					,	
Enter this figu	re in box below and carry it forw	ard to Summary	y Evalua	ation.			
Score carried	forward to Summary Evaluation.					•	
*Any score of 1	in positions in Col (1) marked with an	asterisk must be re	esolved b	efore	the final sco	ore is o	calculated.
Date		Visit Condu	icted hi	,			

### PART 2. SCORE FOR PERSONNEL

Name of Labor	ratory	<u>-</u>				<del></del>	
,			(1) Score	x	(2) Weight	= E	(3) Extension
Question 1.	Supervisor Training				1		
Question 2.	Supervisor Experience*				2		
	Job Descriptions				1		
Question 4.	Training Program				2		
•	Turnover Rate* General Morale				2 2		
Question 6.	General Morale		•			_	
	ī	TOTAL					
Calculation:							
	$\frac{\text{Total Col (3)}}{50}  X  20$						·
Enter this figu	re in box below and carry it forward to	o Summary	Evaluat	ion.			
Score carried f	orward to Summary Evaluation. 🗆						
	·						
						•	
Any score of 1 in	positions in Col (1) marked with an asterisk	k must be res	olved befo	ore ti	ne final scor	e is calc	ulated.

### PART 3. SCORE FOR LABORATORY SPACE AND FACILITIES

Name of Labo	ratory					
		(1) Sco		(2) Weight	=	(3) Extension
Question 1.	General Characteristics of					
	Space and Facilities			1		
Question 2.	Office Space			1		
Question 3.	Laboratory Space*			2		
Question 4.	Bench-top Space			1		
Question 5.	Hood Space and Operation			1		
Question 6.	Storage Space - Chemicals			1		
Question 7.	Sample Storage Space*			2		
Question 8.	Controlled Space*			2		
Question 9.	Library			1		
Question 10.	Safety Equipment/Procedures*			2		
Question 11.	Distilled/Deionized Water*			2		
Question 12.	Glassware Supply and Washing*			2		
Question 13.	Housekeeping			1		
Question 14.	Data Processing Equipment					
<b>X</b>	and Logistic Services			1		
	u			•	•	
		TOTAL				
Calculation:						
	$\frac{\text{Total Col (3)}}{100}  X  10$					
Enter this figu	re in box below and carry it forward	l to Summary Eva	luation			
Score carried f	orward to Summary Evaluation. 🛘					
	,					
		•				
*Any score of 1	in positions in Col (1) marked with an ast	erisk must be resolve	ed before	the final sc	ore is c	alculated.
Date		Visit Conducted	hv			

### PART 4. SCORE FOR ANALYTICAL METHODS

Name of Labo	oratory				· .
			(1) >	( (2) Weight	= (3) Extension
Question 1.	Reference Methods or Approved Alternates*			1	
Question 2.	Reagent and Media Preparation			1	
Question 3.	Performance According to Standard			2	
		TOTAL			
Calculation:					
	$\frac{\text{Total Col (3)}}{20}  X  10$				
Score carried	forward to Summary Evaluation.				•
		•			
					•
*Any score of 1	in positions in Col (1) marked with an	asterisk must be	resolved befo	re the final sc	ore is calculated.
Date		Visit Cond	ucted by		

### PART 5. SCORE FOR INSTRUMENTS

Name of Laboratory					
		(1) Score	X (2) Weight	= (3) Extension	
Question 1. Question 2.	Required Instrumentation* Function Tests and Standardization of		1		
Question 3.	Instruments Calibration Equipment		2 1		
		TAL			
Calculation:					
	$\frac{\text{Total Col (3)}}{20}  X  10$				
Score carried f	orward to Summary Evaluation.				
				·	
*Any score of 1	*Any score of 1 in positions in Col'(1) marked with an asterisk must be resolved before the final score is calculated.				
Date	Vis	it Conducted by			

### PART 6. SCORE FOR INTERNAL AND EXTERNAL CONTROLS

Name of Labo	ratory				<del></del>	1 2 2 2 1 5
		(1) Score	X	Weight	=	(3) Extension
Question 1.	Responsibility for			o i statule a a Geografia		
	Calibration			1 y		
Question 2.	Adequacy of Calibration Logs			1		
Question 3.	Adequacy of Calibration and Maintenance Practices*			2		
Question 4.	Sampling Plans and Sampling Equipment			1		
Question 5.	Sample Collection and Preservation:					
Question 6.	Identification and Storage	٠.	•	۷.,	4 200	
Question o.	of Samples *			1		
Question 7.	Laboratory Handling of			. 1	** r	
Question 7.	Samples			1		
Question 8.	Chain of Custody*			2		
Question 9.	Field Control of Sampling			1		
Question 10.	Control of Monitoring					
	Activities			1		
Question 11.	Clarity of QC Policy			1		
Question 12.	Written Program/Manual*			1		
Question 13.	Responsibility for Quality*			1		
Question 14.	Training in QC			1		
Question 15.	Control of Chemicals and Reagents			1		
Question 16.	Internal Checks: Precision and					
Overtion 17	Accuracy Internal Checks: Routine			1		
Question 17.				2		
Question 18.	Duplicates, Blanks, Spikes* Statistical Methods			1		
Question 19.	Inter-lab Proficiency Tests*			1		
Question 19.	Record System			1		
Question 21.	Report System			i		
•	TOTAL					
Calculation:						
	Total Col (3) X 30					
Enter this figu	re in the box below and carry it forward to Sumn	nary Eva	aluat	ion.		
Score carried	Forward to Summary Evaluation.					
*Any score of 1	in positions in Col (1) marked with an asterisk must be re	solved b	efore	the final so	ore is	calculated.
Date	Visit Conduc	cted by				

### SUMMARY OF LABORATORY EVALUATION

		Score
		Score
Part 1.	General Information	
Part 2.	Personnel	
Part 3. Part 4.	Lab Space and Facilities  Tochnical Services (Applytical Methods)	
Part 5.	Technical Services (Analytical Methods)  Lab Equipment	
Part 6.	Internal and External Controls	
	TOTAL	
Inadequacies	marked by * in the score sheets have not been resolved and above	e is a tentative score. F
	marked by in the score sheets have not been resolved and abov	c is a tentative score.
Final Score	-E. P. P. P. P. P. P. P. P. P. P. P. P. P.	

Evaluation Completed by

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#### APPENDIX

#### MAJOR EQUIPMENT REQUIREMENTS FOR EACH ANALYTICAL METHOD

#### General Analytical Methods

- Alkalinity as CaCO<sub>3</sub> (mg CaCO<sub>3</sub>/liter)
  - (a) Electrometric Titration, Manual
    - pH meter, Type I or II as defined in ASTM D1293
  - (b) Electrometric Titration, Automated
    - An automatic titrimeter meeting the pH meter specifications in (a).
  - (c) Automated, Methyl Orange
    - Technicon AutoAnalyzer with
      - (1) Sampler I
      - (2) Continuous filter
      - (3) Manifold
      - (4) Proportioning pump
      - (5) Colorimeter with 15 mm tubular flow cell and 550 nm filters
      - (6) Recorder with range expander
- 2. Biochemical Oxygen Demand (B.O.D.) 5-day, 20°C (mg.liter)
  - (a) Modified Wrinkler with Full-Bottle
    - B.O.D. incubation bottles
  - (b) Probe Method
    - No specific probe is recommended as superior in the 1974 EPA Methods Manual, but ones evaluated and found reliable were Weston and Stack DO Analyzer Model 30, Yellow Springs Instrument (YSI) Model 54, and the Beckman Fieldlab Oxygen Analyzer.

- 3. Chemical Oxygen Demand (C.O.D.) (mg/liter)
  - (a) No special equipment, other than standard laboratory glassware.
- 4. Total Solids (Total Residue) (mg/liter)
  - (a) Gravimetric, dried at 103-105°C
    - Blender (if samples contain oil or grease)
    - · Porcelain, vycor, or platinum evaporating dishes
    - Muffle furnace, 550°C
    - Steam bath or 98°C oven
    - Drying oven, 103-105°C
    - Dessicators
      - Analytical balance; 200 g capacity, weighing to 0.1 mg
- 5. Total Dissolved Solids (Total Filterable Residue) (mg/liter)
  - (a) Glass fiber filtration, dried at 180°C
    - Glass fiber filter discs: Reeve Angel 934A, 984-H, Gelman type A, or equivalent
    - Filter holder, membrane filter funnel, or Gooch crucibles and adapter
    - Suction flask
    - Porcelain, vycor, or platinum evaporating dishes
    - Muffle furnace, 550°C
    - Steam bath
    - Drying oven, 180°C
    - Dessicators
    - Analytical balance, 200 g capacity, weighing to 0.1 mg
- 6. Total Suspended Solids (Total Non-Filterable Residue) (mg/liter)
  - (a) Glass fiber filtration, dried at 103-105°C
    - Same as (5), except drying oven is at 103-105°C and steam bath, muffle furnace, and evaporating dishes are not required.

- Total Volatile Solids (Volatile Residue) (mg/liter)
  - (a) Gravimetric, dried at 550°C
    - Same as (5)
- 8. Ammonia (as N) (mg/liter)
  - (a) Distillation and titration
    - All glass distillation apparatus (Kjeldahl)
    - Standard titration apparatus
  - (b) Distillation and nesslerization
    - All-glass distillation apparatus (Kjeldahl)
    - Nessler tubes, 50 ml, matched set, APHA standard
    - Spectrophotometer or filter photometer for use at 425 nm with light path > 1 cm.
  - (c) Distillation and ammonia electrode
    - All-glass distillation apparatus (Kjeldahl)
    - Electrometer (pH meter) with expanded mV scale or specific ion meter
    - Ammonia selective electrode, such as Orion Model 95-10 or EIL Model 8002-2
    - Magnetic stirrer, thermally-insulated, and Teflon-coated stirring bar
  - (d) Automated colorimetric phenate method
    - Technicon AutoAnalyzer (AAI or AAII) with
      - (1) Sampler
      - (2) Manifold (AAI) or Analytical Cartridge (AAII)
      - (3) Proportioning pump
      - (4) Heating bath with double delay coil (AAI)
      - (5) Colorimeter with 15 mm tubular flow cell and 630-660 nm filters
      - (6) Recorder
      - (7) Digital printer for AAII (optional)

- 9. Total Kjeldahl Nitrogen (as N) (mg/liter)
  - (a) Digestion, distillation, and titration
    - Same as 8(a) with suction takeoff to remove SO<sub>3</sub> fumes during digestion
  - (b) Digestion, distillation, and nesslerization
    - Same as 8(b) with suction takeoff to remove SO<sub>3</sub> fumes during digestion
  - (c) Digestion, distillation, and ammonia electrode
    - Same as 8(c) with suction takeoff to remove SO<sub>3</sub> fumes during digestion
  - (d) Automated phenate method
    - Technicon AutoAnalyzer with
      - (1) Sampler II with continuous mixer
      - (2) Two proportioning pumps
      - (3) Manifolds I and II
      - (4) Continuous digester
      - (5) Planetary pump
      - (6) Five-gal. Carboy fume trap
      - (7) Heating bath, 80°C
      - (8) Colorimeter equipped with 50 mm tubular flow cell and 630 nm filters
      - (9) Recorder with range expander
      - (10) Vacuum pump
  - (e) Automated selenium method
    - Technicon AutoAnalyzer with
      - (1) Sampler
      - (2) Two manifolds (as in EPA Manual)
      - (3) Two proportioning pumps
      - (4) Continuous digester
      - (5) Two 5-gal. Carboys
      - (6) Colorimeter with 15 or 50 mm flow cell and 630 or 650 nm filter
      - (7) Recorder
      - (8) Vacuum pump

#### 10. Nitrate (as N) (mg/liter)

- (a) Cadmium Reduction Method (Nitrate Nitrate)
  - Glass fiber or membrane filters and associated apparatus
  - Copper/cadmium reduction column
  - Spectrophotometer or filter photometer for use at 540 nm with light path > 1 cm.
- (b) Automated Cadmium Reduction Method (Nitrate Nitrate)
  - Glass fiber or membrane filters and associated apparatus
  - Copper/cadmium reduction column
  - Technicon AutoAnalyzer (AAI or AAII) with
    - (1) Sampler
    - (2) Manifold (AAI) or Analytical Cartridge (AAII)
    - (3) Colorimeter with 15 or 50 mm tubular flow cell and 540 nm filters
    - (4) Recorder
    - (5) Digital printer for AAII (optional)

#### (c) Brucine Method

- Spectrophotometer or filter photometer for use at 410 nm
- Water bath at 100°C (Temperature control is critical: all sample tubes must be held at the same temperature, and temperature must not drop significantly when tubes are immersed in bath.)
- Water bath at 10-15°C
- Neoprene-coated wire rack for holding sample tubes in baths
- Glass sample tubes (40-50 ml)

#### 11. Total Phosphorus (as P) (mg/liter)

- (a) Single Reagent (Ascorbic Acid Reduction Method)
  - Spectrophotometer or filter photometer for use at 650 nm (less sensitive) or 880 nm
  - Acid-washed, detergent-free glassware
  - Hotplate or autoclave (for persulfate digestion)

- (b) Automated Colorimetric Ascorbic Acid Reduction Method
  - Acid-washed, detergent-free glassware
  - Hotplate or autoclave (for persulfate digestion)
  - Technicon AutoAnalyzer with
    - (1) Sampler
    - (2) Manifold (AAI) or Analytical Cartridge (AAII)
    - (3) Proportioning pump
    - (4) Heating bath, 50°C
    - (5) Colorimeter with 15 or 50 mm tubular flow cell and 650-660 or 880 nm filter
    - (6) Recorder
    - (7) Digital printer for AAII (optional)
- 12. Acidity (mg CaCO<sub>3</sub>/liter)
  - (a) Hydrogen peroxide digestion and electrometric titration
    - pH meter, Type I or II as defined in ASTM D1293
  - (b) Hydrogen peroxide digestion and phenolphthalein end-point titration
    - No special equipment, other than standard laboratory glassware
- 13. Total Organic Carbon (T.O.C.) (mg/liter)
  - (a) Combustion and infrared method  $(CO_2)$  or flame ionization method  $(CH_4)$ 
    - Waring or other blender
    - Apparatus for total and dissolved organic carbon (No specific model is recommended, but several have been found reliable: Dow-Beckman Carbonaceous Analyzer Model #915 (infrared), Dohrmann Envirotech DC-50 Carbon Analyzer (flame ionization), Oceanographic International Total Carbon Analyzer).
- 14. Total Hardness (mg CaCO<sub>3</sub>/liter)
  - (a) EDTA titration
    - No special equipment, other than standard laboratory glassware

(b) Automated colorimetric

Technicon AutoAnalyzer with

- (1) Sampler I
- (2) Continuous filter
- (3) Manifold
- (4) Proportioning pump
- (5) Colorimeter equipped with 15 mm tubular flow cell and 520 nm filters
- (6) Recorder with range expander
- (c) Atomic absorption (Ca + Mg)

(See atomic absorption section below)

- 15. Nitrate (as N) (mg/liter)
  - (a) Manual colorimetric diazotization

Spectrophotometer for use at 540 nm with cells  $\geq$  1 cm.

Nessler tubes or volumetric flasks, 50 ml

(b) Automated colorimetric diazotization

Glass fiber or membrane filters and associated apparatus

Technicon AutoAnalyzer (AAI or AAII) with

- (1) Sampler
- (2) Manifold (AAI) or Analytical Cartridge (AAII)
- (3) Colorimeter with 15 or 50 mm tubular flow cell and 540 nm filters
- (4) Recorder
- (5) Digital printer for AAII (optional)

#### Analytical Methods for Trace Metals: Atomic Absorption Methods

For each parameter listed, EPA specifies atomic absorption as at least one of the reference methods. The required equipment in each case will include (1) an atomic absorption spectrophotometer, (2) the hollow cathode (or electric discharge) lamp for each metal, and (3) the fuels and other apparatus specified below. Design features of some common atomic absorption spectrophotometers (as of June, 1972) are discussed in the EPA Handbook for Analytical Quality Control in Water and Wastewater Laboratories. If extraction procedures are to be used, special reagents are required but no special equipment other than standard laboratory glassware. Results are reported in mg/liter.

Fuels Acetylene Nitrous oxide Parameter Air Other X X X Aluminum Antimony X X Arsenic (Gaseous Argon-hydrogen flame Hydride) X X Barium Beryllium | Х Х Cadmium X X Calcium X X or X Nitrous oxide more sensitive Chromium VI X X X Nitrous oxide more or sensitive; extraction with APDC required for separation of Cr VI from Cr III Chromium, total X X or X Nitrous oxide more sensitive X X Cobalt X Х Copper X X Iron X X Lead X Magnesium X X X Manganese Flameless atomic Mercury (Cold Vapor) absorption: details below X Molybdenum X Nickel X X

#### Continued

00110211100			Fuels	
Parameter	Acetylene	Air	Nitrous oxide	Other
Potassium	Х	x		Osram potassium vapor discharge lamp also may be used.
Selenium (Gaseous Hydride)	ν.			Argon-hydrogen lamp
Silver	X	X		
Sodium	X	X		
Thallium	x	X		
Tin	X	X		
Titanium	X		X	
Vanadium	X		X	
Zinc	x	Х		

#### Other Reference Methods for Metals

#### 16. Aluminum (mg/liter)

- (a) Eriochrome cyanine R colorimetric method
  - Spectrophotometer for use at 535 nm, or
  - Filter photometer with 525-535 nm filters (green, or
  - Nessler tubes, 50 ml

#### 17. Arsenic (mg/liter)

- (a) Gaseous Hydride Silver Diethyldithiocarbamate Colorimetric Method
  - Arsine generator and absorption tube
  - Spectrophotometer for use at 535 nm, or
  - Filter photometer with 530-540 nm filter (green)

- 18. Beryllium (mg/liter)
  - (a) Aluminon method
    - Spectrophotometer or filter photometer for use at 515 nm with 5 cm cells
- 19. Boron (mg/liter)
  - (a) Curcumin method
    - Spectrophotometer or filter photometer for use at 540 nm with cells > 1 cm.
    - Vycor or platinum evaporating dishes, 100-150 ml
    - Water bath,  $55 \pm 2^{\circ}C$
    - Ion exchange column, 50 cm x 1.3 cm (diameter)
- 20. Cadmiun (mg/liter)
  - (a) Dithizone Colorimetric Method
    - Spectrophotometer or filter photometer for use at 515 nm
- 21. Calcium (mg/liter)
  - (a) EDTA Titration
    - No special equipment
- 22. Chromium VI (mg/liter)
  - (a) Diphenylcarbazide colorimetric
    - Membrane or sintered glass filter
    - Spectrophotometer or filter photometer for use at 540 nm with cells > 1 cm.
- 23. Chromium, total (mg/liter)
  - (a) Oxidation and diphenylcarbazide colorimetric
    - Membrane or sintered glass filter.
    - Spectrophotometer or filter photometer for use at 540 nm with cells > 1 cm.

#### 24. Copper (mg/liter)

- (a) Neocuproine colorimetric
  - Spectrophotometer for use at 457 nm with cells ≥ 1 cm, or
  - Filter photometer with narrow-band violet filter (max. transmittance at 450-460 nm) and cells > 1 cm, or
  - Nessler tubes, 50 ml.

#### 25. Iron (mg/liter)

- (a) o-Phenanthroline colorimetric
  - Spectrophotometer or filter photometer for use at 510 nm with cells > 1 cm, or
  - Nessler tubes, 100 ml

#### 26. Lead (mg/liter)

- (a) Dithizone colorimetric
  - Spectrophotometer or filter photometer for use at 520 nm with cells > 1 cm
  - pH meter
- 27. Magnesium (mg/liter)
  - (a) Gravimetric
    - No special equipment
- 28. Mercury (mg/liter)
  - (a) Manual Cold Vapor Technique (Water or Sediment)
    - Commercially available mercury analyzer employing this technique, or
    - Atomic absorption spectrophotometer with open sample presentation area for mounting 10 cm absorption cell
    - Mercury hollow cathode lamp: Westinghouse WL-22847, argonfilled, or equivalent
    - Recorder: multi-range, variable speed, compatible with UV detection system

- Absorption cell, 10 cm, quartz end windows, vapor inlet and outlet ports
- Air pump, peristaltic, 1 liter/min.
- Flowmeter
- Aeration tubing and drying tube (or incandescent lamp to warm cell)
- Autoclave (optional, for digestion procedure)
- (b) Automated Cold Vapor Technique
  - Technicon AutoAnalyzer with
    - (1) Sampler II with provision for sample mixing
    - (2) Manifold
    - (3) Proportioning Pump II or III
    - (4) High temperature heating bath with two distillation coils in series
  - Vapor-liquid separator
  - Absorption cell, 10 cm, quartz end windows
  - Atomic absorption spectrophotometer with open sample presentation area for mounting 10 cm cell (or commercially available analyzer employing this technique)
  - Mercury hollow cathode lamp: Westinghouse WL-22847, argonfilled, or equivalent
  - Recorder: multi-range, variable speed, compatible with UV detection system
  - Cooling water for mixing coil and connector and heat lamp for absorption cell
- 29. Nickel (mg/liter)
  - (a) Heptoxime colorimetric method
    - Spectrophotometer or filter photometer for use at 445 nm with cells > 1 cm.
- 30. Potassium (mg/liter)
  - (a) Colorimetric
    - Spectrophotometer for use at 425 nm with cells > 1 cm, or

- Filter photometer with violet filter (max. transmittance near 425 nm) and  $\geq$  1 cm cells, or
- Nessler tubes, 100 ml
- Centrifuge and 25 ml. centrifuge tubes

#### (b) Flame photometric

• Flame photometer, direct-reading or internal-standard, and associated equipment for measurement at 768 nm

#### 31. Sodium (mg/liter)

- (a) Flame photometric
  - Flame photometer, direct-reading or internal-standard, and associated equipment for measurement at 589 nm
  - For low-solids water, air filter and blower for burner housing, oxyhydrogen flame, and polyethylene or Teflon cups, bottles, etc.

#### 32. Vanadium (mg/liter)

- (a) Colorimetric (Catalysis of gallic acid oxidation)
  - Spectrophotometer or filter photometer for use at 415 nm with 1-5 cm cells
  - Water bath,  $25 \pm 0.5$ °C

#### 33. Zinc (mg/liter)

- (a) Dithizone colorimetric method
  - Spectrophotometer or filter photometer for use at 535 or 620 nm with 2 cm cells, or
  - . Nessler tubes, matched
  - pH meter

#### Analytical Methods for Nutrients, Anions, and Organics

- 34. Organic Nitrogen (as N) (mg/liter)
  - (a) Kjeldahl Nitrogen minus Ammonia Nitrogen
    - See (8) and (9) above.

- 35. Orthophosphate (as P) (mg/liter)
  - See (11) above
- 36. Sulfate (as SO<sub>4</sub>) (mg/liter)
  - (a) Gravimetric
    - Analytical balance, weighing to 0.1 mg
    - Steam bath
    - Drying oven, 180°C
    - Muffle furnace, 800°C
    - Appropriate filters or crucibles
  - (b) Trubidimetric
    - Nephelometer or
    - Spectrophotometer or filer photometer for use at 420 nm with 4-5 cm cells
    - Magnetic stirrer with timer or stopwatch
  - (c) Automated colorimetric barium chloroanilate
    - Technicon AutoAnalyzer with
      - (1) Sampler I
      - (2) Continuous filter
      - (3) Manifold
      - (4) Proportioning pump
      - (5) Colorimeter with 15 mm tubular flow cell and 520 nm filters
      - (6) Recorder
      - (7) Heating bath, 45°C
    - Magnetic stirrer
- 37. Sulfide (as S) (mg/liter)
  - (a) Titrimetric iodine
    - No special equipment, other than standard laboratory glassware.

- 38. Sulfite (as  $SO_3$ ) (mg/liter)
  - (a) Titrimetric iodide-iodate
    - No special equipment, other than standard laboratory glassware
- 39. Bromide (mg/liter)
  - (a) Titrimetric iodide-iodate
    - No special equipment, other than standard laboratory glassware
- 40. Chloride (mg/liter)
  - (a) Silver nitrate
    - No special equipment, other than standard laboratory glassware
  - (b) Mercuric nitrate
    - No special equipment, other than standard laboratory glassware
  - (c) Automated colorimetric ferricyanide
    - Technicon AutoAnalyzer with
      - (1) Sampler I
      - (2) Continuous filter
      - (3) Manifold
      - (4) Proportioning pump
      - (5) Colorimeter with 15 mm tubular flow cell and 480 nm filters
      - (6) Recorder
- 41. Cyanide, total (mg/liter)
  - (a) Distillation and silver nitrate titration
    - . Cyanide distillation apparatus
    - Koch microburet, 5 ml.
  - (b) Distillation and pyridine-pyrazolone (or pyridine-barbituric acid) colorimetric
    - Cyanide distillation apparatus

• Spectrophotometer or filter photometer for use at 578 or 620 nm with > 1 cm cells.

#### 42. Fluoride (mg/liter)

- (a) Distillation SPADNS
  - Simple Bellack distillation apparatus
  - Spectrophotometer for use at 570 nm with  $\geq$  1 cm cells, or
  - Filter photometer with green-yellow filter (max. transmittance 550-580 nm) and > 1 cm cells
- (b) Automated complexone method
  - Technicon AutoAnalyzer with
    - (1) Sampler I
    - (2) Manifold
    - (3) Proportioning pump
    - (4) Continuous filter
    - (5) Colorimeter with 15 mm tubular flow cell and 650 nm filters
    - (6) Recorder with range expander
- (c) Fluoride electrode
  - Electrometer
  - Fluoride ion activity electrode
  - Reference electrode, single junction, sleeve-type
  - Magnetic mixer
- 43. Chlorine, total residual (mg/liter)
  - (a) Starch-iodide titration
    - No special equipment, other than standard laboratory glassware
  - (b) Amperometric titration
    - Amperometric end-point detection apparatus, consisting of noble metal electrode, salt bridge, and silver - silver chloride reference electrode cell unit connected to microammeter with appropriate electrical accessories.
    - Agitator

- 44. Oil and Grease (mg/liter)
  - (a) Gravimetric
    - Separatory funnels or soxhlet apparatus
    - Vaccuum
  - (b) Infrared
    - Spearatory funnels
    - Infrared spectrophotometer, double beam, with 1, 5, and 10 cm cells
- 45. Phenols (mg/liter)
  - (a) Colorimetric (4-AAP method with distillation)
    - Phenols distillation apparatus
    - Spectrophotometer or filter photometer for use at 460 nm (following chloroform extraction) or 510 nm and 1-10 cm cells
    - pH meter
  - (b) Automated 4-AAP method
    - Technicon AutoAnalyzer (I or II) with
      - (1) Sampler
      - (2) Manifold
      - (3) Proportioning pump II or III
      - (4) Heating bath with distillation coil
      - (5) Distillation head
      - (6) Colorimeter with 50 mm flow cell and 505 or 520 nm filter
      - (7) Recorder
- 46. Surfactants (mg/liter)
  - (a) Methylene blue colorimetric
    - Spectrophotometer or filter photometer for use at 625 nm with > 1 cm cells
- 47. Algicides (mg/liter)
  - (a) Gas chromatography

• There is no reference procedure for algicides as a class, and, therefore, detailed equipment requirements cannot be specified. For general discussion of gas chromatography and its application in environmental monitoring, see the EPA Training Manual for Pesticide Residue Analysis in Water and the EPA Methods Manual for Analysis of Pesticide Residues in Human and Environmental Samples.

#### 48. Benzidine (mg/liter)

- (a) Diazotization and colorimetric
  - Spectrophotometer, scanning, 510-370 nm
  - Cells, 1-5 cm pathlength, 20 ml max. volume
- 49. Chlorinated organic compounds (except pesticides) (mg/liter)
  - (a) Gas chromatography
    - There is no reference procedure for chlorinated organic com compounds as a class, and, therefore, detailed equipment requirements cannot be specified. Gas chromatography with electron capture, microcoulometry, or electrolytic conductivity detection may be appropriate for individual compounds or groups of compounds. For general discussions of gas chromatography and its application in environmental monitoring, see the EPA Training Manual for Pesticide Residue Analysis in Water and the EPA Methods Manual for Analysis of Pesticide Residues in Human and Environmental Samples.

#### 50. Pesticides (µg/liter)

• There is no single reference procedure for pesticides as a class. However, specific reference procedures for several sub-classes are available from EMSL, USEPA, Cincinnati, Ohio. To be qualified in this parameter, the laboratory should be equipped to analyze for all specified sub-classes. The analysis of pesticides at the levels normally found in wastewater and other environmental sources requires special expertise and experience, in addition to up-to-date, well-maintained, calibrated instrumentation and apparatus. The equipment lists below are based on the EMSL methods; for further information on the equipment and methodology of pesticide analysis, see the EPA Training Manual for Pesticide Residue Analyses in Water and the EPA Methods Manual for Analysis of Pesticide Residues in Human and Environmental Samples.

#### (a) Organochlorine pesticides

- Gas chromatograph with
  - (1) Glass-lined injection port
  - (2) One or more of the following detectors:

    Electron capture, radioactive (H<sup>3</sup> or Ni<sup>63</sup>)

    Microcoulometric titration

    Electrolytic conductivity
  - (3) Recorder, potentiometric, 10" strip chart
  - (4) Appropriate Pyrex gas chromatographic columns
- Snyder columns, 3-ball (macro) and 2-ball (micro), and other K-D glassware
- Appropriate columns for liquid-solid partition chromatography
- Blender
- Special materials, such as PR Grade Florisil and pesticide standards
- (b) Organophosphorus pesticides
  - Gas chromatograph with
    - (1) Glass-lined injection port
    - (2) One or more of the following detectors:

      Flame photometric, 526 nm phosphorus filter
      Electron capture, radioactive (H<sup>3</sup> or Ni<sup>63</sup>)
    - (3) Recorder, potentiometric, 10" strip chart
    - (4) Appropriate Pyrex gas chromatographic columns
  - Snyder columns, 3-ball (macro) and 2-ball (micro), and other K-D glassware
  - Appropriate columns for liquid-solid partition chromatography
  - Blender
  - Special materials, such as PR Grand Florisil, Woelm neutral aluminá, and pesticide standards
- (c) Polychlorinated biphenyls (PCB's)
  - Gas chromatograph with
    - (1) Glass-lined injection part
    - (2) One or more of the following detectors:

Electron capture, radioactive (H<sup>3</sup> or Ni<sup>63</sup>) Microcoulometric titration Electrolytic conductivity

- (3) Recorder, potentiometric, 10" strip chart
- (4) Appropriate Pyrex gas chromatographic columns
- Snyder column, 3-ball (macro)
- Appropriate columns for liquid-solid partition chromatography
- Low-pressure regulator (0-5 psig) with low-flow needle valve
- Blender
- Special materials, such as PR Grade Florisil, high-quality silica gel, and Aroclor (PCB) standards

#### (d) Triazine pesticides

- Gas chromatograph with
  - (1) Glass-lined injection part
  - (2) Electrolytic conductivity detector
  - (3) Recorder, potentiometric, 10" strip chart
  - (4) Appropriate Pyrex gas chromatographic column
- Snyder columns, 3-ball (macro) and 2-ball (micro), and other K-D glassware
- Appropriate columns for liquid-solid partition chromatography
- Blender
- Special materials, such as PR Grade Florisil and pesticide standards

#### (e) 0-Aryl carbamate pesticides

- Thin layer chromatography plates, 200 x 200 mm, coated with Silica Gel G, 0.25 mm
- Associated TLC apparatus, including spotting template, developing chamber, and sprayer (20 ml)

- 51. Specific Conductance (mho/cm @ 25°C)
  - (a) Wheatstone bridge
    - Commercial conductivity meter, or
    - Apparatus consisting of
      - (1) Wheatstone bridge (reading to 1% accuracy or better)
      - (2) Appropriate source of electrical current
      - (3) Specific conductance cell
      - (4) Water bath, 25°C, with racks
- Turbidity (Jackson units)
  - (a) Turbidimeter method
    - Nephelometric turbidimeter, such as Hach Model 2100 or 2100A or equivalent
- Streptococci bacteria, fecal (number/100 ml)
  - (a) MPN
    - Autoclave (to 121°C)
    - Inoculation tubes
    - Incubator, 35 + 0.5°C
  - (b) Membrane filter
    - Autoclave (to 121°C)
    - Filter membranes
    - Petri culture dishes
    - Incubator, 35 + 0.5°C, ca. 90% relative humidity
    - Low-power (10-15X), binocular, wide-field, dissecting microscope and light source
  - (c) Plate count
    - Autoclave (to 121°C)
    - Petri culture dishes
    - Incubator,  $35 \pm 0.5^{\circ}C$

- Microscope and light source, or
- Colony counter
- 54. Specific Conductance (mho/cm @ 25°C)
  - (a) Wheatstone bridge
    - Commercial conductivity meter, or
    - Apparatus consisting of
      - (1) Wheatstone bridge (reading to 1% accuracy or better)
      - (2) Appropriate source of electrical current
      - (3) Specific conductance cell
      - (4) Water bath, 25°C, with racks
- 55. Turbidity (Jackson units)
  - (a) Turbidimeter method
    - Nephelometric turbidimeter, such as Hach Model 2100 or 2100A or equivalent
- 56. Streptococci bacteria, fecal (number/100 ml)
  - (a) MPN
    - Autoclave (to 121°C)
    - Inoculation tubes
    - Incubator, 35 + 0.5°C
  - (b) Membrane filter
    - Autoclave (to 121°C)
    - Filter membranes
    - Petri culture dishes
    - Incubator, 35 ± 0.5°C, ca. 90% relative humidity
    - Low-power (10-15X), binocular, wide-field, dissecting microscope and light source
  - (c) Plate count
    - Autoclave (to 121°C)

- Petri culture dishes
- Incubator, 35 + 0.5°C
- Microscope and light source, or
- Colony counter
- 57. Coliform bacteria, fecal (number/100 ml)
  - (a) MPN
    - Autoclave (to 121°C)
    - · Inoculation tubes
    - Incubator,  $35 \pm 0.5$ °C
    - Water bath, 44.5 + 0.2°C
  - (b) Membrane filter
    - Autoclave (to 121°C)
    - Filter membranes
    - Petri culture dishes
    - Water bath,  $44.5 \pm 0.2$ °C
    - Low-power (10-15X), binocular, wide-field, dissecting microscope and light source
- 58. Coliform bacteria, total (number/100 ml)
  - (a) MPN
    - Same as 56 (a)
  - (b) Membrane filter
    - Same as 56 (b)

#### Radiological Parameters:

The analysis of radiological parameters requires special expertise and experience, in addition to up-to-date, well-maintained, calibrated instrumentation and apparatus.

- 59. Alpha, total (pCi/liter)
  - Windowless Gas-Flow Proportional Counter and associated equipment, or
  - Thin Window Gas-Flow Proportional Counter and associated Equipment, or
  - Alpha Scintillation Counter and associated equipment, or
  - Alpha Spectrometer (Surface Barrier Type) System and associated equipment
- 60. Alpha counting error (pCi/liter)
  - Same as 59.
- 61. Beta, total (pCi/liter)
  - Windowless Gas-Flow Proportional Counter and associated equipment, or \
  - Thin Window Gas-Flow Proportional Counter and associated equipment, or
  - Beta Scintillation Counter and associated equipment, or
  - Liquid Scintillation Counter and associated equipment
- 62. Beta counting error (pCi/liter)
  - Same as 61.
- 63. Radium, total (pCi/liter)
  - Windowless Gas-Flow Proportional Counter and associated equipment, or
  - Thin Window Gas-Flow Proportional Counter and associated equipment, or
    - Alpha Scintillation Counter and associated equipment, or
    - Alpha Spectrometer (Surface Barrier Type) System and associated equipment, or
    - Radon Gas Counting System and associated equipment

#### Other Parameters

- 64. Temperature
  - Good quality mercury-filled or dial type centigrade thermometer, or a thermistor
- 65. pH
- pH meter (electrometer using either glass electrode and reference, such as saturated calomel, or a combination glass and reference electrode)

#### Air Parameters

- 66. Sulfur Dioxide ( $\mu g/m^3$  or ppm)
  - (a) Pararosamiline Method
    - Absorber
    - Pump
    - Air flowmeter or critical orifice
    - Spectrophotometer for use at 548 nm, band width
       15 nm, with 1 cm cells
- 67. Suspended Particulates  $(\mu g/m^3)$ 
  - (a) High-Volume Method
    - High-volume Sampler
    - Shelter for Sampler
    - Flow measurement equipment, including:
      - (1) Rotameter
      - (2) Orifice Calibration Unit
      - (3) Differential manometer
      - (4) Positive Displacement Meter
    - Barometer
    - Environment for conditioning filters
    - Analytical balance: chamber to hold unfolded 8" x 10" filters, sensitivity = 0.1 mg

- Glass fiber filters
- Acceptable alternative equipment for flow measurement (3-6): Exhaust orifice meter, interfaced with a circular chart recorder.
- 68. Carbon monoxide ( $\mu g/m^3$  or ppm)
  - (a) Non-dispersive Infrared Spectrometry
    - Carbon monoxide analyzer (for example, Intech NDIR-CO Analyzer)
    - Pump, flow control value, and flowmeter
    - In-line filter for particles (2-10 µm)
    - Moisture control (refrigeration unit, or drying tube)
- Photochemical Oxidant  $(0_3)$   $(\mu g/m^3 \text{ or ppm})$ 
  - (a) Chemiluminescence, continuous
    - Commercial photochemical oxidant (03) analyzer, or
    - Apparatus consisting of:
      - (1) Detector cell

      - (2) Flowmeters (air and ethylene)(3) Air Inlet Filter (Teflon, 5 m)
      - (4) Photomultiplier tube
      - (5) High Voltage Power Supply(6) Direct Current Amplifier

      - (7) Recorder
      - (8) Ozone Source (low pressure Hg lamp/quartz tube) and Dilution System
    - Apparatus for Calibration (KI $\longrightarrow$  I<sub>2</sub> spectrophotometric method)
- 70. Total Hydrocarbons (corrected for methane) GC FID
  - (a) Method
    - Commercially Available THC, CH4, and CO Analyzer
    - Pump, flow control valves, automatic switching valves, and flowmeter
    - In-line filter (3-5 μm)

- Stripper or Precolumn
- Oven (for column and catalytic converter)
- 71. Nitrogen Dioxide ( $\mu g/m^3$  or ppm)
  - (a) Arsenite 24-Hour Sampling Method
    - Sampling train (Bubbler, trap, membrane filter, 27-gauge hypodermic needle, air pump, calibration equipment)
    - Standard glassware (volumetrics, pipets, graduated cylinders, etc.)
    - Spectrophotometer or colorimeter for use at 540 nm.
  - (b) Continuous Chemiluminescent Method

    - Calibration apparatus (gas-phase titration method): generally including air flow controller, air flowmeters, pressure regulator for NO cylinder, NO flowmeters, capillary restriction, ozone generator, reaction chamber and mixing bulb, sample manifold, NO detector, iodometric calibration apparatus.
  - (c) Griess-Saltzman Colorimetric, Continuous
    - Sampling train
    - Colorimeter for use at 550 nm

TECHNICAL REPORT DATA (Please read Instructions on the reverse before completing)					
1. REPORT NO.	2.	3. RECIPIENT'S ACCESSION NO.			
EPA-600/4-78-017					
4. TITLE AND SUBTITLE		5, REPORT DATE			
PROCEDURE FOR THE EV	ALUATION OF ENVIRONMENTAL	March 1978 issuing date			
MONITORING LABORATOR	6. PERFORMING ORGANIZATION CODE				
7. AUTHOR(S)	<del></del>	8. PERFORMING ORGANIZATION REPORT NO.			
Charles A. Bicking,	Steven Olin and Peter King				
9. PERFORMING ORGANIZATION	NAME AND ADDRESS	10. PROGRAM ELEMENT NO.			
Tracor Jitco, Inc.		1 HD 621			
1776 E. Jefferson St	reet	11. CONTRACT/GRANT NO.			
Rockville, MD 20852		Contract No. 68-03-2171			
12. SPONSORING AGENCY NAME	AND ADDRESS	13. TYPE OF REPORT AND PERIOD COVERED			
Environmental Monito	ring and Support Laboratory-	Cin., OH Contract 1/10/75 to 1/10/76			
Office of Research a	nd Development	14. SPONSORING AGENCY CODE			
U.S. Environmental P	rotection Agency	EPA/600/06			
Cincinnati OH 4526	8	EPA/000/00			
15. SUPPLEMENTARY NOTES					

#### 16. ABSTRACT

A procedure was developed for the on-site evaluation of environmental laboratories in such media as air, water, radiation and pesticides. The procedure includes registration and preliminary questionnaire forms, on-site visits checklist, evaluator's guide and a scoring system for assessment of the laboratory's management, personnel, facilities, analytical methodology and instruments and its quality control procedures.

This research report is not an official EPA manual. Rather, it is a report which is but one of a series being used as input to develop EPA Manuals and Guidelines for Certification Programs.

17. KEY WORDS AND DOCUMENT ANALYSIS		
a. DESCRIPTORS	b.IDENTIFIERS/OPEN ENDED TERMS	c. COSATI Field/Group
Laboratories*, Evaluation*, Acceptability*, Assessments*, Inspection*.	Testing Laboratories, Scoring System, On-Site Checklist, Preliminary Forms, Evaluator's Guide, Grading.	43F 68Ø 91A
18. DISTRIBUTION STATEMENT Release to Public	19. SECURITY CLASS (This Report) Unclassified	21. NO OF PAGES
	20. SECURITY CLASS (This page)  Unclassified	22. PRICE