

GENERAL STANDARD
FOR
WATER POLLUTION CONTROL

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0. INTRODUCTION

The sources of pollution and T.L.V (Threshold Limit Value) of these pollutants in wastewater are described in IPS-E-SF-880 and IPS-E-PR-725. An effort has been made to present the most appropriate methods and equipment that apply as generally as possible for monitoring surface water, groundwaters, cooling or circulating water, boiler water, boiler feed water, wastewater effluents after varying degrees of treatment, and untreated municipal or industrial wastewaters. Here again, the effort has been made to present methods and equipment of the widest possible application, but when monitoring of sample of highly unusual composition are encountered, the equipment of this manual may require modification, or may be wholly inappropriate.

1. SCOPE

This Standard specifies the minimum requirements for material, operation, tests, inspection and covers site, process and laboratory measurements and calibration of instruments.

Note:

For further information refer to IPS-E-SF-880 "Water Pollution Control" and IPS-E-PR-735 "Process Design of Plant Waste Treatment and Recovery System".

2. REFERENCES

Throughout this Standard the following standards and codes are referred to. The editions of these standards and codes that are in effect at time of publication of this Standard shall, to the extent specified herein, form a part of this Standard. The applicability of changes in standards and codes that occur after the date of this Standard shall be mutually agreed upon by the Company and the Vendor/Consultant.

BSI (BRITISH STANDARD INSTITUTE)

BS 6068 6.2	"Water Sampling"
BS 2690 Pt 9 (1970)	"Appearance (Color and Turbidity) Odor, Suspended and Dissolved Solids and Electrical Conductivity"

ASTM (AMERICAN SOCIETY FOR TESTING AND MATERIALS)

ASTM Volume 11.02	"Water"
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IPS (IRANIAN PETROLEUM STANDARDS)

IPS-G-SF-860	"Air Pollution Control"
IPS-G-SF-110	"Protection Against Radioactive Sealed Sources"
IPS-E-SF-880	"Engineering Standard for Water Pollution"
IPS-E-PR-725	"Engineering, Standard for Process Design of Plant Control Wastewater Sewer Systems"
IPS-E-CE-340	"Water Resources and Distribution Systems"

Publication:

- J.M Krajca, Water Sampling, John Wiley & Sons, 1989.
- K.H. Mancy, Instrumental Analysis for Water Pollution Control, an Arbor Science, 1987.
- M.C. Rand, A.E. Green Berg, M.J. Taras, Standard Methods for the Examination of Water and Wastewater, American Public Health Association, 1987.
- H.D. Parker and G.D. Pitt Pollution Control Instrumentation for Oil and Effluents, Graham and Trotman, 1987.
- J. Rodier, L'Analyse De L'Eau, Eaux Naturelles, Eaux Residuares, Eau De Mer, Dunod Technique, 1990.
- W, Horwitz, Official Methods of Analysis, Association of Official Analytical Chemists, 1975.
- D.A Skoog, Principles of Instrumental Analysis, Holt, Rinehart and Winston, Inc., 1981.

3. DEFINITIONS AND TERMINOLOGY

3.1 Biochemical Oxygen Demand (BOD)

The mass concentration of dissolved oxygen consumed under specified conditions by the biological oxidation of organic and/or inorganic matter in water.

3.2 Chemical Oxygen Demand (COD)

Chemical Oxygen Demand (COD) is the equivalent amount of oxygen consumed under specified conditions in the chemical oxidation of the organic and oxidizable inorganic matter contained in a wastewater corrected for the influence of chlorides. In American practice, unless otherwise specified, the chemical oxidizing agent is hot acid dichromate.

3.3 Chemical Treatment

A process involving the addition of chemicals to achieve a specific result.

3.4 Contractor

Contractor is the person, firm or company whose tender has been accepted by the Company and includes the contractor's personnel representative, successors and permitted assigns.

3.5 Dissolved Oxygen (DO)

Dissolved Oxygen (DO) is the oxygen dissolved in sewage, water, or other liquid, usually expressed in milligrams per liter or percent of saturation. It is the test used in BOD determination.

3.6 Drinking Water, Potable Water

Water of a quality suitable for drinking purposes.

3.7 Effluent

Water or wastewater discharge from a containing space such as a treatment plant, industrial process.

3.8 Employer Company Owner

Refers to any of the related affiliated companies of the Petroleum Industries of Iran such as National Iranian Oil Company (NIOC), National Iranian Gas Company (NIGC), National Petrochemical Company (NPC), etc., as a part of the Ministry of Petroleum.

3.9 Ground Water

Water that fills all of the unlocked of material underlying the water table within the upper limit of saturation.

3.10 Industrial Water

Any water used for industrial purpose.

3.11 Oxygen Consumed

Oxygen consumed is the quantity of oxygen taken up from potassium permanganate in solution by a liquid containing organic matter commonly regarded as an index of the carbonaceous matter present. Time and temperature must be specified.

3.12 Pollution

Contamination of water with actively or potentially toxic or otherwise harmful materials.

3.13 Rain Water

Water arising from atmospheric perception, which has not yet collected soluble matter from the earth.

3.14 Raw Water

Water taken from natural sources, i.e. water wells or surface water.

3.15 Sewage Effluent

Treated sewage discharge from a sewage treatment works.

3.16 Site

Works or plant where sampling is to be carried out.

3.17 Sampling Point

A specific location on a sampling line at which an individual sample is extracted.

3.18 Standard Water

Water with known amount of component.

3.19 Surface Water

Water which flows over, or rests on the surface of a land mass.

3.20 Threshold Limit Values (TLVs)

Refers to maximum concentration of substances which could be discharged in wastewater issued by the national standard of environmental protection agency for industrial waste.

3.21 Total Organic Carbon (TOC)

TOC is a measure of the amount of carbon in a sample originating from organic matter only. The test is run by burning the sample and measuring the CO₂ produced.

3.22 Wastewater

Water discharged after being used in, or produced by a process and is of no further immediate value to that process.

3.23 Wet Chemical Method

Method based on chemical reagent not instrumental.

4. UNITS

This Standard is based on International System of Units (SI), except where otherwise specified.

5. ON SITE PORTABLE INSTRUMENTS

All parameters which have been mentioned in IPS-E-SF-880 (Maximum Effluent Standard for Industrial Waste) could be monitored by this portable instrument excluding organic and radioactive substances.

This equipment has been developed to meet the need for a simplified, convenient, accurate means of testing water in the field. The colorimetric tests are made with colorimeter using precalibrated meter scales for direct read-out.

The volumetric tests are conducted by titration using a unique buret and titration stand with a precision screw plunger. This device is shown at the right of the kit in Fig. 1. It dispenses titration solution so that accurate, reliable results are obtained from test to test.

5.1 Alternative Current Colorimeter

5.1.1 As mentioned above the basic of measurement is colorimetric which some reagent could be added to water sample to produce colored solution. This should be tested later. Typical type of portable instrument could be seen in Fig. 1. See also the following Technical Data:

TECHNICAL DATA

ALKALINITY

Titration for phenolphthalein and total alkalinity
Enough of each indicator for 100 tests.
Enough titrant for 100 average (125 ppm) tests.

CARBON DIOXIDE

Standard Titration procedure-
Enough reagent for approximately 100 tests.

CHLORIDE

Mercuric Nitrate Titration-
Enough reagent for approximately
100 tests.

CHLORINE

Improved Orthotolidine Method
Colorimetric Range: 0-1 ppm
Enough reagent for 60 tests.

IRON

Simplified Phenanthroline Method
Colorimetric Range: 0-3 ppm
Enough reagent for approximately 100 tests.

MANGANESE

Cold Periodate Oxidation Method
Colorimetric Range: 1-10 ppm
Enough reagent for approximately 100 tests.

NITRATE, NITROGEN

Cadmium Reduction-Diazotization Method
Colorimetric Range: 0-1.5 ppm N,
0-15 ppm N. 0-150 ppm N
Enough reagent for approximately 100 tests.

NITRITE, NITROGEN

Diazotization Method
Colorimetric Range: 0-0.2 ppm N,
Enough reagent for approximately 100 tests.

CHROMATE

Diphenylcarbohydrazide Method
 Colorimetric Range: 0-1.5 ppm
 Enough reagent for approximately 100 tests.

COLOR

Colorimetric Range: 0-500 APHA
 platinum-cobalt units.
 No reagents required.

COPPER

Cuprethol Method
 Colorimetric Range: 0-3 ppm
 Enough reagent for approximately 100 tests.

FLUORIDE

SPADNS Method
 Colorimetric Range: 0-2 ppm
 Enough reagent for approximately 10 tests.

HARDNESS, CALCIUM

EDTA Titration method
 Enough reagent for approximately 100 tests.

HARDNESS, TOTAL

EDTA Titration method
 100 tests.

HYDROGEN SULFIDE

Screen test color chart comparison
 . . . 0.1-5 ppm
 Enough test paper for 100 tests.

OXYGEN DISSOLVED

Improved Winkler method, Alkaline-Iodide-Azide Modification
 Titration...1 drop=1 ppm,
 1 drop=.2 ppm.
 Enough reagent for approximately 100 tests.

pH, WIND RANGE

Colorimetric Range: 4.0 -10,0
 Enough reagent for approximately 100 test.

PHOSPHATE, ORTHO & META

Stannous Reduction Method
 Colorimetric Range 0-2 ppm and 0-8 ppm.
 Enough reagent for approximately 100 tests

SILICA

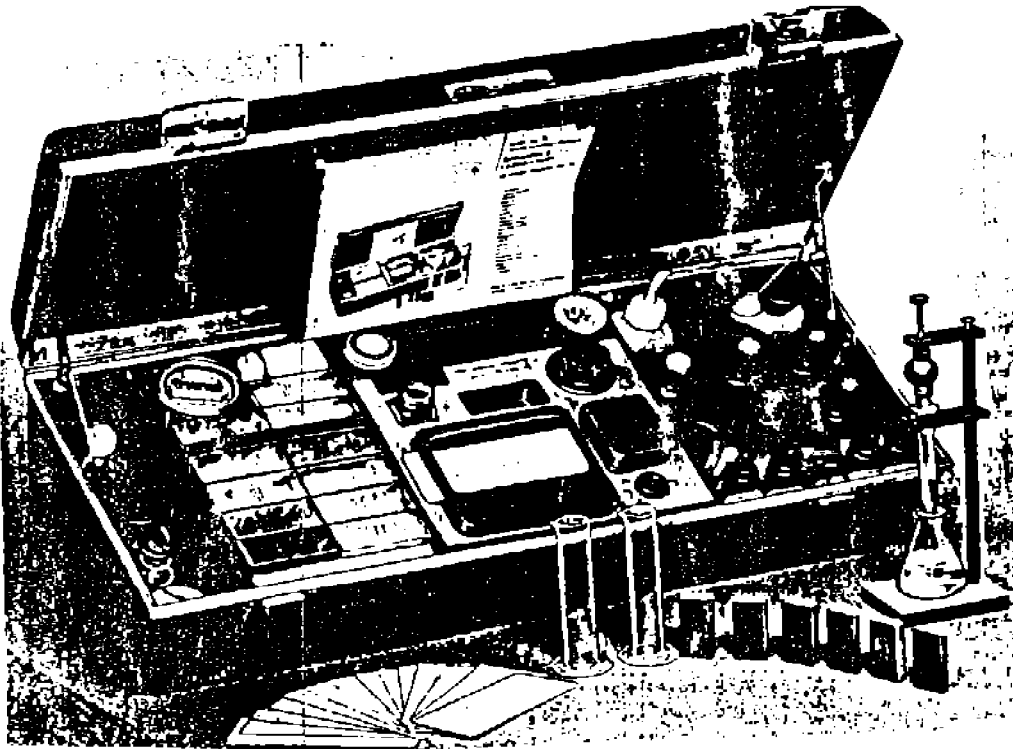
Heteropoly Blue Method
 Colorimetric Range: 0-3 ppm
 Enough reagent for approximately 100 tests.

SULFATE

Turbidimetric Method
 Range: 0-300 ppm
 Enough reagent for approximately 100 tests.

TURBIDITY

Absorption method
 Range: 0-500 JTU
 No reagents required.



ALTERNATIVE CURRENT OPERATED COLORIMETER

Fig. 1

5.1.2 For new equipment See catalog of laboratory equipment issued in August 1990, Environmental Science and Technology or any Relevant Journal.

5.2 Calibration and Inspection

5.2.1 In general this type of instrument is factory calibrated, but can be calibrated by competent person in accordance with the manufacturer's recommended standard.

5.2.2 All reagent and chemical must be freshed or in case of changing color and precipitation replaced by new one.

6. ON LINE FIXED MEASUREMENT OR CONTINUOUS MONITORING

The use of continuous monitoring equipment will permit the frequent measurement of several water quality parameters and the recording of this data at the measurement site or the transmitting of it to another location.

6.1 Continuous Water Sampling and Clarification System

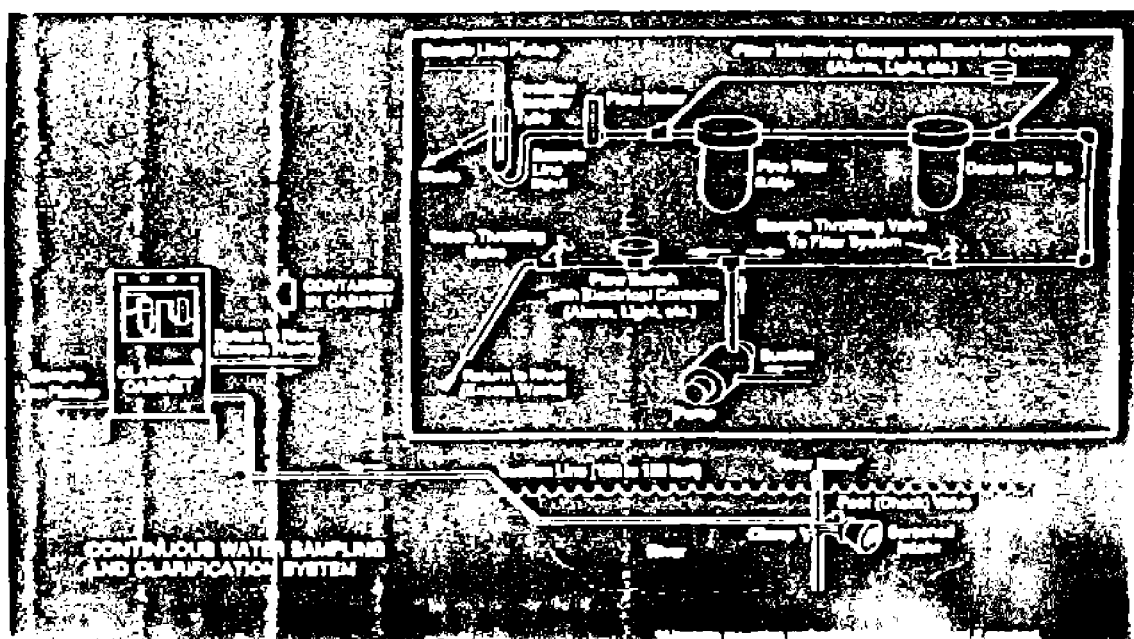
6.1.1 The monitoring instruments should contain a sampling module, the sensors, a signal conditioning module, and a data logging or transmission module. A continuous flow of sample must be provided to the sensors by a submersible pump.

Each sensor in the instrument should be supplied with its own signal conditioner which converts the input to a standard electrical output. For more detail on sensors and ion selective electrode. See K.H. Mancy, Instrumental Analysis for water pollution control, and Ann Arbor Science, 1987.

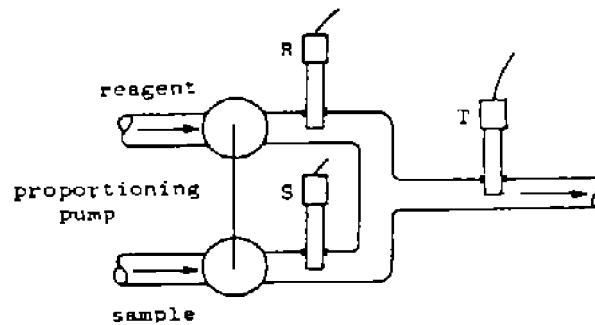
6.1.2 A typical continuous matter sampling and electrode placement for continuous monitoring is shown in Figs. 2 and 3. Further information has been given hereunder (see also ASTM Volume 11.01, 1989).

ELECTRODES, ION-SELECTIVE

Ion Determined	Recommend Reelectrode
Ammonia	Not applicable
Ammonium	Calomel
Barium	Calomel
Bromide	Double junction
Cadmium	Calomel
Calcium	Calomel
Chloride	Double junction
Copper	Calomel
Cyanide	Double junction
Fluoride	Double junction
Fluoroborate	Not applicable
Iodide	Double junction
Lithium	Double junction
Nitrate	Double junction
Oxygen	Not applicable
Potassium	Double junction
Silver/Sulphide	Double junction
Sodium	Double junction
Sulphur	Not applicable



CONTINUOUS WATER SAMPLING AND CLARIFICATION SYSTEM
Fig. 2



ELECTRODE PLACEMENT FOR CONTINUOUS MONITORING
Fig. 3

6.2 Calibration and Inspection

6.2.1 Calibration of the sensor signals should be carried out by adjustment of the controls on the signal conditioner.

6.2.2 The sample intake should be shielded with a screen and must be inspected according to manufacturer's instructions to prevent debris from clogging the system or damaging the pump.

7. LABORATORY INSTRUMENTS

All parameters which are summarized in the following sections could be monitored in laboratory using instrumental or wet chemical methods.

7.1 Collection and Preservation of Samples

7.1.1 Sampling of different sources

The summary of different kind of sampling is mentioned here, for more details See J.M. Kragca, Water Sampling, John wiley, 1989 and BS 6068, Water Sampling.

7.1.2 Sampling of atmospheric precipitation

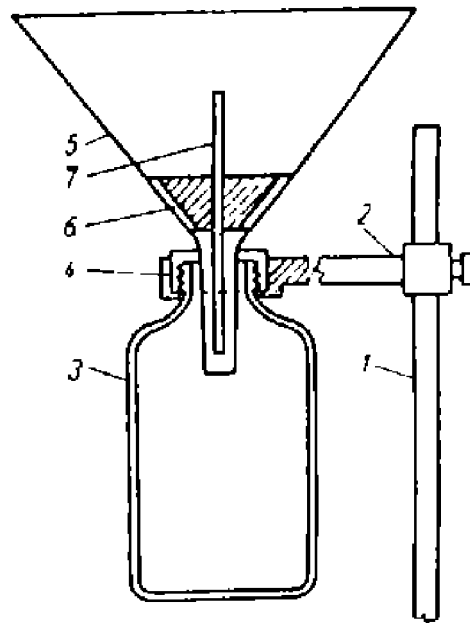
Atmospheric precipitation usually occurs in discrete events of limited duration.

7.1.2.1 Sampling precipitation equipment

Equipment for sampling precipitation, includes, a polyethylene polypropylene bucket and a stand.

7.1.2.2 Construction

The basic of this equipment is shown in Fig. 4. The equipment could be operated manually or automatically.



Funnel system for precipitation water sampling:

- 1) Stand
- 2) Adjustable support
- 3) Sample container
- 4) Threaded cap
- 5) Collecting funnel
- 6) Truncated grooved cone (fly trap)
- 7) Vent tube

SAMPLING OF ATMOSPHERIC PRECIPITATION

Fig. 4

7.1.2.3 Inspection

The equipment must be designed to minimize pollution during its operation. It should be manufactured from chemically inert materials to avoid contamination.

7.1.3 Sampling surface water

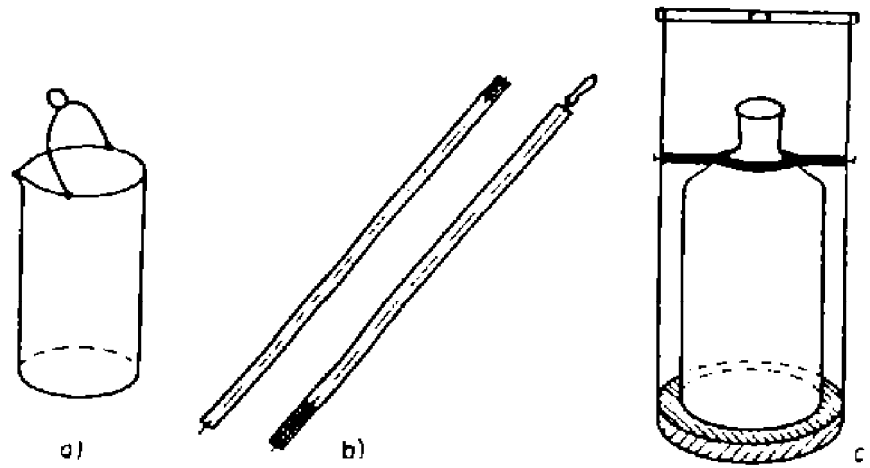
Surface water samples are usually taken directly into the sample container. It is not possible to take the sample by submerging the container by hand, laboratory forceps or a holder with a sliding sleeve should be used.

7.1.3.1 Surface water container

For such sampling the container, which should have a volume of at least two liters, is usually made of stainless steel.

7.1.3.2 Construction

The typical container is shown in Fig. 5.



- a) Metal can;
- b) segmented rod;
- c) submersible bottle with ballast and frame.

SIMPLE AIDS FOR SURFACE WATER SAMPLING

Fig. 5

7.1.4 Soil water sampling

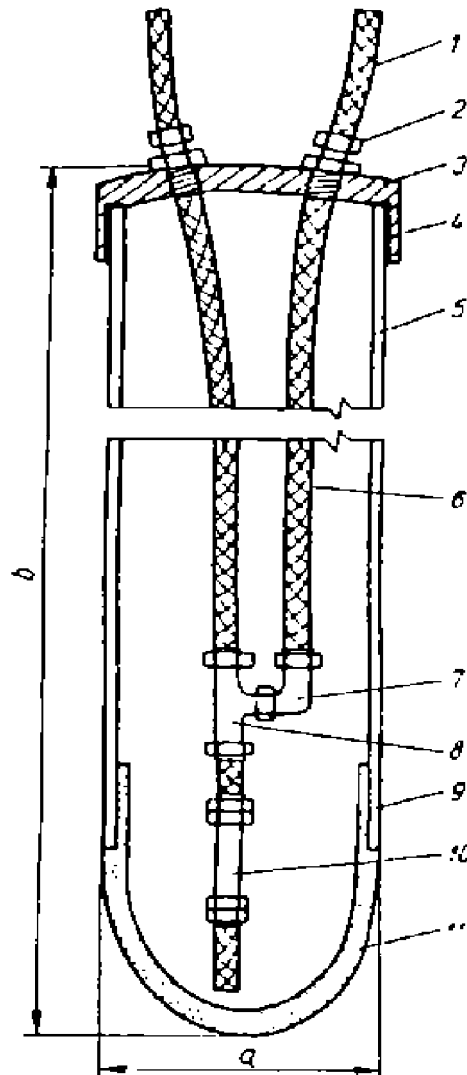
Soil water is defined as being all of the water contained in the soil.

7.1.4.1 Soil water sampling system

A common soil sampling system comprises a ceramic vessel with two pressure (vacuum) hoses inserted into it. The system operates by developing a vacuum through one hose, which causes the soil water to be drawn in through the porous walls of the vessel.

7.1.4.2 Construction

A schematic of soil water sampler is shown in Fig. 6.



- 1) And 6, braided, flexible pressure hose
- 2) Hose connector
- 3) PVC sampler cover
- 4) Seal between cover and sample pot
- 5) Upper wall of sample container
- 6) Elbow connector
- 7) T-piece connector
- 8) Glued connection between upper wall of sample container (PVC) and porous ceramic pot
- 9) Non-return valve
- 10) Porous ceramic pot. Typical dimensions _ a = 50 mm, b= 500 mm.

SOIL WATER SUCTION SAMPLER

Fig. 6

7.1.5 Groundwater sampling

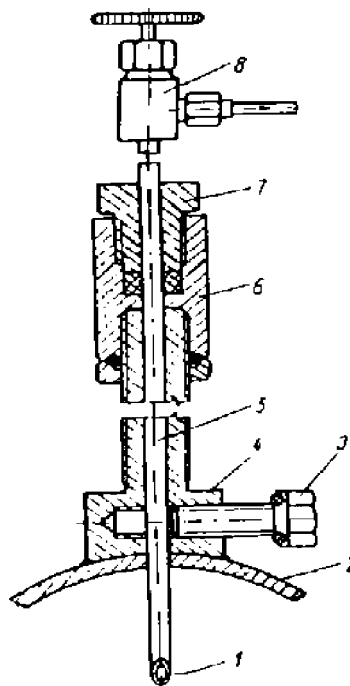
The sampling of groundwater is usually carried out with devices and equipment very similar to that used for surface waters.

7.1.5.1 Ground water sampling system

Similar containers are used for groundwater, mainly glass or polythene bottles.

7.1.5.2 Construction

A typical diagram of transferable sampler for use on wastewater pipelines is shown in Fig. 7.



- 1) Inlet (aperture cut obliquely across tube)
- 2) Pipeline wall
- 3) Sample point closing valve
- 4) Sampler inlet guide, welded to pipeline wall
- 5) Sampling tube
- 6) Connecting bush with o-ring seal
- 7) Locking nut
- 8) Needle valve

TRANSFERABLE SAMPLER FOR USE ON PIPELINES

Fig. 7

7.1.6 Different types of sampling

7.1.6.1 Grab or catch samples

A sample collected at a particular time and place.

7.1.6.2 Composite samples

In most cases the term composite sample refers to a mixture of grab samples collected at the same sampling point at different times.

7.1.6.3 Integrated samples

For certain, the information needed is provided best by analysis of mixtures of grab samples collected from different points simultaneously, or as nearly so as possible.

8. PHYSICAL EXAMINATION

8.1 Color

Color in water may result from the presence of metallic ion humus and peat materials, plankton, weed and industrial waste.

The unit of color produced by 1 mg/L platinum in the form of the chloroplatinate ion.

For color monitoring see ASTM Volume 11.02 and BS 2690: Part 9, 1970. Maximum concentration tabulated in IPS-E-SF-880.

8.2 Conductivity

Conductivity is a numerical expression of the ability of a water sample to carry an electric current and is defined as μ moh/cm². This number depends on the total concentration of the ionized substances dissolved in the water. See also BS 2690: Part 9 1970.

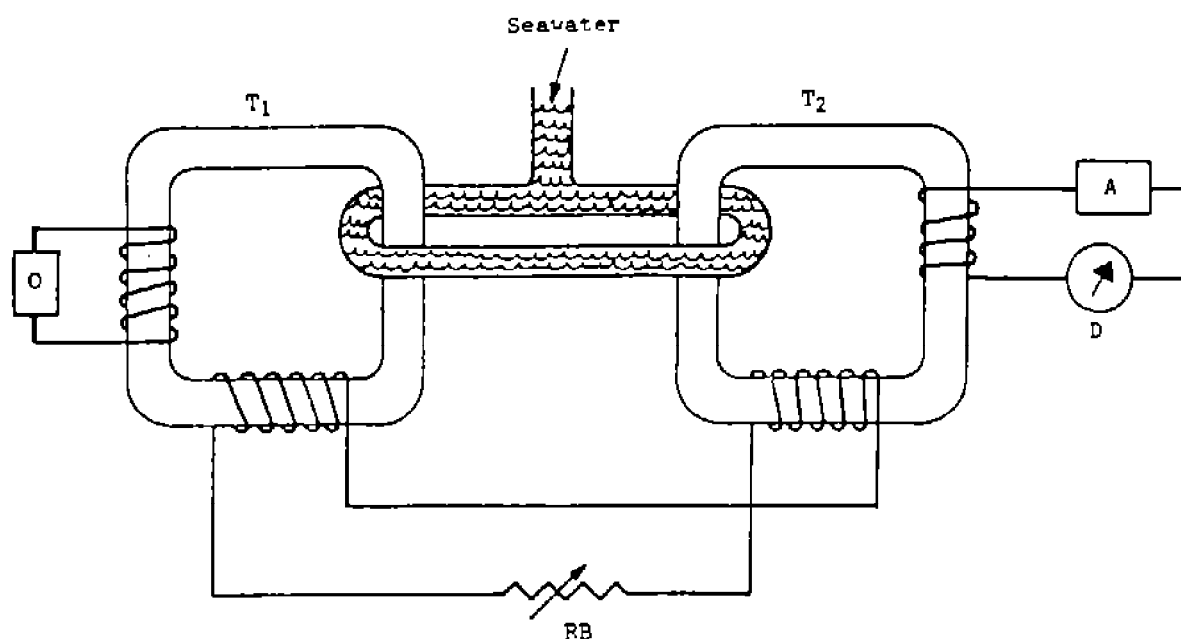
8.2.1 Conductivity instruments

Measurement of electrical conductance is usually based on the use of conductivity bridges (whetstone bridge). The bridge circuit should always be arranged so that at the balance point the detector indicates either zero or minimum potential. Basic of instrument is shown in Fig. 8.

The commercial form is an electrode connected to electrical circuit. Best result can be obtained with the following technical data:

	<u>TECHNICAL</u>	<u>DATA</u>
Ranges	0-1000	mS
	0-199.9	mS
	0-19.99	mS
	0-1999	μ S
	0-199.9	μ S
	0-19.99	μ S
	0-100.0	°C
	(automatic selection)	
Resolution	1	mS
	0.1	mS
	0.01	mS
	1	μ S

	0.1 μS
	0.01 μS
	0.1 $^{\circ}\text{C}$
Accuracy	$\pm 0.5\%$
Temperature compensation range	0 to 100°C
Slope auto temp comp	Variable from 0.5 to $4\%/^{\circ}\text{C}$
Cell constant	0.05 to 19.99 digitally settable
Outputs	Analogue 1 mV/digit, RS 232
Display	Conductivity 20 mm LED
	Temperature 10 mm LED
Overall dimensions (H \times W \times D)	120 \times 300 \times 180 mm
Mass	2kg
Power requirements	115/230 V ac $\pm 20\%$ 50/60 Hz



- T_1 First electrical circuit
- T_2 Second electrical circuit
- RB Variable resistance
- A Ampere
- D Conductivity meter

" INDUCTIVE-TYPE" CONDUCTIVITY CELL
Fig. 8

8.2.2 Calibration

Calibration should be carried out with standard water.

8.3 Odor

Odor and taste are known as the "chemical senses" since they depend on actual contact of the stimulating substance with the appropriate human receptor cell.

For odor monitoring See M.C. Rand, A.E. Greenberg, standard methods for the examination of water and wastewater, American Public Health Association, 1987 and BS 2690 Part 9 1970. For maximum odor see IPS-E-SF-880.

8.4 Residue

The term residue refers to solid matter suspended or dissolved in water or wastewater.

8.5 Salinity

It is defined as the total solid in water after all carbonates have been converted to oxides, all bromide, and iodide have been replaced by chloride and all organic matter oxidized.

8.6 Specific Gravity

Specific gravity is determined by comparing the weight of a volume of the sample of mud, sludge, or industrial waste with that of an equal volume of distilled water.

8.7 Taste

Taste, like odor, is one of the chemical senses.

8.8 Temperature

Normally, temperature measurement may be made by mercury-filled Celsius thermometer with scale marked for every 0.1°C.

8.9 Turbidity

Turbidity is an expression of the optical property that causes light to be scattered and absorbed rather than transmitted in straight lines through sample.

Turbidity in water is caused by the presence of suspended matter, such as clay, silt, finely divided organic and inorganic matter, plankton, and other microscopic organisms. See also BS 2690: Part 9, 1970.

Note:

For Clauses 9.4 to 9.9 the above reference, or ASTM 11.02 should be seen. For maximum characteristic see IPS-E-SF-880.

9. DETERMINATION OF METALS (as defined in Chemical Periodic Table)

The presence of metals in potable water, domestic wastewater, industrial effluents is a matter of serious concern. Metals could be monitored by atomic absorption spectroscopy, polarography, ICP and colorimetric methods.

9.1 Atomic Absorption Spectroscopy

In atomic absorption the sample is atomized into a flame, producing atomic vapor of the elements in question. The atoms absorb the radiation from the light source. The amount of light absorbed is proportional to the amount of the element.

9.1.1 Atomic absorption instrument

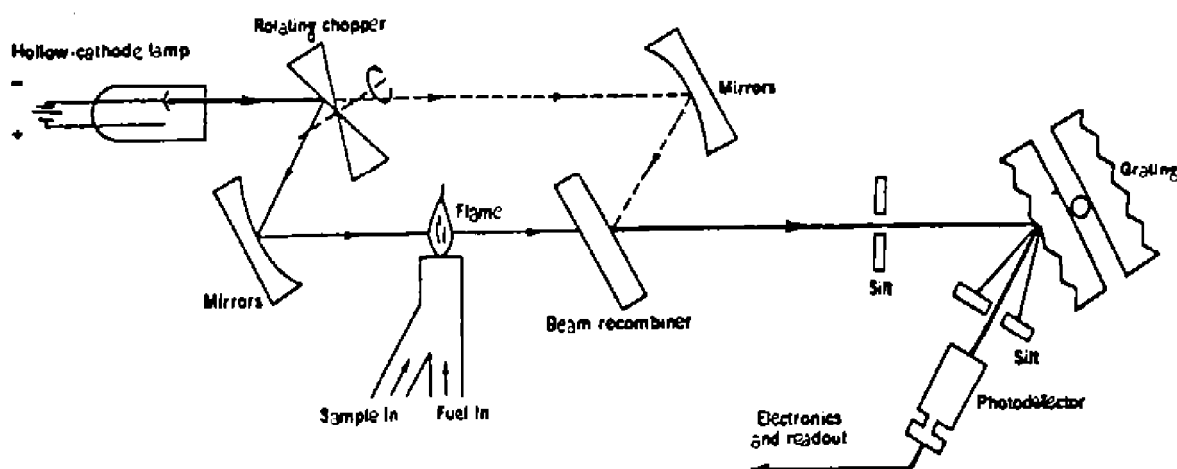
Fig. 9 shows schematic diagram of an instrument. A mechanical modulated chopper alternately passes and reflects the light beam. One beam by-passes the sample and its intensity is measured as I_0 the sample's beam is measured as I_1 . The absorbed light is as $I_0 - I_1$. See also the following technical data:

TECHNICAL DATA

Monochromator	:	Focal length 267 mm.
Wavelength	:	Range 190-870 nm. 1800.
Lines/mm		
Dispersion	:	1.6 nm/min
Photometer	:	Double-beam
Deuterium background corrector	:	Internally mounted correction for molecular absorption
Wavelength drive	:	5 nm/min
Burner	:	Stainless steel
Power requirements	:	105-125 or 200-240 Volts, 50/60 Hz, 150 Watts

9.1.2 Calibration

The instrument should be calibrated according to manufacturer's manual book.



ATOMIC ABSORPTION SPECTROMETER

Fig. 9

For more detail see D.A. Skoog, Principles of instrumental analysis, Holt Rinehart and Winston, Inc., 1981 the toxic pollutants such as Pb, As, Hg with low concentration (PPb) part per billion should be monitored by special device (vapor Generation and Hydride Generation) which could be installed on atomic absorption instrument. Detail has been written in Atomic Absorption manual book.

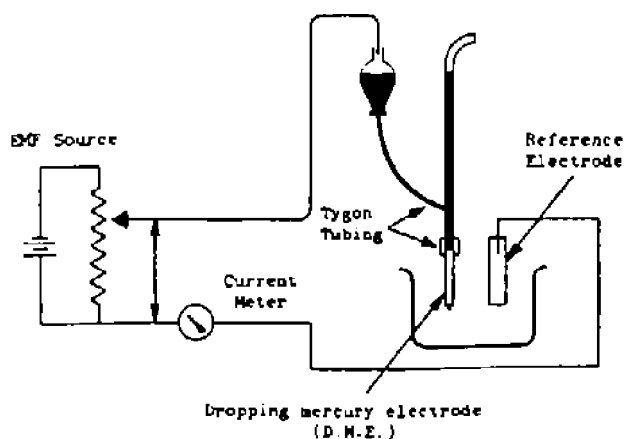
9.2 Polarography

Electrochemical methods are used for analysis of Raw water and wastewaters.

9.2.1 Polarography instrument

A schematic diagram of a classical polarograph is shown in Fig. 10.

Polarographic measurement is based on determining the time-averaged current of the Dropping Mercury Electrode (DME) under diffusion condition.



SCHEMATIC DIAGRAM OF CLASSICAL POLAROGRAPH
Fig. 10

For more details see D.A. Skoog, Principles of Instrumental Analysis, Holt, Rinehart and Winston, Inc., 1981 and Technical Data.

TECHNICAL DATA

Concentration range:	10^{-2} to 10^{-6} M (Mole)
Initial voltage:	Infinitely variable -5 and +5
Voltage range:	-0.75V, -1.5V, -3V
Voltage speed:	Adjustable from 1 to 500 mv/S
Current range:	Adjustable from 20 nA to 10 mA

9.2.2 Calibration

It should be calibrated according to instrument's manual book.

9.3 UV-Visible Spectroscopy

For UV-VIS spectroscopy See IPS-G-SF-860.

9.4 ICP Spectrometry

For inductively coupled argon plasma spectrometry See IPS-G-SF-860.

10. DETERMINATION OF INORGANIC NON-METALLIC CONSTITUENTS

10.1 Acidity and Alkalinity

For above chemicals See ASTM 11.02 and J.Rodier, L'Analyse De L'Eau Faux naturelles, Eaux Residuaire, Eau De Mer, Dunod Technique, 1990.

10.2 Arsenic

Arsenic could be monitored by wet chemical method or hydride generation technic. Detail is described in M.R. WPCF, "Standard Methods for The Examination of Water and Wastewater, American Public Health Association, 1987."

10.3 Boron

Monitored by UV-VIS spectroscopy see 9.3.

10.4 Carbon Dioxide

Measured by titrimetric method see ASTM Volume 11.02.

10.5 Chloride

Chloride, in the form of Cl Ion, is one of the major inorganic anions in water and wastewater. Chloride could be monitored by titrimetric, potentiometric and Ion selective electrode. More detail could be obtained in ASTM Volume 11.02 or W, Horwitz, Official Methods of Analysis, Association of Official Analytical Chemists 1975.

10.6 Chlorine (Residual)

The chlorination of water supplies and polluted water serves to destroy or deactivate disease, producing micro-organisms. It could improve water quality by reaction with ammonia, Iron, Manganese, Sulfide and some organic substances. It should be monitored by titrimetric, colorimetric techniques. More detail should be obtained in ASTM Volume 11.02.

10.7 Cyanide, Fluoride, Iodide

These three pollutants should be measured by titrimetric and/or colorimetric methods (see IPS-G-SF-860, UV-VIS spectroscopy) and Ion selective electrodes.

Detail about Ion selective electrode could be provided in D.A. Skoog, D.M. Wes, principles of instrumental analysis, Holt Rinehart Winston, 1981.

For cyanide See also BS 2690: Part II, 1971.

10.8 Nitrogen (Ammonia, Nitrate, Organic)

Nitrogen group should be monitored by titrimetric and/or colorimetric methods (see IPS-G-SF-860), and for ion selective electrodes see Clause 11.7.

10.9 Oxygen

Oxygen should be monitored by specific electrode for ISE. See clause 11.7 or titrimetric method as detailed in ASTM Volume 11.02.

10.10 Ozone

Ozone should be measured by titrimetric method as given in ASTM Volume 11.02.

10.11 pH Value

The pH value of a solution is defined as the logarithm of the reciprocal of the hydrogen-Ion concentration. Detail on effect of pH could be provided in API, manual on disposal of refinery wastes volume on liquid wastes, Chapter 2. pH should be monitored by electronic pH meter.

10.11.1 pH meter instrument

The pH meter consists of electronic part, glass electrode, reference electrode. See also the following Technical Data.

TECHNICAL DATA

Digital pH meter range	:	0-00 to 14.00 pH
mV range	:	-1999 to +1999 mV
°C range	:	-50.0 to +199.9 °C
Accuracy	:	Better than 0-01 pH
Output, analogue	:	1 mv/Digit

10.11.2 Calibration

It should be calibrated with standard buffer solution with known pH.

10.12 Phosphate

Phosphate content should be monitored by colorimetric method. For detail see ASTM 11.02 and BS 6068 Section 2.28 1986.

10.13 Silica

Silica concentration could be measured by calorimetric, gravimetric method. For detail See ASTM 11.02.

10.14 Sulfate

Sulfate could be measured by gravimetric method. For detail See M.R. WPCF, standard methods for the examination of water and wastewater, American Public Health Association, 1985.

10.15 Sulfide

For sulfide monitoring see Clause 10.14 above, but in general it should be measured by colorimetric and titrimetric methods.

11. DETERMINATION OF ORGANIC CONSTITUENTS

11.1 Grease and Oil

11.1.1 Solvent extraction infrared absorption

In this sort of monitoring the oil is extracted from sample and the infrared absorption of the oil in solution is then measured.

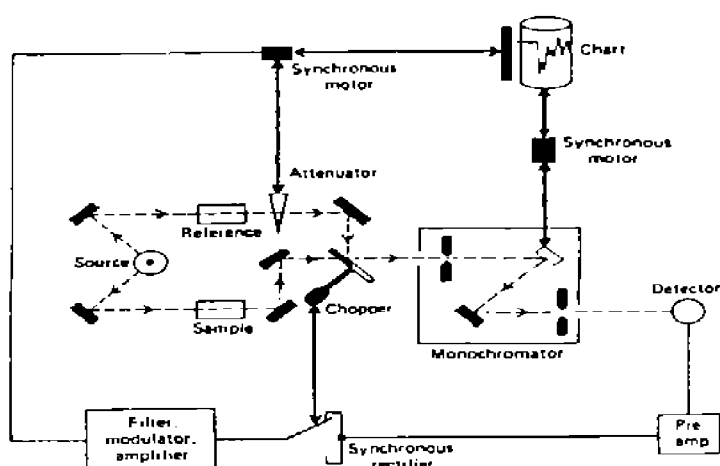
11.1.1.1 Infrared spectrophotometers instrument

Radiation from the source is split into two beams, half passing into the sample cell and the other half into the reference cell. The reference beam then passes through the attenuator and onto the chopper. After dispersion by the prism or grating, the alternating beams fall on the detector and are converted to an electrical signal.

Schematic diagram is shown in Fig. 11. See also the following Technical Data:

TECHNICAL DATA

Accuracy:	$\pm 6 \text{ Cm}^{-1}$ from 4000 to 2000 Cm^{-1} $\pm 3 \text{ Cm}^{-1}$ from 2000 to 600 Cm^{-1}
Repeatability:	$\pm 1 \text{ Cm}^{-1}$ from 4000 to 2000 Cm^{-1} $\pm 0.5 \text{ Cm}^{-1}$ from 2000 to 600 Cm^{-1}
Resolution:	5.6 Cm^{-1} at 3000 Cm^{-1}
Spectral slit width:	2.8 Cm^{-1} at 1000 Cm^{-1}
Stray light %T:	Less than 0.3 %
Noise:	0.8 %T



Heavy dark line= mechanical linkage;

Light line = electrical linkage;

Dotted line = radiation path

SCHEMATIC DIAGRAM OF A DOUBLE-BEAM SPECTROPHOTOMETER

Fig. 11

11.1.1.2 Calibration

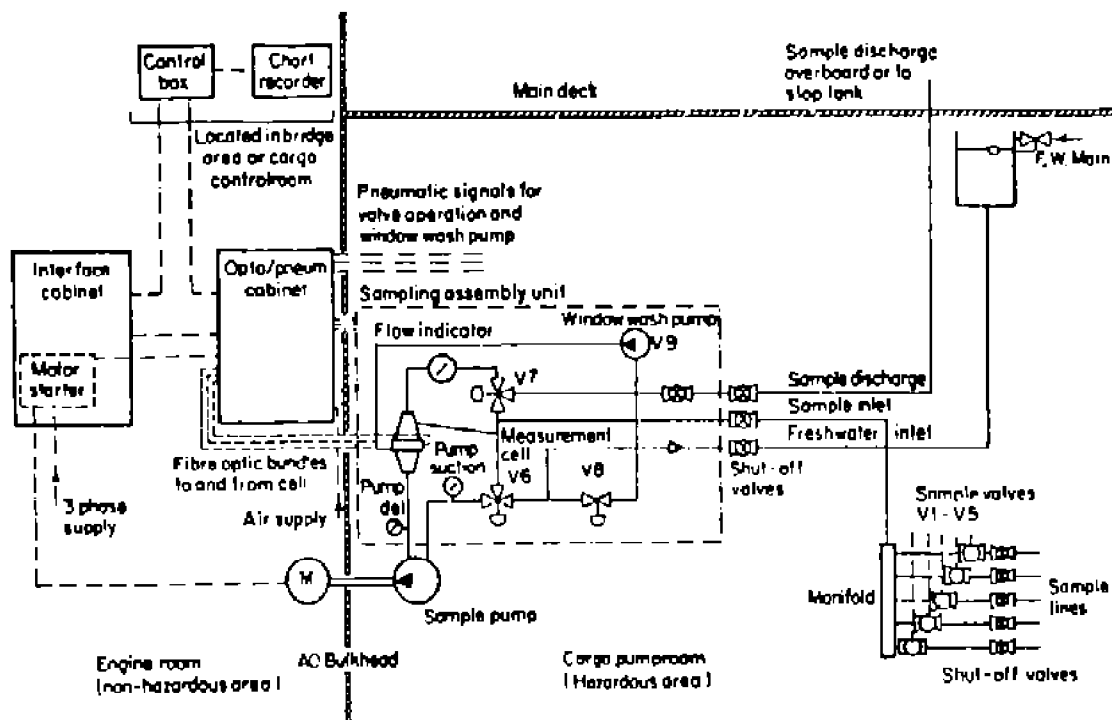
Calibration of instrument should be according to manufacturer's manual book.

11.1.2 Optical fiber

This method is the use of bared optical fiber immersed in the discharge stream.

11.1.2.1 Optical fiber monitoring instrument

Water has a refractive index of 1.33-1.38 and oil or hydrocarbon have near 1.5 so radiation would be lost from the fiber when oil droplets became attached. For radiation helium-neon laser could be used. Schematic diagram of monitoring by this system is shown in Fig. 12.



A fibre optic link joins the measurement cell with the electronics box.

SCHEMATIC DIAGRAM OF A TANKER'S BALLAST MONITORING SYSTEM

Fig. 12

11.1.2.2 Calibration

Calibration should be according to manufacturer's manual book.

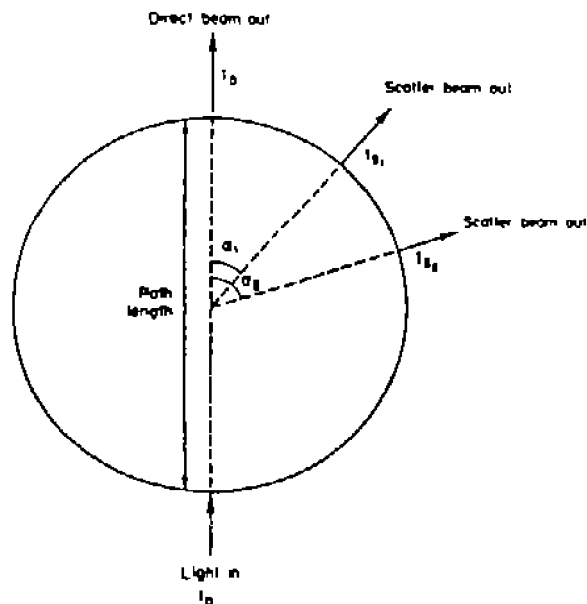
11.1.3 Scatter and turbidity

Oil concentration could be monitored by light scatter from oil droplets in the sample.

11.1.3.1 Optical scatter instrument

The light source should preferably have a wave length at which the water absorption is low, white light sources or near infrared emitters at about 0.9 μm can be used.

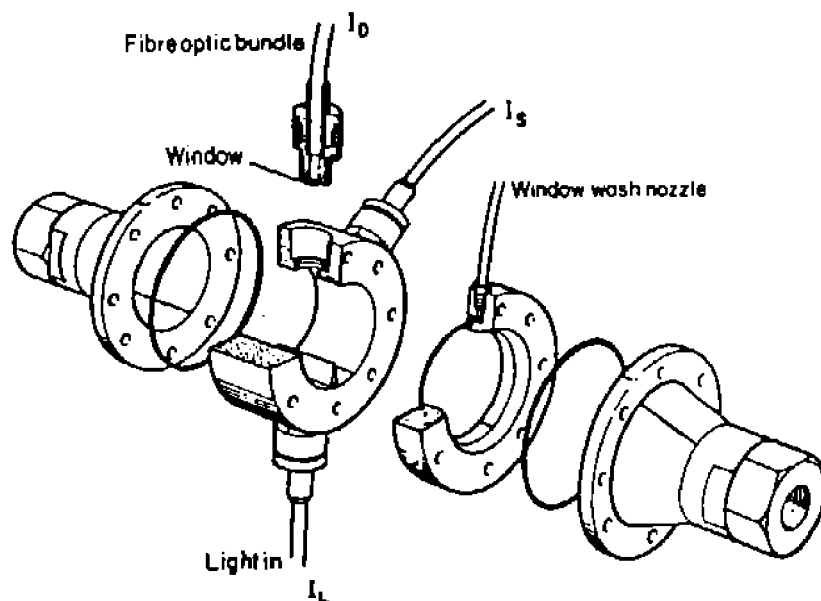
Basic of this method is shown in Fig. 13.



SCATTER DIAGRAM, SCHEMATIC REPRESENTATION OF A TYPICAL SCATTER CELL WITH THE FLOW PERPENDICULAR TO THE PAGE

Fig. 13

Also schematic diagram of an optical scatter cell is provided in Fig. 14. More detail is provided in H.D. Parker and G.D. PITT, Pollution Control Instrumentation for Oil and Effluents, Graham and Trotman, 1987.



The device is designed for incorporation of optical fibres.

The windows are cleaned by periodically passing a jet of clean water through adjacent slots.

SCHEMATIC DIAGRAM OF AN OPTICAL SCATTER CELL WITH DETECTION AT ONE FORWARD ANGLE

Fig. 14

11.1.3.2 Calibration

Calibration should be according to manufacturer's manual book.

11.2 Methane

11.2.1 Volumetric method

For volumetric method. See ASTM Volume 11.02.

11.2.2 Gas chromatograph

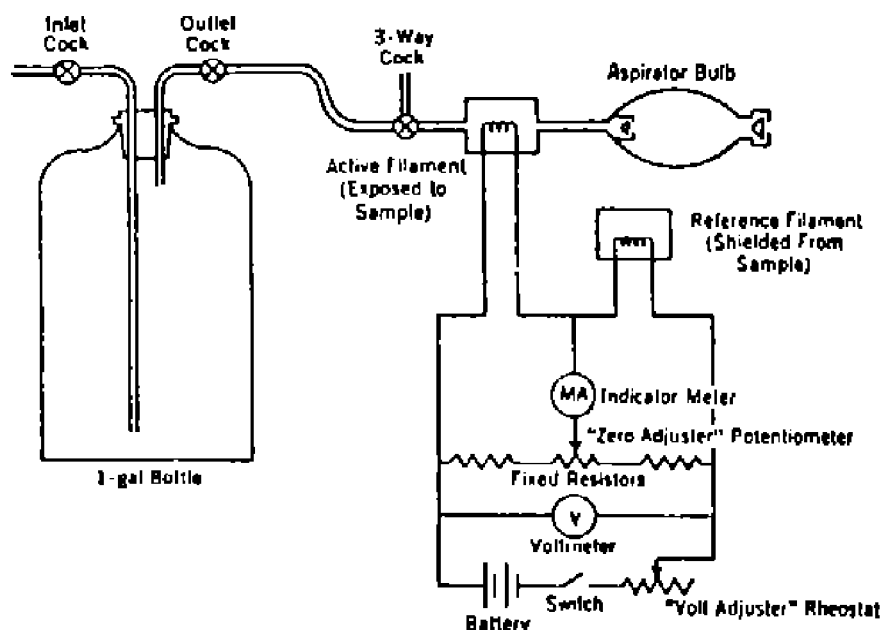
For equipment and basic of gas chromatograph. See IPS-G-SF-860.

11.2.3 Combustible gas indicator

An equilibrium is established between methane in solution and the partial pressure of methane in the gas phase above the solution. The partial pressure of methane could be determined with a combustible gas indicator.

11.2.3.1 Combustible-gas indicator instrument

The heat generated by the oxidation of the gas increases the electrical resistance of the filament. The resulting imbalance of the electrical circuit causes deflection of a milliammeter. A diagrammatic view of the apparatus is shown in Fig. 15.



COMBUSTIBLE-GAS INDICATOR CIRCUIT AND FLOW DIAGRAM

Fig. 15

11.2.3.2 Calibration

The milliammeter should be calibrated in terms of the percentage of methane.

11.3 Organic Acid and Volatile Acids

Organic acid and volatile acids should be monitored by chromatographic and steam distillation. See ASTM Volume 11.02 and IPS-G-SF-860.

11.4 Organic Carbon (Total)

Organic Carbon TOC could be monitored by carbon analyzer in the range of 1 to 150 mg/L in water and wastewater.

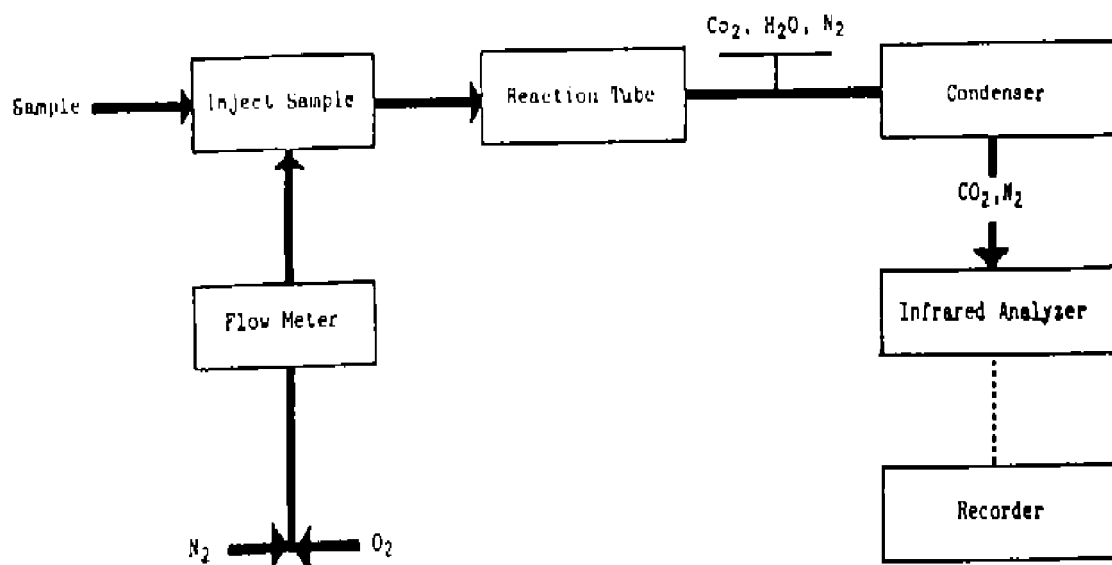
11.4.1 Total carbon analyzer instrument

Microportion of sample is injected into a heated packed tube in a stream of oxygen or purified air.

The water is vaporized and the organic matter is oxidized to carbon dioxide, which is measured by means of a nondispersive type of infrared analyser. Flow diagram is shown in Fig. 16. See also the following Technical Data:

TECHNICAL DATA

Sampler	:	Quartz boat
Combustion temperature	:	800 °C
Carrier gas	:	N ₂
Combustion gas	:	O ₂
Measuring range	:	1 g ^c /l
Sensitivity	:	0.1 mg ^c /l
Voltage	:	220 V, 50 Hz
Power input	:	2.35 kVA



TOTAL CARBON ANALYZER FLOW DIAGRAM

Fig. 16

11.4.2 Calibration

The instrument should be calibrated according to manufacturer's manual book.

11.5 Oxygen Demand (Biochemical)

The Biochemical Oxygen Demand (BOD) is a test in which standardized laboratory procedures should be used to determine the relative oxygen requirements of wastewaters, effluents, and polluted waters.

For this procedures see ASTM Volume 11.02. See also BS 6068 Section 2.3 1984.

11.6 Oxygen Demand (Chemical)

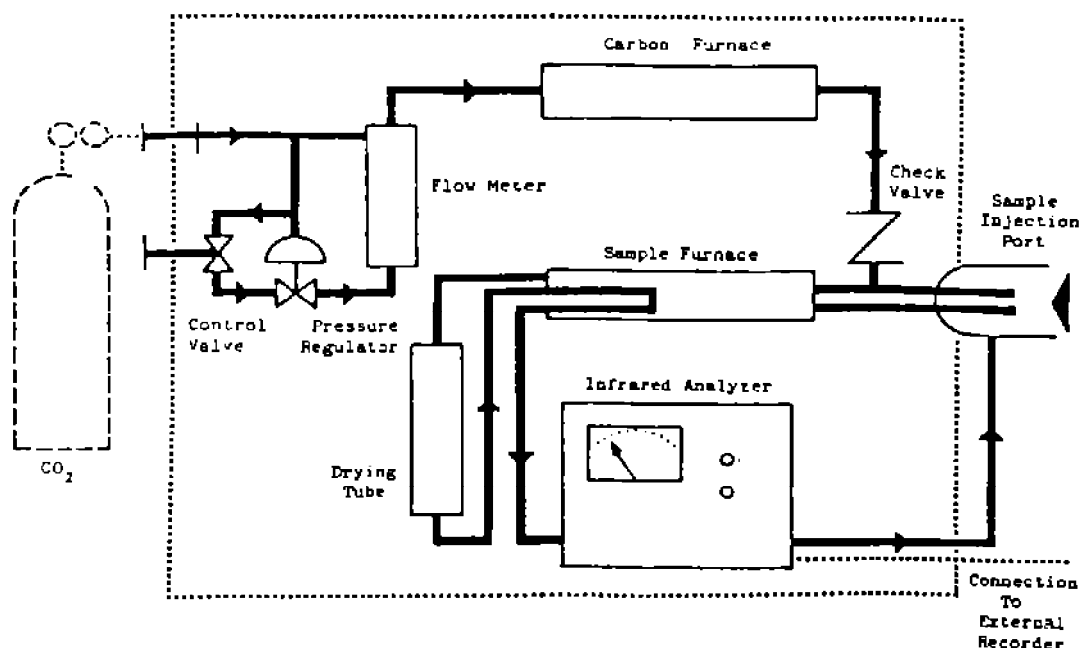
COD is an important measured parameter for stream and industrial waste control.

11.6.1 COD analyzer instrument

By using dry CO_2 to carry the organic matter through a platinum catalytic combustion furnace which oxidizes it to CO and H_2O . Following water removal and passage through a second catalytic treatment, the CO concentration is measured using an infrared analyser. A schematic of instrument is shown in Fig. 17. See also the following Technical Data:

TECHNICAL DATA

Sampler	:	Quartz boat
Combustion temperature	:	800 °C
Carrier gas	:	CO_2
Measuring range	:	1 g O/l
Sensitivity	:	0.1 mg O/l
Voltage	:	220 V, 50 Hz
Power input	:	2.35 KVA



COD ANALYZER FLOW DIAGRAM
Fig. 17

11.6.2 Calibration

Calibration should be according to instrument's manual book.

11.7 Pesticides

Gas chromatographic procedure should be used for quantitative determination of pesticides which is fully described in IPS-G-SF-860.

11.8 Phenols

Phenols, defined as hydroxy derivatives of benzene and its condensed nuclet, may occur in domestic and industrial wastewaters. It should be monitored by photometric and gas chromatographic methods which are described in IPS-G-SF-860. For column packing, carrier gas refer to ASTM Volume 11.02 1985.

11.9 Surfactants

One of the pollutant which must be controlled in wastewater is surfactants (Synthetic Detergents). This should be monitored by infrared equipment or spectrophotometer. For infrared description see 12.1.1 and for spectrophotometer, IPS-G-SF-860. For surfactants monitoring. See also BS 6068: Section 2.24, 1968.

12. EXAMINATION OF WATER AND WASTEWATER RADIOACTIVITY

The radioactivity in water and wastewater originates from natural and artificial or man-made sources. Artificial sources of radioactivity include fission, fusion, or particle acceleration, giving rise largely to alpha, beta and gamma radioactivity. For more detail . See IPS-G-SF-110 "Protection Against Radioactive Sealed Sources" and ASTM 11.02.

12.1 Counting Room

The room should be free of dust and fumes that may affect the electrical stability of instrument. The background could be stabilized and lowered considerably by making the walls, floor, and ceiling out of several centimeters of concrete. Generally, temperature could be constant within 3°C and should not exceed 30°C.

Samples containing appreciable activity should be stored at a distance so as not to affect instrument background counting rate.

12.2 Alpha Particle Counter Instrument

Alpha particle counter, consisting of either a proportional detector or a scintillation detector, and a scaler conforming to the following requirements. For method, see ASTM 11.02 1989 and Technical Data.

12.2.1 Proportional detector

This may be one of several types commercially available. The material used in the construction of the detector should be free from detectable radioactivity.

The manufacturer should supply voltage plateau and background counting rate data. Voltage plateau data should show the threshold voltage, slope, and length of plateau for a particular input sensitivity.

12.2.2 Scintillation detector

It should consist of an "activated" zinc sulfide phosphor having a minimum effective diameter of 36.5 mm. The phosphor should be mounted so that it could be attached and optically coupled to a multiplier photo tube.

12.2.3 Scaler

Mechanical register, power supply, and amplifier are contained in a single chassis, generally termed the scaler.

12.2.4 Sample mounting disks or dishes

Having a flat bottom of a diameter slightly less than the inside diameter of the detector. Flat disks are preferred, but dishes could be used that have 3.2 mm high side walls. Platinum and stainless steel have been used for this purpose.

12.2.5 Calibration and standardization for general monitoring

Place a known amount of alpha standard into a volume of water sufficient to dissolve salts equivalent to those of the test samples and prepare for counting.

12.3 Beta Particle Radioactivity Instrument

Beta particle radioactivity of water and wastewater in general monitored by beta particle counter which consist of the following components:

12.3.1 Detector

The end-window Geiger-Muller tube and the internal or external proportional gas-flow chambers are the two most commercially available types of detector.

12.3.2 Detector shield

The detector assembly shall be surrounded by an external radiation shield of massive metal equivalent to approximately 51 mm of lead and lined with 3.2 mm thick aluminum.

12.3.3 Scaler

Described in 12.2.3.

12.3.4 Sample mounting dishes

Described in 12.2.4.

12.3.5 Calibration for general monitoring

See 12.2.5. For more technical data see ASTM Volume 11.02, 1989.

12.4 Gamma-Ray Monitoring

This section covers the monitoring equipment for gamma-ray emitting radio nuclides in water or wastewater by means of gamma-ray spectrometry.

12.4.1 Gamma-Ray instrument

Gamma-ray spectra are measured with modular equipment consisting of a detector, and analyzer, memory and a permanent data storage device. Lithium-drifted detectors, P-Type or N-Type, are used. A multichannel pulse-height analyzer should be used to determine the amplitude of each pulse originating in the detector.

12.4.2 Shield

See 12.3.2.

12.4.3 Calibration and standardization

See 12.2.5. For more technical data see ASTM Volume 11.02, 1989.

13. MICROBIOLOGICAL EXAMINATION OF WATER

In this section there should be suitable procedures in making microbiological examinations of samples of water to determine sanitary quality and suitability for general use. For more detail about these procedures following standards could be seen; ASTM, Volume 11.02, 1989 and standard methods for the examination of water and wastewater, American Public Health Association, 1985. See also Appendix A and IPS-E-CE-340 Water Resources and Distribution Systems.

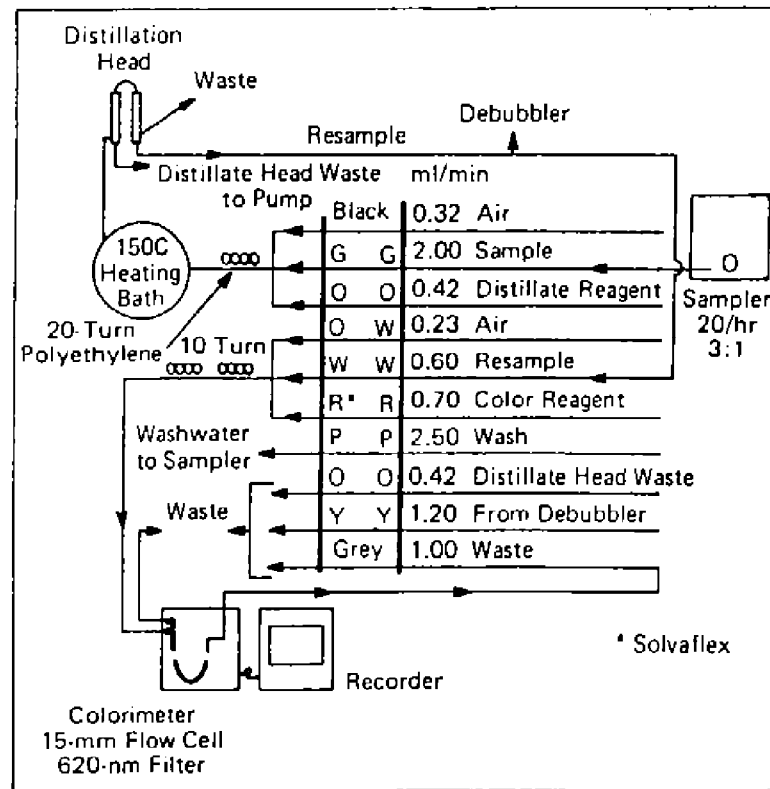
13.1 Physical and chemical characteristics of potable water. Appendix A defines the maximum and desired values of physical and chemical characteristics in separate tables.

14. AUTOMATED LABORATORY EQUIPMENT FOR MONITORING WATER AND WASTEWATER

Automated instrument are available and in use to monitor individual samples at rate of 10 to 60 samples/hr. The same instruments could be modified to make analysis for multiple constituents simultaneously from one sample. The read-out system includes sensing elements with indicators, alarms, and recorders.

Appropriate methodology could be supplied by the manufacturer for many of the common constituents of water and wastewater.

Monitoring of fluoride as a typical example could be totally automated. The basic of measurement is shown in Fig. 18. Uptaking of sample is automated and all valves which are indicated by different colors (B,O,Y,etc.) operated by computer in measuring time.



FLUORIDE MANIFOLD

FLUORIDE MANIFOLD

Fig. 18

15. MARKING

Each equipment shall be marked using stamp and cast figure or metal nameplate with letters not less than 8 mm in height.

- 1) Name of manufacturer or identifying symbol.
- 2) Distinctive catalogue designation.
- 3) Date of manufacture.

16. SHIPMENT

Shipment of all equipment should be under warranty and unpacking must be within this period (time of insurance for shipment).

It must be supplied and rapped in moisture proof material and also it shall bear in a clearly visible manner, with appropriate instruction, for storage.

17. SPARE PARTS

Packaging list should be with instrument which include, instrument + spare parts and two manual books. Spare parts must be supplied for at least two years.

18. PRE-INSTALLATION AND INSTALLATION

Pre-installation and installation must be done by manufacturer or representative of supplier and purchaser. Representatives should witness tests carried out for installation and commissioning of equipment.

19. WARRANTY

The instruments should be under warranty for at least one year after test run.

20. SERVICE AND INSPECTION

Instruments should be inspected by qualified persons assigned by the manufacturer or user according to letter of agreement.

APPENDICES

APPENDIX A

PHYSICAL CHARACTERISTIC OF POTABLE WATER

CHARACTERISTIC	DESIRED THRESHOLD LIMIT VALUE (T.L.V)	MAXIMUM VALUE
Color	5 units	5 units
Odor	2 "	3 "
Turbidity	5 "	25 "
pH	7-8.5	6.5-9.2
Chemical characteristic of potable water mg/L		
CHARACTERISTIC	DESIRED (T.L.V)	MAXIMUM VALUE
As	0	0.05
Cd	0	0.01
Cn	0	0.05
Pb	0	0.1
Hg	0	0.001
Se	0	0.01
Cr	0	0.05
Ba	0	1
Ag	0	0.05
B	0	1
Hardness	150	500
Ca	75	200
Mg	50	150
Mn	0.05	0.5
Fe	0.3	1
Zn	5	15
Cr	0.5	1.5
So ₄	200	400
Cl	200	600
N	0.002	0.05
Detergent	0.1	0.2
P	0.1	0.2
Total Dissolved Solid	500	1500