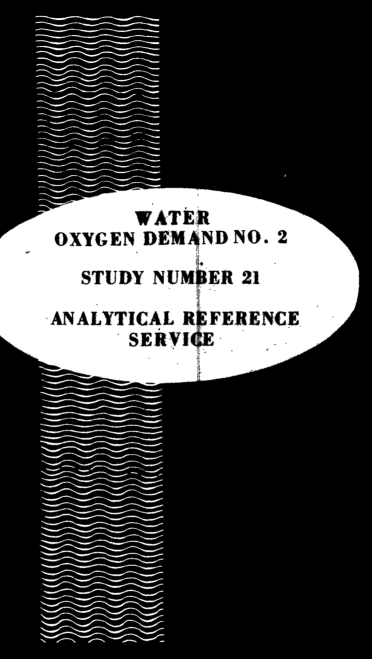
ENVIRONMENTAL HEALTH SERIES Water Supply and Pollution Control



U.S. DEPARTMENT OF HEALTH, EDUCATION, AND WELFARE
Public Health Service

WATER OXYGEN DEMAND NO. 2

Study Number 21

Report of a Study Conducted by the

ANALYTICAL REFERENCE SERVICE

TRAINING PROGRAM

U. S. DEPARTMENT OF HEALTH, EDUCATION, AND WELFARE
Public Health Service
Bureau of State Services
Division of Water Supply and Pollution Control

Robert A. Taft Sanitary Engineering Center Cincinnati, Ohio

1965

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FOREWORD

The Analytical Reference Service is conducted by the Training Program of the Robert A. Taft Sanitary Engineering Center for the evaluation of laboratory methods in the environmental field. Cooperative studies by member organizations, through analysis of identical samples and critical review of methodology, provide the mechanism for:

- Evaluation of analytical procedures, including precision and accuracy, by comparison of the procedures and results reported by participating laboratories.
- 2. Exchange of information regarding method characteristics.
- Improvement or replacement of existing methods by development of more accurate procedures and by development of new methodology for determination of new pollutional components.

Samples are designed to contain measured amounts of selected constituents. Decisions as to qualitative makeup are made by the ARS staff, the membership, and consultants. Notice of each study is sent to the entire membership.

A portion of the study sample with accompanying data forms for reporting numerical values, a critique of the procedures used, comments on modifications, sources of error, difficulties encountered, or other pertinent factors, is then shipped to each of those who expresses a desire to participate. The results and comments of each study are compiled and a report is prepared.

Initially directed toward examination of water, studies now include air, milk, and food. Some studies are periodically repeated, for the advantage of new members, to evaluate new methods or to reevaluate existing methods.

The selection of studies is guided by the responses to questionnaires periodically circulated among the membership which now includes 198 federal, state, and municipal agencies; industries; universities; consulting firms; and foreign agencies.

> James P. Sheehy Director, Training Program

STUDIES ON WHICH REPORTS HAVE BEEN COMPLETED

Water-Minerals Calcium, magnesium, hardness, sulfate, chloride, alkalinity, nitrite, nitrate, sodium,

and potassium. Studies completed in 1956,

1958, and 1961.

Water-Metals Lead, copper, cadmium, aluminum, chromium,

iron, manganese, and zinc. Studies completed

in 1957 and 1962.

Water-Fluoride Fluoride in the presence and absence of inter-

ferences, with and without distillation using a specified procedure. Studies completed in

1958 and 1961.

Water-Radioactivity Studies completed in 1959, 1961, and 1963. The

first two studies were designed to determine gross beta activity, while the third study was concerned with gross beta and strontium-90

activity.

Water-Surfactant Surfactant in various waters. Studies com-

pleted in 1959 and 1963.

Water-Oxygen Demand Biochemical oxygen demand and chemical oxy-

gen demand study completed in 1960; COD

study completed in 1965.

Water-Trace Elements Arsenic, boron, selenium, and beryllium.

Study completed in 1962.

Freshwater Plankton Evaluation of the precision and accuracy ob-

tainable by the use of various methods of plankton counting and identification. Study

completed in 1964.

Air-Inorganics Chloride, sulfate, fluoride, and nitrate in

aqueous solution and on glass fiber Hi-Vol filter mats. Study completed in 1958.

Air-Lead Filter paper tape impregnated with lead.

Study completed in 1961.

Air-Sulfur Dioxide Determination of sulfur dioxide in air using a

specified method. Study completed in 1963.

Milk-DDT Residue DDT in milk. Study completed in 1962.

PARTICIPANTS IN THIS STUDY

Alabama Water Improvements Commission

Alexander Orr, Jr., Water Treatment Plant, Miami Florida

American Cyanamid Company, Bound Brook, New Jersey

Arizona State Department of Health

British Coke Research Association, Chesterfield, Derbyshire, England

California State Department of Public Health, Berkeley

California State Department of Public Health, Los Angeles

Connecticut State Department of Health

Delaware Water Pollution Commission

Department of Municipal Laboratories, Hamilton, Ontario, Canada

Department of Water Resources, Durham, North Carolina

Dow Chemical Company, Midland, Michigan

Erie County Laboratory, Buffalo, New York

Florida State Board of Health (Division of Sanitary Engineering, Jacksonville)

Florida State Board of Health (Pensacola)

Florida State Board of Health (Winter Haven)

General Electric Company, Louisville, Kentucky

Georgia Institute of Technology, Department of Applied Biology

HALL Laboratories Division, Calgon Corporation, Pittsburgh, Pennsylvania

Hawaii State Department of Health

Health Department, Beaumont, Texas

Idaho Department of Health

Illinois State Water Survey Division

Indiana State Board of Health

Industrial Chemicals, Incorporated, South Bend, Indiana

Kentucky State Department of Health

Los Angeles Department of Public Works, Hyperion Treatment Plant

Los Angeles Department of Water and Power

Louisiana State Board of Health

Louisville Water Company, Incorporated

Maryland State Department of Health

Massachusetts Department of Public Health

Metropolitan Utilities District, Omaha, Nebraska

Metropolitan Water District of Southern California

Michigan Water Resources Commission

Minnesota Department of Health

Missouri Department of Public Health and Welfare

Monsanto Chemical Company, St. Louis, Missouri

Montana State Board of Health

NALCO Chemical Company, Chicago, Illinois

National University of Colombia, Bogota, Colombia, South America

Nebraska State Department of Health

Nevada State Department of Health

New Hampshire Water Pollution Commission

New Jersey State Department of Health

New York State Conservation Department

North Carolina Department of Water Resources, Raleigh Ohio State Department of Health Oregon State Board of Health Pennsylvania Department of Health Rensselaer Polytechnic Institute, New York Roy F. Weston, Incorporated St. Louis County Water Company Scientific Research Council, Kingston, Jamaica, West Indies Sixth U.S. Army Medical Laboratory, Fort Baker, California Tennessee Valley Authority (Stream Pollution Control) Chattanooga Texas State Department of Health 2793D U.S. Air Force Hospital, Regional Environmental Health

Laboratory, McClelland Air Force Base, California
2794th U.S. Air Force Dispensary - Class B, Kelly Air Force Base, Texas
United Kingdom Atomic Energy Authority, Didcot, Berks., England
U.S. Army Environmental Hygiene Agency, Edgewood Arsenal, Maryland
U.S. Industrial Chemicals Company, Tuscola, Illinois
University of Beograd, Civil Engineering Faculty, Beograd, Yugoslavia
University of Kansas, School of Engineering and Architecture
University of Leeds, Houldsworth School of Applied Science,
Leeds, England

University of North Carolina, Chapel Hill
Virginia State Water Control Board
Washington State Department of Health
Washington State University, Division of Industrial Research
Water Department, Charlotte, North Carolina
Water Department, Long Beach, California
Water Research Association, Marlow, Buckinghamshire, England
West Virginia State Water Resources Commission
Wisconsin State Board of Health

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ABSTRACT

This study consisted of four samples which 74 participating laboratories were instructed to dilute to a specified volume and analyze by both the Standard Method for Chemical Oxygen Demand and by the Mercuric Sulfate modification.

The results from this study indicate that the two procedures produce similar precision and accuracy when no interfering materials are present. When interferences due to high concentrations of chloride are present, the standard method will produce equal precision and accuracy only if the appropriate corrective techniques are applied.

The Mercuric Sulfate modification is the method of choice for COD measurement since with less manipulation it effectively removes the interference due to chloride oxidation and is less time consuming.

WATER-OXYGEN DEMAND NUMBER 2

PURPOSE OF THE STUDY

The chemical oxygen demand (COD) of a liquid sample is one of the oldest analytical parameters of pollution and is used quite frequently. Many oxidants and variations in procedure have been used in the past, but the dichromate procedure, as described in the 11th edition of Standard Methods for the Examination of Water and Wastewater, ¹ is used by most analysts today.

A brief evaluation of the precision and accuracy of this test was made in the previous Water-Oxygen Demand study, which involved a sample containing no interfering material and having an oxidizability that varied slightly from the theoretical.

It is generally recognized that chloride ions in the sample prevent a true measurement of the COD. Several remedial measures involving a mathematical correction, either alone or in conjunction with the use of silver sulfate, are offered in <u>Standard Methods</u>. On the other hand, the mercuric sulfate modification developed by Dobbs and Williams, of the Robert A. Taft Sanitary Engineering Center, seemed to remove effectively interference by chlorides. A comparative evaluation of the two methods, therefore, appeared timely.

DESIGN OF THE STUDY

To achieve a sound evaluation of accuracy, a search was undertaken for an oxidizable material that would consistently exert a COD that is 100 % of the theoretical and that would be stable for several months. Potassium acid phthalate was found to meet these requirements in the concentrations used for the samples as shipped.

The study consisted of four samples, designated as A, B, C, and D, which the recipient was instructed to dilute to a specified volume and analyze in triplicate by both the standard method and the mercuric sulfate modification. Instructions for the latter procedure were supplied to the participants.

Sample A, designed as a control, contained 8.5020 g/liter potassium acid phthalate in sterile distilled water with no interference added. When 10 ml of this sample was diluted to 500 ml, the resulting solution produced a COD of 200 mg/liter.

Sample B, designed to simulate a fairly average sample, contained 6.8016 g/liter potassium acid phthalate and 8.2440 g/liter sodium chloride. When 10 ml of this sample was diluted to 500 ml,

the resulting solution had a COD of 160 mg/liter and a chloride concentration of 100 mg/liter.

Sample C, represented the type of sample most likely to cause erroneous COD values. This sample contained 6.3764 g/liter potassium acid phthalate and 82.4400 g/liter sodium chloride. When 10 ml of this sample was diluted to 500 ml, the theoretical COD of the resulting solution was 150 mg/liter and the chloride content was 1,000 mg/liter.

Sample D was designed to evaluate the overall efficiency of the methods in analyzing a variety of substrates. This sample contained 3. 4008 g/liter potassium acid phthalate, and 5 ml was to be diluted to 500 ml with surface water or wastewater (Solution D_1) collected by the participants. The resulting solution (D_2) would then contain 40 mg/liter COD in addition to the COD of the diluent (D_1).

Many participants provided additional analytical data on the water used as Sample $\mathrm{D}_{1}\,.$

TREATMENT OF DATA

The mean of the results reported by each participant was plotted on probability paper to determine the distribution. Values showing a deviation from the normal distribution line were rejected as non-representative because of errors in calculation, dilution, or other indeterminate factors. The rejected values are circled on the probability plot. These rejected values were not included in the development of statistical parameters. In several instances in which a mathematical error was noted, the corrected values were submitted in the interim report to the participant for his approval. If the corrected values were approved by the participant, they were then used in the report.

Calculation of the standard deviations was based on the difference between the average result submitted by each participant and the overall mean value reported for each method. The average reported values are also used in the bar charts. The results obtained by use of the standard method were grouped according to the method employed for correction of chloride interference and are identified as follows:

Technique (1) Mathematical correction

Technique (2) Silver sulfate added before reflux

Technique (3) Silver sulfate added after 20 minutes boiling

Technique (4) Silver sulfate dissolved in the sulfuric acid

Statistical terms, as used in this report, are defined in the glossary.

DISCUSSION

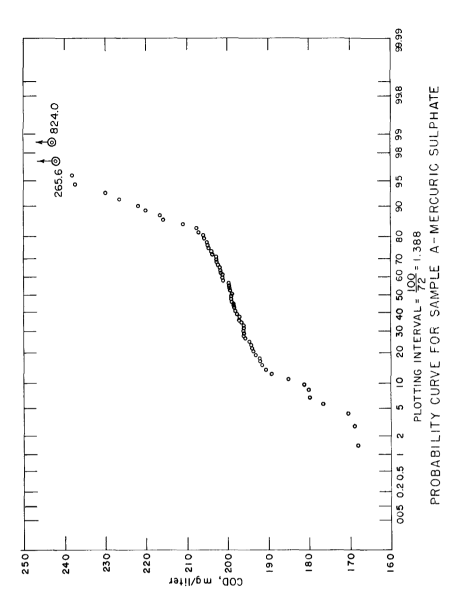
SAMPLE A (200 mg/liter COD in distilled water)

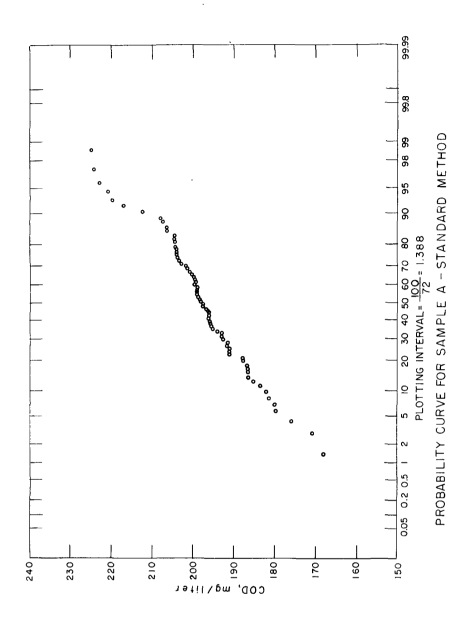
Sample A provided for evaluation of the precision and accuracy of the two methods on waters containing no interfering substances. This sample functioned as a control, and the two methods were expected to produce results of equal accuracy and precision.

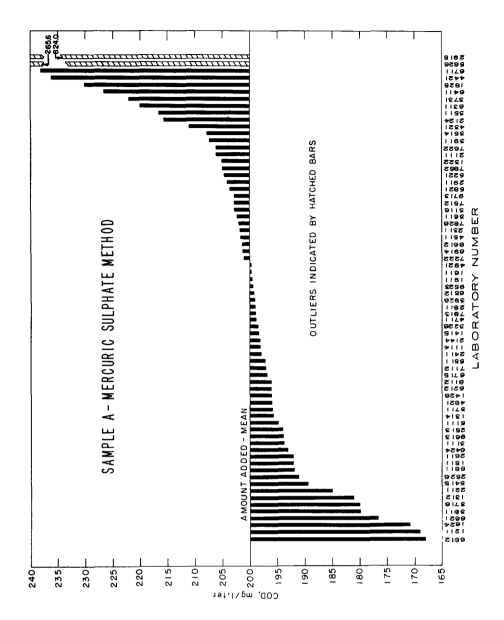
The mercuric sulfate method seems to exhibit slightly more accuracy and a little less precision (Table 1) than the standard method, when the technique for chloride correction is not considered. (See accuracy and precision in Glossary.) The laboratories that used the silver sulfate correction (technique 2) seemed, according to the mean, to achieve better accuracy than those using the other techniques, but the precision was adversely affected. Also, the 50 % range shows that the mean is misleading, since the better half of the results were still twice as far from the theoretical as the results obtained by the other chloride correction techniques. The best standard method values were obtained by the combination of techniques 1 and 3, although no chloride interference was present. Thus, this sample provided an evaluation of procedures rather than efficiency of chloride tie-up. In general, the results indicate that there is very little difference between the two methods when applied to this type of sample.

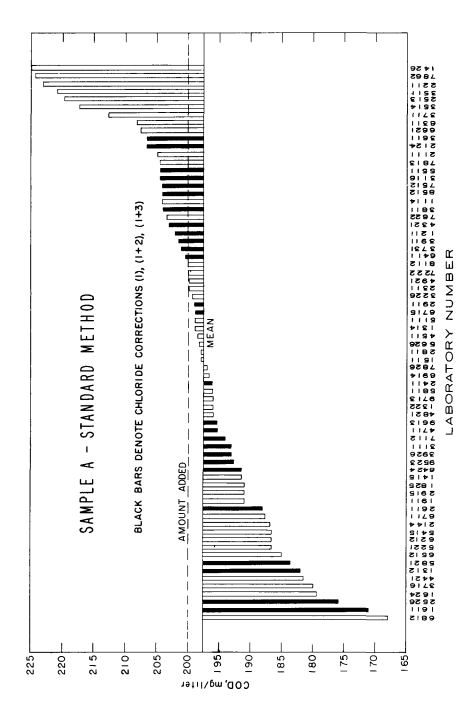
Table 1. SUMMARY OF STATISTICAL DATA ON SAMPLE A

Method	Mean	Standard deviation	Median	50% range	Number
Mercuric sulfate	200.2	13.1	198.9	+ 4.6	69
Standard method, Overall	197.5	11.4	198.2	<u>+</u> 4. 8	71
Standard method (1)	194.9	9. 2	195.3	+ 4.8	15
Standard method (2)	199.8	14.4	199.2	+ 8.6	27
Standard method (3)	190.0	13.2	196.8	$\frac{-}{+}$ 3. 2	5
Standard method (4)	194.9	5.8	198.4	+ 1.6	5
Standard method (1 + 2)	200. 2			_ 	2
Standard method (1 + 3)	196.0	10.5	200. 4	<u>+</u> 4. 4	9









SAMPLE B (160 mg/liter COD in distilled water containing 100 mg/liter chloride)

Sample B was designed to contain a moderate amount of interference, such as might be found in many surface waters. This amount of chloride would tend to produce a high COD result.

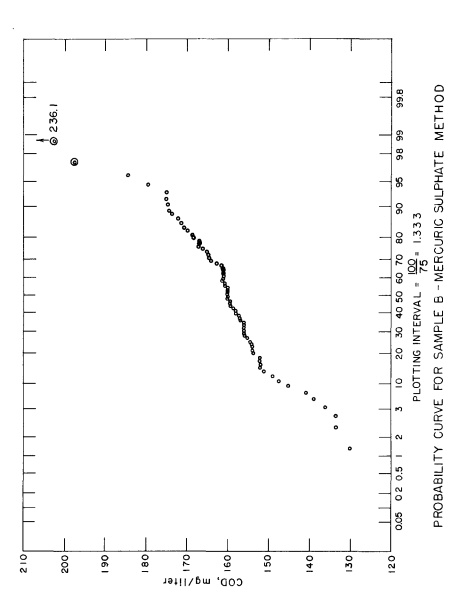
The mercuric sulfate method produced good accuracy as shown by the close agreement (Table 2) between the mean, median, and amount added. The 50% range shows that the better half of the results had an error of 5 mg/liter or less. Surprisingly, the precision as indicated by the standard deviation was a little better than on Sample A.

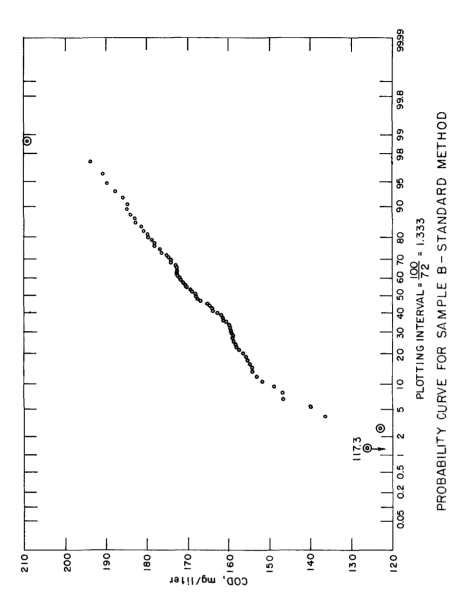
The standard method (overall) showed substantially less accuracy. The mean and median are in close agreement, but are about 8 mg/liter higher than the amount added. The $50\,\%$ range of 11 mg/liter also indicates substantial inaccuracy in even the better half of the results. The results show a normal distribution and a precision nearly as good as on the control sample A.

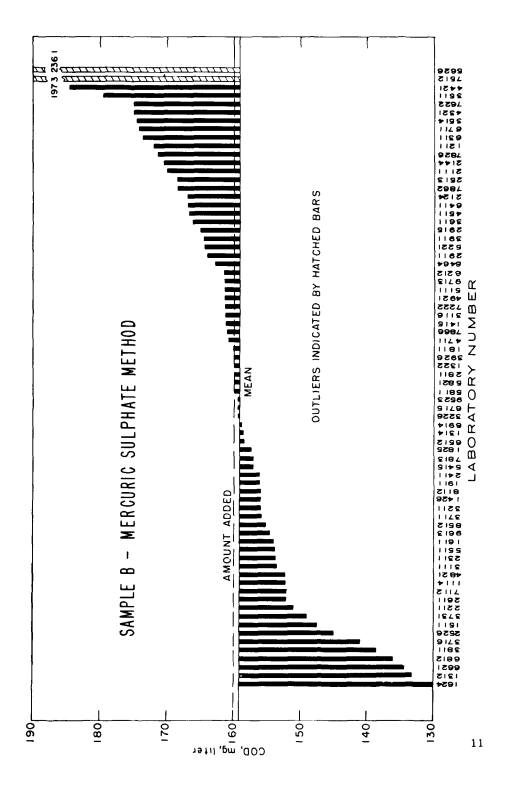
It is evident that on this type of sample substantially equal accuracy and precision can be obtained by use of either the mercuric sulfate method or the standard method when the mathematical correction is applied in conjunction with the procedure of refluxing for 20 minutes before adding silver sulfate (technique 1 and 3).

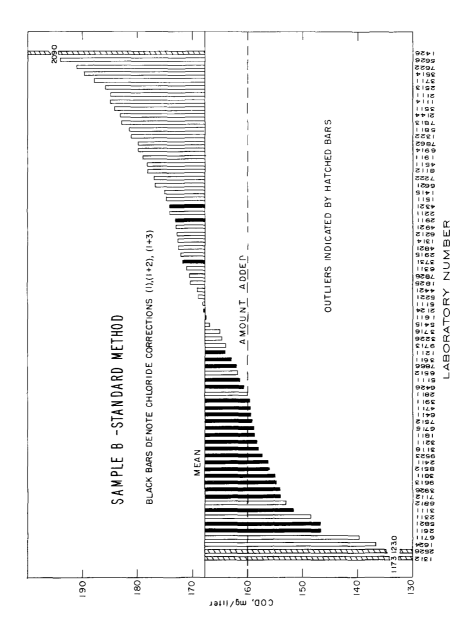
Table 2. SUMMARY OF STATISTICAL DATA ON SAMPLE B

Method	Mean	Standard deviation	Median	50% range	Number
Mercuric sulfate	159.1	10.4	159.9	+ 5.0	72
Standard method, Overall	167.7	12.4	168.0	<u>+</u> 11.0	71
Standard method (1)	154.0	14.7	157.2	+ 5.3	17
Standard method (2)	175.0	11.0	176.6	+ 16.8	27
Standard method (3)	154.0	11.6	153.0	$\frac{-}{+}$ 10.4	5
Standard method (4)	174.0	4.7	172.6	+ 12.6	5
Standard method (1 + 2)	161.0				2
Standard method (1 + 3)	159.0	11.8	159.3	+ 1.4	9









SAMPLE C (150 mg/liter COD in distilled water containing 1,000 mg/liter chloride)

The greatest difficulty with the COD test is experienced in analysis of waters of high chloride content, such as some rivers in the Southwest and estuary waters. Sample C was designed to represent a sample of this type.

The mercuric sulfate method produced results that were normally distributed and a precision that was nearly the same as on the control sample A. The values, however, showed a tendency to be slightly high (Table 3). The mean and median were in close agreement, but were approximately 6 mg/liter higher than the amount added; therefore, chloride oxidation may not have been completely inhibited in all cases.

The overall performance of the standard method was dramatically inferior. The general tendency was toward very high results with a mean value of 219.2 mg/liter and a median value of 200.8 mg/liter. The precision, as indicated by a standard deviation of 89.0, was poor.

Examination of the data showed that the results could be divided into two main groups. One group contained all values to which a mathematical correction for chloride interference had been applied - either combined with the use of silver sulfate added initially, added after 20 minutes of refluxing, or not used at all. The other group of values then included all results to which no mathematical correction had been applied, but which had involved the use of silver sulfate in some manner. Separate bar graphs show the results in these groups. One laboratory reported that no correction for chloride interference was used; therefore, this result appears only in the overall presentation. Statistical data showing further breakdown of the results into six separate groups according to the specific technique used for chloride correction appear in Table 3.

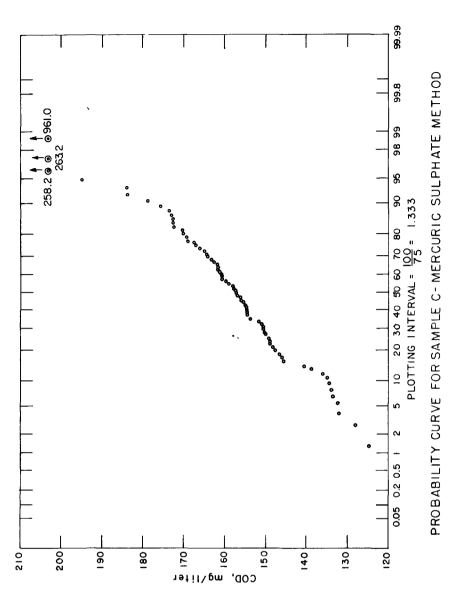
It is evident that, whether or not silver sulfate is used, a mathematical correction is mandatory for this level of chloride interference if the standard method is used. Although two laboratories reported adding silver sulfate initially, in conjunction with a mathematical correction (techniques 2 and 1), these two techniques are incompatible since the silver sulfate would produce a precipitate of silver chloride, which is only partially oxidized by the procedure, making the standard mathematical correction inapplicable. In spite of this, the results submitted by the two participants show much better accuracy than expected and are tabulated but unexplained.

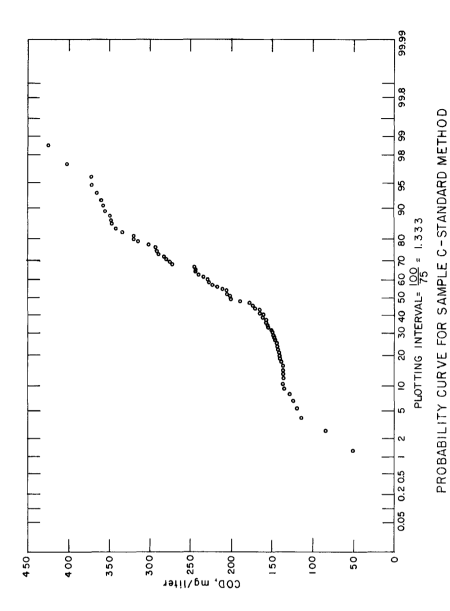
Again, the technique that produced the best accuracy and precision for the standard method was the application of a mathematical correction in conjunction with 20 minutes refluxing before the addition of silver sulfate (techniques 1 and 3).

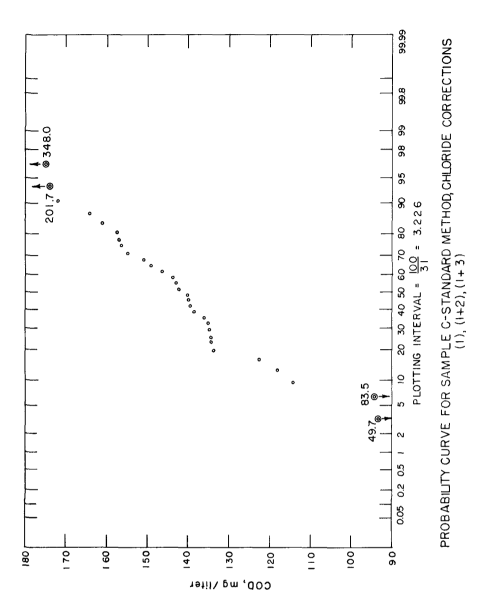
The use of silver sulfate as a catalyst was not necessary for any of the samples in this study. As a result, the participants that used only the mathematical correction achieved essentially the same accuracy as the group using both the mathematical correction and the silver sulfate after 20 minutes reflux. The latter technique, however, would have been required if the sample had contained materials such as straight-chain alcohols and acids that require the catalyst for complete oxidation.

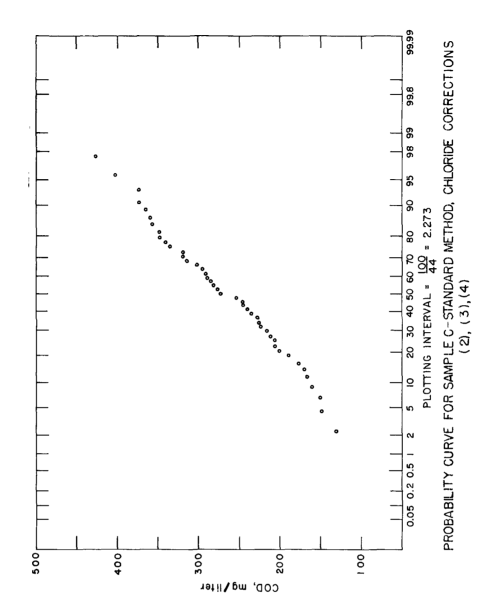
Table 3. SUMMARY OF STATISTICAL DATA ON SAMPLE C

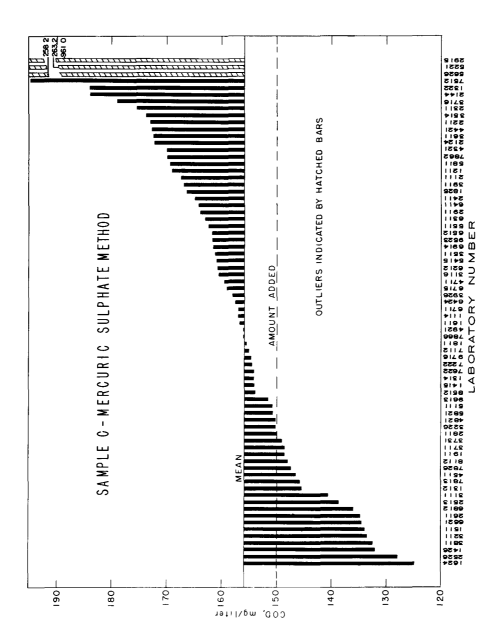
Method	Mean	Standard deviation	Median	50% range	Number
Mercuric sulfate	155.9	13.9	156.8	+ 11.6	71
Standard method, Overall	219.2	89.0	200.8	<u>+</u> 55.3	74
Standard method (1)	147.6	58.0	143.4	+ 11.8	18
Standard method (2)	282.8	66.0	285.8	$\frac{1}{7}$ 132.3	30
Standard method (3)	221.6	112.9	147.9	+ 23.0	5
Standard method (4)	233.8	75.5	224.9	+ 61.0	8
Standard method (1 + 2)	138.7				2
Standard method (1 + 3)	146.6	22.6	139.8	+ 10.2	10
Standard method (1) $+ (1 + 2) + (1 + 3)$	143,0	13.8	141.2	<u>+</u> 10.3	26
Standard method (2) + (3) + (4)	266.6	76.2	270.9	+ 120.9	43

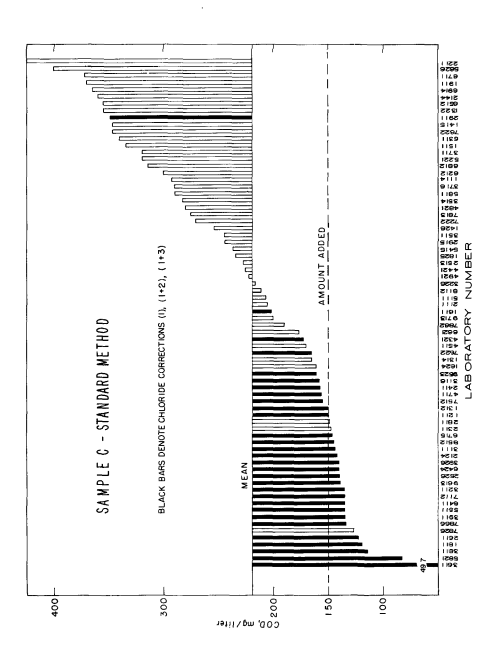


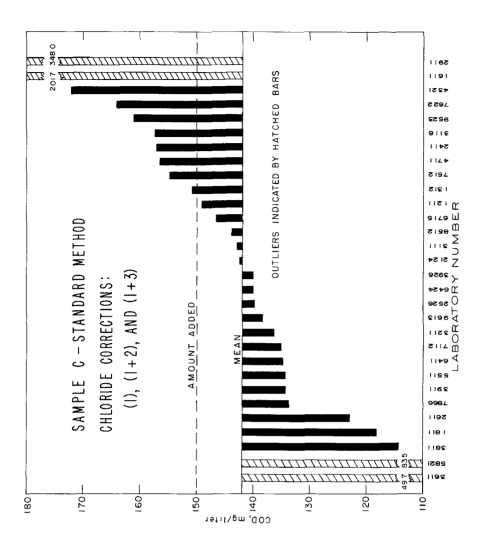


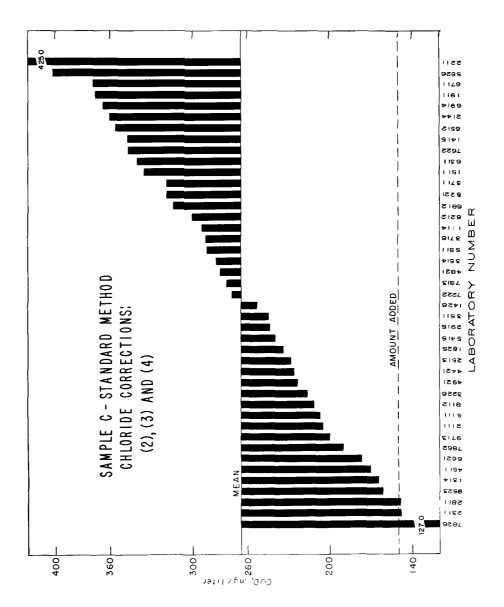












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SAMPLE D (40 mg/liter COD to be added to water sample collected by participant)

Sample D was designed to provide an overall estimate of the efficiency of the COD methods in analyzing a variety of collected samples (D_1) containing various amounts of chemically oxidizable material and interferences.

In preparing sample $D_2,\,5$ ml of the sample D concentrate was made up to 500 ml by adding 495 ml of the collected sample (D_1). Since before analysis D_1 was not similarly diluted, sample D_1 contained 1% more of the collected sample than did D_2. That is to say, to make the samples strictly comparable, 495 ml of the collected sample should have been diluted with 5 ml of distilled water and this sample then called D_1.

To minimize confusion, additional manipulations, and possibly mathematical errors, it was decided to ignore the dilution factor in the instructions to participants. The submitted results were then corrected by the Analytical Reference Service staff to compensate for the dilution. This was done by adding 1 % of the COD value for sample D_1 to the difference obtained by subtracting the initial COD value of D_1 from the COD value of D_2 . This, in effect, raised the difference value to the value that would have been obtained if there had been no dilution of D_1 in the preparation of D_2 . The theoretical value for $\mathrm{D}_2\text{-}\mathrm{D}_1$ was therefore adjusted to 40.0 mg/liter COD in every case. Statistical calculations were thereby simplified, and presentation of the data in a manner comparable to that used in the other parts of this study was made possible.

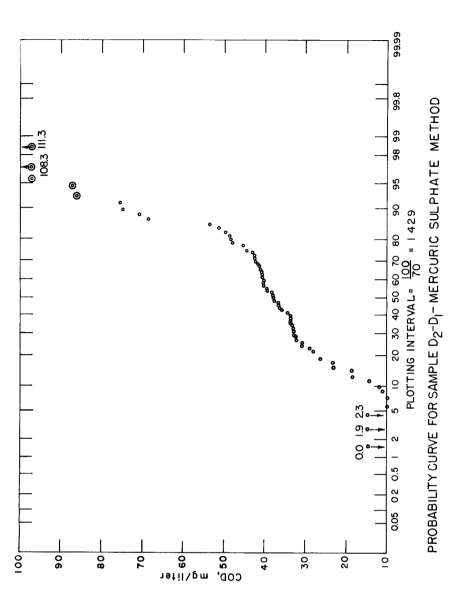
Since so many participants used a water having a very low COD for sample D_1 , it is unfortunate that the instructions referred only to the high COD procedures, although many participants did use the low COD, procedure. Without doubt, many results would have been improved by using the N/40 reagents rather than the N/4 $\,$ In addition, the varied composition of the water used as sample D_1 precludes the use of statistical parameters in the usual manner, and they are, therefore, presented only for the purpose of aiding in the discussion of this sample and should not be used to predict the precision and accuracy of results that might be obtained from a different sample.

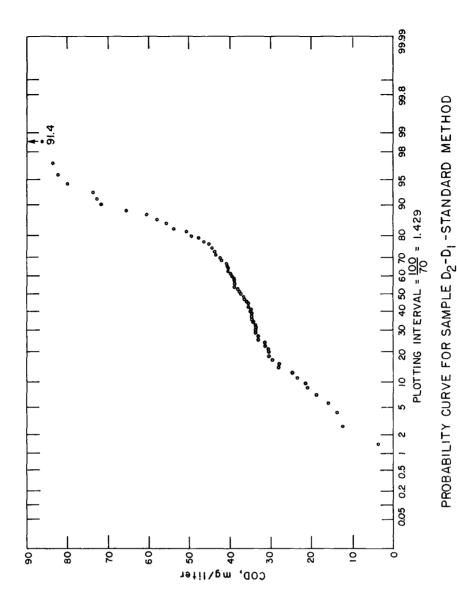
Many of the participants provided considerable analytical data on the water used for sample D $_{\rm L}$. Unfortunately, no significant correlation was found between the quality of the water and the results obtained. Errors, such as neglecting to correct for high chlorides in the standard method, were self-canceling and did not affect the D $_2$ -D $_1$ difference. Difficulties such as an endpoint obscured by color or turbidity undoubtedly caused inaccurate results, but were not reported in sufficient number to evaluate.

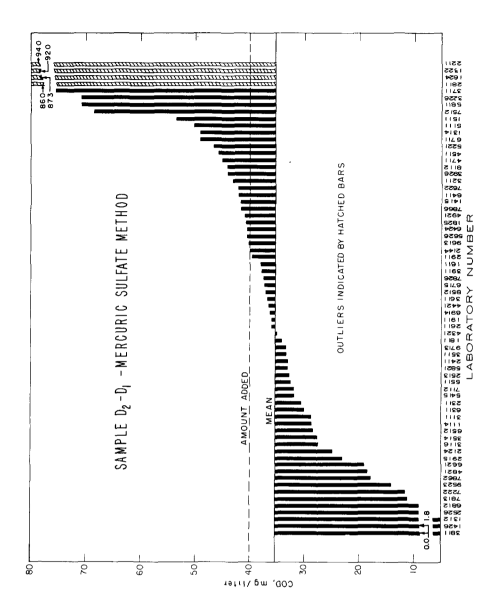
The data indicate (Table 4) that the mercuric sulfate procedure is slightly more precise than the standard method, as shown by the standard deviations of 15.5 and 17.0. The accuracy, however, was less, as shown by the deviation of the mean from the theoretical. On the other hand, the 50% range shows that the better half of the results and the median of all of the results obtained by either method are in close agreement. The differences are not considered significant, and it is the opinion of the Analytical Reference Service staff that either method would be suitable for most of the $\rm D_1$ samples.

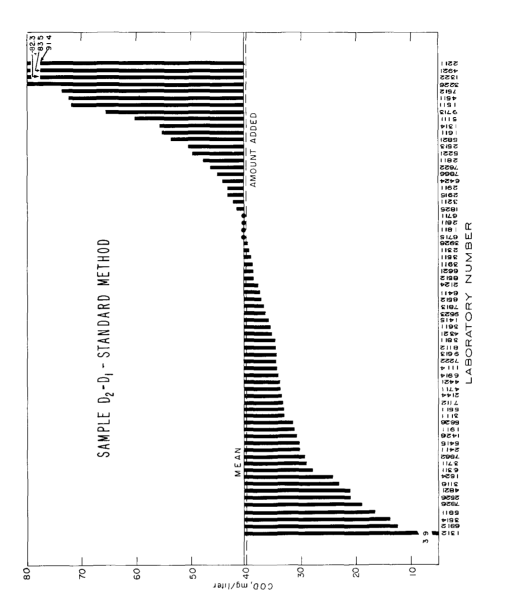
Table 4. SUMMARY OF STATISTICAL DATA ON SAMPLE D

Table 1, Schille-101 of Striff Striff De-11 of Striff III S						
Method	Mean	Standard deviation	Median	50 % range	Number	
Mercuric sulfate	35. 4	15.5	36.4	+ 7.4	64	
Standard method Overall	40.5	17.0	37.2	_ 6, 9	69	









COMMENTS OF PARTICIPANTS

The most frequent comment concerned the precipitate formed when mercuric sulfate was added to Sample C. A few participants found this troublesome in detecting the endpoint of the titration, but the majority experienced no difficulty. Several suggestions were made to use pumice stone or porcelain chips instead of glass beads to control bumping. The general opinion of the participants who used a mathematical correction for chloride in the standard method was that the mercuric sulfate method was quicker and just as good. Those who did not perform a chloride analysis found the weighing of mercuric sulfate a time-consuming extra step. Many suggestions were made to use more than the recommended 2-3 drops of indicator to improve the color change at the endpoint of the titration. Some participants took exception to the statement in the instructions for the mercuric sulfate method that the COD calculation was the same as that for the standard method.

In preparing the instructions it was assumed that since the mercuric sulfate method purports to eliminate chloride interference, it would be evident that the COD calculation would not include a correction factor for chloride.

It was suggested that quantitative control of excess mercuric sulfate might give better results for high chloride samples. This undoubtedly would be true for extremely saline waters which would require more than the specified 1 gram of mercuric sulfate to complex the chloride completely. This type of sample, however, would be a special case requiring further investigation.

A comment that the sample should be added last to the cooled acid-dichromate mixture to avoid loss of volatile fractions may have some virtue in special cases. This technique could then incorporate another suggestion that the mercuric sulfate be dissolved first in the sulfuric acid. A sample containing high chlorides and requiring the catalytic action of silver sulfate might not, however, be completely oxidized because of some of the silver sulfate precipitating with the chloride, if this technique were used.

SUMMARY AND CONCLUSIONS

The results from this study indicate that the two procedures produce similar precision and accuracy when no interfering materials are present. When high concentrations of chloride are present, the mercuric sulfate procedure will effectively remove the interference due to chloride oxidation. The standard method may produce equal precision and accuracy only if the proper technique is used; namely, the measurement of chloride concentration in order to be able to apply the mathematical correction (technique 1), and refluxing the sample with the acid-dichromate mixture for 20 minutes before adding silver sulfate if the catalyst is required (technique 3). Many inaccurate COD values are entirely due to inaccurate chloride analysis.

An alarmingly large number of participants were apparently unaware of the effect of chloride in the measurement of COD by the standard method. This is evident from the many comments expressing concern over the great difference in COD values produced by the two methods in the analysis of Sample C.

Undoubtedly, much of the difficulty is due to a lack of clarity in the procedure as written in the 11th edition of <u>Standard Methods</u>. It is hoped that the forthcoming 12th edition will be improved in this respect.

The Analytical Reference Service staff members feel that the mercuric sulfate modification is the best method for COD measurement because it is less time-consuming than the correctly performed standard method and will provide at least equivalent precision and accuracy regardless of interferences present. Additional benefits, described by Dobbs and Williams, ² but not evaluated in this study, are the elimination of inaccuracies due not only to the series of cyclic changes from chlorine to chloride through the formation of chloramines in wastewater containing chlorides and a high concentration of ammonia, organic amine, or nitrogenous matter, but also the reaction of chlorine, produced by the oxidation of chlorides, with organic matter in the sample. The latter can materially affect the COD.

A statistical summary of the results is presented in Table 5.

Table 5. STATISTICAL SUMMARY					
	Sample A	Sample B	Sample C	Sample D	
Mg/liter COD added	200	160	150	40	
Mg/liter chloride added	0	100	1000	0	
Mean					
Mercuric sulfate	200.2	159.1	155.9	35, 4	
Standard method, overall Standard method (1) +	197.5	167.7	219.2	40, 5	
(1+2)+(1+3)			143.0		
Standard method (2) + (3) + (4)			266.6		
Median					
Mercuric sulfate	198.9	159.9	156.8	36, 4	
Standard method, overall	-	168.0	200.8	37, 2	
Standard method (1) + $(1+2)$ + $(1+3)$			141.2		
Standard method (2) + (3) + (4)			270.9		
Standard deviation					
Mercuric sulfate	13.1	10.4	13.9	15.5	
Standard method, overall Standard method (1) +	11.4	12.4	89.0	17.0	
(1+2)+(1+3)			13.8		
Standard method (2) + (3) + (4)			76.2		
50% range					
Mercuric sulfate	+ 4.6	+ 5.0	+ 11.6	+ 7.4	
Standard method, overall	$\frac{-}{4}$ 4.8	$\frac{-}{+}11.0$	$\frac{\pm}{\pm}$ 11.6 \pm 55.3	$\frac{+}{+}$ 7. 4 $\frac{+}{+}$ 6. 9	
Standard method (1) + $(1 + 2) + (1 + 3)$			<u>+</u> 10.3		
Standard method (2) + (3) + (4)			<u>+</u> 120.9		

LITERATURE REFERENCES

- Standard Methods for the Examination of Water and Wastewater 11th edition. APHA, AWWA, WPCF. New York, 1960.
- 2. Dobbs, R. A. and Williams, R. T. Elimination of Chloride Interference in the Chemical Oxygen Demand Test. Anal. Chem. Vol. 35, p. 1064, 1963.

APPENDICES

APPENDIX A.

TABULATION OF RESULTS

Table A-1. SAMPLE A (Amount added = 200 mg/liter COD)

Mercuric sulfate method Standard method					
	Mercuric sulf	ate metnod	Stai	ndard meth	
Laboratory	D 11-	7. 6	D . 11	3.7	Chloride
number	Results	Mean	Results	Mean	correction
					method
	199.3		205.7		
1114	196.9	198.1	201.7	204.0	2
	198.1		204.5		
	177.0		200.0		
1211	138.0	169.0	207.0	202.0	1 and 3
	193.0	-	200.0		
1312		181.2		182.0	1
					_
	196.4		200.0		
1314	194.4	195.7	200.0	198.7	4
	196.4		196.0		
	204.0		200.0		
1322	208.0	205.0	192.0	196.0	none made
	204.0		196.0		110110 111040
1415		198.3		191.4	2
	204.0		330 A		
1426	204.0 196.0	196.0	$228.0 \\ 224.0$	225.0	4
1420	188.0	190.0	224.0	22 5. 0	4
	100.0		44.U		
	192.0		195.2		
1511	192.0	192.0	199.2	197.8	2
	192.0		199.2		
	195.8		170.0		
1611	195.8	199.7	172.8	170.9	1 and 3
	207.4		170.0		
	174.8		182.0		
1624	168.8	170.8	178.0	179.3	2
	168.8	. • •	178.0		_
	201.5		189.0		
1911	197.5	199.5	191.0	191.0	2
1011	199.5	100.0	192.9	202.0	_

Table A-1 SAMPLE A (continued)

	Table A-1	SAMPL	E A (contin	ued)	
	Mercuric sulf	ate method	Sta:	ndard metl	nod
Laboratory number	Results	Mean	Results	Mean	Chloride correction method
2111	206.0 206.0 206.0	206.0	206.0 204.0 204.0	204.7	2
2124	217.1 214.7 215.1	215.6	208.0 203.2 208.0	206.4	1 and 2
2144	196.8 198.8 198.8	198.1	188.0 188.0 184.4	186.8	2
2211	185 185 184	185	247 214 207	223	2
2311	199.1 207.3 199.1	201.8	202.5 202.5 194.2	199.7	3
2411	198.8 196.1 198.8	197.9	198.0 191.4 199.2	196.2	1 and 3
2513	192.0 196.8 193.6	194.1	216.0 225.6 217.6	219.7	2
2526	189.4 198.3 185.0	190.9	183.1 177.0 167.2	175.8	1
2611	192.0 192.0 192.0	192.0	188.0 188.0 188.0	188.0	1
2811	200.5 198.4 197.8	198.9	200.5 198.6 194.7	197.9	3

Table A-1. SAMPLE A (continued)

Table A-1. SAMPLE A (continued)					
	Mercuric sulfa	ite method	Star	ndard meth	nod
Laboratory number	Results	Mean	Results	Mean	Chloride correction method
2911	208 204 200	204	199 199 199	199	1
2915	832 816 824	824	188 196 188	191	4
3111	193.7 195.6 191.7	193.7	197.1 193.1 189.1	193.1	1
3116	202.3 202.3 203.5	202.7	204.9 203.9 204.5	204. 4	1 and 3
3226	197.1 199.0 199.0	198.4	199.2 199.2 199.2	199.2	2
3511	217.6 217.6 214.4	216.5	222. 4 217. 6 222. 4	220.8	2
3514	204. 2 211. 2	207, 7	212.0 222.3	217.2	2
3611	204.8 201.1 201.1	202, 3	211. 4 204. 1 204. 1	206.5	1
3711	192.2 198.6 196.5	195,8	210.9 218.6 208.1	212.5	2
3716	181 181 179	180	179 179 181	180	2

Table A-1. SAMPLE A (continued)

	Mercuric sulfa		Standard method		
Laboratory number	Results	Mean	Results	Mean	Chloride correction method
3731	216. 0 228. 0 222. 0	222.0	196.0 208.0 200.0	201.0	1
3811	172.6 182.7 178.4 185.8	179.9	204.0 200.0 208.9 202.7	203.9	1
3911	209.6 209.6 202.4	207.2	206.8 200.1 197.4	201.4	1 and 3
3926	197 196 205	199	195 195 190	193	1 and 3
4321	210 213 210	211	196 204 210	203	1
4421	237.1 246.2 225.1	236.1	180.2 189.3 175.0	181.5	2
4511	201.6 201.6 201.6	201.6	198. 4 198. 4 198. 4	198.4	4
4711	198.8 196.8 200.7	198.8	195. 2 197. 2 193. 2	195.2	1
4821	195.9 195.9 195.9	195.9	195.9 195.9 195.9	195.9	2

Table A-1. SAMPLE A (continued)

	Mercuric sulfa		E A (contin	idard meth	od
Laboratory number	Results	Mean	Results	Mean	Chloride correction method
4921	199.7 199.7 199.7	199.7	199.7 199.7 199.7	199,7	2
5111	196.0 196.0 192.0	194.7	200.0 196.0 200.0	198.7	2
5221	206.6 204.6 202.6	204.6	186.6 186.6 186.6	186,6	2
5415	189.3 185.4 193.2	189.3	182.8 190.3 186.7	186.6	4
5511	197. 2 197. 2 197. 2	197.2	201.1 207.1 205.1	204.4	1 and 3
5626		265.6		198, 2	None
5811	194.9 190.2 190.2	191.8	199.5 194.9 194.9	196.1	2
5821	203.6 203.6 203.6	203.6	186.3 182.3 182.3	183.6	1
6212	192.0 200.0 196.0	196.0	182.0 186.0 192.0	186.6	2
6311	230.6 210.7 218.7	220.0	186.9 242.5 194.8	208.1	2

Table A-1. SAMPLE A (continued)

	Table A-1.	SAMPLI	A (continu		
	Mercuric sulfat	e method	Stand	lard method	
Laboratory	Results	Mean	Results	Mean	Chloride correction method
	226.6		200.4		
6411	226.6	226.6	196.4	200.4	1 and 3
	226.6		204.4		
	194.3		192.7		
6424	192.4	193.0	188.7	191.4	1 and 3
	192.4		192.7		
	199.2		187.1		
6512	197.2	199.2	185.1	185.1	2
	201.2		183.1		
	169.0		207.4		
6621	184.3	176.6	207.4	207.4	2
	176.6		207.4		
	239.2		187.7		
6711	235.5	238.0	187.7	187.7	3
	239.2		187.7		
	197.0		199.4		
6715	197.0	196.7	199.4	198.7	1
	196.0		197.4		
	169		169		
6812	165	168	169	168	3
	169		165		
	199.2		198.0		
6914	203.2	201.2	194.0	196.7	2
			198.0		
	195		193		
7112	200	197	194	194	1 and 2
	196		194		
	205.2		199.1		
7222	199.1	201.1	201.2	198.8	2
	199.1		199.1		

Table A-1	SAMPLE A	(continued)
TADTE T-T.	DUMETT U	(COmminger)

	ercuric sulfa		Stan_	dard metho	od
Laboratory number	Results	Mean	Results	Mean	Chloride correction method
	208.0		206.8		
7512	200.0	202.7	198.8	204.1	1
	200.0		206.8		
	208.8		204.6		
7622	204.6	206.0	200.4	203.2	2
	204.6		204.6		
	197.4		205.8		
7813	199.5	198.8	205.8	204.4	2
	199.5		201.6		
	202. 0		196.8		
7826	202.0	202.0		196.8	3
	202.0				
	204. 4		208.8		
7862	205, 2	204.8	226.3	224. 3	2 and 4
	204.8		237.8		
	192		199		
8112	200	196	199	200	4
	196		203		
	202.0		210.0		
8512	199.6	201.2	198.0	204.0	1
	202.0				
	198.8		190.5		
9523	200.7	199.4	195.0	192.7	1
	198.8		192.6		
	192.4		195.1		
9613	193.9	193.8	194.7	195.3	1
	195.1		196.2		
	200.0		200.0		
9713	208.0	202.7	196.0	196.0	2
	200.0		192.0		
	226.4		190.1		
1825	211.7	230.1	191.4	191.0	2
	222.3		191.4		

Table A-2. SAMPLE B
(Amount added = 160 mg/liter.COD + 100 mg/liter chloride)

	Mercuric sulf		COD + 100 mg/liter chloride) Standard method		
Laboratory number	Results	Mean	Results	Mean	Chloride correction method
1114	151. 4 154. 2 151. 0	152.2	187. 2 184. 0 182. 8	184.7	2
1211	157 168 190	172	164 167 162	164	1 and 3
1312		133.2		117.3	1
1314	158. 4 158. 4 158. 4	158. 4	173.9 171.9 171.9	172.6	4
1322	160 160 160	160	180 180 184	181	none made
1415		161.1		174.9	2
1 426	164 156 148	156	212 212 204	209	4
1511	147. 4 147. 4 147. 4	147. 4	175. 5 179. 3 169. 2	174.6	2
1611	153.6 156.4 151.7	153, 9	160. 8 168. 8 172. 8	167.5	1 and 3
1624	130.8 130.8 128.8	130.1	142.0 146.0 122.0	136.7	2
1811	161. 4 160. 3 158. 6	160.1	158. 1 159. 0 158. 6	158.6	1 and 3

Table A-2. SAMPLE B (continued)

	Monounio gulfe		E B (Contin		
	Mercuric sulfa	ile method	Stan	dard metho	Chloride
Laboratory number	Results	Mean	Results	Mean	correction method
	153.6		176.9		
1911	158.4	156.1	178.8	178.8	2
	156.4		180.8		
	170.0		184.0		
2111	170.0	170.0	188.0	184.7	2
	170.0		182.0		
	169.7		166.7		
2124	163.7	167.0	166.7	167.9	1 and 2
	167.7		170.3		
	172.4		183.6		
2144	170.4	170.4	183.6	182.9	2
	168. 4		181.6		
	148		174		
2211	157	151	172	174	2
	148		175		
	153.9		153.8		
2311	150, 2	153.7	146.0	148.6	3
	157.1		146.0		
	156.6		154, 2		
2411	155.1	156.1	157.0	156.2	1 and 3
	156.6		157.3		
	164.8		179.2		
2513	172.8	168.5	182.4	185.6	2
	1.68.0		195.2		
	144.3		114.8		
2 526	150.2	145.0	127.4	123.0	1
	140.6		126.7		
	152.0		146.6		
2611	152.0	152.0	146.6	146.6	1
	152.0		146.6		

Table A-2. SAMPLE B (continued)

	Table A-2		E B (contin		
	Mercuric sulfa	te method	Stan	dard meth	
Laboratory number	Results	Mean	Results	Mean	Chloride correction method
2811	160, 2 160, 5 159, 0	159.9	160, 3 160, 0 159, 2	159.8	3
2911	164 164 164	164	173 173 173	173	1
2915	166 168 160	165	172 172 172	172	4
3111	156.1 152.2 152.2	153. 5	151.6 151.6 151.6	151.6	1
3116	161.2 161.2 161.2	161. 2	157, 5 158, 3 158, 3	158. 0	1 and 3
3211	157.2 155.3 155.3	155, 9	156. 2 158. 2 160. 2	158 . 2	1
3226	158.0 158.0 161.9	159. 3	173.7 160.0 160.0	164.6	2
3511	176.6 180.8 180.8	179.4	185, 6 185, 6 180, 8	184.0	2
3514	174.6 180.5 168.5	174.5	189, 5 185, 3 193, 5	189. 4	2
3611	162.5 166.2 169.9	166. 2	156. 7 171. 4 160. 4	162.8	1

Table A-2. SAMPLE B (continued)

	Mercuric sulfa	ate method	Stan	dard metho	od
Laboratory number		Mean	Results	Mean	Chloride correction method
3711	156.3 155.1 155.9	155.8	187. 1 190. 6 185. 3	187. 7	2
3716	140 140 142	141	165 165 164	165	2
3731	165.0 171.0 112.0	149.0	153.7 161.7 199.7	171.7	1
3811	137.0 140.5 138.2	138.6	148. 9 150. 0 165. 8	154.9	1
3911	165. 2 166. 8 161. 6	164.5	161.6 158.0 158.8	159,5	1 and 3
3926	163 160 156	160	152 158 151	154	1 and 3
4321	174 178 174	175	171 171 181	174	1
4421	213.9 151.7 187.8	184.5	173. 0 166. 7 167. 5	169.1	2
4511	173. 4 165. 3 161. 3	166.7	182. 2 182. 2 170. 0	178. 1	4
4711	161. 4 157. 4 163. 3	160.7	160, 0 160, 0 158, 0	159.3	1

Table A-2. SAMPLE B (continued)

Table A-2. SAMPLE B (continued)							
	Mercuric sulfa	te method	Stan	dard method			
Laboratory number	Results	Mean	Results	Mean	Chloride correction method		
4821	152.2 152.2 152.2	152. 2	176. 4 172. 3 168. 3	172. 3	2		
4921	157.4 161.3 165.1	161.3	176.6 172.8 169.0	172. 8	2		
5111	164.0 164.0 156.0	161.3	168.0 168.0 168.0	168.0	2		
5221	164.4 164.4 164.4	164. 4	168.3 168.3 169.3	168. 8	2		
5415	158.3 154.5 158.3	157.0	164.3 171.7 164.3	166.8	4		
5511	153.8 153.8 153.8	153.8	163. 3 161. 3 159. 3	161. 3	1 and 3		
56 2 6		236.1		193. 8	None		
5811	159.8 159.8 159.8	159.8	181. 4 181. 4 181. 4	181.4	2		
5821	163.9 159.9 155.9	159.9	146.6 146.6 146.6	146.6	1		
6212	158.0 164.0 162.0	161. 4	174.0 172.0 172.0	172. 7	2		

	Table A-2.	SAMPL	E B (contin	ued)	
	Mercuric sulfate	method	Star	dard method	
Laboratory number	Results	Mean	Results	Mean	Chloride correctio method
6911	174. 9	173 6	174.9	171 0	9

	Mercuric sullat	emeniou	Stal	luar u memou	_
Laboratory number	Results	Mean	Results	Mean	Chloride correction method
6311	174.9 171.0 174.9	173.6	174. 9 167. 0 171. 0	171.0	2
6411	166. 2 168. 2 166. 2	166,9	160.0 160.0 158.0	159.3	1 and 3
6424	164.8 162.8 160.8	162.8	162. 6 158. 6 160. 6	160.6	1 and 3
6512	158.3 158.3 158.3	158.3	162. 3 164. 3 158. 3	161.7	2
6621	142.1 126.7 134.4	134. 4	176.6 176.6 176.6	176, 6	2
6711	173.0 173.0 176.6	174. 2	139. 8 139. 8 139. 8	139.8	3
6715	159.3 159.3 159.3	159.3	159. 3 157. 5 159. 3	158.7	1
6812	137 133 137	136	153 153 153	153	3
6914	161. 4 157. 4 157. 4	158.7	178. 4 180. 3 180. 3	179.7	2
7112	152 152 152	152	154 154 154	154	1 and 2

Table A-2, SAMPLE B (continued)

	Table A-	2. SAMPL	E B (contin		
	Mercuric sulfa	ate method	Sta	indard metl	nod
Laboratory number	Results	Mean	Results	Mean	Chloride correction method
7222	166.6 156.5 160.5	161.2	180.8 178.8 170.7	176.8	2
7512	168.0 200.0 224.0	197, 3	159. 0 159. 0 159. 0	159. 0	1
7622	167. 0 179. 0 179. 0	175.0	185. 0 196. 6 190. 9	190.8	2
7813	159. 9 155. 5 155. 5	157.0	182. 8 184. 9 180. 7	182.8	2
78 2 6	171.3 171.3 171.3	171.3	165. 2 177. 2 169. 2	170. 5	3
7862	162. 4 174. 3	168, 4	178. 6 177. 0 183. 4	179.7	2 and 4
7866	152. 4 165. 0 165. 4	160.9	165. 0 161. 5 159. 1	161.9	1
8112	152 160 156	156	177 177 180	178	4
8512	155. 2 155. 2	155.2	159. 8 151. 8	155. 8	1
9523	159. 4 159. 4 159. 4	159.4	152. 1 159. 9 159. 5	157. 2	1

Table A-2. SAMPLE B (continued)

	Mercuric sulf	ate method	d Sta	ndard met	hod
Laboratory number	Results	Mean	Results	Mean	Chloride correction method
9613	154. 0 155. 1 154. 8	154, 6	153. 4 153. 8 156. 9	154.7	1
9713	172.0 152.0 160.0	161.3	168. 0 168. 0 156. 0	164. 0	2
1825	157. 2 157. 2	157. 2	167.9 172.7	170.3	2

Table A-3. SAMPLE C (Amount added = 150 mg/liter COD + 1000 mg/liter chloride)

	Mercuric sulfa	te method	Stand	dard metho	
Laboratory number	Results	Mean	Results	Mean	Chloride correction method
1114	154.2 153.8 163.0	157.0	287.4 291.5 298.3	292.4	2
1211	182 168 166 160	169	169 147 141 138	149	1 and 3
1312		145.4		150.8	1
1314	156.1 154.1 152.2	154.1	164.0 164.0 164.0	164.0	4
1322	184 192 176	184	352 356 356	355	none made
1415		153.9		347.3	2
1426	128 140 128	132	252 256 252	253	4
1511	137.7 129.9 133.8	133.8	334.3 330.5 338.2	334.4	2
1611	163.2 157.4 149.8	156.8	195.0 203.0 207.0	201.7	1 and 3
1624	128.8 118.8 126.8	124.8	164.0 130.0 190.0	161.3	
1811	154.6 155.0 156.6	155.4	116.0 118.3 120.0	118.1	1 and 3

Table A-3. SAMPLE C (continued)

	Table A-3.		E C (continu		
	Mercuric sulfa	te method	Star	idard meth	od Chloride
Laboratory number	Results	Mean	Results	Mean	correction method
1911	144.0 149.9	148.6	373.7 367.7	370. 4	2
1911	151.8	140.0	36 9. 7	310.4	2
0111	166.0	107.0	200.0	005.0	•
2111	168.0 168.0	167.3	204.0 212.0	205.3	2
	170.9		143.7		
2124	177.3 168.5	172.2	143.7 139.7	142.4	1 and 2
	181.2		364.8		
2144	186.0 184.4	183.9	354.4 360.4	359.9	2
	185		423		
2211	167 167	173	426 426	425	2
	179.0		150.0		
2311	179.0 168.0	175.3	146.0 146.0	147.3	3
	164.4		156.9		
2411	164.4 165.6	164.8	156.9 156.9	156.9	1 and 3
	144.0		236.8		
2513	137.6 134.4	138.7	217.6 228.8	227.7	2
	137.6		139.7		
2526	121.8 124.3	127.9	141.2 138.2	139.7	1
	132.0		119.6		
2611	132.0 140.0	134.7	127.6 119.6	122.3	1

Table A-3. SAMPLE C (continued)

	Table A-3		E C (contin		
	Mercuric sulfa	ate method	Stan	dard meth	od
Laboratory number	Results	Mean	Results	Mean	Chloride correction method
2811	149.8 150.6 149.8	150.1	144.7 149.7 149.3	147.9	3
2911	164 164 164	164	348 348 348	348	1
2915	960 964 960	961	228 256 244	243	4
3111	140.1 140.1 142.0	140.7	142.9 142.9 142.9	142.9	1
3116	159.7 160.9 160.9	160, 5	158.6 156.5 156.5	157.2	1 and 3
3211	133. 4 133. 4	133.4	135.5 137.5 135.5	136.2	1
3226	148.3 148.3 154.1	150, 2	216.7 216.7 216.0	216.5	2
3511	158.4 164.8 160.0	161.1	244.8 244.8 241.6	243.7	2
3514	179, 5 166, 7 175, 2	173.8	273.2 290.0 283.8	282.3	2
3611	173.6 173.6 169.9	172.4	49.1 49.1 50.9	49.7	1

Table A-3, SAMPLE C (continued)

			EC (contin		
	Mercuric sulfa	te method	Stan	dard meth	
Laboratory number	Results	Mean	Results	Mean	Chloride correction method
3711	148.6 147.8 149.5	148.6	292.2 324.1 340.9	319.1	2
3716	179 179 179	179	290 290 290	290	2
3731	168.0 168.0 112.0	149.0			1
3811	143.2 133.9 122.4 129.8	132.3	130.1 104.2 108.3	114.2	1
3911	164.0 160.8 176.0	166.9	135.8 134.2 132.6	134. 2	1 and 3
3926	161 159 155	158	142 138 140	140	1 and 3
4321	170 172 168	170	173 169 173	172	1
4421	166.2 177.8 174.2	172.7	225.1 225.9 227.2	226.1	2
4511	145.2 149.2 145.2	146.5	170.0 174.1 166.0	170.0	4
4711	159.4 161.4 157.4	159.4	136.9 175.9	156.4	1

Table A-3. SAMPLE C (continued)

Mercuric sulfa		Table A-3. SAMPLE C (continued)					
Mercuric sulfate method		Standard method Chlo		Chloride			
Results	Mean	Results	Mean	correction method			
150.2 156.2 144.2	150.2	352.2 277.3 210.5	280.0	2			
153.6 157.4 157.4	156.1	222.7 215.0 230.4	222.7	2			
152.0 152.0 148.0	150.7	208.0 208.0 204.0	206.7	2			
296.4 237.1 256.2	263.2	325.0 302.0 328.8	318.6	2			
158.3 158.3 166.2	160.9	235.2 242.5 238.8	238.8	4			
165.6 161.7 159.7	162.3	138.8 130.9 132.9	134.2	1 and 3			
261.8 257.4 255.3	258.2	401.3 401.3 401.3	401.3	4			
164.0 184.0 160.0	169.3	320.0 284.0 264.0	289.3	2			
152.0 148.0 152.0	150.7	86.2 78.2 86.2	83.5	1			
160.0 158.0 164.0	160.7	304.0 300.0 298.0	300.6	2			
163.0 163.0 163.0	163.0	345.9 330.0 345.9	340.6	2			
	150. 2 156. 2 144. 2 153. 6 157. 4 157. 4 152. 0 152. 0 148. 0 296. 4 237. 1 256. 2 158. 3 158. 3 166. 2 165. 6 161. 7 159. 7 261. 8 257. 4 255. 3 164. 0 184. 0 160. 0 152. 0 148. 0 160. 0 152. 0 148. 0 163. 0 163. 0 163. 0	150.2 156.2 144.2 153.6 157.4 152.0 152.0 152.0 152.0 150.7 148.0 296.4 237.1 263.2 256.2 158.3 158.3 160.9 166.2 165.6 161.7 162.3 159.7 261.8 257.4 258.2 255.3 164.0 184.0 169.3 160.0 152.0 148.0 150.7 169.3 160.0 152.0 148.0 169.3 160.0 150.7	150.2 352.2 156.2 150.2 277.3 144.2 210.5 153.6 222.7 157.4 156.1 215.0 157.4 230.4 152.0 208.0 152.0 150.7 208.0 148.0 204.0 296.4 325.0 237.1 263.2 302.0 256.2 328.8 158.3 160.9 242.5 166.2 238.8 165.6 138.8 161.7 162.3 130.9 159.7 132.9 261.8 401.3 257.4 258.2 401.3 255.3 401.3 164.0 320.0 184.0 169.3 284.0 160.0 264.0 152.0 86.2 148.0 150.7 78.2 152.0 86.2 148.0 150.7 78.2 152.0 86.2 148.0 150.7 78.2 152.0	150.2			

Table A-3. SAMPLE C (continued)

	Table A-S	B. SAMPL	E C (contin		
	Mercuric sulfa	te method	Star	dard meth	od
Laboratory number	${ m Results}$	Mean	Results	Mean	Chloride correction method
6411	162.2 164.2 166.2	164.2	134.6 134.6 134.6	134.6	1 and 3
6424	158.9 158.9 154.9	157.6	141.8 137.8 139.8	139.8	1 and 3
6512	163.0 163.0 159.0	161.7	355.9 353.9 357.8	355. 9	2
6621	142.1 126.7 134.4	134.4	169.0 184.3 176.6	176.6	2
6711	161.9 154.6 154.6	157.0	368.0 375.0 372.0	371.7	3
6715	155.3 160.7 161.1	159.0	145.9 147.7 145.9	146.5	1
6812	137 133 137	136	314 314 314	314	3
6914	159.4 163.3 161.4	161.4	366.5 362.6 364.6	364.6	2
7112	156 152 156	155	135 139 131	135	1 and 2
7222	158.5 152.4 152.4	1 54 . 4	286.5 266.2 260.1	270.9	2

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Table A-3.	SAMPLE C	(continued)

	Table A-3	. SAMPL	E C (contin		
	Mercuric sulfa	te method	Stan	dard meth	od
Laboratory					Chloride
·	Results	Mean	Results	Mean	correction
number					method
					- Incured
	200.0		149.4		
7512	184.0	194.7	165.4	154.7	1
	200.0		149.4		
	150.2		341.8		
7622	154.1	154.1	335.9	347.1	2
	158.1	201, 1	363,6	011	2
	100.1		500.0		
7690				104 0	1
7622				164.0	1
	145.0		272.9		
7813	145.0	145.7	260.3	275.0	2
	147.1		291.8		
	148.0		131.0		
7826	150.0	147.3	123.0	127.0	3
1020	144.0	11.,0	127.0	12.,0	O
	111.0		121.0		
	100.2				
7000	168.3	100.0		100.0	0 1 1
7862	147.3	169.8	183.4	189.6	2 and 4
	193.7		195.7		
	154.0		135.4		
7866	157.1	156.1	131.9	133.6	1
	157.1		133.4		
	148		211		
8112	148	148	211	211	4
	148		212		
	153.6		145.4		
8512	154.0	153.8	142.2	143.8	1
0012		100.0	112.2	140.0	1
	162.1		153.1		
0532		101 0		101 0	
9523	165.3	161.6	163.9	161.0	1
	157.4		165.9		
	150,1		138.2		
9613	151.3	151.5	136.7	138.2	1
	153.2		139.8		

Table A-3. SAMPLE C (continued)

	1001011 0.	~~~~~	(0011011	.ucu,					
	Mercuric Sulfate	e method	Stan	Standard method					
Laboratory number	Results	Mean	Results	Mean	Chloride correction method				
	164.0		188.0						
9713	148.0	154.6	208.0	200.0	2				
	152.0		204.0						
	175.9		233.8						
1825	157.2	166.3	225.2	233.9	2				
	165.7		242.8						

Table A-4. SAMPLE D (continued)

Chloride correction method	1	က	1	4	1	1 and 3	T	23	2	63	1	23	 1	1 and 3	1 and 3	1	83
Standard method Diff. Corrected Diff. Diff.	40.4	47.7	43, 2	43.2	33.0	23, 2	42,3	80.0	39, 1	13,8	3 5, 6	29.0	34.7	38,8	39,6	35, 4	33, 8
andard Diff.	40.0	47.5	43.0	39.0	32,9	22.7	42, 3	79.9	38.9	13.2	34,8	28.4	34, 5	36.9	39.0	35.0	33, 8
Mean D 2	78.6	63,1	0.09	461,0	40,9	72,9	42,3	89, 7	59,2	70.0	111,9	85.9	52,1	224,0	95.0	76.0	35, 4
Mean D ₁	38.6	15.6	17.0	422.0	8.0	50.2	0	8 6	20.3	56.8	77.1	57.6	17.6	187,1	56.0	41.0	1.6
Corrected Diff.	36.4	87.8	40,4	23,4	30,6	27.9	43.0	75.4	33, 5	28.0	37, 7	86,8	0.0	39, 7	44, 5	35, 4	37, 5
method Diff.	36.0	87.3	40.0	19.0	30, 5	27.4	43.0	75.4	33, 3	27.6	36.7	86.0	0.0	37.9	44.0	35.0	37.3
Sulfate Mean	81,3	134,6	76.0	464.0	41.8	72.6	43.0	79.2	49.6	68, 4	141.5	166.0	0.0	213,9	94.0	78.0	56,1
$ \begin{array}{ccc} \textit{Mercuric sulfate method} \\ \textit{Mean} & \textit{Mean} \\ \textit{D}_1 & \textit{D}_2 \\ \end{array} \text{ Diff.} $	45, 3	47, 3	36.0	445.0	11,3	45, 2	0	3, 8	16,3	40,8	104.8	80.0	0.0	176.0	50.0	43.0	18,8
Laboratory	2611	2811	2911	2915	3111	3116	3211	3226	3511	3514	3611	3711	3811	3911	3926	4321	4421

Table A-4. SAMPLE D (continued)

Chloride orrection method		4		2	2	2	2	4	က	v	. 2		2	က	က	2	2	c:
2	correction method								1 and	none				1 and	1 and			
nethod	Corrected Diff.	72.2	33.8	21.2	83.5	60.3	49.7	30.3	33.1	31.5	16.6	53.7	27.8	37.4	44.2	37.2	38.6	40.7
Standard method	Diff.	70.2	33.2	21.1	75.1	0.09	49.7	25.9	32.8	30.8	12.0	53.6	27.8	36.8	44.1	36.9	38.4	40.5
St	Mean D_2	269.9	69.7	28.6	916.6	85.3	53.5	462.7	63.5	96.9	469.3	61.7	27.8	97.1	53.1	67.8	61.4	58.9
	Mean D_1	199.7	36.5	7.5	841.5	25.3	3.8	436.8	30.7	66.1	457.0	8.1	0	60.3	0.6	30.9	23.0	18.4
	Corrected Diff.	48.1	47.9	18.4	41.5	53.5	48.6	32.6	33.0	40.7	75.0	33.4	32.0	42.2	40.9	28.9	23.0	49.4
nethod	Diff.	45.7	47.6	18.4	40.2	53.3	48.5	28.3	32.7	39.6	70.7	33.1	31,8	41.6	40.8	28.7	23.0	49.1
sulfate r	Mean D_2	284.9	79.1	18.4	166.2	69.3	57.4	456.0	67.0	152.4	504.0	62.0	55.7	100.5	54.5	45.2	23.0	76.1
Mercuric sulfate method	Mean $^{ m D}_1$	239.2	31.5	0.0	126.0	16.0	8.9	427.7	34.3	112.8	433.3	28.9	23.9	58.9	13.7	16.5	0.0	27.0
	Laboratory number	4511	4711	4821	4921	5111	5221	5415	5511	5626	5811	5821	6311	6411	6424	6512	6621	6711

Table A-4. SAMPLE D (continued)

	Caloride correction method	-	က	83	1 and 2	87	1	87	87	က	2 and 4	П	4	П	1	1	87	62
nethod	Corrected Diff.	40.3	12.5	34.1	33.3	34.5	73.4	46.3	36.7	18.9	29.3	45.1	34.6	38.6	36.5	34.5	65.6	41.7
Standard method	Diff.	40.2	0.6	34.0	33.1	33.9	73.3	46.1	36.6	15.2	29.0	44.9	33.0	38.5	36.4	34.4	65.3	41.5
St	Mean D ₂	46.8	356.0	41.8	50.0	93.5	88.1	67.2	43.3	382.6	62.3	64.7	188.0	53.2	43.0	43.8	93.3	57.5
	$\substack{\text{Mean}\\ D_1}$	9.9	347.0	7.8	16.9	59.6	14.8	21.1	6.7	367.4	33.3	19.8	155.0	14.7	9.9	9.4	28.0	16.0
	Corrected Diff.	38.1	9.6	36.5	32.6	11.8	6.07	42.3	11.1	39.5	18.2	41.9	45.1	37.7	14.1	40.5	33.6	40.7
method	Diff.	38.0	0.6	36.5	32.5	11.5	7.07	42.1	11.0	36.0	17.9	41.7	44.0	37.5	14.0	40.4	33.3	40.7
Mercuric sulfate method	Mean D ₂	45.3	68.0	40.5	40.5	44.0	7.06	64.5	17.3	390.4	51.9	58.6	156.0	62.0	19.7	48.8	0.09	41.6
Mercuri	Mean D ₁	7.3	59.0	4.0	8.0	32.5	20.0	22.4	6.3	354.4	34.0	16.9	112.0	24.5	5.7	8.4	26.7	6.0
	Laboratory number	6715	6812	6914	7112	7222	7512	7622	7813	7826	7862	7866	8112	8512	9523	9613	9713	1825

APPENDIX B.

MERCURIC SULFATE PROCEDURE

- A. Measure 50 ml of sample or aliquot diluted to 50 ml with distilled water, and place in a standard reflux flask, then add:
 - (1) 1 gram mercuric sulfate
 - (2) 5 ml concentrated $\rm H_2SO_4$ swirl to dissolve mercuric salt
 - (3) 25 ml 0, 25N $K_2 Cr_2 O_7$
 - (4) 70 ml concentrated $\mathrm{H_2SO_4}$ (cautiously)
 - (5) 0.75 gram Ag_2SO_4
 - (6) Several glass beads or porcelain chips
- B. Mix well by swirling flask.
- C. Connect flask to condenser and reflux for 2 hours.
- D. Wash down the condenser with distilled water and cool to room temperature.
- E. Add 10 drops of o-phenanthroline ferrous indicator, and titrate to a red endpoint with standardized ferrous ammonium sulfate (approx. 0.25N).

NOTE: Reagents, equipment, and calculations are the same as in Standard Methods. 1

APPENDIX C.

GLOSSARY OF STATISTICAL TERMS

A glossary of statistical terms with definitions of their meaning as used in these reports is presented to insure uniform and complete understanding.

Arithmetic mean The sum of a series of test results divided

by the number in the series.

Median The value above and below which an equal

number of observations lie.

Accuracy Accuracy is the correctness of a measurement.

or the degree of correspondence between the

result and the true value.

Accuracy Data Measurements that relate to the difference

between the average test results and the true result when the latter is known or assumed.

The following measures apply:

50% Range - The maximum deviation from the true amount for the more accurate half of the

mean results reported.

Average deviation from true concentration -The average difference without regard to sign between each laboratory mean and the true

value.

Average percent deviation from established concentration (or amount added) - The average of the differences between a laboratory's replicate results and the established concentration expressed as percentages of the established

concentration.

Precision Precision is a measure of the reproducibility

of measurements, or the degree to which the measurements correspond to one another.

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Precision Data

Measurements that relate to the variation among the test results themselves, i.e., the scatter or dispersion of a series of test results without assumption of any prior information. The following measures apply:

Variance - The sum of squares of deviations of the average test results from the mean of the series divided by one less than the total number of average test results.

Standard deviation - The square root of the variance.

<u>Coefficient of variation</u> - The standard deviation of the laboratories' means as a percentage of the mean of this series.

Range - The difference in magnitude between the highest and lowest laboratory mean.

Average percent deviation within laboratory - The average of the differences between a laboratory s replicate results and their mean, expressed as percentages of their mean.

Confidence limits - Limits within which the true mean of the series will lie with a given probability.

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- Bennett, C. A. and Franklin, N. L. Statistical Analysis in Chemistry and the Chemical Industry. John Wiley & Sons, Inc. New York. 1954.
- Bauer, E. L. A Statistical Manual for Chemists. Academic Press. New York. 1960.
- 4. Dixon, W. J. and Massey, F. J. Introduction to Statistical Analysis. McGraw-Hill Book Co., Inc. New York. 1951.
- 5. Standard Methods for the Examination of Water and Wastewater. 11th edition. APHA, AWWA, WPCF. New York, 1960.

APPENDIX D

PROBABILITY EXPLANATION

This section deals with the description and use of normal probability paper. The bell-shaped normal distribution curve can be reduced to a straight line on probability paper. Obviously, it is easier to deal with this straight line than with the complicated bell-shaped curve.

Construction of Normal Probability Paper

In an ideal sample, frequencies of the measurements plotted against their magnitudes give a bell-shaped normal distribution curve. This curve is symmetric about its mean, 0, and the percentage of the area lying between any two points on the curve can be found. For example, 68.26% of the area under the curve lies between + one standard deviation (Figure D-1).

The first step in the construction of normal probability paper is the transformation of the bell-shaped distribution curve to the probability summation curve. This curve gives the summation of the area from left to right under the bell-shaped curve up to any deviation. For example, taking the point, 0, on the X-axis of the summation curve, we read from the Y axis that 50% of the total area lies below this zero value, which is the mean. The summation of area up to a deviation of -1 σ below the mean is 50 - 34.13, or 15.87%, while the summation of area up to +1 σ above the mean is 50 + 34.13, or 84.13%. Similarly, other transformations can be made to form a complete, smooth ogee or s-shaped curve (Figure D-2).

The next step is to reduce the probability summation curve to a straight line. This is done by projecting the points on the ogee curve vertically to the linear X-axis scale and by writing the corresponding percentage opposite each point. For example, project point a to a', and write its probability summation (15.87%); project b to b', and write 84.13%, etc. Such a scale, with the percentages shown, constitutes the X-axis of normal probability paper.

Plotting on Normal Probability Paper

Since the probability summation is the cumulative area from left to right, it represents probability of a measurement equal to or less than a given value. Arranging the data in ascending order

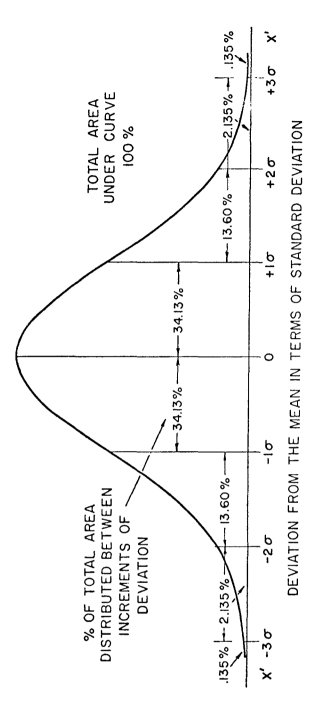
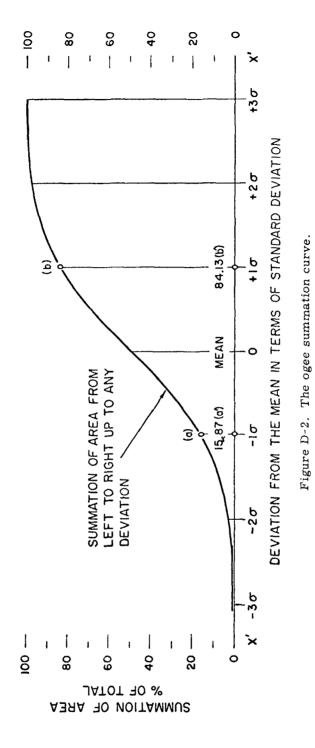


Figure D-1. The bell-shaped distribution curve.



of magnitude automatically places them in the order of their position on the probability summation scale. This arrangement permits ready determination of the probability not exceeding a certain magnitude of measurement.

In the development of the probability paper many difficulties were encountered in choosing a formula for plotting the position of the measurements on the probability summation scale (or X-axis), but the problem was solved in the following way. Ideally, the plotting position for the mean of any series, regardless of the number of values, is at 50 %; i.e., half of the values above and half below. If the mean is considered as part of the series, the number of plotting positions becomes n + 1. Thus the X-axis, which represents 100 %, is divided into n + 1 intervals. To plot the third point, for example, multiply $\frac{1}{n+1}$ by 3, or $\frac{3}{n+1}$. The formula for the plotting interval, therefore, is $\frac{m}{n+1}$ where

m equals the serial number of the measurements arranged in ascending order of magnitude, and n equals the total number of measured values to be plotted. Multiplication of the resulting value by 100 converts the ratio to a percentage.

In summary, the procedure for plotting data on normal probability paper is as follows:

- 1. Array data in order of ascending magnitude.
- 2. Calculate the plotting position of each value by the expression $\frac{m\,(100)}{n+1}$, which gives the ratio as a percentage. This point designates the percentage of the values that are equal to or less than the plotted value.
- 3. The Y-axis is graduated linearly from the lowest to the highest reported value, while the X-axis is graduated according to the probability scale. Place the first (lowest) value above the calculated plotting interval for m = 1 on the X-axis and at the appropriate value on the Y-axis. Plot the remaining values in a similar manner.

INTERPRETATION

If a straight line develops in the plotting, the data have a normal distribution; that is, in accordance with the theory of probability, this is the expected distribution of results.

If a straight line does not develop in the plotting, a change in the conditions affecting the observed measurements is suspected. It may mean, for example, that the same characteristic has not been measured under the same conditions.

Sometimes the great majority of the data approximate a straight line, but on the ends some results will be either extremely high or low. Just as one of these erratic results is far removed from the others on the bell-shaped normal curve, so it is far removed from the others on the straight-line curve. When this happens, these erratic results are presented in the published report on the probability curve, but the statistics are based on only the normal segment of the distribution.

REFERENCE

Velz, C.J. Graphical Approach to Statistics. Water and Sewage Works. pp. R106-R135. 1950

APPENDIX E.

ANALYTICAL REFERENCE SERVICE MEMBERSHIP

STATE AGENCIES

Alabama Water Improvement Commission

Arizona State Department of Health

Arkansas State Board of Health

California Department of Water Resources

California State Department of Public Health (Berkeley)

California State Department of Public Health (Los Angeles)

Colorado State Department of Public Health

Connecticut State Department of Health

Delaware Water Pollution Commission

Florida State Board of Health (Bureau of Laboratories, Jacksonville)

Florida State Board of Health (Division of Sanitary Engineering,

Jacksonville)

Florida State Board of Health (Pensacola)

Florida State Board of Health (Winter Haven)

Hawaii State Department of Health

Idaho Department of Health

Illinois State Department of Public Health (Chicago)

Illinois State Department of Public Health (Springfield)

Illinois State Water Survey Division

Indiana State Board of Health

Kansas State Board of Health (Sanitary Engineering Laboratories)

Kentucky State Department of Health

Louisiana State Board of Health

Maryland State Department of Health

Maryland State Water Pollution Control Commission

Massachusetts Department of Public Health

Michigan State Department of Health

Michigan Water Resources Commission

Minnesota State Department of Health

Missouri Department of Public Health and Welfare

Montana State Board of Health

Nassau County Department of Health

Nebraska State Department of Health

Nevada State Department of Health (Las Vegas)

Nevada State Department of Health (Reno)

New Hampshire State Department of Health

New Hampshire Water Pollution Commission

New Jersey State Department of Health

New Mexico State Department of Public Health

New York State Conservation Department

North Carolina Department of Water Resources New York State Department of Health North Dakota State Department of Health Ohio State Department of Health Oklahoma State Department of Health Oregon State Board of Health Pennsylvania Department of Health Puerto Rico Institute of Health Laboratories Rhode Island Department of Health South Carolina Water Pollution Control Authority South Dakota Department of Health Tennessee Division of Preventable Diseases Tennessee Stream Pollution Control Board Texas State Department of Health Utah State Department of Health Vermont State Department of Health Vermont State Department of Water Resources Virginia State Department of Health Virginia State Water Control Board Washington State Department of Health West Virginia State Water Resources Commission Wisconsin State Board of Health

MUNICIPAL AGENCIES

Air Pollution Control District, Pasadena, California Alexander Orr Jr., Water Treatment Plant, Miami, Florida City Department of Health, New York, New York City Department of Public Health, Pasadena, California City Department of Water, Dayton, Ohio City Department of Water Resources, Durham, North Carolina City Health Department, Baltimore, Maryland City Health Department, Beaumont, Texas City Health Department, Houston, Texas City Water Department, Charlotte, North Carolina Department of Air Pollution Control, Chicago, Illinois Department of Public Works and Utilities, Flint Water Plant, Flint, Michigan Department of Service and Buildings, Dayton, Ohio Department of Water and Sewers, South District Filtration Plant, Chicago, Illinois Erie County Laboratory, Buffalo, New York Long Beach Water Department, Long Beach, California Los Angeles County Flood Control District Los Angeles Department of Public Works, Hyperion Treatment Plant Los Angeles Department of Water and Power

Louisville Water Company, Incorporated

Metropolitan Utilities District, Omaha, Nebraska

Metropolitan Water District of Southern California

Philadelphia Department of Public Health

Philadelphia Department of Public Health (Occupational Environment Section)

Philadelphia Suburban Water Company

Philadelphia Water Department (Belmont Laboratory)

Philadelphia Water Department (Torresdale Laboratory)

St. Louis County Water Company

St. Louis Department of Health and Hospitals

Water Works, Topeka, Kansas

FEDERAL AGENCIES

Associated Universities, Incorporated, Brookhaven National Laboratory DHEW, PHS, Great Lakes—Illinois River Basins Project

DHEW, PHS, Northeastern Radiological Health Laboratory, Winchester, Massachusetts

DHEW, PHS, Northeast Shellfish Sanitation Research Center, Narragansett, Rhode Island

DHEW, PHS, Off-Site Radiological Safety Program, Las Vegas, Nevada DHEW, PHS, Water Quality Section, Division of Water Supply and Pollution Control, Cincinnati, Ohio

DHEW, PHS, Water Supply Section, Interstate Carrier Branch, Washington, D. C.

Food and Drug Administration, Division of Pharmacology, Washington, D. C.

Fourth U. S. Army Medical Laboratory, Fort Sam Houston, Texas Oak Ridge Institute of Nuclear Studies, Oak Ridge, Tennessee Pearl Harbor Naval Shipyard

Sixth U. S. Army Medical Laboratory, Fort Baker, California Tennessee Valley Authority, Chattanooga, Tennessee (Stream Pollution Control)

Tennessee Valley Authority, Wilson Dam, Alabama (Occupational Health Branch)

2793D U. S. Air Force Hospital, Regional Environmental Health Laboratory, McClelland Air Force Base, California

2794th U. S. Air Force Dispensary - Class B, Kelly AFB, Texas U. S. Air Force Radiological Health Laboratory, Wright-Patterson AFB, Ohio

U. S. Army Environmental Hygiene Agency, Maryland

U. S. Department of the Interior, Bureau of Reclamation, Denver, Colorado

U. S. Department of the Interior, Geological Survey, Columbus, Ohio

U. S. Department of the Interior, Geological Survey, Denver, Colorado

- U. S. Department of the Interior, Geological Survey, Philadelphia, Pennsylvania
- U. S. Department of the Interior, Geological Survey, Sacramento, California

Walter Reed Army Medical Center, Washington, D. C.

UNIVERSITIES

Case Institute of Technology, Department of Civil and Sanitary Engineering Columbia University University of Florida, Department of Chemistry Georgia Institute of Technology, Department of Applied Biology Johns Hopkins University, School of Hygiene and Public Health University of Kansas, School of Engineering and Architecture New Mexico State University, Department of Civil Engineering University of North Carolina, Chapel Hill, North Carolina University of Pittsburgh, Graduate School of Public Health Purdue University, Department of Chemistry Rensselaer Polytechnic Institute, Troy, New York Rutgers - The State University Department of Agricultural Chemistry Department of Environmental Science Washington State University, Division of Industrial Research

INDUSTRY

American Cyanamid Company, Bound Brook, New Jersey Anaconda Company, Grants, New Mexico Atlantic Refining Company, Philadelphia, Pennsylvania Battelle Memorial Institute, Columbus, Ohio Bethlehem Steel Company, Bethlehem, Pennsylvania California Water Service Company, San Jose, California Culligan, Incorporated, Northbrook, Illinois Dearborn Chemical Company, Chicago, Illinois Dow Chemical Company, Midland, Michigan E. I. du Pont de Nemours and Co., Aiken, South Carolina El Paso Natural Gas Products Company, El Paso, Texas Ekroth Laboratories, Incorporated, Brooklyn, New York Fairbanks, Morse, and Company, Research Center, Beloit, Wisconsin Food Machinery and Chemical Corporation, Pocatello, Idaho General Electric Company, Louisville, Kentucky Goodyear Atomic Corporation, Piketon, Ohio HALL Laboratories Division, Calgon Corporation, Pittsburgh, Pennsylvania

Hazleton Nuclear Science Corporation, Palo Alto, California Industrial Chemicals, Incorporated, South Bend, Indiana Ionics, Incorporated, Cambridge, Massachusetts Kennecott Copper Corporation, Salt Lake City, Utah Midwest Research Institute, Kansas City, Missouri Minute Maid Company, Anaheim, California Monsanto Chemical Company, St. Louis, Missouri NALCO Chemical Company, Chicago, Illinois Pacific Gas and Electric Company, Emeryville, California Pacific Gas and Electric Company, San Francisco, California Pan American World Airways, Patrick Air Force Base, Florida Pomeroy and Associates, Pasadena, California Radiation Detection Company, Mountain View, California Reynolds Electrical and Engineering Company, Las Vegas, Nevada Roy F. Weston, Incorporated, Newtown Square, Pennsylvania Sandia Corporation, Sandia Base, Albuquerque, New Mexico Shell Chemical Company, New York, New York Tracerlab, Incorporated, Richmond, California U. S. Industrial Chemicals Company, Tuscola, Illinois Water Service Laboratories, Incorporated, New York, New York

FOREIGN

British Coke Research Association, Chesterfield, Derbyshire, England Central Electricity Research Laboratories, Leatherhead, Surrey, England

Comissao Inter-Municipal de Controle da Poluicao das Aguas E Do Ar, Sao Paulo - Brasil

Department of Health Services and Hospital Insurance, Vancouver, B. C., Canada

Department of Municipal Laboratories, Hamilton, Ontario, Canada Department of National Health and Welfare, Ottawa, Ontario, Canada Department of National Health and Welfare, Occupational Health Division, Ottawa, Ontario, Canada

Department of National Health and Welfare, Vancouver, B. C., Canada Instituto Nacional de Obras Sanitarias, Caracas, Venezuela Metropolitan Water, Sewerage, and Drainage Board, Sydney, Australia National University of Colombia, Bogota, Colombia, South America Permutit Company, Limited, London, England Scientific Research Council, Kingston, Jamaica, West Indies

Taiwan Institute of Environmental Sanitation, PHA, Taiwan Pingtung Air Pollution Laboratory
Pingtung Organic Waste Laboratory
Taichung Water Laboratory
Tainun Water Laboratory
Taipei Milk Laboratory

Taipei Radiological Health Laboratory Taipei Water Laboratory Taitung Water Laboratory United Kingdom Atomic Energy Authority, Didcot, Berks, England University of Beograd, Civil Engineering Faculty, Beograd, Yugoslavia University of Leeds, Houldsworth School of Applied Science, Leeds, England

APPENDIX F.

STAFF AND ACKNOWLEDGMENTS

ANALYTICAL REFERENCE SERVICE STAFF

- E. F. McFarren, Chief R. J. Lishka, Chemist
- R. T. Cope, Statistician
- J. M. Matthews, Chemist
- P. A. Miller, Secretary

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- D. G. Ballinger, Supervisory Chemist Technical Advisory and Investigations Water Supply and Pollution Control
- J. M. Cohen, Chemist, In Charge Engineering Research
- R. C. Kroner, In Charge General Laboratory Services National Water Quality Network

GPO 820-837-7 75



ACCESSION NO. PIRLIOCPAPHIC: Robert A. Tare Sanitary Engineering Center, PATER OXYGEN DEM AND NO. 2. STUDY TEY WORLS NUME II : Public Feilth & ricc Publication No 999 N ≥ -26. 19€ 75 po ABSTRI. In Pris study consists of formsa aples which "a part upating laboratorie were just deted to dilite to a specified volume a danalyze by both the Star and Methor for Connical Dayger Demand and by the Merchane Sulfate modification. The results from this study indicate that the two procedures produce s mitar precision and accuracy when no interfering majerials are present When interferences due to high concentrations of chloride are present, the standard method will produce equal precision and accuracy only if the appropriate corrective techniques are applied. The Mercuric Sulfate mountreatter is the method of choice for COD measi rement since with less (over) BIBLIOGRAPHIC: Robert A. Taft Sanitary Engineering ACCESSION NO. Center. WATER OXYGEN DEMAND NO. 2. STUDY NUMBER 21. Public Health Service Publication KEY WORDS: No. 999-WP-26. 1965. 75 pp. ABSTRACT: This study consisted of four samples which 74 participating laboratories were instructed to dilute to a specified volume and analyze by both the Standard Method for Chemical Oxygen Demand and by the Mercuric Sulfate modification. The results from this study indicate that the two procedures produce similar precision and accuracy when no interfering materials are present. When interferences due to high concentrations of chloride are present, the standard method will produce equal precision and accuracy only if the appropriate corrective techniques are applied. The Mercuric Sulfate modification is the method of choice for COD measurement since with less (over) BIBLIOGRAPHIC: Robert A. Taft Sanitary Engineering ACCESSION NO. Center. WATER OXYGEN DEMAND NO. 2. STUDY NUMBER 21. Public Health Service Publication KEY WORDS: No. 999-WP-26, 1965, 75 pp. ABSTRACT: This study consisted of four samples which 74 participating laboratories were instructed to dilute to a specified volume and analyze by both the Standard Method for Chemical Oxygen Demand and by the Mercuric modification. The results from this study indicate that the two procedures produce similar precision and accur-

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