

# SP-910 Portable Water Analyzer Operation Manual

Rev.B



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## **Standard Limited Warranty**

Pyxis Lab warrants its products for defects in materials and workmanship. Pyxis Lab will, at its option, repair or replace instrument components that prove to be defective with new or remanufactured components (i.e., equivalent to new). The warranty set forth is exclusive and no other warranty, whether written or oral, is expressed or implied.

#### **Warranty Term**

The Pyxis warranty term is thirteen (13) months ex-works. In no event shall the standard limited warranty coverage extend beyond thirteen (13) months from original shipment date.

### **Warranty Service**

Damaged or dysfunctional instruments may be returned to Pyxis for repair or replacement. In some instances, replacement instruments may be available for short duration loan or lease.

Pyxis warrants that any labor services provided shall conform to the reasonable standards of technical competency and performance effective at the time of delivery. All service interventions are to be reviewed and authorized as correct and complete at the completion of the service by a customer representative or designate. Pyxis warrants these services for 30 days after the authorization and will correct any qualifying deficiency in labor provided that the labor service deficiency is exactly related to the originating event. No other remedy, other than the provision of labor services, may be applicable.

Repair components (parts and materials), but not consumables, provided in the course of a repair, or purchased individually, are warranted for 90 days ex-works for materials and workmanship. In no event will the incorporation of a warranted repair component into an instrument extend the whole instrument's warranty beyond its original term.

### Shipping

A Repair Authorization Number (RA) must be obtained from the Technical Support (service@pyxis-lab.com) before any product can be returned to the factory. Pyxis will pay freight charges to ship replacement or repaired products to the customer. The customer shall pay freight charges for returning products to Pyxis. Any product returned to the factory without an RA number will be returned to the customer.



## 1 General Description

### 1.1 Specification

Colorimeter Wavelength: 365/420/455/525/560/570/630 nm
 Turbidity Excitation Wavelength: White/infrared LED/90-degree scattering

Fluorescence Excitation Wavelength: 365/460 nm LED
 Fluorescence Emission Wavelength: 410/520 nm
 Wavelength Accuracy: ±1 nm

Absorbance Reproducibility:
 0.005 au (0 - 1.0 au) (3sigma)

Absorbance Linearity Range: 0 to 1.0 au

PTSA Reproducibility:
 1 ppb PTSA (3 sigma)

PTSA Detection Limit: 1 ppb
PTSA Range: 0 - 300 ppb

Fluorescein Reproducibility: 0.2 ppb or 2% of the value

Fluorescein Detection Limit: 0.1 ppb
 Fluorescein Range: 600 ppb
 Turbidity Reproducibility: 1 NTU (3 sigma)
 Turbidity Detection Limit: 1 NTU

Turbidity Detection Limit: 1 NTU
Turbidity Range: 0 - 200 NTU
Battery: 4 AA alkaline
Typical Battery Life: 3 months

Display: Graphical LCD 160x240 pixels, visible under direct sunlight

Instrument Dimension: L 265mm W 88mm H 62mm
 Instrument Weight: 600 g without batteries
 Storage Temperature Range: 0 to 140°F (-18 - 60°C)
 Operation Temperature Range: 40 to 120 °F (4 - 49°C)
 Humidity: 85% at 106 °F (41 °C)
 Environmental: IP67, dustproof and waterproof

### Note:

- 1. Specifications are subject to change without notice with Pyxis' continuous development.
- The fluorescein range in earlier versions of the SP-910 may be only up to 20 ppb. To extend the upper limit to 600 ppb, please contact Pyxis customer support at service@pyxis-lab.com.



### 1.2 Pyxis Major Features

The SP-910 analyzer shown in Figure 1 is a combination of photometer and fluorometer. It provides colorimetric measurements at 7 LED wavelengths, fluorometric measurement of fluorescent tracer PTSA and fluorescein, and nephelometric turbidity measurement using white LED and infrared LED as the excitation sources. The SP-910 is pre-calibrated for colorimetric measurements of analyses common in industrial water treatment and other water testing in the laboratory or in the field, such as chlorine, phosphate, iron, and copper. Main features include:

- The SP-910 is pre-calibrated for measuring PTSA (pyrenetetrasulfonic acid) in the range of 0 to 300 ppb. The fluorescence PTSA measurement is automatically compensated for sample color and turbidity interference.
- The SP-910 is pre-calibrated for measuring fluorescein in the range of 0 to 600 ppb.
- The SP-910 is pre-calibrated for measuring turbidity in the range of 0 to 200 NTU.
- Automatically select the primary wavelength according to the method selected and switches to the secondary wavelength to extend the primary measurement range.

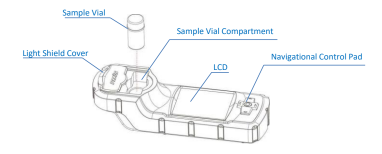


Figure 1. Sample Vial and Major Components

- Display a concentration-time profile curve during the last time period in a colorimetric
  measurement. The user can terminate the timing process and take a reading if the displayed
  concentration reaches a plateau before completing the predefined time period.
- The user can update the calibration parameter of any pre-calibrated colorimetric method by testing a standard solution first and then following a setup procedure to update the calibration parameters.
- Built-in Bluetooth allows easy connection to PC or mobile apps for downloading datalog and adding new colorimetric methods.

## 1.3 Unpackaging the Instrument

Remove the instrument and accessories from the shipping container and inspect each item for any damage that may have occurred during shipping. Verify that all items listed on the packing slip are included. If any items are missing or damaged, please contact Pyxis Customer Service at <a href="mailto:service@pyxis-lab.com">service@pyxis-lab.com</a>



#### 1.4 Standard Accessories

- Sample Vials two 10 ml (Part # MA-24), round, 0.78 inch (20 mm) pathlength, glass vials, which can be used for all measurements including turbidity and fluorescence measurements.
- 4 AA alkaline batteries
- Instrument Manual, also available from www.pyxis-lab.com
- 25 ml sample via (Part # MA-25)
- 16 mm tube adapter (Part # 52214)
- Bluetooth/USB Adapter for Desktop (Part # MA-NEB)

### 1.5 Optional Accessories

- 100 ppb PTSA standard in a 500 ml brown plastic bottle (Part # 21001)
- 50,250 and 500 ppb fluorescein standard in a 500 ml brown plastic bottle (Part #s FLUO50, FLUO250, FLUO500)

## 1.6 Sample Vial Compartment

The sample vial compartment is shown in Figure 1 along with a 10-ml sample vial. When the sample vial is inserted into the sample vial compartment, the triangular mark on the sample vial should be aligned approximately with the 6 o'clock position of the sample vial compartment or any position consistently.

The sample vial compartment can take in a 25 ml sample vial. The light shield cover is not required to be closed if the 25 ml sample via is used.

The 16 mm tube adapter is needed for colorimetric methods using the 16 mm sample tube. The instruction

to us the adapter is provided in section 8.

The sample vial compartment should be kept clean. A small amount foreign material could significantly affect turbidity and fluorescence measurement results. Use a soft cloth or lint free paper tissue to clean sample vial compartment periodically. Remove debris, scale, and deposit promptly.

## 1.7 Light Shield Cover

The light shield cover is shown in Figure 2. The light shield cover can be conveniently slid between the open and closed positions. The light shield cover is held firmly at the rest positions by permanent magnets.

The light shield cover should be in the closed position during storage, transportation, and measurements, especially during the turbidity and fluorescence measurements. When turned on, the SP-910 carries out self-diagnosis including checking the performance of a variety of optical devices. The light shield door shall be at the closed position to shield interference from ambient light during self-diagnosis.

Care should be taken to avoid water or debris being trapped in the track of the light shield door.

**Commented [LR1]:** 还有两个 25ml 的瓶子也是标配 MA-25 COD 适配器也是标配

- •25 ml sample via (Part # MA-25)
- •16 mm tube adapter (Part # 52214)

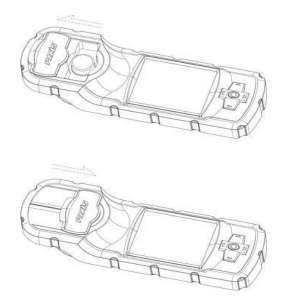


Figure 2 Open and Close the Light Shield Cover

## Warning

Magnetic sensitive devices, including but not limited to, credit cards, watches, hard disks, should be keep at a distance of at least 2 inches from the Light Shield Door to avoid possible damage and/or loss of information recorded.

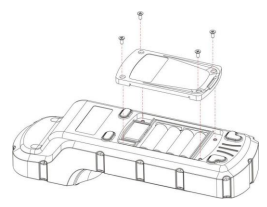
### 2 Start the SP-910

### 2.1. Battery Installation

The SP-910 is powered by four AA-size alkaline batteries. Do not use rechargeable nickel cadmium (NiCad) batteries or any AA-size lithium batteries. A set of batteries typically lasts for three months. When the batteries capacity is low, the SP-910 will prompt a LOW BATTERY warning. Replace all four batteries to resume operation of the SP-910 after the battery warning.

The SP-910 battery compartment, shown in Figure 3, is on the back side of the instrument. Insert a small pad underneath the screen area to make the back-surface level when the instrument is turned upside down. Install batteries as followings:

- 1. Remove the battery compartment cover by loosening four screws.
- 2. Insert four batteries into the battery holder as shown in Figure 3. Make sure the positive battery polarity marker (+) is aligned with the positive marker (+) on the battery holder.
- Replace the battery compartment cover, making sure that the sealing O-ring is lying flat on the battery holder and tighten the four screws.



**Figure 3 Replace Batteries** 

## 2.1 Description of the Navigational Control Pad

The SP-910 navigational control pad consists of five keys as shown in Figure 1. The left, right, up, and down keys are navigational keys that are used to select an icon, a button, or other items in various pages. The center key is the OK key. Press the OK key on a selected item to launch the action associated with the selected item. The OK key is also used to accept the current selection, like the return key in a computer keyboard.

### 2.2 Turning on the SP-910

After new batteries installation, the SP-910 will not be automatically turned on. To turn on SP-910, press the OK key, and release the OK key when the LCD is lit.



You can navigate the main page menu and launch an operation by pressing on an icon. If battery voltage is too low for the instrument to work properly, the SP-910 will show a low battery warning message when it is being turned on If this happens, replace all four batteries.

### 2.3 Main Page

The SP-910 provides intuitive icon driven user operations. On the main page, eight major feature groups are illustrated as below:

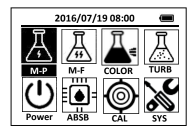


Figure 4. Main Menu

A brief description of each feature group is given in Table 1. Detailed operation instructions can be found in the following chapters.

No. Title Description M-P PTSA measurement 2 M-F Fluorescein measurement 3 **COLOR** Colorimetric measurement methods 4 **TURB** Turbidity measurement 5 Power Turn off SP-910

Absorbance measurements

Calibration routines

Table 1 Feature Groups on Main Menu

## 2.4 Turning off the SP-910

ABSB

CAL

SYS

6

7

8

Turn the SP-910 off by navigating to Power icon and press the OK key. Alternatively, you can turn off the SP-910 by pressing OK key for 5 seconds in any menu.

System and diagnosis information, Bluetooth enabling

### 2.5 The SP-910 Auto Power off

The SP-910 automatically turns itself off with no-key activity for a given period, except for during a measurement. The auto power-off time can be set in **SYS->System Set.** Pressing OK key will wake up the instrument, and the SP-910 will return to the original page if it has any measurement data.

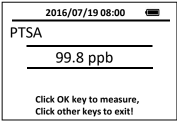
### 2.6 Auto LCD Power Saving

During a colorimetric method measurement, The SP-910 automatically turns LCD backlight off with no-key activity and continues the measurement with the LCD backlight off. The auto LCD power-off time can be set in SYS->System Set. Pressing any key will turn on the LCD backlight. Under normal ambient lighting condition, icons and other contents shown on the LCD screen are readable without backlight being on.

### 3 PTSA Measurement

#### 3.1 PTSA Measurement

- 1. Fill the 10 ml sample vial with the test solution and tightly cap the sample vial.
- Place the sample vial into the sample vial compartment and slide the light shield cover to the closed position.
- 3. Press the M-P on the main page, The SP-910 will start to measure the PTSA concentration
- 4. The SP-910 will display the PTSA concentration in ppb as PTSA.



## 3.2 Figure 5. PTSA Measurement

Deionized water (DI) as the blank calibration solution and the 100 ppb PTSA calibration s

During the fluorescence measurement to determine the PTSA concentration, the SP-910 checks the sample turbidity. If the sample turbidity value detected is greater than 40 NTU, The SP-910 will display a warning. For best results, the sample should be filtered if turbidity exceeds 40 NTU.

Sample color causes a lower PTSA concentration to be measured. The SP-910 automatically compensates for sample color. If the sample color is too intense, The SP-910 will display a warning.

For best results, ensure that the sample vial is clean. Wipe off water on the outside wall of the sample vial using a lint-free tissue paper. Fill the sample vial to the 10 ml mark. If the sample contains air bubbles, tap the sample vial gently to remove the bubbles before placing the sample vial to sample vial compartment.

### 3.3 PTSA calibration

tandard solution are needed.

 Press the CAL on the main page, then choose the M-P and press the OK key to launch the PTSA calibration page.



- Follow the message prompts, insert the DI blank into the sample vial compartment and press the OK kev.
- Follow the message prompts, use the upper and down key to switch between 100 ppb and 200 ppb standard
- 4. Fill the sample vial with the 100 ppb or 200 ppb standard and place the sample vial into the sample vial compartment and press the OK key to start calibration

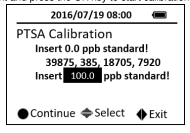


Figure 6. PTSA Calibration

If calibration fails, the followings should be checked:

- The DI blank is being contaminated.
- The 100 ppb PTSA standard solution is decayed or being contaminated.
- The light shield cover is not in the closing position.
- The sample vial compartment is blocked with debris, water, or other materials.

The 100 ppb standard solution shall be stored in a brown or black opaque bottle. Exposing the PTSA standard to light will cause the standard losing the PTSA concentration. Many substances, such as quaternary amine cause a negative interference. Many other substances such laundry detergents that contain optical brightener will cause a significant positive interference.

### 4 Fluorescein Measurement

### 4.1 Fluorescein Measurement

- 1. Fill the 10 ml sample vial with the test solution and tightly cap the sample vial.
- 2. Place the sample vial into the sample vial compartment and slide the light shield cover to the closed position.
- Press the M-F on the main page, then press the OK button, The SP-910 will start to measure the fluorescein concentration in the sample.
- 4. The SP-910 will display the fluorescein concentration in ppb.

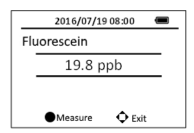


Figure 7. Fluorescein Measurement

For best results, ensure that the sample vial is clean. Wipe off water on the outside wall of the sample vial using a lint-free tissue paper. Fill the sample vial to the 10 ml mark. If the sample contains air bubbles, tap the sample vial gently to remove the bubbles before placing the sample vial to sample vial compartment.

### 4.2 Fluorescein calibration (Firmware version before v1.0r295)

Deionized water (DI) as the blank calibration solution and the 20 ppb fluorescein calibration standard solution are needed.

- 1. Press the CAL on the main page, then choose the **Fluorescein** and press the OK key to launch the fluorescein calibration page.
- Follow the message prompts, insert the DI blank into the sample vial compartment and press the OK key.
- 3. Follow the message prompts and insert the 20 ppb standard into the sample vial compartment and press the OK key.
- 4. Press the OK key to return to the main page.



Figure 8. Fluorescein Calibration

If calibration fails, the followings should be checked:

- The DI blank is being contaminated.
- The 20 ppb fluorescein standard solution is decayed or being contaminated.
- The light shield cover is not in the closing position.
- The sample vial compartment is blocked with debris, water, or other materials.

### 4.3 Fluorescein calibration (Firmware version v1.0r295 and after)

Deionized water (DI) as the blank calibration solution, the 50 ppb fluorescein, the 250 ppb fluorescein and the 500 fluorescein calibration standard solutions are needed.

- Press the CAL on the main page, then choose Fluorescein and press the OK key to launch the fluorescein calibration page.
- Follow the message prompts, insert the DI blank into the sample vial compartment and press the OK key.
- 3. Insert the 50 ppb standard into the sample vial compartment and press the OK key to complete the low range calibration.
- 4. Press the **OK** key to proceed with middle range calibration or press any other keys to return to main page.
- Insert the 250 ppb standard into the sample vial compartment and press the OK key to complete the middle range calibration.
- 6. Press the **OK** key to start proceed with high range calibration or press any other keys to return to main page.
- 7. Insert the 500 ppb standard into the sample vial compartment and press the OK key to complete the high range calibration.

The middle range and high range calibrations from steps 4 to 8 are optional if only low range fluorescein measurement is intended.

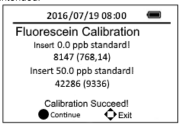


Figure 9. Low Range Fluorescein Calibration



Figure 10. Middle Range Fluorescein Calibration

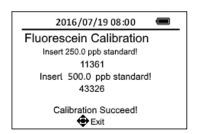


Figure 11. High Range Fluorescein Calibration

The standard solutions shall be stored in a brown or black opaque bottle. Exposing the fluorescein standard to light will cause the standard losing the fluorescein concentration. Many substances, such as quaternary amine cause a negative interference. Many other substances such laundry detergents that contain optical brightener will cause a significant positive interference.

## 5 Colorimetric Measurement

## 5.1 Supported Methods

A wide range of colorimetric methods is supported by the SP-910 analyzer and the number of them keeps increasing with continuous development of Pyxis. See corresponding Hach® methods in Appendix A.

Table 2 List of Supported Colorimetric Methods

Abbreviated Method Name	Method Name	Description	Range
AL	Alumi	Aluminum,Aluminon method	0.8 ppm
ALKLR	Alkalinity	Alkalinity, Total, Low Range	100 ppm as CaCO3
ALKHR	Alkalinity	Alkalinity, Total, High Range	500 ppm as CaCO3
AZOL	Azole	UV digestion for tolyltriazole and benzotriazole	16 ppm
BLCH	Bleach	Direct method measuring sodium hypochlorite concentration	16%
BLCHL	Bleach	Direct method measuring sodium hypochlorite concentration, Low Range	1.50%
Br-T	Bromine	Bromine,DPD method for	4.5 ppm
Ca	Ca	Calcium ,Calmagite method	4 ppm as CaCO3
CaHR	Ca	Calcium hardness ,Murexide method	500 ppm as CaCO3
CaMgL	CaMg	Total hardness, Chlorophosphonazo colorimetric method, Ultra-Low Range	1 ppm as CaCO3
CODLR	COD	Oxygen Demand, Chemical (Reactor Digestion 20 Minutes Method) ,Low range	150 ppm



Abbreviated Method Name	Method Name	Description	Range
CODHR	COD	Oxygen Demand, Chemical (Reactor Digestion 20 Minutes Method), High range	1500 ppm
CODUH	COD	Oxygen Demand, Chemical (Reactor Digestion 20 Minutes Method) ,Ultra-High Range	15000 ppm
CODUL	COD	Oxygen Demand, Chemical (Reactor Digestion 20 Minutes Method) ,Ultra-Low Range	40 ppm
CLLR	CLLR	Turbidimetric method,Low Range	40 ppm
CLMR	CLMR	Turbidimetric method ,Medium Range	400 ppm
CL2HR	CL2High	Total Chlorine, DPD method , High Range	10 ppm
CL2HR	CL2High	Free Chlorine, DPD method , High Range	10 ppm
CL2UH	CL2UH	Free Chlorine, Iodimetr method ,Ultra-High Range	400 ppm
CL-F	F-Chlorine	Free chlorine, DPD method	2.2 ppm
CLTMB	Chlorine, Free	Free chlorine ,TMB method	1.2 ppm
CIO2	CIO2-DPD	DPD method, USEPA accepted for reporting drinking water analysis	5 ppm
CIO2D	CIO2Direct	Direct method for chlorine dioxide,Medium Range	45 ppm
CLO2H	CIO2Direct	Direct method for chlorine dioxid, High Range	1500 ppm
CL-T	T-Chlorine	Total chlorine, DPD method	2.2 ppm
CN	Cyanide	Cyanide, Pyridine-Pyrazalone method	0.24 ppm
COLOR	Color	Color, APHA Platinum-Cobalt Standard Method	500 units
Cr6	Cr6	chromium hexavalent,1,5- Diphenylcarbohydrazide method , USEPA accepted for wastewater analyses	0.6 ppm
CrT	CrTot	Total Chromium ,Alkaline hypobromite Oxidation method	0.6 ppm
CuBi	Cu_Bicinch	Bicinchoninate method, EPA approved for reporting wastewater analysis	5 ppm
CuLR	CuPorp	Copper,Porphyrin method	0.21 ppm
CYAN	CYAN	Cyanuric acid,Turbidimetric method	55 ppm
CYN-F	Cyclohexylamine	Cyclohexylamine,Fluorescent method	1.2 ppm
DEHA	DEHA	Iron Reduction method for N,N- diethylhydroxylamine and other oxygen scavengers	0.5 ppm
DO	DO	Dissolved Oxygen,lodimetry method	10 ppm
F	Floride	Fluoride, SPADNS method	2 ppm
FeMo	FeMo	Total iron method for water containing molybdate	1.8 ppm
FePh	Fe_phenanth	Total iron using 1,10-phenanthroline, USEPA approved for reporting wasterwater analysis	3 ppm
FeSal	Fe-Sal	Total Iron using 5-Sulfosalicylic Acid Dihydrate	5 ppm
FeTp	FeTptz	Total iron using TPTZ	1.8 ppm
FeZi	FeZine	Total iron , FerroZine method	1.3 ppm

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Abbreviated Method Name	Method Name	Description	Range
H2O2	H2O2	Hydrogen peroxide,lodimetry method	500 ppm
H2O2L	H2O2L	Hydrogen peroxide, DPD method,Low Range	1.5 ppm
Mg	Mg	Magnesium,Calmagite method	4 ppm as CaCO3
MnHR	MnHigh	Manganese,Periodate Oxidation method,High range	20 ppm
MnLR	MnLow	Manganese,Periodate Oxidation method,Low range	0.7 ppm
MoHR	Mo_HighRange	Molybdate, Mercaptoacetic Acid method, High range	40 ppm
MoLR	Mo_LowRange	Molybdate, ternary complex method,Low range	3 ppm
N2H4	N2H4	Hydrazine,p-Dimethylaminobenzaldehyde method	0.5 ppm
NH2C	NH2CL	Chloramine mono ,Indophenol method	3 ppm
N-TLR	N-TLR	Nitrogen, Total (Test 'N Tube Method) ,Low range	25 ppm as N
N-THR	N-TLR	Nitrogen, Total (Test 'N Tube Method) ,High range	150 ppm as N
NH3S	NH3Sal	nitrogen, ammonia ,Salicylate method	0.5 ppm as N
NH3-F	NH3-F	Ammonia Nitrogen ,Fluorescent Method	0.07 ppm as N
NH3LR	NH3LR	Nitrogen, Ammonia (Test 'N Tube) - Low range	2.5 ppm as N
NH3HR	NH3HR	Nitrogen, Ammonia (Test 'N Tube) -High range	50 ppm as N
Ni	Ni	Nickel,PAN method	1.2 ppm
NO2D	NO2D	Direct method for nitrite	1000 as NO2
NO2H	NO2H	High range nitrite, ferrous sulfate method	150 as NO2
NO2L	NO2L	Low range nitrite, diazotization method, USEPA approved for reporting wastewater and drinking water analysis	0.35ppm as NO2
NO3HR	NO3H	Nitrate ,Cadmium Reduction method,High range	30 ppm as N
NO3MR	NO3M	Nitrate ,Cadmium Reduction method,Middle range	5 ppm as N
NO3CA	NO3CA	Nitrate, High Range (Test 'N Tube Method)	30 ppm as N
O3	O3	Ozone ,DPD method	2 ppm
PAA	PAA	Peroxyacetic , lodimetry method	500 ppm
OPO4	OPO4	Reactive phosphate using ascorbic acid molybdenum blue method, USEPA accepted for wastewater analysis	2.5 as PO4
OrgP	Phosphonate	UV digestion and ascorbic acid reduction molybdenum blue method	2.5 as PO4
PAmi	OPO4-Amino	Reactive phosphate, amino acid reduction method	30 as PO4
P-TLR	P-TLR	Phosphorus, Total (Test 'N Tube Method) - Low range	3.5 as PO4
P-THR	P-THR	Phosphorus, Total (Test 'N Tube Method) - High range	100 as PO4

Abbreviated Method Name	Method Name	Description	Range
pН	pН	Phenol red method for pH	8.5
PMoV	OPO4-MoV	Reactive phosphate , Molybdovanadate method	45 as PO4
POLY	Polymer	Anionic polymeric dispersant, Turbidimetric method	14 ppm
Sb3+	Sb3+	Antimony Trivalent ,PADAP Method	0.11 ppm
Sb-T	Sb-T	Antimony, Total ,PADAP Method	0.11 ppm
S2-	Sulfide	Methylene blue method for sulfide, USEPA accepted for reporting wastewater analysis	0.7 ppm
SiHR	SiHR	Silica, Silicomolybdate method, High Range	75 as SiO2
SiLR	SiLR	Silica, Heteropoly Blue method, Low Range	5 as SiO2
SO3LR	SO3LR	Sulfite,OPA method ,Low Range	5 ppm
SO3HR	SO3HR	Sulfite,OPA method , High Range	50 ppm
SO4	SO4	Barium sulfate, Turbidimetric method	70 ppm
TOC	TOC	Total Organic Carbon	20 ppm
Urea	Urea	Urea (Reactor Digestion Method)	10 ppm
ZnXO	ZnXO	Zinc ,Xylenol orange method	3 ppm
Zn	Zinc	Zincon method for zinc, USEPA approved for wastewater analysis	3 ppm

### 5.2 Select a Method

Move the icon focus to the method icon **COLOR** using the navigational (left, right, up, or down) keys. Press OK on the icon to launch the first method selection page. The methods shown on the top row of the page are the most frequently selected methods.

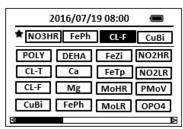


Figure 12. Method Selection

The followings are the operations associated with this page:

- 1. Use the navigational keys and the OK key to select and launch a method.
- Long press the OK key to return to the main page. Press the arrow icon at the lower right corner of the page to display the second method selection page if the device is loaded with more than 23 methods.

Note: Methods shown in the method selection pages include Hach® equivalent methods and Pyxis specific advanced methods. The table in Appendix A provides a brief description of Pyxis method names



and their corresponding Hach® program number. Hach® reagents for 10 ml sample can be used for the test.

### 5.3 Single Timing Step Method

Most of colorimetric methods have only one timing step. As an example, in the DPD free chlorine method, it takes one minute for the DPD powder reagent to completely react with chlorine in the water sample. The DPD free chlorine method has a single one-minute timing step. Figure 13 shows the main page of a method with a single timing step.

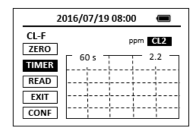


Figure 13. Single Timer Method

### 5.4 Single-Vial Procedure

- 1. Place the sample vial filled with the water sample in the SP-910 sample vial compartment and press the **ZERO** button. The SP-910will display the page shown in Figure 13.
- 2. Take the sample vial out and add the reagent to the sample vial.
- 3. Place sample vial back into the sample vial compartment and press the timer button TMR1. The SP-910 will start to monitor the reaction between the reagent and the species you want to measure in the water sample. The concentration is shown in the chart as a function of time (Figure 14).
- 4. When the timer reaches the preset time and the reaction is complete, the value of concentration will be shown on the top right corner of the page.
- 5. The rate of the reaction is often faster than the standard pre-set time, which will become apparent from the concentration-time plot. You can press the STOP button to stop the timer and terminate the timing step. The last read concentration value will be displayed on the top right corner of the page after you terminate the timing step.

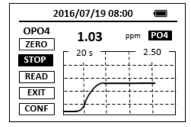


Figure 14. Concentration as a Function of Time

### 5.5 Two-Vial Methods



Some colorimetric methods require using two vials. The water sample is added to two identical vials. One vial is being used to zero the colorimeter, referred as to the prepared blank. A reagent is added to the other vial, referred as to the prepared sample. The absorbance value is determined from the prepared sample.

If the method requires two or more reagents, the prepared blank could be the resulting solution after one or more reagents have been added to the sample.

The following procedure is typical for two-vial methods:

- Place the prepared blank into the SP-910 sample vial compartment and press the ZERO button to zero the instrument.
- Place the prepared sample into the SP-910 sample vial compartment and press the TMR1 button to start the method timer.
- When the timing step is completed, the measured concentration will be displayed on the top of the page. The timing step could be terminated earlier by pressing STOP button.
- 4. Optionally, the SP-910 can be re-zeroed using the prepared blank after the timing step is completed or terminated. The blank reading will be subtracted from the measured concentration value, and the displayed concentration value on the top-right corner will be updated. This step is optional. It is only necessary if the prepared blank changes its color during the timing period.
- Optionally, the prepared sample vial can be put back and read again by pressing the READ button if the blank is re-zeroed after the timing step is completed or terminated. A new concentration value based on the last absorbance value measured will be calculated and displayed.

### 5.6 Multiple Timing Steps Method

Some colorimetric methods have two or three timing steps. The SP-910 shows a count-down timer for the timing steps before the last timing step (Figure 15). During these timing steps, one or more reagents are added to the sample, or operations such as swirling the vial to mix the reagent and the sample are being performed. These methods usually use one vial for the prepared blank and the other for the prepared sample.

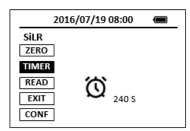


Figure 15. Multiple Timer Method

In order to show the concentration-time curve as shown in Figure 14 during the last timing step, The SP-910 must be zeroed using the prepared blank before the last timing step. Thus, the last timer button will not be selectable until the SP-910 has been zeroed using the prepared blank. Multi-timing step Hach® methods require zeroing the colorimeter using the prepared blank after the last timing step is completed. The SP-910 can optionally be re-zeroed using the prepared blank after the last timing step. The blank



value measured will be subtracted from the concentration value measured at the end of the last timing step. Optionally, the **READ** button could be pressed to read the prepare sample again.

The following procedure is typical for methods having two-timing steps:

- Press the TMR1 button to start the first timer. Complete the necessary operations to prepare the blank and the sample.
- 2. Place the prepared blank into the SP-910 sample vial compartment and press the **ZERO** button
- 3. Place the prepared sample into the SP-910 sample vial compartment and press the **TMR2** button to start the second timer. The SP-910 will display the measured concentration as a function of time as shown in Figure 14.
- When the timing step is completed, the measured concentration will be displayed on the top right of the screen. The timing step could be terminated earlier by pressing STOP button.
- 5. Optionally, The SP-910can be re-zeroed using the prepared blank after the timing step is completed or terminated. The blank reading will be subtracted from the measured concentration value, and the displayed concentration value on the top-right corner will be updated. This step is optional. It is only necessary if the prepared blank changes its color during the timing period.

### 5.7 Advanced Methods

The SP-910 provides 7 LED wavelengths and can measure absorbance values at multiple LED wavelengths. Consequently, the SP-910 can provide many predefined advanced methods that traditionally require complex and often expensive lab testing procedures.

### 5.7.1 Low range, direct reading chlorine dioxide, 0 to 35.0 ppm

The maximum absorption bank of aqueous chlorine dioxide is around 360nm. The SP-910 has a 365nm UV LED and can be used to directly measure chlorine dioxide. It offers a much lower detection limit (0.2 ppm) than direct methods available from other portable colorimeters having only light sources in the visible range.

Select CIO2D in the method selection page and carry out the following steps to measure chlorine dioxide:

- Place a vial filled with deionized water into the vial compartment and press the ZERO button to zero the SP-910.
- Discard the deionized water and fill the same vial with the sample. Place the vial into the vial compartment and press READ button to read. The measured chlorine dioxide concentration will be displayed in the top of the method page.

### 5.7.2 Turbidimetric Anionic Polymer Method

- 1. Add polymer reagent 1 to 10 ml sample and inverse the sample vial 5 times to mix the reagent with the sample. Place the sample via to the sample vial compartment.
- 2. Press on ZERO.
- 3. Add polymer reagent 2 and press on TMR1 to start the five minutes timer.



- Gently inverse the sample via for 10 times and place the sample vial to the sample vial compartment.
- Polymer concentration will be measured and displayed when the five-minute timer is reached.The polymer concentration is shown as ppm PAA (polyacrylic acid) equivalent.

### 5.7.3 Direct Reading Bleach Percent Method, 0 to 15%

The SP-910 has a 365nm UV LED and other deep blue LEDs that can be used to directly measure bleach concentration in the range of 0 to 15%. No reagent is required for the method and the displayed result is the sodium hypochlorite concentration in percentage.

Select **BLCH** in the method selection page and carry out the following steps:

- Press the OK key to enter the temperature input interface. Enter the temperature of the sample.
- 2. Place a vial filled with deionized water into the vial compartment and press the **ZERO** button to zero the SP-910.
- Discard the deionized water and fill the same vial with the bleach sample. Place the vial into the vial compartment and press READ button to read. The measured bleach concentration will be displayed in the top of the method page.

### 5.8 Method Setup and Calibration

Press the SETUP button in the method result page to launch the method setup and calibration page.

### 5.8.1 Set up the method parameters

Press the **FORM** button to select a concentration form from the list of forms that are available for this specific method (Figure 16).

Press the **UNIT** button to select a concentration unit among the list of ppb, ppm, mg/L, ug/L and No Unit (Figure 17).

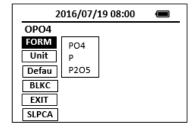


Figure 16. Method from Selection

Commented [LR3]: 目前有温度补偿关于漂水测试,要更新这部分内容

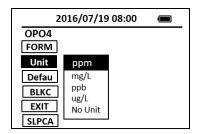


Figure 17. Method Unit Selection

### 5.8.2 Slope Calibration

If the method has been calibrated prior to shipping, there is no need to calibrate unless a calibration check indicates that the method needs a calibration. The following steps are used to calibrate a method:

- Use a calibration standard of known concentration. Follow the steps required by the method and note the value reported by the SP-910.
- If the measured value differs from the known standard value, Press the CONFG button to launch the method configuration page.
- 3. Press the slope calibration button **SlpCal**. A numeric keyboard will be displayed.
- 4. Enter the concentration value and press the OK key on the enter key in the numeric keyboard to return to the configuration page.
- Press the EXIT button. Press the OK key to accept the calibration or other key to cancel the calibration.

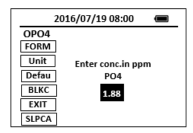


Figure 18. Slope Calibration

For best results, the concentration of the standard solution should be less than the maximum concentration for the method (Table 2) and greater than the half of the maximum concentration. For example, to calibrate total chlorine, the chlorine concentration in the standard solution should be between 1.1 and 2.2 ppm.

The corresponding calibration parameters will be updated and saved in the memory as the working calibration parameter set. Note that this set of calibration parameters are not the same as the default set. You can use **Default** button to copy the default calibration parameters to the working set.

### 5.8.3 LowC Calibration

Commented [LR4]: 目前已修改为 LowC 低点校准



Some methods have a non-zero intercept value in the calibration equation. For these methods, a proper non-zero intercept value is pre-loaded in the SP-910 prior to shipping. The following steps are used to carry out a reagent blank calibration:

- 1. Follow the normal steps to carry out a measurement on a deionized water sample.
- 2. Press the CONFG button to launch the method configuration page.
- 3. Press the LowC calibration button LowC
- 4. Press the OK key to save when exiting from the configuration page or press other keys to cancel.

### 5.8.4 Resume to Default Calibration Parameters

Pressing the **Default** button will copy the default calibration intercept and slope to the working intercept and slope, respectively. If the default calibration parameters were created prior to shipping, this button action is to restore the working calibration parameters to the original factory loaded calibration parameters.

## 6 Turbidity Measurement

### 6.1 Operation

Follow the following steps to measure turbidity:

- 1. Fill the 10 ml sample vial to above the 10 ml mark.
- 2. Insert the sample vial to the sample vial compartment.
- 3. Slide the light shield cover to the closed position.
- Press the TURB on the main page, then press the OK key, The SP-910 will start to measure the turbidity in the sample.

### 6.2 Turbidity Calibration

- 1. Fill the 10 ml sample vial to above 10 ml mark with the deionized water.
- 2. Insert the sample vial to the sample vial compartment.
- 3. Slide the light shield cover to the closed position.
- Press the CAL on the main page, then choose the Turbidity calibration and press the OK button to launch the Turbidity calibration page. (Figure 19)
- 5. Press the **OK** key to measure the deionized water
- Fill the 10 ml sample vial to above 10 ml mark with the 50 NTU standard. Insert the sample vial to the sample vial compartment.
- 7. Press the **OK** key to measure the 50 NTU standard. Low range turbidity calibration is successful
- 8. Press the **OK** key to continue high range turbidity calibration. If high range turbidity calibration not required, press any keys to exit. (Figure 20)
- Fill the 10 ml sample vial to above 10 ml mark with the 100 or 200 NTU standard. Insert the sample vial to the sample vial compartment.



- Follow the message prompts, use the upper or down key to switch the standard between 100 NTU and 200 NTU.
- 11. Press the **OK** key to measure the selected standard. High range turbidity calibration is successful. (Figure 21)
- 12. Press any keys to exit.

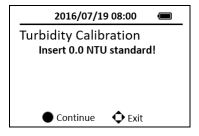


Figure 19. Turbidity Calibration-1

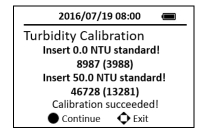


Figure 20. Turbidity Calibration-2

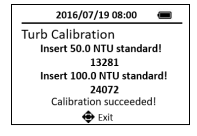


Figure 21. High Range Turbidity Calibration

## 7 Absorbance Measurement

The following steps are used to measure the absorbance values of a sample:

1. Press the ABS to launch the absorbance measurement page.



- Place a vial filled with the blank sample in the sample vial compartment. Press the ZERO button to zero the method.
- 3. Place a vial filled with the sample in the sample vial compartment. Press the READ button to read absorbance. The absorbance values of first 6 wavelengths (Table 3) will be shown. Press the READ button again to show the absorbance values of the last three wavelengths.

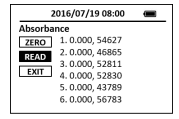


Figure 22. Absorbance Measurement

Press **EXIT** to return to the main page. Timing function for absorbance measurement may not be available for some models.

Table 3 Wavelength of each channel

Channel	Wavelength (nm)
1	560
2	570
3	Not used
4	Not used
5	455
6	525
7	365
8	630
9	420

Note that the absorbance values measured with the SP-910 is generally smaller than those measured with a spectrophotometer equipped with a monochromatic light source or detector. The SP-910 absorbance values should, however, linearly correlate with the absorbance values measured with the spectrophotometer. Thus, for any colorimetric system, The SP-910 absorbance follows Lambert-Beer law.

## 8 Bluetooth Interface

The SP-910 equipped with a Bluetooth interface, which allows a user to connect to SP-910 with a computer or a mobile device to do the following tasks:

- Configure device
- Add user defined colorimetric methods



- Upgrade device firmware
- Download saved datalog

With the Bluetooth interface, the user can calibrate an inline fluorometer directly from SP-910 in the field. Below sections describe how to connect and communicate with your SP-910 via a computer and uPyxis apps.

### 8.1 Install Software

Download uPyxis software from www.pyxis-lab-lab/supports/, unzip and install uPyxis, The Bluetooth adapter driver will be installed as well. Plugin the Bluetooth adapter shipped along with your SP-910 device, open uPyxis app.

## 8.2 Turn on SP-910 Bluetooth

The SP-910 Bluetooth function is normally switched off in order to reduce power consumption, to turn on Bluetooth, select SYS in the main menu and click BTLE in the SYS screen.

## 8.3 Connect uPyxis to SP-910

Click **Device** tap on the top right of uPyxis app. Select UBS-Bluetooth in the dropdown menu. uPyxis will scan nearby Pyxis Bluetooth devices including SP-910 units. Click the discovered SP-910 to connect.If the SP-910 is being automatically powered off during connection, please push the OK key to power on the SP-910 again. The SP-910 will automatically turn on the Bluetooth and repeat the connection steps again in uPyxis.



Figure 23. uPyxis Scans Bluetooth Devices

### 8.3.1 Upgrade Firmware



When connected, click **System** tap to view the device information. The user can upgrade the SP-910 firmware to the latest. The latest version of the SP-910 firmware can be obtained from service@pyxis-lab.com



Figure 24. System Information

### 8.3.2 Setup Product

The SP-910 will display PTSA in ppb unit as the default in the PTSA measurement. A product name can be assigned to allow the SP-910 to display the product name instead of PTSA. The product/PTSA ratio can be set up according to Figure 25. For products containing 0.1% PTSA, the ratio is 1000, which means that 100 ppb PTSA equals to 100 ppm product. For products containing 0.2% PTSA, the ratio is 2000, which means that 200 ppb PTSA equals to 100 ppm product.



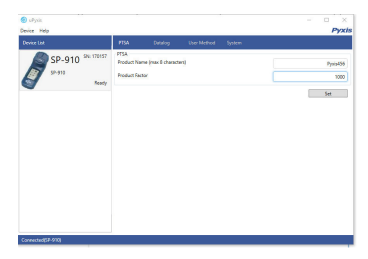


Figure 25. Setup Product

### 8.3.3 Add User Defined Colorimetric Methods

Click **User Method** tap and click **Read from Device** to read all current default or user defined methods into uPyxis. Select a method and clone a new method from the selected. The parameters in the method can be edited and are defined as:

No.: a sequence number, the user does not need to enter

Name: the method name

**LED:** the LED wavelength in nm and can be selected from the dropdown menu.

**T1:** the first reaction period of the method

 ${\bf T2:}$  the second period of the method

T3: the third period of the method

DP0, DP1, and DP2: calibration coefficients as shown in the following equation, where A is absorbance

 $ppm = DP0*A^2 + DP1*A + DP2.$ 

Max: the maximum range of the method.

Form1: the default display form of the method, such as PO4 for a phosphate method.

Factor1: it is always 1 If the method has just one display formp.

FormId: default is 0 and the user does not need to change it.



**UnitId:** default is 0 and the user does not need to change it.

**DecNum:** the number of decimal places in the displayed concentration

After a user defined method is created, click Write to Device to save the method.

### 8.3.4. Download Datalog

Click **Datalog** tap to open the datalog page. Click **Read Datalog List** to retrieve the datalog list from the SP-910 (Figure 28). Select a datalog entry from the list and then click **Read Datalog** to load the measurement values to uPyxis (Figure 29). The datalog can be saved as a CSV file.

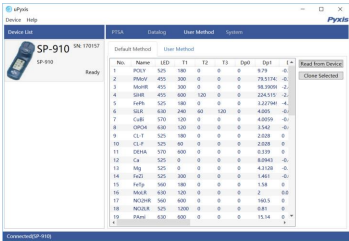


Figure 26. Load the default Methods

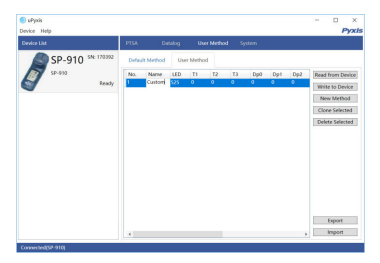


Figure 27. Add a User Defined Method



Figure 28. Datalog List

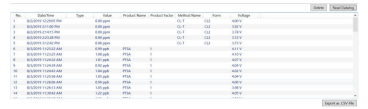


Figure 29. Detailed Measurement Data

## 9 Calibrate a ST-500 with SP-910

The SP-910 can be used to verify the result of inline Pyxis ST-500 and other probes by measuring the sample water took from the inline probe sample line. The SP-910 can then be used to calibrate the inline probes over the Bluetooth connection.



Choose the **CAL** in main menu and select **Inline Device**, the following interface then appears in the screen. SP-910 starts to scan devices via Bluetooth interface.

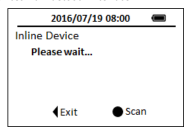


Figure 30. Scan Inline Device

Active inline probes will be listed in the following screen, use **Up** and **Down** key to select the device you want to pair with, click **OK** key to connect.

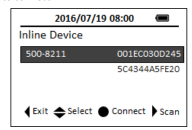


Figure 31. Pair Inline Device

Once the connection is established, the SP-910 will read the latest reading from the connected ST-500 and display the reading as shown in Figure 32.

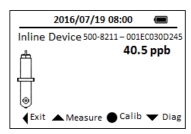


Figure 32. Read Inline Device

Use the SP-910 to measure the sample water by clicking **Up** key, as shown in Figure 33

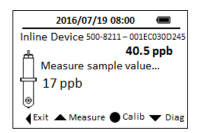


Figure 33. SP-910 Measures Sample Water

Click **OK** key to send the calibration instruction to the ST-500 via Bluetooth connection. After that, the connected ST-500 will be calibrated to the value measured by the SP-910. The SP-910 will keep reading ST-500 every 4 seconds to verify if the calibration is successful. Please note that it takes about a minute for the ST-500 to approach to the calibrated reading.

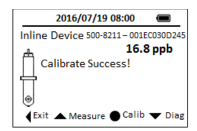


Figure 34. Calibration Success

Click **Down** key to start diagnose ST-500 probe. As in Figure 35, a range of ST-500 operation parameters will be displayed. Furthermore, click **OK** button in diagnosis page to check whether ST-500 is fouled.

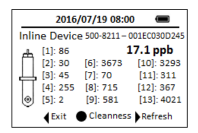


Figure 35. Inline Device Diagnosis Data

In the cleanliness page, please put the ST-500 probe into DI water and then click the cleanliness button again to conduct cleanness check.

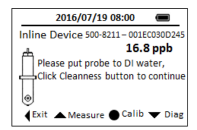


Figure 36. Cleanliness Check

Figure 37 shows a probe may be fouled according to its diagnosis operational parameters.

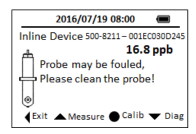


Figure 37. Probe is Fouled

## 10 Maintenance

Use a soft cloth or lint free paper tissue to clean the sample vial compartment periodically. Remove debris, scale, and deposit promptly.

Although The SP-910 is protected from water damage, it is a good practice to avoid water entering the sample vial compartment and becoming trapping underneath the navigational control pad. Deposits left behind when the water is evaporated could affect Pyxis performance.

The SP-910 should be stored in the temperature range of 0 to 140°F (-18 to 60°C) and relative humidity less than 85% at 106 °F (41 °C). Do not leave the SP-910 in a parked vehicle. The temperature inside a parked vehicle can reach above 150 °F in summer and -20 °F in winter. Exposing the SP-910 to extreme temperature or humidity will cause a gradual decay in performance of fluorescence measurements and require more frequent calibrations.

During storage and transportation, do not leave a sample vial in the sample vial compartment. Close the lid of the sample vial compartment during storage and transportation.

Replace batteries when the SP-910 displays a warning message indicating LOW BATTERY voltage. Remove batteries from the SP-910 battery compartment if the SP-910 is going to be placed in storage for a long period time.



When the SP-910 is shipped, a desiccant pack is included in the desiccant compartment underneath the cover of the battery compartment. It is recommended that a new desiccant pack is replaced each time the batteries are replaced.

## 11 Troubleshooting

The SP-910 will prompt a warning message if it detects an abnormal condition or operation. On screen prompts direct the user to take appropriate corrective actions in most cases.

If an unspecific error occurs or the SP-910 cannot be turned on, reboot the instrument by taking a battery out of the battery holder and re-install the battery.

If the SP-910 has been idle for more than two months and cannot be turned on, replace all four batteries with four new AA alkaline batteries.

A diagnostics page can be launched by press the **SYS** icon in the main page. The software version and its associated hash code can be found in the diagnosis page. Contact Pyxis professionals at <a href="mailto:service@pyxis-lab.com">service@pyxis-lab.com</a> and provide with following information to ensure high quality technical support.

**Table 4 Contact Information** 

Items	Note
Contact Name	
Phone	
Email	
Customer Name	
Product Number (P/N)	Can be found on the product label on back of product
Serial Number (S/N)	Can be found on the product label on back of product
Firmware version	Can be found in diagnosis page
Problem Description	Capture warning message if applicable

## 12 Appendix A.

## Pyxis Method and Hach® Method Number (PRMP) Cross Reference

Abbreviated Method Name	Method Name	Corresponding Hach © method	Hach Method Number
AL	Alumi	Aluminon Method for Aluminum	8012
ALKLR	Alkalinity	Alkalinity, Total, Low Range	N/A
ALKHR	Alkalinity	Alkalinity, Total, High Range	N/A
AZOL	Azole	Benzotriazole, UV Photolysis	8079
BLCH	Bleach	Direct method measuring sodium hypochlorite concentration	N/A
BLCHL	Bleach	Direct method measuring sodium hypochlorite concentration, Low Range	N/A
Br-T	Bromine	Bromine, DPD Method	8016
Ca	Ca	Calcium: Calmagite Colorimetric Method	8030
CaHR	Ca	Calcium hardness ,Murexide method	N/A
CaMgL	CaMg	Total hardness, Chlorophosphonazo colorimetric method , Ultra-Low Range	8374
CODLR	COD	Oxygen Demand, Chemical (Reactor Digestion 20 Minutes Method) ,Low range	10259
CODHR	COD	Oxygen Demand, Chemical (Reactor Digestion 20 Minutes Method) , High range	10259
CODUH	COD	Oxygen Demand, Chemical (Reactor Digestion 20 Minutes Method) ,Ultra-High Range	8000
CODUL	COD	Oxygen Demand, Chemical (Reactor Digestion 20 Minutes Method) ,Ultra-Low Range	8000
CLLR	CLLR	Turbidimetric method,Low Range	N/A
CLMR	CLMR	Turbidimetric method ,Medium Range	N/A
CL2HR	CL2High	High Range DPD Chlorine	10070
CL2UH	CL2UH	Free Chlorine, Iodimetr method ,Ultra-High Range	N/A
CL-F	F-Chlorine	Chlorine, Free, DPD	8021
CLTMB	Chlorine, Free	Free chlorine ,TMB method	N/A
CIO2	CIO2-DPD	Chlorine Dioxide, DPD	10126
CIO2D	CIO2Direct	Chlorine Dioxide, Direct Reading	8345
CLO2H	ClO2Direct	Chlorine Dioxide, Direct Reading, High Range	N/A
CL-T	T-Chlorine	Chlorine, Total, DPD	8167
CN	Cyanide	Pyridine-Pyrazalone Method for Cyanide	8027
COLOR	Color	Color, APHA Platinum-Cobalt Standard Method	8025
Cr6	Cr6	Hexavalent chromium, 1,5- Diphenylcarbohydrzaide Method	8023
CrT	CrTot	Chromium total Alkaline Hypobromite Oxidation Method,	8024



Abbreviated Method Name	Method Name	Corresponding Hach © method	Hach Method Number
CuBi	Cu_Bicinch	Copper, Bicinchoninate	8506
CuLR	CuPorp	Copper,Porphyrin method	8143
CYAN	CYAN	Turbidimetric method for cyanuric acid	8139
CYN-F	Cyclohexylamine	Cyclohexylamine,Fluorescent method	N/A
DEHA	DEHA	DEHA, Iron Reduction Method for Oxygen Scavengers	8140
DO	DO	Dissolved Oxygen,lodimetry method	N/A
F	Floride	SPADNS 2 Method for Fluoride	8029
FeMo	FeMo	Iron, for cooling water with molybdenum- based treatment	8365
FePh	Fe_phenanth	Iron, 1,10 phenanthroline	8008
FeSal	Fe-Sal	Total Iron using 5-Sulfosalicylic Acid Dihydrate	N/A
FeTp	FeTptz	Iron, TPTZ	8112
FeZi	FeZine	Iron, FerroZine	8147
H2O2	H2O2	Hydrogen peroxide, lodimetry method	N/A
H2O2L	H2O2L	Hydrogen peroxide, DPD method, Low Range	N/A
Mg	Mg	Calcium: Calmagite Colorimetric Method	8030
MnHR	MnHigh	High Range Manganese, Periodate Oxidation Method	8034
MnLR	MnLow	Low Range Manganese PAN Method	8149
MoHR	Mo_HighRange	Molybdenum, High Range, Mercaptoacetic Acid	8036
MoLR	Mo_LowRange	Molybdenum, Low Range, Ternary Complex	8169
N2H4	N2H4	P-Dimethylaminobenzaldehyde Method for Hydrazine	8141
NH2C	NH2CL	Indophenol Method for MonoChloramine	10171
N-TLR	N-TLR	Nitrogen, Total (Test 'N Tube Method) ,LR TNT Persulfate Digestion Method	10071
N-THR	N-TLR	Nitrogen, Total (Test 'N Tube Method) ,HR TNT Persulfate Digestion Method	10072
NH3S	NH3Sal	Ammonia Salicylate Method	8155
NH3-F	NH3-F	Ammonia Nitrogen ,Fluorescent Method	N/A
NH3LR	NH3LR	Nitrogen, Ammonia (Test 'N Tube) - Low range, Salicylate Method	10023
NH3HR	NH3HR	Nitrogen, Ammonia (Test 'N Tube) -High range, Salicylate Method	10031
Ni	Ni	PAN method for nickel	8150
NO2D	NO2D	Direct method for nitrite	N/A
NO2H	NO2H	Nitrite, High Range, Ferrous Sulfate	8153
NO2L	NO2L	Nitrite, Low Range, Diazotization	8507
NO3HR	NO3H	High range nitrate	8039
NO3MR	NO3M	Middle range nitrate	8171
NO3CA	NO3CA	NITRATE, High Range, Test 'N Tube	10020



Abbreviated Method Name	Method Name	Corresponding Hach © method	Hach Method Number
		Chromotropic Acid Method	
O3	O3	Ozone ,DPD method	N/A
PAA	PAA	Peroxyacetic , lodimetry method	N/A
OPO4	OPO4	Phosphorus, Reactive, Orthophosphate Ascorbic Acid	8048
OrgP	Phosphonate	Phosphonates, Persulfate UV Oxidation	8007
PAmi	OPO4-Amino	Phosphorus, Reactive, Amino Acid	8178
P-TLR	P-TLR	Phosphorus, Total (Test 'N Tube Method) -LR PhosVer 3 with Acid Persulfate Digestion	8190
P-THR	P-THR	Phosphorus, Total (Test 'N Tube Method) - HR-Molybdovanadate Method with Acid Persulfate Digestion	10127
pН	pН	Phenol red method for pH	10076
PMoV	OPO4-MoV	Phosphorus, Reactive, Molybdovanadate	8114
POLY	Polymer	Turbidimetric method for anionic polymers	N/A
Sb3+	Sb3+	Antimony Trivalent ,PADAP Method	N/A
Sb-T	Sb-T	Antimony, Total ,PADAP Method	N/A
S2-	Sulfide	Methylene Blue Method for Sulfide	8131
SiHR	SiHR	Silica, High Range, Silicomolybdate	8185
SiLR	SiLR	Silica, Low Range, Heteropoly Blue	8186
SO3LR	SO3LR	Sulfite,OPA method ,Low Range	N/A
SO3HR	SO3HR	Sulfite,OPA method , High Range	N/A
SO4	SO4	Sulfate	8051
TOC	TOC	Total Organic Carbon	10129
Urea	Urea	Urea (Reactor Digestion Method)	N/A
ZnXO	ZnXO	Zinc ,Xylenol orange method	N/A
Zn	Zinc	Zincon Method for Zinc	8009

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