

COD3 Plus

Colorimeter

Operator's Manual

Firmware Version 2.10 and greater

1925-MN-V3

12.06.19

Warning! This set contains chemicals that may be harmful if misused. Read cautions on individual containers carefully. Not to be used by children except under adult supervision.



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CONTENTS

COD3 Plus

| | |
|--|---|
| ▪ Kit Contents..... | 5 |
| ▪ Accessories..... | 5 |
| ▪ General Testing Procedures..... | 6 |
| ▪ Testing with LaMotte Pre-Programmed Tests..... | 6 |
| ▪ Calibrating LaMotte Pre-Programmed Tests..... | 8 |

SET UP

| | |
|-------------------------------|----|
| ▪ Setting the Clock..... | 11 |
| ▪ Logging Data..... | 12 |
| ▪ Factory Setup..... | 13 |
| ▪ Setting Power Save..... | 13 |
| ▪ Setting Backlight Time..... | 14 |
| ▪ Selecting A Language..... | 15 |
| ▪ Looping Menus..... | 16 |

COMPUTER CONNECTION

| | |
|----------------------------|----|
| ▪ Output..... | 17 |
| ▪ Computer Connection..... | 17 |

BATTERY

| | |
|-----------------------------|----|
| ▪ Battery/AC Operation..... | 17 |
| ▪ Battery Replacement..... | 18 |

MAINTENANCE

| | |
|-----------------------|----|
| ▪ Cleaning..... | 18 |
| ▪ Repairs..... | 18 |
| ▪ Meter Disposal..... | 19 |

GENERAL OPERATING INFORMATION

| | |
|--------------------------------------|----|
| ▪ Overview..... | 19 |
| ▪ General Operating Information..... | 19 |
| ▪ The Keypad..... | 20 |
| ▪ The Display and Menus..... | 20 |
| ▪ Tubes and Chambers..... | 22 |
| ▪ Sample Holders..... | 23 |
| ▪ Sample Dilution Techniques..... | 23 |

GENERAL INFORMATION

| | |
|-------------------------------|----|
| ■ Packaging and Delivery..... | 24 |
| ■ General Precautions | 24 |
| ■ Safety Precautions | 24 |
| ■ Limits of Liability | 24 |
| ■ Specifications | 25 |
| ■ CE | 26 |
| ■ IP67 | 26 |
| ■ Warranty..... | 26 |
| ■ Register Your Meter | 26 |

TROUBLE SHOOTING GUIDE

| | |
|------------------------------|----|
| ■ Error Messages..... | 27 |
| ■ Calibration..... | 27 |
| ■ Stray Light | 27 |
| ■ Troubleshooting Guide..... | 28 |

TEST PROCEDURES (ALPHABETICAL ORDER)

■ CONTENTS AND ACCESSORIES

Contents

| | |
|------------------------------|--------------|
| COD3 Plus Colorimeter | |
| Test Tubes, with Caps | Code 0290 |
| COD/UDV Adapter | Code 1724 |
| USB Wall Adapter | Code 1721 |
| USB Cable | Code 1720-01 |
| COD3 Plus Colorimeter Manual | |

Accessories

| | |
|---|------------------|
| Test Tubes, with Caps | Code 0290-6 |
| Replacement Chamber | Code 3-0038 |
| USB Cable | Code 1720-01 |
| USB Wall Adapter | Code 1721 |
| COD/UDV Adapter | Code 1724 |
| Car Charger | Code 5-0132 |
| Small Field Carrying Case [37.5 x 27.5 x 13.75 cm] | Code 1910-GCS150 |
| Large Field Carrying Case [45 x 32.5 x 20 cm] | Code 1910-GCS440 |

WARNING: Only use the USB Cable [Code 1720-01] that is supplied with the kit. Make no substitutions.

■ GENERAL TESTING PROCEDURES

The following is a step-by-step example of how to run tests from the Testing Menu. These test procedures are designed to be used with LaMotte SMART Reagent Systems.

LaMotte Company continuously updates the list of pre-programmed tests as the calibrations become available. Call LaMotte Technical Services at 1-800-344-3100 [410-778-3100 outside the USA] or email at tech@lamotte.com for a current list of available calibrations.

■ TESTING WITH LaMOTTE PRE-PROGRAMMED TESTS

| | | |
|---|--------------|---|
| 1. Press and briefly hold  to turn the meter on. The LaMotte logo screen will appear for about 3 seconds and the Main Menu will appear. | Main Menu | |
| | Testing Menu | |
| | Editing Menu | |
| | 12:00:00 | 001/500  |

| | | |
|--|-------------------|---|
| 2. Press  to select Testing Menu . | All Tests | |
| | 006 Ammonia-N LRF | |
| | 007 Ammonia-N LRS | |
| | 008 Ammonia-N HR | |
| | 014 Boron |  |
| | 12:00:00 | 001/500  |

| | | |
|--|-------------------|---|
| 3. Press  or  to scroll to the desired test. | All Tests | |
| | 006 Ammonia-N LRF |  |
| | 007 Ammonia-N LRS | |
| | 008 Ammonia-N HR |  |
| | 014 Boron | |
| | 12:00:00 | 001/500  |

| | | |
|--|-------------------|---|
| 4. Press  to select the test. | 007 Ammonia-N LRS | |
| | | |
| | Scan Bank |  |
| | Scan Sample | |
| | 12:00:00 | 001/500  |

| | |
|---|--|
| <p>5. Insert the blank into the chamber. Close the lid. Press ENTER to scan the blank. The screen will display Scan Blank Blank Done for about 1 second and then return to the test menu.</p> | <p>007 Ammonia-N LRS</p> <p>Scan Blank</p> <p>Scan Sample</p> <p>12:00:00 001/500 </p> |
|---|--|

| | |
|--|--|
| <p>6. Insert the reacted sample into the chamber. Close the lid. Press ENTER to scan the sample. The screen will display Scan Sample Sample Done for about 1 second. The result will appear on the screen.</p> | <p>007 Ammonia-N LRS</p> <p>1.00 ppm</p> <p>Scan Blank</p> <p>Scan Sample</p> <p>12:00:00 001/500 </p> |
|--|--|

| | |
|---|---|
| <p>To repeat the test, press ENTER to scan the sample again. The last blank scanned is used by the colorimeter for repeated scans. A different blank can be used by pressing ▲ or ▼ to scroll to Scan Blank and then scanning another blank. Scroll with ▲ or ▼ and make another selection with ENTER. The %T or Absorbance of the last test can be viewed by scrolling down and choosing %T/Abs. Press EXIT to escape to previous menus.</p> <p>NOTE: The menu loop in this screen so either ▲ or ▼ will lead to the menu selection needed.</p> | <p>007 Ammonia-N LRS</p> <p>1.00 ppm</p> <p>Scan Bank</p> <p>Scan Sample</p> <p>12:00:00 001/500 </p> |
|---|---|

■ CALIBRATING LaMOTTE PRE-PROGRAMMED TESTS

The LaMotte Pre-Programmed Tests have been pre-calibrated. Recalibration of the pre-programmed tests by the user is not possible. However, a procedure to standardize the calibration can be performed to obtain the most accurate readings or to meet regulatory requirements.

The LaMotte Pre-Programmed tests are standardized with one standard solution. To standardize over the full range of the test, the concentration of the standard should be chosen from the high end of the range. Alternatively, if samples do not cover the full range of the test, a standard should be chosen that is close to the concentration of the samples.

The standardization procedure should be followed as often as required by regulations and laws for compliance monitoring.

In the example below, the Aluminum calibration will be standardized.

Prepare a standard solution to be tested. In this example, 0.30 ppm ammonia nitrogen.

| | | |
|---|--------------|---|
| 1. Press and briefly hold  to turn the meter on. The LaMotte logo screen will appear for about 3 seconds and the Main Menu will appear. | Main Menu | |
| | Testing Menu |  |
| Editing Menu | | |
| | 12:00:00 | 001/500  |

| | | |
|--|-------------------|---|
| 2. Press  to select Testing Menu . | Testing Menu | |
| | 006 Ammonia-N LRF |  |
| 007 Ammonia-N LRS | | |
| | 008 Ammonia-N HR | |
| | 014 Boron | |
| | 12:00:00 | 001/500  |

| | | |
|---|-------------------|---|
| 3. Press  or  to scroll to the desired test factor. | All Tests | |
| | 006 Ammonia-N LRF |  |
| 007 Ammonia-N LRS | | |
| | 008 Ammonia-N HR | |
| | 014 Boron | |
| | 12:00:00 | 001/500  |

| | | |
|---|-----------------------|---|
| 4. Press ENTER to select the test. | 007 Ammonia-N LRS | |
| | Scan Blank | ↓ |
| | Scan Sample | |
| | 12:00:00 001/500 | |

| | | |
|---|-----------------------|---|
| 5. Follow the test procedure in the manual to test the prepared standard. Insert the blank into the chamber. Close the lid. Press ENTER to scan the blank. The screen will display Scan Blank Blank Done for about 1 second and then return to the Test Menu . | 007 Ammonia-N LRS | |
| | Scan Blank | ↑ |
| | Scan Sample | ↓ |
| | 12:00:00 001/500 | |

| | | |
|--|-----------------------|---|
| 6. Insert the reacted standard solution into the chamber. Close the lid. Press ENTER to scan the sample. The screen will display Scan Sample Sample Done for about 1 second. The result will appear on the screen. | 007 Ammonia-N LRS | |
| | 0.28 ppm | ↑ |
| | Scan Blank | |
| | Scan Sample | |
| | 12:00:00 001/500 | |

| | | |
|---|-----------------------|---|
| 7. The displayed result can now be standardized. Press ▲ or ▼ to scroll to Calibrate . | 007 Ammonia-N LRS | |
| | 0.28 ppm | ↑ |
| | %T/Abs | |
| | Calibrate | |
| | 12:00:00 001/500 | |

| | | |
|---|-----------------------|--|
| 8. Press ENTER to select Calibrate . A reverse font [light background with dark characters] will appear to indicate that the reading can be adjusted. | 007 Ammonia-N LRS | |
| | 0.28 ppm | |
| | ^, v=Edit, ENTER=Save | |
| | ^ +ENTER=Default | |
| | 12:00:00 001/500 | |

| | | |
|---|-----------------------|---|
| <p>9. Press  or  to adjust the value shown to the concentration of the prepared standard, 0.30 in this example.</p> <p>NOTE: A maximum adjustment of 25% is possible.</p> | 007 Ammonia-N LRS | |
| | 0.30 ppm | |
| | ^, v=Edit, ENTER=Save | |
| | ^ +ENTER=Default | |
| | 12:00:00 | 001/500  |

| | | |
|---|---|---|
| <p>10. Press  to save the value.</p> <p>To leave the Calibration procedure without saving the adjustment, press .</p> <p>Press  and  at any time to return to the default value.</p> <p>The calibration has now been standardized and can be used for testing. Scroll to Scan Blank and begin testing.</p> | 007 Ammonia-N LRS | |
| | 0.30 ppm  | |
| | %T/Abs | |
| | Calibrate | |
| | 12:00:00 | 001/500  |

■ SETTING THE CLOCK

Setting the clock allows the correct time and date stamp to be stored with each reading in the data logger.

| | | | |
|---|----------------|---|--|
| <p>1. From the Editing Menu, press  or  to scroll to Set Clock.</p> | Editing Menu | | |
| | Set Clock | |   |
| | Logging | | |
| | Factory Setup | | |
| | Set Power Save | | |
| 12:00:00 | 001/500 |  | |

| | | | |
|---|--------------------|---------|---|
| <p>2. Press ENTER to select Set Clock. The year is displayed. Press  or  to scroll to the appropriate character. Press ENTER to select the character. The month, day, hour, format hour, minute, second, AM/PM will be displayed. Repeat for each.</p> | Set Time | | |
| | Year: 20 <u>00</u> | | |
| | 12:00:00 | 001/500 |  |

| | | | |
|---|----------------|---|--|
| <p>3. Press ENTER to select the final character. The time and date will be saved and the meter will return to the Editing Menu.</p> | Editing Menu | | |
| | Set Clock | |   |
| | Logging | | |
| | Factory Setup | | |
| | Set Power Save | | |
| 12:00:00 | 001/500 |  | |

■ LOGGING DATA

The default setting for the data logger is enabled. The meter will log the last 500 data points. The counter in the center bottom of the display will show how many data points have been logged. The display will show 500+ when the data logger has exceeded 500 points and the data points are being overwritten.

| | | |
|--|----------------|---|
| 1. From the Editing Menu , press  or  to scroll to Logging . | Editing Menu | |
| | Set Clock | ↑ |
| | Logging | |
| | Factory Setup | ↓ |
| | Set Power Save | |
| 12:00:00 001/500  | | |

| | | |
|---|------------------|--|
| 2. Press  to select Logging . | Logging | |
| | Display Test Log | |
| | Logging Enabled | |
| | Logging Disabled | |
| | Erase Log | |
| 12:00:00 001/500  | | |

| | | |
|--|------------------|--|
| 3. Press  or  to scroll to desired function. | Logging | |
| | Display Test Log | |
| | Logging Enabled | |
| | Logging Disabled | |
| | Erase Log | |
| 12:00:00 001/500  | | |

| | | |
|---|----------------|---|
| 4. Press  . The screen will display Storing... for about 1 second and return to the Editing Menu . | Editing Menu | |
| | Set Clock | ↑ |
| | Logging | |
| | Factory Setup | ↓ |
| | Set Power Save | |
| 12:00:00 001/500  | | |

■ FACTORY SETUP

The Factory Setup menu is used in manufacturing of the colorimeter. This menu is not for use by the operator in the field.

■ SETTING POWER SAVE

The power saving Auto Shutoff feature will turn the meter off when a button has not been pushed for a set amount of time. The default setting is disabled. To change the setting:

| | | | | | |
|---|--|-----------|--------|---------|---------------|
| 1. From the Editing Menu , press  or  to scroll to Set Power Save . | Editing Menu | | | | |
| | <table border="1"> <tr><td>Set Clock</td><td rowspan="4" style="text-align: center;">↑ ↓</td></tr> <tr><td>Logging</td></tr> <tr><td>Factory Setup</td></tr> <tr><td>Set Power Save</td></tr> </table> | Set Clock | ↑ ↓ | Logging | Factory Setup |
| Set Clock | ↑ ↓ | | | | |
| Logging | | | | | |
| Factory Setup | | | | | |
| Set Power Save | | | | | |
| | 12:00:00 001/500  | | | | |

| | | | | | |
|--|---|---------|--|-----------|------------|
| 2. Press  to select Set Power Save . | Set Power Save | | | | |
| | <table border="1"> <tr><td>Disable</td><td rowspan="4" style="text-align: center;"> </td></tr> <tr><td>5 Minutes</td></tr> <tr><td>15 Minutes</td></tr> <tr><td>30 Minutes</td></tr> </table> | Disable | | 5 Minutes | 15 Minutes |
| Disable | | | | | |
| 5 Minutes | | | | | |
| 15 Minutes | | | | | |
| 30 Minutes | | | | | |
| | 12:00:00 001/500  | | | | |

| | | | | | |
|--|---|---------|--|-----------|------------|
| 3. Press  or  to scroll to desired function. | Set Power Save | | | | |
| | <table border="1"> <tr><td>Disable</td><td rowspan="4" style="text-align: center;"> </td></tr> <tr><td>5 Minutes</td></tr> <tr><td>15 Minutes</td></tr> <tr><td>30 Minutes</td></tr> </table> | Disable | | 5 Minutes | 15 Minutes |
| Disable | | | | | |
| 5 Minutes | | | | | |
| 15 Minutes | | | | | |
| 30 Minutes | | | | | |
| | 12:00:00 001/500  | | | | |

| | | | | | |
|---|--|-----------|--------|---------|---------------|
| 4. Press  . The screen will display Storing.... for about 1 second and the meter will return to the Editing Menu . | Editing Menu | | | | |
| | <table border="1"> <tr><td>Set Clock</td><td rowspan="4" style="text-align: center;">↑ ↓</td></tr> <tr><td>Logging</td></tr> <tr><td>Factory Setup</td></tr> <tr><td>Set Power Save</td></tr> </table> | Set Clock | ↑ ↓ | Logging | Factory Setup |
| Set Clock | ↑ ↓ | | | | |
| Logging | | | | | |
| Factory Setup | | | | | |
| Set Power Save | | | | | |
| | 12:00:00 001/500  | | | | |

■ SETTING THE BACKLIGHT TIME

The backlight illuminates the display for enhanced viewing. The default setting is 10 seconds. If Button Control is chosen the backlight button on the key pad will act as an on/off switch and the backlight will remain on or off when the meter is being used. When one of the other settings – 10, 20 or 30 seconds – is chosen, the display will be illuminated for the specified amount of time after any button is pressed.

NOTE: The backlight feature uses a significant amount of power. The longer the backlight is on, the more frequently the battery will have to be charged if the USB/Wall Adapter is not being used.

| <p>1. From the Editing Menu, press  or  to scroll to Backlight Time.</p> | <table border="1"> <thead> <tr> <th colspan="3">Editing Menu</th> </tr> </thead> <tbody> <tr> <td>Logging</td> <td></td> <td rowspan="4">   </td> </tr> <tr> <td>Factory Setup</td> <td></td> </tr> <tr> <td>Set Power Save</td> <td></td> </tr> <tr> <td>Set Backlight Time</td> <td></td> </tr> <tr> <td>12:00:00</td> <td>001/500</td> <td></td> </tr> </tbody> </table> | Editing Menu | | | Logging | |   | Factory Setup | | Set Power Save | | Set Backlight Time | | 12:00:00 | 001/500 |  |
|--|---|--|--|--|---------|--|--|---------------|--|----------------|--|--------------------|--|----------|---------|---|
| Editing Menu | | | | | | | | | | | | | | | | |
| Logging | |   | | | | | | | | | | | | | | |
| Factory Setup | | | | | | | | | | | | | | | | |
| Set Power Save | | | | | | | | | | | | | | | | |
| Set Backlight Time | | | | | | | | | | | | | | | | |
| 12:00:00 | 001/500 |  | | | | | | | | | | | | | | |

| <p>2. Press  to select Set Backlight Time.</p> | <table border="1"> <thead> <tr> <th colspan="3">Set Backlight Time</th> </tr> </thead> <tbody> <tr> <td>Button Control</td> <td></td> <td></td> </tr> <tr> <td>10 seconds</td> <td></td> <td></td> </tr> <tr> <td>20 seconds</td> <td></td> <td></td> </tr> <tr> <td>30 seconds</td> <td></td> <td></td> </tr> <tr> <td>12:00:00</td> <td>001/500</td> <td></td> </tr> </tbody> </table> | Set Backlight Time | | | Button Control | | | 10 seconds | | | 20 seconds | | | 30 seconds | | | 12:00:00 | 001/500 |  |
|--|---|---|--|--|----------------|--|--|------------|--|--|------------|--|--|------------|--|--|----------|---------|---|
| Set Backlight Time | | | | | | | | | | | | | | | | | | | |
| Button Control | | | | | | | | | | | | | | | | | | | |
| 10 seconds | | | | | | | | | | | | | | | | | | | |
| 20 seconds | | | | | | | | | | | | | | | | | | | |
| 30 seconds | | | | | | | | | | | | | | | | | | | |
| 12:00:00 | 001/500 |  | | | | | | | | | | | | | | | | | |

| <p>3. Press  or  to scroll to desired option.</p> | <table border="1"> <thead> <tr> <th colspan="3">Set Backlight Time</th> </tr> </thead> <tbody> <tr> <td>Button Control</td> <td></td> <td></td> </tr> <tr> <td>10 seconds</td> <td></td> <td></td> </tr> <tr> <td>20 seconds</td> <td></td> <td></td> </tr> <tr> <td>30 seconds</td> <td></td> <td></td> </tr> <tr> <td>12:00:00</td> <td>001/500</td> <td></td> </tr> </tbody> </table> | Set Backlight Time | | | Button Control | | | 10 seconds | | | 20 seconds | | | 30 seconds | | | 12:00:00 | 001/500 |  |
|---|---|---|--|--|----------------|--|--|------------|--|--|------------|--|--|------------|--|--|----------|---------|---|
| Set Backlight Time | | | | | | | | | | | | | | | | | | | |
| Button Control | | | | | | | | | | | | | | | | | | | |
| 10 seconds | | | | | | | | | | | | | | | | | | | |
| 20 seconds | | | | | | | | | | | | | | | | | | | |
| 30 seconds | | | | | | | | | | | | | | | | | | | |
| 12:00:00 | 001/500 |  | | | | | | | | | | | | | | | | | |

| <p>4. Press . The screen will display Storing... for about 1 second and the meter will return to the Editing Menu.</p> | <table border="1"> <thead> <tr> <th colspan="3">Editing Menu</th> </tr> </thead> <tbody> <tr> <td>Logging</td> <td></td> <td rowspan="4">   </td> </tr> <tr> <td>Factory Setup</td> <td></td> </tr> <tr> <td>Set Power Save</td> <td></td> </tr> <tr> <td>Set Backlight Time</td> <td></td> </tr> <tr> <td>12:00:00</td> <td>001/500</td> <td></td> </tr> </tbody> </table> | Editing Menu | | | Logging | |   | Factory Setup | | Set Power Save | | Set Backlight Time | | 12:00:00 | 001/500 |  |
|---|---|--|--|--|---------|--|--|---------------|--|----------------|--|--------------------|--|----------|---------|---|
| Editing Menu | | | | | | | | | | | | | | | | |
| Logging | |   | | | | | | | | | | | | | | |
| Factory Setup | | | | | | | | | | | | | | | | |
| Set Power Save | | | | | | | | | | | | | | | | |
| Set Backlight Time | | | | | | | | | | | | | | | | |
| 12:00:00 | 001/500 |  | | | | | | | | | | | | | | |

■ SELECTING A LANGUAGE

There are seven languages available: English, Spanish, French, Portuguese, Italian, Chinese, and Japanese.

| | | |
|--|--------------------|---|
| 1. From the Editing Menu , press  or  to scroll to Select Language . | Editing Menu | |
| | Factory Setup |  |
| Set Power Save | | |
| | Set Backlight Time | |
| | Select Language | |
| | 12:00:00 | 001/500  |

| | | |
|---|-----------------|---|
| 2. Press  to select Select Language . | Select Language | |
| | English |  |
| Spanish | | |
| French | | |
| Portugese | | |
| | 12:00:00 | 001/500  |

| | | |
|--|-----------------|--|
| 3. Press  or  to scroll to desired language. | Select Language | |
| | English |  |
| Spanish | | |
| French | | |
| Portugese | | |
| | 12:00:00 | 001/500  |

| | | |
|--|-----------------|---|
| 4. Press  . The screen will display Storing... for about 1 second and the meter will return to the Editing Menu . | Editing Menu | |
| | Set Power Save |  |
| Set Backlight Time | | |
| | Bluetooth Menu | |
| | Select Language | |
| | 12:00:00 | 001/500  |

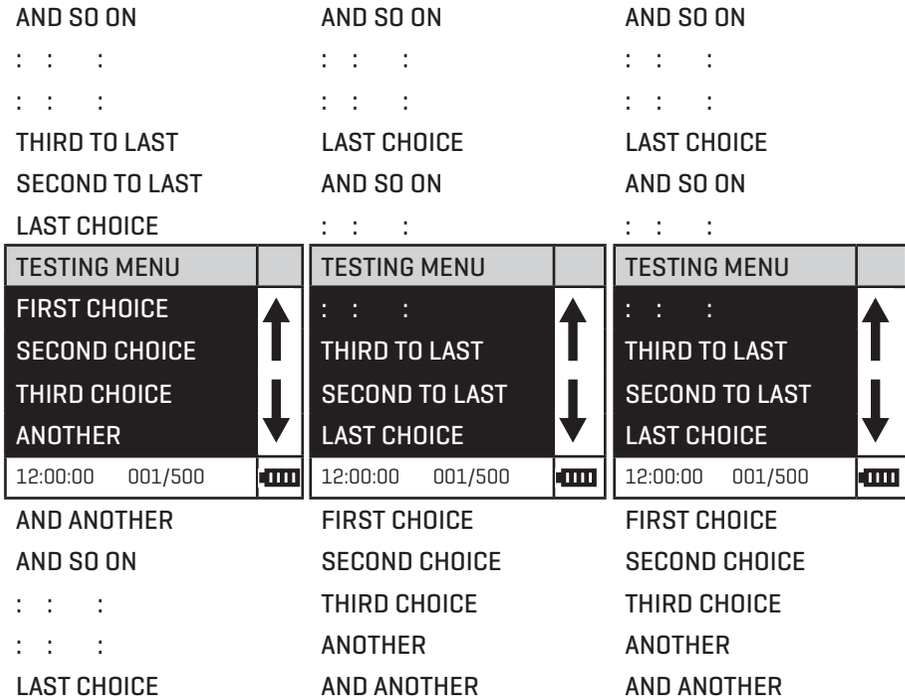
NOTE: If meter unintentionally switches to another language, use the procedure above to reset the meter to the desired language. For example, to reset the meter to English:

Turn meter on.

1. Press  one time. Press .
2. Press  five times. Press .
3. Press .

■ LOOPING MENUS

Long menus, such as the Testing Menu, incorporate a looping feature which allows the user to quickly reach the last choice in the menu from the first choice. In a looping menu the last choices in the menu are above the first choice and scrolling upward moves through the menu in reverse order. Scrolling downward moves through the menu from first choice to last but the menu starts over following the last choice. So all menu choices can be reached by scrolling in either direction. The diagrams below demonstrate a looping menu.



The feature called **Looping Menu** can be turned on and off in the **Editing Menu**. The default setting is ON.

COMPUTER CONNECTION

■ OUTPUT

USB

■ COMPUTER CONNECTION

USB Type A, USB mini B, Order Cable Code 1720.

■ WATERLINK CONNECT 2

The meter may be interfaced with any Windows-based 64 bit computer by using the LaMotte WaterLink Connect 2 program and a USB cable. The program will store test information and results in a local database, and allow for exporting this data to a comma separated value [CSV] file. The meter will send the following data: time/date stamp, name of test, sample value, sample units, meter name, and location. To download WaterLink 2 go to <http://www.waterlinkconnect.com/>. Then select "Products" > "WaterLink Connect 2 Application" > "Download".

BATTERY

■ BATTERY/AC OPERATION

The colorimeter may be operated on battery power using a USB wall adapter or USB computer connection. If using the meter as a bench top unit, use the wall adapter if possible to extend the battery life. The meter will remain on when the USB adapter is used.

To charge the battery with the wall adapter, plug the smaller end of the USB cable [USB mini B connector] into the meter and the larger end of the USB cable [USB type A connector] into the wall adapter. Plug the wall adapter into an AC outlet. Reinsert the rubber USB port plug after charging. To charge the battery from a computer, plug the smaller end of the USB cable [USB mini B connector] into the meter and the larger end of the USB cable [USB Type A connector] into a USB port on a computer. The battery icon will show no bars and flash when the unit first turns on. Then the indicator will indicate the battery status by showing 0, 1, 2, 3, or 4 bars.

It will take 6 hours to fully charge a low battery. The battery icon will flash when the battery is charging. The battery icon will show four bars and stop flashing when it is fully charged. The charging circuit will automatically switch to a float charge when the battery is fully charged. The charger may remain connected. Some computers will NOT supply power to their USB ports during standby operation. The wall adapter will charge the unit continuously.

Storing the meter above ambient room temperature will decrease the battery charge more quickly than storage at room temperature. If the meter does not turn on, it means that the battery is at a very low charge. Charging the battery with the wall adapter in this state may take up to 10 hours. At low temperatures, approaching 0 °C,

the battery will charge more slowly. It will not charge at all below 0 °C. The battery icon will show no bars and continuously flash if the battery is getting low but the unit will still operate normally. A “Low Battery” message on the status bar of the display will replace the time when the battery voltage is too low for proper operation and accuracy may be degraded. A “Shutdown Low Batt” message on the display will appear for a few seconds before the power is switched off when the battery is too low to operate the unit. When the battery icon simultaneously flashes bars 1 and 2 followed by bars 3 and 4, it is an indication that the battery is damaged and technical support should be contacted.

To extend the battery life:

- Shut down the unit with the power switch when not taking measurements or use the power save option to have the unit automatically turn off after 5 minutes.
- Store the unit at a moderate temperature.
- Fully charge the battery before storing the unit for extended periods of time.
- Fully charge the battery at least once per year. Failure to do so may result in a permanently drained battery.
- Limit backlight use. The unit consumes 3X normal power with the backlight on. Set the backlight time option to 10 seconds, or select “Button Control” and keep the backlight off.

BATTERY REPLACEMENT

The lithium ion battery used in this unit should last for many years with normal use. When it no longer powers the unit long enough to meet testing requirements it will need to be replaced. Lithium ion batteries that are properly charged and stored do not usually lose all capacity; they just have less capacity after hundreds of charge cycles. This unit uses a custom battery assembly that is only available from LaMotte Company. Battery replacement must be performed at a LaMotte authorized repair facility. The water resistant housing of this meter should not be opened by the user. Contact LaMotte Company by phone [1-800-344-3100] or email [tech@lamotte.com] for a return authorization number.

MAINTENANCE

.....

■ CLEANING

Clean the exterior housing with a damp, lint-free cloth. Do not allow water to enter the light chamber or any other parts of the meter. To clean the light chamber and optics area, point a can of compressed air into the light chamber and blow the pressurized air into the light chamber. Use a cotton swab dampened with Windex® window cleaner to gently swab the interior of the chamber. Do not use alcohol; it will leave a thin residue over the optics when dry.

■ REPAIRS

Should it be necessary to return the meter for repair or servicing, pack the meter

carefully in a suitable container with adequate packing material. A return authorization number must be obtained from LaMotte Company by calling 800-344-3100 (US only) or 410-778-3100, faxing 410-778-6394, or emailing tech@lamotte.com. Often a problem can be resolved over the phone or by email. If a return of the meter is necessary, attach a letter with the return authorization number, meter serial number, a brief description of problem and contact information including phone and FAX numbers to the shipping carton. This information will enable the service department to make the required repairs more efficiently.

■ METER DISPOSAL

Waste Electrical and Electronic Equipment [WEEE]

Natural resources were used in the production of this equipment. This equipment may contain materials that are hazardous to health and the environment. To avoid harm to the environment and natural resources, the use of appropriate take-back systems is recommended. The crossed out wheeled bin symbol on the meter encourages the use of these systems when disposing of this equipment.



Take-back systems will allow the materials to be reused or recycled in a way that will not harm the environment. For more information on approved collection, reuse, and recycling systems contact local or regional waste administration or recycling services.

GENERAL OPERATING INFORMATION

■ OVERVIEW

The meter is a portable, microprocessor controlled, direct reading colorimeter. It has a graphical liquid crystal display and 6 button keypad. These allow the user to select options from the menu driven software, to directly read test results or to review stored results of previous tests in the data logger. The menus can be displayed in seven different languages.

The LaMotte tests are precalibrated for LaMotte reagent systems. The colorimeter displays the result of these tests directly in units of concentration.

The optics feature a colored LED. The LED has a corresponding silicon photoiode with an integrated interference filter. The interference filter selects a narrow band of light from the corresponding LED for the colorimetric measurements.

A USB wall adapter, USB computer connection or lithium battery powers the colorimeter.

■ GENERAL OPERATING INFORMATION

The operation of the colorimeter is controlled by the menu driven software and user interface. A menu is a list of choices. This allows a selection of various tasks for the colorimeter to perform, such as scan blank and scan sample. The keypad is used to make menu selections that are viewed on the display.

■ THE KEYPAD

The keypad has 6 buttons which are used to perform specific tasks.

| | |
|--|---|
|  | This button will scroll up through a list of menu selections. |
|  | The button is used to select choices in a menu viewed on the display. |
|  | This button controls the backlight on the display. |
|  | This button will scroll down through a list of menu selections. |
|  | This button exits to the previous menu. |
|  | This button turns the meter on or off. |

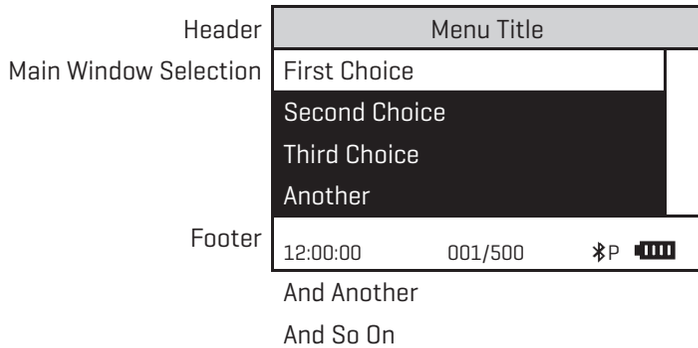


■ THE DISPLAY AND MENUS

The display allows menu selections to be viewed and selected. These selections instruct the colorimeter to perform specific tasks. The menus are viewed in the display using two general formats that are followed from one menu to the next. Each menu is a list of choices or selections.

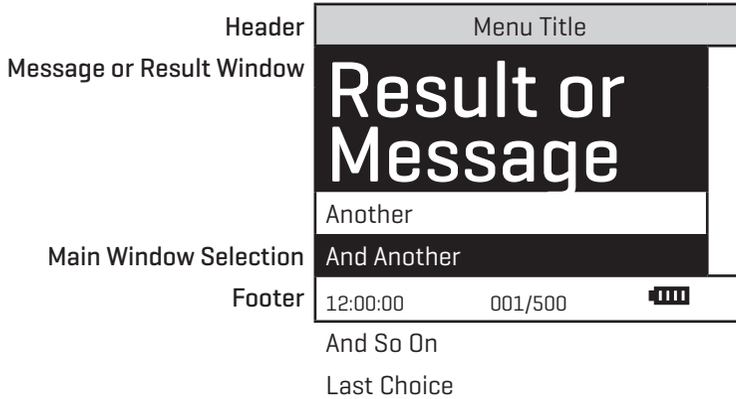
The display has a header line at the top and a footer line at the bottom. The header displays the title of the current menu. The footer line displays the time and the date, the data logger status, the bluetooth/printer status and the battery status. The menu selection window is in the middle of the display between the header and the footer.

The menu selection window displays information in two general formats. In the first format only menu selections are displayed. Up to 4 lines of menu selections may be displayed. If more selections are available they can be viewed by pressing the arrow buttons   to scroll the other menu selections into the menu selection window. Think of the menu selections as a vertical list in the display that moves up or down each time an arrow button   is pressed. Some menus in the colorimeter are looping menus. The top and bottom menu choices are connected in a loop. Scrolling down past the bottom of the menu will lead to the top of the menu. Scrolling up past the top of the menu will lead to the bottom of the menu.



A light bar will indicate the menu choice. As the menu is scrolled through, the light bar will highlight different menu choices. Pressing the  button will select the menu choice that is indicated by the light bar.

In the second format the menu choice window takes advantage of the graphical capabilities of the display. Large format graphic information, such as test results or error messages or the LaMotte logo is displayed. The top two lines of the display are used to display information in a large, easy to read format. The menus work in the same way as previously described but two lines of the menu are visible at the bottom of the display.



As described previously, the **EXIT** button allows an exit or escape from the current menu and a return to the previous menu. This allows a rapid exit from an inner menu to the main menu by repeatedly pushing the **EXIT** button. Pushing  at any time will turn the colorimeter off.

The display may show the following messages:

| | |
|---|---|
|  | Battery Status |
|  | More choices are available and can be viewed by scrolling up and/or down through the display. |
| Header | Identifies the current menu and information on units and reagent systems if applicable. |
| Footer | In the data logging mode the number of the data point is displayed and the total number of data points in the memory will be shown. The footer also shows current time and battery status |

■ TUBES AND CHAMBERS

The colorimeter uses one type of tube [Code 0290] for all test factors.

The handling of the tubes is of utmost importance. Tubes must be clean and free from lint, fingerprints, dried spills and significant scratches, especially the central zone between the bottom and the sample line.

Scratches, fingerprints and water droplets on the tube can cause stray light interference leading to inaccurate results. Tubes that have been scratched in the light zone through excessive use should be discarded and replaced with new ones.

Tubes should always be washed on the inside and outside with mild detergent prior to use to remove dirt or fingerprints. The tubes should be allowed to air-dry in an inverted position to prevent dust from entering the tubes. Dry tubes should be stored with the caps on to prevent contamination.

After a tube has been filled and capped, it should be held by the cap and the outside surface should be wiped with a clean, lint-free absorbent cloth until it is dry and smudge-free. Handling the tube only by the cap will avoid problems from fingerprints. Always set the clean tube aside on a clean surface that will not contaminate the tube. It is imperative that the tubes and light chamber be clean and dry. The outside of the tubes should be dried with a clean, lint-free cloth or disposable wipe before they are placed in the meter chamber.

Tubes should be emptied and cleaned as soon as possible after reading a sample to prevent deposition of particulates on the inside of the tubes.

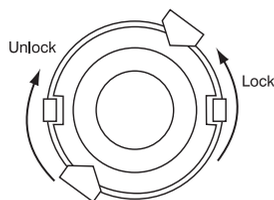
Variability in the geometry of the glassware and technique is the predominate cause of variability in results. Slight variations in wall thickness and the diameter of the tubes may lead to slight variations in the test results. To eliminate this error the tubes should be placed in the chamber with the same orientation each time.

Chambers which have been scratched through excessive use should be discarded and replaced with a new one.

■ SAMPLE HOLDERS

The sample chamber is designed for 25 mm round tubes. An adapter to hold 16 mm COD tubes and 10 mm square UDV cuvettes is included. 10 cm cuvettes can be scanned only at 525 nm and 568 nm. The light path is blocked at 428 nm and 635 nm with the use of the adapter. COD tubes can be scanned at all wavelengths.

Position the COD/UDV Adapter [Code 1724] so that the notches in the adapter fit around the posts on the chamber. Turn the adapter counterclockwise until the arrows are at the top and bottom of the chamber and the adapter is locked into place. Turn the adapter clockwise to unlock the adapter and remove it from the chamber.



■ SAMPLE DILUTION TECHNIQUES

If a test result is out of the range of the meter, it must be diluted. The test should then be repeated on the diluted sample. The following table gives quick reference guidelines for dilutions of various proportions.

| Amount of Sample | Deionized Water to Bring Final Volume to 10 mL | Multiplication Factor |
|------------------|--|-----------------------|
| 10 mL | 0 mL | 1 |
| 5 mL | 5 mL | 2 |
| 2.5 mL | 7.5 mL | 4 |
| 1 mL | 9 mL | 10 |
| 0.5 mL | 9.5 mL | 20 |

All dilutions are based on a final volume of 10 mL, so several dilutions will require small volumes of the water sample. Graduated pipets should be used for all dilutions. If volumetric glassware is not available, dilutions can be made with the colorimeter tube. Fill the tube to the 10 mL line with the sample and then transfer it to another container. Add 10 mL volumes of deionized water to the container and mix. Transfer 10 mL of the diluted sample to the colorimeter tube and follow the test procedure. Repeat the dilution and testing procedures until the result falls within the range of

the calibration. Multiply the test result by the dilution factor. For example, if 10 mL of the sample water is diluted with three 10 mL volumes of deionized water, the dilution factor is four. The test result of the diluted sample should be multiplied by four.

GENERAL INFORMATION

■ PACKAGING AND DELIVERY

Experienced packaging personnel at LaMotte Company assure adequate protection against normal hazards encountered in transportation of shipments.

After the product leaves LaMotte Company, all responsibility for safe delivery is assured by the transportation company. Damage claims must be filed immediately with the transportation company to receive compensation for damaged goods.

■ GENERAL PRECAUTIONS

READ THE INSTRUCTION MANUAL BEFORE ATTEMPTING TO SET UP OR OPERATE THE METER. Failure to do so could result in personal injury or damage to the meter. The meter should not be used or stored in a wet or corrosive environment. Care should be taken to prevent water from wet tubes from entering the meter chamber.

NEVER PUT WET TUBES IN THE METER.

■ SAFETY PRECAUTIONS

Read the labels on all LaMotte reagent containers prior to use. Some containers include precautionary notices and first aid information. Certain reagents are considered hazardous substances and are designated with a * in the instruction manual. Safety Data Sheets [SDS] can be found at www.lamotte.com. Read the SDS before using these reagents. Additional emergency information for all LaMotte reagents is available 24 hours a day from the National Poison Control Center 1-800-222-1222 or by contacting the 24 hour emergency line for ChemTel 1-800-255-3924 [USA, Canada, Puerto Rico]; locations outside the North American Continent 813-248-0585 [call collect]. Be prepared to supply the name and four-digit LaMotte code number found on the container label or at the top of the SDS or in the contents list of the procedure. LaMotte reagents are registered with a computerized poison control information system available to all local poison control centers.

Keep equipment and reagent chemicals out of the reach of young children.

Ensure that the protection provided by this equipment is not impaired. Do not install or use this equipment in a manner that is not indicated in this manual.

■ LIMITS OF LIABILITY

Under no circumstances shall LaMotte Company be liable for loss of life, property, profits, or other damages incurred through the use or misuse of its products.

■ SPECIFICATIONS

INSTRUMENT TYPE: Colorimeter

| | |
|--------------------------|--|
| Readout | 160 x 100 backlit LCD, 20 x 6 line graphical display |
| Wavelengths | 428 nm, 525 nm, 568 nm, 635 nm |
| Wavelength Accuracy | ±2% FS |
| Readable Resolution | Determined by reagent system |
| Wavelength Bandwidth | 10 nm typical |
| Photometric Range | -2 to +2 AU |
| Photometric Precision | ± 0.001 AU at 1.0 AU |
| Photometric Accuracy | ±0.005 AU at 1.0 AU |
| Sample Chamber | Accepts 25 mm diameter flat-bottomed test tubes, 10 mm square cuvettes†, 16 mm COD test tubes |
| Light Sources | 4 LEDs |
| Detectors | 4 silicon photodiodes |
| Modes | Pre-programmed tests, absorbance, %T |
| Pre-Programmed Tests | YES, with automatic wavelength selection |
| User Defined Tests | Up to 25 user tests can be input |
| Languages | English, Spanish, French, Portuguese, Italian, Chinese, Japanese |
| Temperature | Operation: 0-50 °C; Storage: -40-60 °C |
| Operation Humidity Range | 0-90 % RH, non-condensing |
| USB Port | Mini B |
| Power Requirements | USB wall adapter, USB computer connection or lithium ion rechargeable battery |
| Battery | Charge Life: Approximately 380 tests with backlight on to 1000 tests with backlight off. [Signal averaging disabled]. Battery Life: Approximately 500 charges. |
| Electrical Rating | Rated voltage [5V], Rated power of input current [1.0A] at mini-USB input port |
| Data Logger | 500 test results stored for download to a PC |
| Waterproof | IP67 with USB port plug in place |
| Dimensions [LxWxH] | 3.5 x 7.5 x 2.5 inches, 8.84 x 19.05 x 6.35 cm |
| Weight | 13 oz, 362 g [meter only] |

†525 nm and 568 nm only

■ CE COMPLIANCE

The Smart3 Colorimeter has earned the European CE Mark of Compliance for electromagnetic compatibility and safety. The Declaration of Conformity for the Smart3 colorimeter is available at www.lamotte.com.

■ IP67 CERTIFICATION

The Smart3 meets IP67 standards for protection against dust and immersion only when the USB port plug is in place. Documentation is available at www.lamotte.com.

■ WARRANTY

LaMotte Company warrants this instrument to be free of defects in parts and workmanship for 2 years from the date of shipment. If it should become necessary to return the instrument for service during or beyond the warranty period, contact our Technical Service Department at 1-800-344-3100 for a return authorization number or visit www.lamotte.com for troubleshooting help. The sender is responsible for shipping charges, freight, insurance and proper packaging to prevent damage in transit. This warranty does not apply to defects resulting from action of the user such as misuse, improper wiring, operation outside of specification, improper maintenance or repair, or unauthorized modification. LaMotte Company specifically disclaims any implied warranties or merchantability or fitness for a specific purpose and will not be liable for any direct, indirect, incidental or consequential damages. LaMotte Company's total liability is limited to repair or replacement of the product. The warranty set forth above is inclusive and no other warranty, whether written or oral, is expressed or implied.

■ REGISTER YOUR METER

To register your meter with the LaMotte Service Department, go to www.lamotte.com and choose SUPPORT on the top navigation bar.

TROUBLESHOOTING

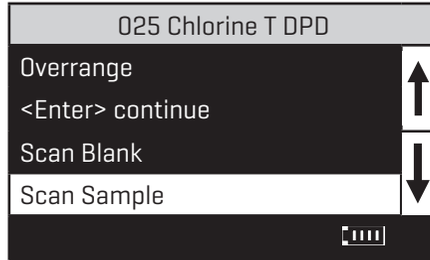
■ ERROR MESSAGES

Over Range

If the message **OVERRANGE** is displayed when scanning a sample, the sample may be over range or under range. If the sample is over range the sample should be diluted and tested again [see Sample Dilution Techniques and Volumetric Measurements, page 23].

If overrange is displayed, press **ENTER** to continue testing on diluted samples.

Note: After pressing **ENTER**, the overrange concentration will be displayed. This concentration is an **approximation only**.



■ CALIBRATION

As with all pre-calibrated meters, it is highly recommended, even if not required by regulations, that the user periodically verify the performance of the meter by running standards with a predetermined concentration. Results outside of specification are an indication that the meter needs to be adjusted. This can be done following the user calibration described on page 6. If the user calibration fails to properly adjust the meter then the meter should be returned to LaMotte Company for recalibration. [See page 8].

■ STRAY LIGHT

The colorimeter should have no problems with stray light. Make sure that the sample compartment lid is always fully closed.

■ TROUBLESHOOTING GUIDE

| <i>PROBLEM</i> | <i>REASON</i> | <i>SOLUTION</i> |
|--|---|---|
|  Flashing | Low battery. Readings are reliable. | Charge battery or use USB wall/computer adapter. |
| “Low Battery” | Battery voltage is very low. Readings are not reliable. | Charge battery or use USB wall/computer adapter. |
| “Shut Down Low Batt” Shut Down | Battery is too low to operate the unit. | Charge battery or use USB wall/computer adapter. |
| “Overrange” | Sample is outside of acceptable range. | Dilute sample and test again. |
| Unusually large negative or positive readings when performing calibration | Incorrect standards used to calibrate meter. | Use fresh 0.0 standard in clean tube. Reset meter to factory default settings. Recalibrate meter. |

COD3 Plus

Colorimeter

Test Procedures

Firmware Version 2.10 and greater

COD3 Plus COLORIMETER REAGENT SYSTEMS

COD3 Plus REAGENT SYSTEMS LIST

LaMotte Company continuously updates the list of pre-programmed tests as the calibrations become available.

Call LaMotte Technical Services at 1-800-344-3100 (410-778-3100 outside the USA) or email at tech@lamotte.com for a current list of available calibrations and more information.

| Test Factor [Test #] | Range [ppm] | MDL | Test Method [# of Reagents] | # of Tests |
|---|-------------|------|----------------------------------|------------|
| Ammonia Nitrogen-Low Range, Fresh Water [006] | 0.00-1.00 | 0.05 | Salicylate [3] | 25 |
| Ammonia Nitrogen-Low Range, Salt Water [007] | 0.00-1.00 | 0.10 | Salicylate [3] | 25 |
| Ammonia Nitrogen-High Range [008] | 0.00-4.00 | 0.05 | Nesslerization [2] | 50 |
| Boron [014] | 0.00-0.80 | 0.05 | Azomethine-H [2] | 50 |
| Cobalt [029] | 0.00-2.00 | 0.04 | PAN [3] | 50 |
| COD-Low Range [031] | 0-150 | 5 | Digestion [1] | 25 |
| COD-Standard Range [032] | 0-1500 | 50 | Digestion [1] | 25 |
| COD-High Range [030] | 0-15000 | 500 | Digestion [1] | 25 |
| Color [033] | 0-1000 cu | 20 | Platinum Cobalt [0] | - |
| Copper-Cuprizone [035] | 0.00-2.00 | 0.03 | Cuprizone [2] | 50 |
| Copper-Thiocarbamate [036] | 0.00-6.00 | 0.10 | Diethyldithiocarbamate [1] | 50 |
| Cyanuric Acid [039] | 5-200 | 10 | Melamine [1] | 50 |
| Dissolved Oxygen [043] | 0.0-11.0 | 0.6 | Winkler Colorimetric [3] | 100 |
| Fluoride [045] | 0.0-2.0 | 0.1 | SPADNS [2] | 50 |
| Hydrazine [049] | 0.00-1.00 | 0.01 | P-dimethyl-aminobenzaldehyde [2] | 50 |
| Molybdenum-High Range [063] | 0.0-50.0 | 0.6 | Thioglycolate [3] | 50 |
| Nickel [064] | 0.00-8.00 | 0.15 | Dimethylglyoxime [6] | 50 |
| Ozone-Low Range [071] | 0.00-0.40 | 0.01 | Indigo Trisulfonate [3] | 100 |
| Ozone-High Range [072] | 0.00-2.50 | 0.05 | Indigo Trisulfonate [3] | 20 |
| Phosphate-Low Range [081] | 0.00-3.00 | 0.05 | Ascorbic Acid Reduction [2] | 50 |
| Phosphate-High Range [080] | 0.0-70.0 | 0.5 | Vanodomolybd-phosphoric Acid [1] | 25 |
| Potassium [085] | 0.0-10.0 | 0.8 | Tetraphenylboron [2] | 100 |
| Silica-Low Range [087] | 0.00-4.00 | 0.05 | Heteropoly Blue [4] | 50 |
| Silica-High Range [086] | 0-75 | 1 | Silicomolybdate [3] | 50 |
| Sulfate-High Range [089] | 0-100 | 3 | Barium Chloride [1] | 50 |

| Test Factor [Test #] | Range [ppm] | MDL | Test Method [# of Reagents] | # of Tests |
|-------------------------|-------------|------|------------------------------------|------------|
| Sulfide-Low Range [090] | 0.00-1.50 | 0.06 | Methylene Blue [3] | 50 |
| Tannin [093] | 0.0-10.0 | 0.1 | Tungsto-molybdophosphoric Acid [2] | 50 |
| Turbidity [095] | 0-400 FAU | 3 | Absorption [0] | - |
| Zinc-Low Range [097] | 0.00-3.00 | 0.05 | Zincon [6] | 50 |

AMMONIA NITROGEN - LOW RANGE

SALICYLATE METHOD · CODE 3659-01-SC

| QUANTITY | CONTENTS | CODE |
|----------|-------------------------------|---------|
| 60 mL | *Salicylate Ammonia #1 | *3978-H |
| 10 g | *Salicylate #2 Reagent | *7457-D |
| 2 x 5 g | *Salicylate #3 Reagent Powder | *7458-C |
| 1 | Spoon, 0.1 g, plastic | 0699 |
| 1 | Spoon, 0.15 g, plastic | 0727 |
| 1 | Pipet, 1.0 mL, plastic | 0354 |

***WARNING:** Reagents marked with an * are considered to be potential health hazards. To view or print a Safety Data Sheet (SDS) for these reagents go to www.lamotte.com. Search for the four digit reagent code number listed on the reagent label, in the contents list or in the test procedures. Omit any letter that follows or precedes the four digit code number. For example, if the code is 4450WT-H, search 4450. To obtain a printed copy, contact LaMotte by email, phone or fax.

Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

Ammonia nitrogen is present in various concentrations in many surface and ground water supplies. Any sudden change in the concentration of ammonia nitrogen in a water supply is cause for suspicion. A product of microbiological activity, ammonia nitrogen is sometimes accepted as chemical evidence of pollution when encountered in natural waters.

Ammonia is rapidly oxidized in natural water systems by special bacterial groups that produce nitrite and nitrate. This oxidation requires that dissolved oxygen be available in the water. Ammonia is an additional source of nitrogen as a nutrient which may contribute to the expanded growth of undesirable algae and other forms of plant growth that overload the natural system and cause pollution.

| | |
|-------------------------------|--|
| APPLICATION: | Low concentrations of ammonia in fresh, brackish and salt water; fresh and salt water aquariums. |
| RANGE: | 0.00 - 1.00 ppm Ammonia-Nitrogen |
| MDL: | 0.05 ppm Fresh Water 0.10 ppm Salt Water |
| METHOD: | Salicylate and ammonia react at high pH in the presence of a chlorine donor and an iron catalyst to form a blue indophenol dye, the concentration of which is proportional to the ammonia concentration in the sample. |
| SAMPLE HANDLE & PRESERVATION: | Ammonia solutions tend to be unstable and should be analyzed immediately. Samples may be stored for 24 hours at 4°C or 28 days at -20°C. |
| INTERFERENCES: | There are few interferences in most natural waters. High concentrations of reducing agents, such as hydrazine, react with the chlorine donor and can result in negative interferences. Color and turbidity can also interfere. |

PROCEDURE - FRESH WATER

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** [or another sequence containing **006 Ammonia-N LRF**] from **Testing Menu**.
4. Scroll to and select **006 Ammonia-N LRF** from menu.
5. Rinse a clean tube [0290] with sample water. Fill to the 10 mL line with sample.
6. Insert tube into chamber, close lid and select **Scan Blank**. [See Note.]
7. Remove tube from colorimeter. Use the 1.0 mL plastic pipet [0354] to add 2.0 mL of *Salicylate Ammonia #1 [3978]. Cap and mix.
8. Use the 0.15 g spoon [0727] to add two measures of *Salicylate #2 Reagent [7457]. Cap and mix until dissolved. Wait 1 minute.
9. At end of 1 minute waiting period use 0.1 g spoon [0699] to add two measures of *Salicylate #3 Reagent Powder [7458]. Cap and shake vigorously for at least 30 seconds and all solid has dissolved. Wait 12 minutes for maximum color development.
10. At the end of the 12 minute waiting period, immediately mix and insert tube into chamber, close lid and select **Scan Sample**. Record result.
11. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

CALCULATIONS:

To express results as Unionized Ammonia [NH_3]:

$$\begin{aligned} \text{ppm Unionized Ammonia } [\text{NH}_3] &= \\ \text{ppm Ammonia-Nitrogen } [\text{NH}_3\text{-N}] &\times 1.2 \end{aligned}$$

To express results as Ionized Ammonia [NH_4]:

$$\begin{aligned} \text{ppm Ionized Ammonia } [\text{NH}_4^+] &= \\ \text{ppm Ammonia-Nitrogen } [\text{NH}_3\text{-N}] &\times 1.3 \end{aligned}$$

To determine the percentages of Unionized and Ionized Ammonia-Nitrogen, consult the Appendix.

NOTE: It is strongly suggested that a reagent blank be determined to account for any contribution to the test result by the reagent system. To determine the reagent blank, follow the above test procedure to scan a distilled or deionized water blank. Then follow the above procedure to perform the test on a distilled or deionized water sample. This test result is the reagent blank. Subtract the reagent blank from all

subsequent test results of unknown samples. It is necessary to determine the reagent blank only when a new lot number of reagents is obtained.

PROCEDURE - SALT WATER

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** [or another sequence containing **007 Ammonia-N LRS**] from Testing Menu.
4. Scroll to and select **007 Ammonia-N LRS** from menu.
5. Rinse a clean tube [0290] with sample water. Fill to the 10 mL line with sample.
6. Insert tube into chamber, close lid and select **Scan Blank**. [See Note.]
7. Remove tube from colorimeter. Use the 1.0 mL plastic pipet [0354] to add 2.0 mL of *Salicylate Ammonia #1 [3978]. Cap and mix.
8. Use the 0.15 g spoon [0727] to add two measures of *Salicylate #2 Reagent [7457]. Cap and mix until dissolved. Wait 1 minute.
9. At end of 1 minute waiting period use 0.1 g spoon [0699] to add two measures of *Salicylate #3 Reagent Powder [7458]. Cap and shake vigorously for at least 30 seconds and all solid has dissolved. Wait 20 minutes for maximum color development.
10. At the end of the 20 minute waiting period, immediately mix and insert tube into chamber, close lid and select **Scan Sample**. Record result.
11. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

CALCULATIONS:

To express results as Unionized Ammonia [NH_3]:

$$\begin{aligned} \text{ppm Unionized Ammonia } [\text{NH}_3] &= \\ \text{ppm Ammonia-Nitrogen } [\text{NH}_3\text{-N}] &\times 1.2 \end{aligned}$$

To express results as Ionized Ammonia [NH_4]:

$$\begin{aligned} \text{ppm Ionized Ammonia } [\text{NH}_4^+] &= \\ \text{ppm Ammonia-Nitrogen } [\text{NH}_3\text{-N}] &\times 1.3 \end{aligned}$$

To determine the percentages of Unionized and Ionized Ammonia-Nitrogen, consult the Appendix.

NOTE: It is strongly suggested that a reagent blank be determined to account for any contribution to the test result by the reagent system. To determine the reagent

blank, follow the above test procedure to scan a distilled or deionized water blank. Then follow the above procedure to perform the test on a distilled or deionized water sample. This test result is the reagent blank. Subtract the reagent blank from all subsequent test results of unknown samples. It is necessary to determine the reagent blank only when a new lot number of reagents is obtained.

AMMONIA NITROGEN - HIGH RANGE

NESSLERIZATION METHOD · CODE 3642-SC

| QUANTITY | CONTENTS | CODE |
|-----------|------------------------------|-----------|
| 30 mL | Ammonia Nitrogen Reagent #1 | V-4797-G |
| 2 x 30 mL | *Ammonia Nitrogen Reagent #2 | *V-4798-G |
| 1 | Pipet, 1 mL, plastic | 0354 |

***WARNING:** Reagents marked with an * are considered to be potential health hazards. To view or print a Safety Data Sheet [SDS] for these reagents go to www.lamotte.com. Search for the four digit reagent code number listed on the reagent label, in the contents list or in the test procedures. Omit any letter that follows or precedes the four digit code number. For example, if the code is 4450WT-H, search 4450. To obtain a printed copy, contact LaMotte by email, phone or fax.

Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

Ammonia nitrogen is present in various concentrations in many surface and ground water supplies. Any sudden change in the concentration of ammonia nitrogen in a water supply is cause for suspicion. A product of microbiological activity, ammonia nitrogen is sometimes accepted as chemical evidence of pollution when encountered in natural waters.

Ammonia is rapidly oxidized in natural water systems by special bacterial groups that produce nitrite and nitrate. This oxidation requires that dissolved oxygen be available in the water. Ammonia is an additional source of nitrogen as a nutrient which may contribute to the expanded growth of undesirable algae and other forms of plant growth that overload the natural system and cause pollution.

| | |
|---------------------------------|--|
| APPLICATION: | Drinking, surface, and saline waters; domestic and industrial wastes. |
| RANGE: | 0.00–4.00 ppm Ammonia Nitrogen |
| MDL: | 0.05 ppm |
| METHOD: | Ammonia forms a colored complex with Nessler's Reagent in proportion to the amount of ammonia present in the sample. Rochelle salt is added to prevent precipitation of calcium or magnesium in undistilled samples. |
| SAMPLE HANDLING & PRESERVATION: | Ammonia solutions tend to be unstable and should be analyzed immediately. Sample may be stored for 24 hours at 4°C or 28 days at –20°C. |
| INTERFERENCES: | Sample turbidity and color may interfere. Turbidity may be removed by a filtration procedure. Color interference may be eliminated by blanking the instrument with a sample blank. |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Scroll to and select **All Tests** [or another sequence containing **008 Ammonia-N HR**] from **Testing Menu**.
4. Scroll to and select **008 Ammonia-N HR** from menu.
5. Rinse a clean tube [0290] with sample water. Fill to the 10 mL line with sample.
6. Insert tube into chamber, close lid and select **Scan Blank**. [See Note]
7. Remove tube from colorimeter. Add 8 drops of Ammonia Nitrogen Reagent #1 [V-4797]. Cap and mix. Wait 1 minute.
8. Use the 1.0 mL pipet [0354] to add 1.0 mL of *Ammonia Nitrogen Reagent #2 [V-4798]. Cap and mix. Allow 5 minutes for maximum color development.
9. At end of the 5 minute waiting period, immediately mix, insert tube into chamber, close lid and select **Scan Sample**. Record result.
10. Press  to turn the colorimeter off or press the  exit to a previous menu or make another menu selection.

CALCULATIONS:

To express results as Unionized Ammonia [NH_3]:

$$\begin{aligned} \text{ppm Unionized Ammonia } [\text{NH}_3] &= \\ \text{ppm Ammonia-Nitrogen } [\text{NH}_3\text{-N}] &\times 1.2 \end{aligned}$$

To express results as Ionized Ammonia [NH_4]:

$$\begin{aligned} \text{ppm Ionized Ammonia } [\text{NH}_4^+] &= \\ \text{ppm Ammonia-Nitrogen } [\text{NH}_3\text{-N}] &\times 1.3 \end{aligned}$$

To determine the percentages of Unionized and Ionized Ammonia-Nitrogen, consult the Appendix.

NOTE: It is strongly suggested that a reagent blank be determined to account for any contribution to the test result by the reagent system. To determine the reagent blank, follow the above test procedure to scan a distilled or deionized water blank. Then follow the above procedure to perform the test on a distilled or deionized water sample. This test result is the reagent blank. Subtract the reagent blank from all subsequent test results of unknown samples. It is necessary to determine the reagent blank only when a new lot number of reagents is obtained.

BORON

AZOMETHINE-H METHOD · CODE 4868-01

| QUANTITY | CONTENTS | CODE |
|----------|-----------------------------|---------|
| 120 mL | *Boron Buffer | *4869-J |
| 10 g | *Boron Indicator Powder | *4870-D |
| 1 | Pipet, plastic, 1.0 mL | 0354 |
| 1 | Spoon, 0.15 g | 0727 |
| 1 | Dark storage chamber, brown | 0108 |

***WARNING:** Reagents marked with an * are considered to be potential health hazards. To view or print a Safety Data Sheet [SDS] for these reagents go to www.lamotte.com. Search for the four digit reagent code number listed on the reagent label, in the contents list or in the test procedures. Omit any letter that follows or precedes the four digit code number. For example, if the code is 4450WT-H, search 4450. To obtain a printed copy, contact LaMotte by email, phone or fax.

Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

Small amounts of boron are necessary for plant growth but large amounts can be toxic. In humans, boron aids in the uptake of calcium and the production of strong bones. An excess of boron can affect the central nervous system resulting in a syndrome known as borism. Some natural waters may contain small amounts of boron. Large concentrations may be due to industrial effluent entering waterways. Boron compounds are used in cleaning compounds, paper and paints, fertilizers, glass and ceramics, fire retardants and the production of alloys. In the atomic energy field, boron is a component of neutron shields and nuclear reactors. Some swimming pools use boron buffering systems.

| | |
|---------------------------------|--|
| APPLICATION: | Surface and saline waters, hydroponic solutions, industrial waste, swimming pools. |
| RANGE: | 0.00–0.80 ppm Boron |
| MDL: | 0.05 |
| METHOD: | Azomethine-H and borate form a yellow complex at pH 6 in proportion to the concentration of boron present. |
| SAMPLE HANDLING & PRESERVATION: | Store samples in polyethylene bottles. Do not use borate detergents or glassware. |
| INTERFERENCES: | Interferences in drinking water are unlikely. Manganese, zirconium, chromium, titanium, copper, vanadium, aluminum, beryllium and iron may cause high results. |

PROCEDURE

1. This test requires a Reagent Blank. Rinse a tube [0290] with clear, colorless, boron free water. Fill to 10 mL line with clear, colorless, boron free water.
2. Use the 1.0 mL pipet [0354] to add 2 mL of *Boron Buffer [4869]. Cap and mix.
3. Use the 0.15 g spoon [0727] to add one level measure of *Boron Indicator Powder [4870]. Press full spoon against side of jar to compress powder. Scrape off excess powder on inside neck of bottle. Tap excess off spoon handle.
4. Cap and shake vigorously for 30 seconds.
5. Insert the tube into meter chamber. Close lid.
6. Start a timer set for 30 minutes. Do not open the lid during the waiting time. The reaction is photosensitive.
7. Rinse a clean tube [0290] with Sample Water. Fill to the 10 mL line with sample water. Repeat steps 2–4.
8. Insert the tube into the Dark Storage Chamber [0108]. Close top.
9. Start a second timer set for 30 minutes. Do not open the chamber during the waiting time. The reaction is photosensitive.
10. When 2 minutes remain on the first timer [Reagent Blank], press and hold ON button until colorimeter turns on.
11. Press and hold  until colorimeter turns on.
12. Press  to select **Testing Menu**.
13. Select **All Tests** [or another sequence containing **014 Boron**] from **Testing Menu**.
14. Scroll to and select **014 Boron** from menu. At the end of the Reagent Blank 30 minute waiting period, remove Reagent Blank tube from meter chamber. Invert several times to mix.
15. Insert the tube into meter chamber, close lid and select **Scan Blank**.
16. Remove the tube from colorimeter.
17. At the end of the Sample Water 30 minute waiting period, remove Sample Water tube from Dark Storage Chamber. Invert several times to mix.
18. Insert tube into meter chamber, close lid and select **Scan Sample**. Record result in ppm boron.
19. Press  to turn colorimeter off or press the  to exit to a previous menu or make another menu selection.

COBALT

PAN METHOD · CODE 4851-01

| QUANTITY | CONTENTS | CODE |
|----------|---------------------------|---------|
| 60 mL | *Cobalt Buffer | *4852-H |
| 60 mL | *Cobalt Indicator Reagent | *4853-H |
| 30 mL | Stabilizer Solution | 4854-G |
| 2 | Pipet, 1.0 mL, plastic | 0354 |
| 1 | Pipet, 0.5 mL, plastic | 0353 |

***WARNING:** Reagents marked with an * are considered to be potential health hazards. To view or print a Safety Data Sheet [SDS] for these reagents go to www.lamotte.com. Search for the four digit reagent code number listed on the reagent label, in the contents list or in the test procedures. Omit any letter that follows or precedes the four digit code number. For example, if the code is 4450WT-H, search 4450. To obtain a printed copy, contact LaMotte by email, phone or fax.

Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

Cobalt rarely occurs in natural water. It is used in the manufacture of alloys to increase corrosion resistance and strength. It is found in wastewaters as a corrosion by-product.

| | |
|---------------------------------|--|
| APPLICATION: | Industrial wastewater. |
| RANGE: | 0.00–2.00 ppm Cobalt |
| MDL: | 0.04 ppm |
| METHOD: | PAN [1-[2-Pyridylazo]-2-Naphthol] forms a greenish complex with Cobalt [Co ⁺²] at a pH of 5. |
| SAMPLE HANDLING & PRESERVATION: | Store samples in acid-washed plastic bottles. Adjust pH to less than 2 with nitric acid. Adjust sample pH to 5 before testing. |
| INTERFERENCES: | Iron [+2] and high concentrations of heavy metals. |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** (or another sequence containing **029 Cobalt**) from **Testing Menu**.
4. Scroll to and select **029 Cobalt** from menu.
5. Rinse a tube (0290) with sample water. Fill to 10 mL with sample.
6. Insert the tube into chamber, close lid and select **Scan Blank**.
7. Remove the tube from colorimeter.
8. Use the 1.0 mL pipet (0354) to add 1 mL of *Cobalt Buffer (4852). Cap and mix.
9. Use the other 1.0 mL pipet (0354) to add 1 mL of *Cobalt Indicator Reagent (4853). Cap and mix.
10. Wait 3 minutes.
11. Use the 0.5 mL pipet (0353) to add 0.5 mL Stabilizer Solution (4854). Cap and invert 15 times to thoroughly mix.
12. Wait 5 minutes. DO NOT MIX.
13. Insert tube into chamber, close lid and select **Scan Sample**. Record result in ppm cobalt.
14. Press  to turn the colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTE: For best possible results, a reagent blank should be determined to account for any contribution to the test result by the reagent system. To determine the reagent blank, follow the above test procedure to scan a distilled or deionized water blank. Then follow the above procedure to perform the test on a distilled or deionized water sample. This test result is the reagent blank. Subtract the reagent blank from all subsequent test results of unknown samples. It is necessary to determine the reagent blank only when a new lot number of reagents is obtained.

COD – LOW RANGE

MERCURY FREE DIGESTION METHOD · CODE 0072-SC MERCURY DIGESTION METHOD · CODE 0075-SC

| QUANTITY | CONTENTS | CODE |
|----------|-----------------------------------|----------|
| 25 | *COD Low Range Mercury Free Tubes | *0072-SC |
| or 25 | *COD Low Range Mercury Tubes | *0075-SC |

***WARNING:** Reagents marked with an * are considered to be potential health hazards. To view or print a Safety Data Sheet [SDS] for these reagents go to www.lamotte.com. Search for the four digit reagent code number listed on the reagent label, in the contents list or in the test procedures. Omit any letter that follows or precedes the four digit code number. For example, if the code is 4450WT-H, search 4450. To obtain a printed copy, contact LaMotte by email, phone or fax.

Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

COD Low Range Mercury Free Tubes are not USEPA approved.

COD Low Range Mercury Tubes are USEPA approved.

Equipment needed but not supplied:

| | | |
|------|----------------------------|------------|
| 1 | COD Adapter | 5-0087 |
| 1 | COD Reactor, 12 tube, 110V | 5-0102 |
| or 1 | COD Reactor, 12 tube, 230V | 5-0102-EX2 |
| 1 | Measuring Pipet, 1.0 mL | 2-2110 |
| 1 | Pipet Bulb | 2-2164 |

Chemical Oxygen Demand (COD) is a measure of the amount of organic matter in water which is susceptible to oxidation by chemical oxidants. COD can be empirically related to the Biological Oxygen Demand (BOD) and organic carbon content of a specific source of water. This correlation must be determined experimentally for each source of water.

| | |
|---------------------------------|--|
| APPLICATION: | Domestic and industrial wastes. |
| RANGE: | 0–150 mg/L COD |
| MDL: | 7.5 mg/L |
| METHOD: | Dichromate in the presence of silver salts, at high temperature in a closed system, oxidizes most organic compounds to 95–100% of the theoretical amount. This process is called digestion. As dichromate oxidizes the organic compounds, the amount of yellow color is reduced. The remaining yellow color is measured colorimetrically at the 420 nm and is directly proportional to the COD of the sample. |
| SAMPLE HANDLING & PRESERVATION: | Collect samples in glass and test as soon as possible. If samples must be stored, preservation is accomplished by the addition of concentrated H ₂ SO ₄ to adjust the pH below 2. Samples with suspended solids should be homogenized in a blender [100 mL for 30 seconds] and then stirred gently with a magnetic stirrer. |
| INTERFERENCES: | <p>Volatile organic compounds are not oxidized to the extent that they are in the vapor above the digestion solution. Therefore, they do not contribute to the COD reading. Chloride concentrations above 10% of COD interfere with the mercury free tubes. Chloride above 2000 ppm will interfere with the mercury tubes. Nitrite gives a positive interference of 1.1 ppm O₂ per ppm NO₂-N which is insignificant unless nitrite concentrations are very high. Other reduced inorganic compounds are stoichiometrically oxidized, causing a positive interference. Corrections can be made for these compounds based upon their stoichiometry and concentrations.</p> <p>When scanning samples in 16 mm tubes, such as COD, the sample chamber lid can not be closed. Use the COD adapter to minimize stray light interference. To further reduce stray light interference, do not Scan Sample in direct sunlight.</p> |

PROCEDURE

Use COD/UDV adapter.

1. Homogenize sample if necessary.
2. Preheat COD heater block to $150\pm 2^{\circ}\text{C}$.
3. Remove cap from COD tube. Hold tube at a 45° angle. Use a volumetric pipet, to carefully add 2.0 mL sample water allowing the sample to run down the side of the tube.
4. Cap and mix thoroughly.
5. Rinse the outside of the tube with distilled water. Wipe dry with a paper towel.
6. Repeat steps 3 through 5 using 2.0 mL distilled water. This is the reagent blank.
7. Place tubes in preheated COD block heater and maintain temperature at $150\pm 2^{\circ}\text{C}$ for two hours.
8. At the end of the heating period turn the heater off. Wait 20 minutes for the tubes to cool to 120°C or less.
9. Remove tubes from block heater. Invert several times to mix.
10. Allow to cool to room temperature.
11. Press and hold  until colorimeter turns on.
12. Press  to select **Testing Menu**.
13. Select **All Tests** [or another sequence containing **031 COD LR**] from **Testing Menu**.
14. Scroll to and select **031 COD LR** from menu.
15. Wipe the blank tube with a damp towel to remove fingerprints and smudges. Wipe with a dry towel.
16. Insert reagent blank tube into chamber. Select **Scan Blank**.
17. Remove tube from colorimeter.
18. Insert digested water sample tube into chamber. Select **Scan Sample**. Record result. For the most accurate results, take three readings on each sample and average the results.
19. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTES: Reagents are light sensitive. Unused reagents should be stored in the shipping container, and in the refrigerator if possible, until needed.

A reagent blank should be run with each set of samples and with each lot of reagents.

The reacted blank will be stable if stored in the dark.

To eliminate error caused by contamination, wash all glassware with 20% sulfuric acid.

For greater accuracy, a minimum of three repetitions should be performed and the results averaged.

Some samples may be digested completely in less than two hours. The concentration may be measured at 15 minute intervals while the vials are still hot until the reading remains unchanged. The vials should be cooled to room temperature before the final measurement is taken.

COD – STANDARD RANGE

MERCURY FREE DIGESTION METHOD · CODE 0073-SC MERCURY DIGESTION METHOD · CODE 0076-SC

| QUANTITY | CONTENTS | CODE |
|----------|--|----------|
| 25 | *COD Standard Range Mercury Free Tubes | *0073-SC |
| or 25 | *COD Standard Range Mercury Tubes | *0076-SC |

***WARNING:** Reagents marked with an * are considered to be potential health hazards. To view or print a Safety Data Sheet [SDS] for these reagents go to www.lamotte.com. Search for the four digit reagent code number listed on the reagent label, in the contents list or in the test procedures. Omit any letter that follows or precedes the four digit code number. For example, if the code is 4450WT-H, search 4450. To obtain a printed copy, contact LaMotte by email, phone or fax.

Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

COD Standard Range Mercury Free Tubes are not USEPA approved.

COD Standard Range Mercury Tubes are USEPA approved.

Equipment needed but not supplied:

| | | |
|------|----------------------------|------------|
| 1 | COD Adapter | 5-0087 |
| 1 | COD Reactor, 12 tube, 110V | 5-0102 |
| or 1 | COD Reactor, 12 tube, 230V | 5-0102-EX2 |
| 1 | Measuring Pipet, 1.0 mL | 2-2110 |
| 1 | Pipet Bulb | 2-2164 |

Chemical Oxygen Demand (COD) is a measure of the amount of organic matter in water which is susceptible to oxidation by chemical oxidants. COD can be empirically related to the Biological Oxygen Demand (BOD) and organic carbon content of a specific source of water. This correlation must be determined experimentally for each source of water.

| | |
|---------------------------------|--|
| APPLICATION: | Domestic and industrial wastes. |
| RANGE: | 0–1500 mg/L COD |
| MDL: | 50 mg/L |
| METHOD: | Dichromate in the presence of silver salts, at high temperature in a closed system, oxidizes most organic compounds to 95–100% of the theoretical amount. This process is called digestion. As dichromate oxidizes the organic compounds, a green complex is formed. The concentration of the green complex is measured at 605 nm and is directly proportional to the COD of the sample. |
| SAMPLE HANDLING & PRESERVATION: | Collect samples in glass and test as soon as possible. If samples must be stored, preservation is accomplished by the addition of concentrated H_2SO_4 to adjust the pH below 2. Samples with suspended solids should be homogenized in a blender (100 mL for 30 seconds) and then stirred gently with a magnetic stirrer. |
| INTERFERENCES: | <p>Volatile organic compounds are not oxidized to the extent that they are in the vapor above the digestion solution. Therefore, they do not contribute to the COD reading. Chloride concentrations above 10% of COD interfere with the mercury free tubes. Chloride above 2000 ppm will interfere with the mercury tubes. Nitrite gives a positive interference of 1.1 ppm O_2 per ppm NO_2-N which is insignificant unless nitrite concentrations are very high. Other reduced inorganic compounds are stoichiometrically oxidized, causing a positive interference. Corrections can be made for these compounds based upon their stoichiometry and concentrations.</p> <p>When scanning samples in 16 mm tubes, such as COD, the sample chamber lid can not be closed. Use the COD adapter to minimize stray light interference. To further reduce stray light interference, do not Scan Sample in direct sunlight.</p> |

PROCEDURE

Use COD/UDV adapter.

1. Homogenize sample if necessary.
2. Preheat COD heater block to $150\pm 2^{\circ}\text{C}$.
3. Remove cap from COD tube. Hold tube at a 45° angle. Use a volumetric pipet, to carefully add 2.0 mL sample water allowing the sample to run down the side of the tube.
4. Cap and mix thoroughly.
5. Rinse the outside of the vial with distilled water. Wipe dry with a paper towel.
6. Repeat steps 2 through 5 using 2.0 mL distilled water. This is the reagent blank.
7. Place tubes in preheated COD block heater and maintain temperature at $150\pm 2^{\circ}\text{C}$ for two hours.
8. At the end of the heating period turn the heater off. Wait 20 minutes for the tubes to cool to 120°C or less.
9. Remove tubes from block heater. Invert several times to mix.
10. Allow to cool to room temperature.
11. Press and hold  until colorimeter turns on.
12. Press  to select **Testing Menu**.
13. Select **All Tests** [or another sequence containing **032 COD SR**] from **Testing Menu**.
14. Wipe the blank tube with a damp towel to remove fingerprints and smudges. Wipe with a dry towel.
15. Scroll to and select **032 COD SR** from menu.
16. Insert reagent blank tube into chamber. Select **Scan Blank**.
17. Remove tube from colorimeter.
18. Insert digested water sample tube into chamber. Select **Scan Sample**. Record result. For the most accurate results, take three readings on each sample and average the results.
19. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTES: Reagents are light sensitive. Unused reagents should be stored in the shipping container, and in the refrigerator if possible, until needed.

A reagent blank should be run with each set of samples and with each lot of reagents.

The reacted blank will be stable if stored in the dark.

To eliminate error caused by contamination, wash all glassware with 20% sulfuric acid.

For greater accuracy, a minimum of three repetitions should be performed and the results averaged.

Some samples may be digested completely in less than two hours. The concentration may be measured at 15 minute intervals while the vials are still hot until the reading remains unchanged. The vials should be cooled to room temperature before the final measurement is taken.

COD – HIGH RANGE

MERCURY FREE DIGESTION METHOD · CODE 0074-SC MERCURY DIGESTION METHOD · CODE 0077-SC

| QUANTITY | CONTENTS | CODE |
|----------|------------------------------------|----------|
| 25 | *COD High Range Mercury Free Tubes | *0074-SC |
| or 25 | *COD High Range Mercury Tubes | *0077-SC |

***WARNING:** Reagents marked with an * are considered to be potential health hazards. To view or print a Safety Data Sheet [SDS] for these reagents go to www.lamotte.com. Search for the four digit reagent code number listed on the reagent label, in the contents list or in the test procedures. Omit any letter that follows or precedes the four digit code number. For example, if the code is 4450WT-H, search 4450. To obtain a printed copy, contact LaMotte by email, phone or fax.

Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

COD High Range Mercury Free Tubes and COD High Range Mercury Tubes are not USEPA approved.

Equipment needed but not supplied:

| | | |
|------|----------------------------|------------|
| 1 | COD Adapter | 5-0087 |
| 1 | COD Reactor, 12 tube, 110V | 5-0102 |
| or 1 | COD Reactor, 12 tube, 230V | 5-0102-EX2 |
| 1 | Measuring Pipet, 1.0 mL | 2-2110 |
| 1 | Pipet Bulb | 2-2164 |

Chemical Oxygen Demand (COD) is a measure of the amount of organic matter in water which is susceptible to oxidation by chemical oxidants. COD can be empirically related to the Biological Oxygen Demand (BOD) and organic carbon content of a specific source of water. This correlation must be determined experimentally for each source of water.

| | |
|--------------------------------|--|
| APPLICATION: | Domestic and industrial wastes. |
| RANGE: | 0–15000 mg/L COD |
| MDL: | 500 mg/L |
| METHOD: | Dichromate in the presence of silver salts, at high temperature in a closed system, oxidizes most organic compounds to 95–100% of the theoretical amount. This process is called digestion. As dichromate oxidizes the organic compounds, a green complex is formed. The concentration of the green complex is measured at 605 nm and is directly proportional to the COD of the sample. |
| SAMPLE HANDLING & RESERVATION: | Collect samples in glass and test as soon as possible. If samples must be stored, preservation is accomplished by the addition of concentrated H ₂ SO ₄ to adjust the pH below 2. Samples with suspended solids should be homogenized in a blender (100 mL for 30 seconds) and then stirred gently with a magnetic stirrer. |
| INTERFERENCES: | <p>Volatile organic compounds are not oxidized to the extent that they are in the vapor above the digestion solution. Therefore, they do not contribute to the COD reading. Contains mercury sulfate to prevent interference from chloride. Nitrite gives a positive interference of 1.1 ppm O₂ per ppm NO₂-N, which is insignificant unless nitrite concentrations are very high. Other reduced inorganic compounds are stoichiometrically oxidized, causing a positive interference. Corrections can be made for these compounds based upon their stoichiometry and concentrations.</p> <p>When scanning samples in 16 mm tubes, such as COD, the sample chamber lid can not be closed. Use the COD adapter to minimize stray light interference. To further reduce stray light interference, do not Scan Sample in direct sunlight.</p> |

PROCEDURE

Use COD/UDV adapter.

1. Homogenize sample if necessary.
2. Preheat COD heater block to $150\pm 2^{\circ}\text{C}$.
3. Remove cap from COD tube. Hold tube at a 45° angle. Use a graduated pipet, to carefully add 0.2 mL sample water allowing the sample to run down the side of the tube.
4. Cap and mix thoroughly.
5. Rinse the outside of the tube with distilled water. Wipe dry with a paper towel.
6. Repeat steps 3 through 5 using 0.2 mL distilled water. This is the reagent blank.
7. Place tubes in preheated COD block heater and maintain temperature at $150\pm 2^{\circ}\text{C}$ for two hours.
8. At the end of the heating period turn the heater off. Wait 20 minutes for the tubes to cool to 120°C or less.
9. Remove tubes from block heater. Invert several times to mix.
10. Allow to cool to room temperature.
11. Press and hold  until colorimeter turns on.
12. Press  to select **Testing Menu**.
13. Select **All Tests** [or another sequence containing **030 COD HR**] from Testing Menu menu.
14. Wipe the blank tube with a damp towel to remove fingerprints and smudges. Wipe with a dry towel.
15. Scroll to and select **030 COD HR** from menu.
16. Insert reagent blank tube into chamber. Select **Scan Blank**.
17. Remove tube from colorimeter.
18. Insert digested water sample tube into chamber. Select **Scan Sample**. Record result. For the most accurate results, take three readings on each sample and average the results.
19. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTES: Reagents are light sensitive. Unused reagents should be stored in the shipping container, and in the refrigerator if possible, until needed.

A reagent blank should be run with each set of samples and with each lot of reagents.

The reacted blank will be stable if stored in the dark.

To eliminate error caused by contamination, wash all glassware with 20% sulfuric acid.

For greater accuracy, a minimum of three repetitions should be performed and the results averaged.

COLOR

PLATINUM COBALT METHOD NO REAGENTS REQUIRED

Color in water may be attributed to humus, peat, plankton, vegetation, and natural metallic ions, such as iron and manganese, or industrial waste. Color is removed to make water suitable for domestic and industrial use. Color may have to be removed from industrial waste before it is discharged to a waterway.

| | |
|---------------------------------|---|
| APPLICATION: | Potable water and water with color due to natural materials. |
| RANGE: | 0–1000 color units |
| MDL: | 20 cu |
| METHOD: | Color is determined by a meter that has been calibrated with colored standards of known platinum cobalt concentration. True color, the color of water in which the turbidity has been removed, is measured. |
| SAMPLE HANDLING & PRESERVATION: | Collect all samples in clean glassware. Determine color as soon as possible to avoid biological or chemical changes that could occur in the sample during storage. |
| INTERFERENCES: | Turbidity will interfere. Filter before testing. |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** (or another sequence containing **033 Color**) from **Testing Menu**.
4. Scroll to and select **033 Color** from menu.
5. Rinse a tube (0290) with color-free water (distilled or deionized water). Fill to 10 mL line with color-free water.
6. Insert the tube into chamber, close lid and select **Scan Blank**.
7. Remove tube from colorimeter. Empty tube.
8. Rinse tube with sample water. Fill to 10 mL line with water sample.
9. Insert tube with sample water, close lid and select **Scan Sample**. Record result in color units.
10. Press  to turn the colorimeter off or press  to exit to a previous menu or make another menu selection.

COPPER

CUPRIZONE METHOD · CODE 4023

| QUANTITY | CONTENTS | CODE |
|----------|-----------|-----------|
| 15 mL | *Copper A | *P-6367-E |
| 15 mL | *Copper B | *P-6368-E |

***WARNING:** Reagents marked with an * are considered to be potential health hazards. To view or print a Safety Data Sheet [SDS] for these reagents go to www.lamotte.com. Search for the four digit reagent code number listed on the reagent label, in the contents list or in the test procedures. Omit any letter that follows or precedes the four digit code number. For example, if the code is 4450WT-H, search 4450. To obtain a printed copy, contact LaMotte by email, phone or fax.

Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

The copper content of drinking water generally falls below 0.03 parts per million, but copper levels as high as 1.0 part per million will give water a bitter taste. Waters testing as high as 1.0 part per million copper have probably been treated with a copper compound, like those used in the control of algae, or have become contaminated from untreated industrial wastes. The addition of copper sulfate to lakes causes an increase in the copper content of the sediments. Acid waters and those high in free carbon dioxide may cause the corrosion or “eating away” of copper, brass and bronze pipes and fittings. This corrosion results in the addition of copper to the water supply.

| | |
|---------------------------------|---|
| APPLICATION: | Drinking, surface, and domestic waters. Pools and spas. |
| RANGE: | 0.00–2.00 ppm Copper |
| MDL: | 0.03 ppm |
| METHOD: | Copper ions form a blue complex with cuprizone, in a 1 to 2 ratio, at a pH of about 8, in proportion to the concentration of copper in the sample. |
| SAMPLE HANDLING & PRESERVATION: | Copper has a tendency to be adsorbed to the surface of the sample container. Samples should be analyzed as soon as possible after collection. If storage is necessary, 0.5 mL of 20% hydrochloric acid per 100 mL of sample will prevent “plating out”. However, a correction must be made to bring the reaction into the optimum pH range. |
| INTERFERENCES: | Hg ⁺¹ at 1 ppm. Cr ⁺³ , Co ⁺² , and silicate at 10 ppm. As ⁺³ , Bi ⁺³ , Ca ⁺² , Ce ⁺³ , Ce ⁺⁴ , Hg ⁺² , Fe ⁺² , Mn ⁺² , Ni ⁺² and ascorbate at 100 ppm. Many other metal cations and inorganic anions at 1000 ppm. EDTA at all concentrations. |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** [or another sequence containing **035 Cu Cuprizone**] from **Testing Menu**.
4. Scroll to and select **035 Cu Cuprizone** from menu.
5. Rinse a tube [0290] with sample water. Fill to the 10 mL line with sample.
6. Insert the tube into chamber, close lid and select **Scan Blank**.
7. Remove tube from colorimeter and add 5 drops of *Copper A [6367]. Cap and mix.
8. Add 5 drops of *Copper B [6368]. Cap and mix.
9. Wait 5 minutes. Mix.
10. Insert tube into chamber, close lid and select **Scan Sample**. Record result.
11. Press  to turn the colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTE: For best possible results, a reagent blank should be determined to account for any contribution to the test result by the reagent system. To determine the reagent blank, follow the above test procedure to scan a distilled or deionized water blank. Then follow the above procedure to perform the test on a distilled or deionized water sample. This test result is the reagent blank. Subtract the reagent blank from all subsequent test results of unknown samples. It is necessary to determine the reagent blank only when a new lot number of reagents are obtained.

The reaction may stain the tubes. Scrub tubes thoroughly after each use.

COPPER

DIETHYLDITHIOCARBAMATE METHOD · CODE 3646-SC

| QUANTITY | CONTENTS | CODE |
|----------|-----------|---------|
| 15 mL | *Copper 1 | *6446-E |

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Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

The copper content of drinking water generally falls below 0.03 parts per million, but copper levels as high as 1.0 part per million will give water a bitter taste. Waters testing as high as 1.0 part per million copper have probably been treated with a copper compound, like those used in the control of algae, or have become contaminated from untreated industrial wastes. The addition of copper sulfate to lakes causes an increase in the copper content of the sediments. Acid waters and those high in free carbon dioxide may cause the corrosion or "eating away" of copper, brass and bronze pipes and fittings. This corrosion results in the addition of copper into the water supply.

| | |
|---------------------------------|---|
| APPLICATION: | Drinking, surface, and saline waters; domestic and industrial wastes. |
| RANGE: | 0.00–6.00 ppm Copper |
| MDL: | 0.10 ppm |
| METHOD: | Copper ions form a yellow colored chelate with diethyldithiocarbamate around pH 9-10 in proportion to the concentration of copper in the sample. |
| SAMPLE HANDLING & PRESERVATION: | Copper has a tendency to be adsorbed to the surface of the sample container. Samples should be analyzed as soon as possible after collection. If storage is necessary, 0.5 mL of 20% hydrochloric acid per 100 mL of sample will prevent "plating out." However, a correction must be made to bring the reaction into the optimum pH range. |
| INTERFERENCES: | Bismuth, cobalt, mercurous, nickel and silver ions and chlorine [6 ppm or greater] interfere and must be absent. |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** (or another sequence containing **036 Cu Thiocarbamate**) from **Testing Menu**.
4. Scroll to and select **036 Cu Thiocarbamate** from menu.
5. Rinse a clean tube (0290) with sample water. Fill to the 10 mL line with sample.
6. Insert tube into chamber, close lid and select **Scan Blank**.
7. Remove tube from colorimeter and add 5 drops of *Copper 1 (6446). Cap and mix. Solution will turn yellow if copper is present.
8. Insert tube into chamber, close lid and select **Scan Sample**. Record result.
9. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTE: The reaction may stain the tubes. Scrub the tubes thoroughly after each use.

CYANURIC ACID

MELAMINE METHOD-TURBIDITY · CODE 3661-01-SC

| QUANTITY | CONTENTS | CODE |
|------------|-----------------------------|--------|
| 2 x 100 mL | Cyanuric Acid Test Solution | 4856-J |
| 1 | Syringe, 5 mL | 0807 |

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Cyanuric acid is added to swimming pool water as a stabilizing agent for free chlorine residuals. It minimizes the loss of chlorine from the action of ultraviolet rays in sunlight. Cyanuric acid levels in pools should be maintained between 25 and 75 ppm and various public health associations recommend that the concentration should never exceed 100-150 ppm.

| | |
|---------------------------------|---|
| APPLICATION: | Swimming pool waters. |
| RANGE: | 5–200 ppm Cyanuric Acid |
| MDL: | 10 ppm |
| METHOD: | A buffered solution of melamine forms a precipitate with cyanuric acid in proportion to the amount of cyanuric acid present. The amount of particles in suspension is measured turbidimetrically. |
| SAMPLE HANDLING & PRESERVATION: | Cyanuric acid samples should be analyzed as soon as possible after collection. Deterioration of the sample can be minimized by keeping samples in the dark or refrigerated until analysis can be performed. |
| INTERFERENCES: | No known interference from compounds normally found in pool water. Temperature of the sample should be maintained between 70°F and 80°F for best results. Check for stray light interference [see p. 69]. |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** (or another sequence containing **039 Cyanuric Acid L**) from **Testing Menu**.
4. Scroll to and select **039 Cyanuric Acid L** from menu.
5. Rinse a clean tube [0290] with sample water. Fill to the 10 mL line with sample.
6. Insert tube into chamber, close lid and select **Scan Blank**.
7. Remove tube from colorimeter and pour out water. Use a graduated cylinder or similar to measure 5 mL of sample water and pour into colorimeter tube.
8. Use the 5 mL syringe [0807] to add 5 mL of Cyanuric Acid Test Solution [4856]. Cap and mix thoroughly. A precipitate will form if cyanuric acid is present. Wait 1 minute.

NOTE: This reagent bottle has a special fitting which enables the syringe to be inserted into the top of the bottle. With syringe in place, invert bottle and withdraw syringe plunger until 5 mL of reagent is contained in the syringe barrel. Remove syringe from reagent bottle and depress plunger to dispense into the tube.

9. At end of 1 minute waiting period, mix thoroughly, insert tube into chamber, close lid and select **Scan Sample**. Record result.
10. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTE: For the most accurate results, the sample and reagents should be at $25 \pm 4^{\circ}\text{C}$.

DISSOLVED OXYGEN

WINKLER COLORIMETRIC METHOD · CODE 3688-SC

| QUANTITY | CONTENTS | CODE |
|----------|----------------------------------|-----------|
| 30 mL | *Manganese Sulfate Solution | *4167-G |
| 30 mL | *Alkaline Potassium Iodide Azide | *7166-G |
| 30 mL | *Sulfuric Acid 1:1 | *6141WT-G |
| 1 | Sample Tube, screw cap | 29180 |
| 1 | Cap | 28570 |

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Dissolved oxygen is vital to the survival of aquatic organisms. Naturally present, dissolved oxygen enters the water when plants photosynthesize. Wind and wave action also cause oxygen from the air to dissolve into water. Dissolved oxygen is consumed by aquatic animals and by the oxidation, or chemical breakdown, of dead and decaying plants and animals. The concentration of dissolved oxygen in natural waters can range from 0 to 14 ppm and is effected by temperature and salinity.

| | |
|----------------|---|
| APPLICATION: | This method is applicable for the determination of dissolved oxygen in drinking water, all surface waters and wastewater. |
| RANGE: | 0.0–11.0 Dissolved Oxygen |
| MDL: | 0.6 ppm |
| METHOD: | This method uses the azide modification of the Winkler Method with a colorimetric determination of the yellow iodine produced from the reaction with the dissolved oxygen. |
| INTERFERENCES: | The presence of other oxidizing agents may cause positive interferences. Reducing may cause negative interferences. Nitrite interferences are eliminated with the azide modification. |

COLLECTION & TREATMENT OF THE WATER SAMPLE

Steps 1 through 4 below describe proper sampling technique in shallow water. For sample collection at depths beyond arm's reach, special water sampling apparatus is required [e.g. the LaMotte Water Sampling Chamber, Code 1060; Model JT-1 Water Samplers, Code 1077; Water Sampling Outfit, Code 3103; or Water Sampling Bottle, Code 3-0026].

1. To avoid contamination, thoroughly rinse the screw cap Sample Tube [29180] with sample water.
2. Tightly cap Sample Tube and submerge to the desired depth. Remove cap and allow the Sample Tube to fill.
3. Tap the sides of the submerged tube to dislodge any air bubbles clinging to the inside. Replace the cap while the Sample Tube is still submerged.
4. Retrieve Sample Tube and examine it carefully to make sure that no air bubbles are trapped inside. Once a satisfactory sample has been collected, proceed immediately with Steps 5 and 6 to "fix" the sample.

NOTE: Be careful not to introduce air into the sample while adding the reagents in steps 5 and 6. Simply drop the reagents into the sample. Cap carefully, and mix gently.

5. Add 2 drops of *Manganese Sulfate Solution [4167] and 2 drops of *Alkaline Potassium Iodide Azide [7166]. Cap and mix by inverting several times. A precipitate will form. Allow the precipitate to settle below the shoulder of the tube before proceeding.
6. Add 8 drops of *Sulfuric Acid, 1:1 [6141WT]. Cap and gently mix until the precipitate has dissolved. A clear-yellow to brown-orange color will develop, depending on the oxygen content of the sample.

NOTE: It is very important that all "brown flakes" are dissolved completely. If the water has a high DO level this could take several minutes. If flakes are not completely dissolved after 5 minutes, add 2 drops of *Sulfuric Acid 1:1 [6141WT] and continue mixing.

NOTE: Following the completion of step 6, contact between the water sample and the atmosphere will not affect the test result. Once the sample has been "fixed" in this manner, it is not necessary to perform the actual test procedure immediately. Thus, several samples can be collected and "fixed" in the field, and then carried back to a testing station or laboratory where the test procedure is to be performed.

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** (or another sequence containing **043 Dissolved Oxygen**) from **Testing Menu**.
4. Scroll to and select **043 Dissolved Oxygen** from menu.
5. Rinse a clean tube [0290] with untreated sample water. Fill to the 10 mL line with sample. This tube is the BLANK.
6. Insert tube into chamber, close lid and select **Scan Blank**.
7. Fill a second tube [0290] to the 10 line with the treated "Fixed" sample. This tube is the SAMPLE.
8. Remove BLANK from colorimeter, insert SAMPLE tube into chamber, close lid and select **Scan Sample**. Record result.
9. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

FLUORIDE

SPADNS METHOD · CODE 3647-02-SC

| QUANTITY | CONTENTS | CODE |
|-----------|-------------------------------|---------|
| 4 x 30 mL | *Acid Zirconyl SPADNS Reagent | *3875-G |
| 2 x 30 mL | *Sodium Arsenite Solution | *4128-G |
| 1 | Pipet, 0.5 mL, plastic | 0353 |
| 1 | Pipet, 1.0 mL, plastic | 0354 |

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Fluoride may occur naturally in some ground waters or it may be added to public drinking water supplies to maintain a 1.0 mg/L concentration to prevent dental cavities. At higher concentrations, fluoride may produce an objectionable discoloration of tooth enamel called fluorosis, though levels up to 8 mg/L have not been found to be physiologically harmful.

NOTE: This procedure uses the EPA approved Reagent System for fluoride found in method 4500-F-D, 18th Edition of Standard Methods, pp. 1-27.

APPLICATION Drinking and surface waters; domestic and industrial waters.

RANGE: 0.0–2.0 ppm Fluoride

MDL: 0.1 ppm

METHOD: Colorimetric test based upon the reaction between fluoride and zirconium dye lake. The fluoride reacts with the dye lake, dissociating a portion of it into a colorless complex ion and dye. As the fluoride concentration increases, the color produced becomes progressively lighter.

Samples may be stored and refrigerated in plastic containers.

SAMPLE HANDLING & PRESERVATION:

INTERFERENCES: The following substances produce a positive interference at the concentration given:

| | |
|--|-----------|
| Chloride [Cl ⁻] | 7000 mg/L |
| Phosphate [PO ₄ ⁻³] | 16 mg/L |
| [NaPO ₃] ₆ | 1 mg/L |

he following substances produce a negative interference at the concentration given:

| | |
|--|-----------|
| Alkalinity [CaCO ₃] | 5000 mg/L |
| Aluminum [Al ³⁺] | 0.1 mg/L |
| Iron [Fe ³⁺] | 10 mg/L |
| Sulfate [SO ₄ ⁻²] | 200 mg/L |

Color and turbidity must be removed or compensated for in the procedure. Temperature should be maintained within 5°C of room temperature.

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** (or another sequence containing **045 Fluoride**) from **Testing Menu**.
4. Scroll to and select **045 Fluoride** from menu.
5. This test requires a reagent blank. Rinse a clean tube [0290] with clear, colorless, fluoride free water. Fill to the 10 mL line with clear, colorless, fluoride free water.
6. Use the 0.5 mL pipet [0353] to add 0.5 mL of *Sodium Arsenite Solution [4128]. Cap and mix.
7. Use the 1.0 mL pipet [0354] to add 2 measures of *Acid-Zirconyl SPADNS Reagent [3875]. Cap and mix thoroughly. (This is the reagent blank.)
8. Insert tube into chamber, close lid and select **Scan Blank**.
9. Rinse a clean tube [0290] with sample water. Fill to the 10 mL line with sample water. Repeat steps 6 and 7 .
10. Insert tube into chamber, close lid and select **Scan Sample**. Record result.
11. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

HYDRAZINE

p-DIMETHYLAMINO BENZALDEHYDE METHOD · CODE 3656-01-SC

| QUANTITY | CONTENTS | CODE |
|-----------|----------------------------|---------|
| 2 x 60 mL | *Hydrazine Reagent A | *4841-H |
| 10 g | Hydrazine Reagent B Powder | 4842-D |
| 1 | Pipet, 1.0 mL, plastic | 0354 |
| 1 | Spoon, 0.15 g, plastic | 0727 |

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Hydrazine, N_2H_4 , is added to the water in high pressure boilers to reduce corrosion by acting as an oxygen scavenger.

| | |
|---------------------------------|---|
| APPLICATION: | Water and boiler water, industrial waste water. |
| RANGE: | 0.00–1.00 ppm Hydrazine |
| MDL: | 0.01 ppm |
| METHOD: | p-Dimethylaminobenzaldehyde reacts with hydrazine under acidic conditions to form a yellow color in proportion to the amount of hydrazine present. |
| SAMPLE HANDLING & PRESERVATION: | Samples should be analyzed as soon as possible after collection due to the ease with which hydrazine becomes oxidized. Acidification of the sample may increase the time between collection and analysis. |
| INTERFERENCES: | The substances normally present in water do not interfere with the test, with the exception of strong oxidizing agents. |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** [or another sequence containing **049 Hydrazine**] from **Testing Menu**.
4. Scroll to and select **049 Hydrazine** from menu.
5. Rinse a clean tube [0290] with sample water. Fill to the 10 mL line with sample.
6. Insert tube into chamber, close lid and select **Scan Blank**.
7. Remove tube from colorimeter. Use the 1 mL pipet [0354] to add 4 mL of *Hydrazine Reagent A [4841]. Cap and mix.
8. Use the 0.15 g spoon [0727] to add one measure of Hydrazine Reagent B Powder [4842]. Cap and shake vigorously for 10 seconds. Wait 2 minutes for maximum color development. An undissolved portion of Hydrazine Reagent B may remain in bottom of tube without adversely affecting results.
9. At the end of the 2 minute waiting period, mix, insert tube into chamber, close lid and select **Scan Sample**. Record result.
10. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTE: For best possible results, a reagent blank should be determined to account for any contribution to the test result by the reagent system. To determine the reagent blank, follow the above test procedure to scan a distilled or deionized water blank. Then follow the above procedure to perform the test on a distilled or deionized water sample. This test result is the reagent blank. Subtract the reagent blank from all subsequent test results of unknown samples. It is necessary to determine the reagent blank only when a new lot number of reagents is obtained.

MANGANESE – HIGH RANGE

PERIODATE METHOD · CODE 3669-SC

| QUANTITY | CONTENTS | CODE |
|----------|------------------------------|---------|
| 10 g | Manganese Buffer Reagent | 6310-D |
| 15 g | *Manganese Periodate Reagent | *6311-E |
| 1 | Spoon, 0.1 g, plastic | 0699 |
| 1 | Spoon, 0.15 g, plastic | 0727 |

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Manganese is present in ground water in the divalent state due to the lack of oxygen. In surface waters, manganese may be in various oxidation states as soluble complexes or as suspended compounds. Manganese is rarely present in excess of 1 mg/L. It may impart an objectionable taste or cause staining problems in laundry, but manganese levels normally encountered in water seldom produce any health hazards. Manganese is removed from water by various means, including chemical precipitation, pH adjustment, aeration, superchlorination and the use of ion exchange resins.

| | |
|---------------------------------|---|
| APPLICATION: | Drinking and surface waters, domestic and industrial wastewaters. |
| RANGE: | 0.0–15.0 Manganese |
| MDL: | 0.3 ppm |
| METHOD: | Periodate oxidizes soluble manganous compounds into permanganate. |
| SAMPLE HANDLING & PRESERVATION: | Manganese may oxidize readily in a neutral water and precipitate from solution. It may adhere to or be absorbed by container walls, especially glass. Acidified samples can be stored in plastic. |
| INTERFERENCES: | Reducing substances capable of reacting with periodate or permanganate must be removed or destroyed before the periodate oxidation is attempted. |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** (or another sequence containing **059 Manganese HR**) from **Testing Menu**.
4. Scroll to and select **059 Manganese HR** from menu.
5. Rinse a tube (0290) with sample water. Fill to the 10 mL line with sample.
6. Insert tube into chamber, close lid and select **Scan Blank**.
7. Remove tube from colorimeter. Use the 0.1 g spoon (0699) to add two measures of Manganese Buffer Reagent (6310). Cap and mix until powder dissolves.
8. Use the 0.15 g spoon (0727) to add one measure of *Manganese Periodate Reagent (6311). Cap and shake for one minute. An undissolved portion of the reagent may remain in the bottom of the tube without adversely affecting the test results. Wait two minutes for maximum color development. Solution will turn pink if manganese is present.
9. At the end of the two minute waiting period, mix, insert tube into chamber, close lid and select **Scan Sample**. Record result.
10. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NICKEL

DIMETHYLGLYOXIME METHOD · CODE 3663-01-SC

| QUANTITY | CONTENTS | CODE |
|----------|---|-----------|
| 60 mL | *Hydrochloric Acid, 2.5N | *6251PS-H |
| 30 g | *Ammonium Persulfate Reagent | *6566-G |
| 30 mL | *Silver Nitrate Solution, 0.0141N | *6346WT-G |
| 250 mL | Sodium Citrate, 10% | 6253-K |
| 60 mL | *Dimethylglyoxime, 1% | *6254-H |
| 60 mL | *Ammonium Hydroxide, Conc. | *6537-H |
| 3 | Pipets, 1.0 mL, plastic | 0354 |
| 1 | Spoon, 0.1 g, plastic | 0699 |
| 1 | Test tube, 5-10-12.9-15-20-25, glass, w/cap | 0608 |
| 1 | Graduated Cylinder, 10 mL, glass | 0416 |

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Nickel is not usually found in natural waters except as a result of contamination from industrial wastewaters as a corrosion product of stainless steel and nickel alloys. Nickel may also enter surface waters from plating bath process water.

| | |
|---------------------------------|---|
| APPLICATION: | Drinking and surface waters; domestic and industrial wastewater. |
| RANGE: | 0.00–8.00 ppm Nickel |
| MDL: | 0.15 ppm |
| METHOD: | Nickel under basic conditions forms a colored complex with dimethylglyoxime in proportion to the concentration of nickel. |
| SAMPLE HANDLING & PRESERVATION: | Samples may be collected in either plastic or glass containers and preserved by adding 5 mL of concentrated nitric acid per liter. |
| INTERFERENCES: | Organic matter interferes. Cobalt, iron, copper, manganese and chromium do not interfere if each of the concentrations is below 15 ppm. |

PROCEDURE

1. Use the 10 mL graduated cylinder [0416] to measure 10 mL of sample water. Pour into glass test tube [0608].
2. Use the 1 mL pipet [0354] to add 1 mL of *Hydrochloric Acid, 2.5N [6251].
3. Use the 0.1 g spoon [0699] to add 2 measures of *Ammonium Persulfate Reagent [6566]. Add two drops of *Silver Nitrate Solution, 0.0141N [6346WT]. Mix until the powder has dissolved. The solution will be slightly cloudy at this point.
4. Use 10 mL graduated cylinder [0416] to add 5 mL of Sodium Citrate, 10% [6253].
5. Use a second 1 mL pipet [0354] to add 1 mL of *Ammonium Hydroxide, Conc. [6537]. Mix, then dilute to 25 mL with deionized water.
6. Use a third 1 mL pipet [0354] to add 1 mL of *Dimethylglyoxime, 1% [6254]. Mix. Wait 20 minutes for color development.
7. At end of 20 minute waiting period fill a clean tube [0290] to the 10 mL line with the developed test sample.
8. Fill a second clean tube [0290] to 10 mL line with deionized water or untreated sample water. This is the blank.
9. Press and hold  until colorimeter turns on.
10. Press  to select **Testing Menu**.
11. Select **All Tests** [or another sequence containing **064 Nickel**] from **Testing Menu**.
12. Scroll to and select **064 Nickel** from menu.
13. Insert the blank into chamber, close lid and select **Scan Blank**.
14. Insert test sample into chamber, close lid and select **Scan Sample**. Record result.
15. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTE: It is strongly suggested that a reagent blank be determined to account for any contribution to the test result by the reagent system. To determine the reagent blank, follow the above test procedure to scan a distilled or deionized water blank. Then follow the above procedure to perform the test on a distilled or deionized water sample. This test result is the reagent blank. Subtract the reagent blank from all subsequent test results of unknown samples.

OZONE

INDIGO METHOD · CODE 3651-SC

| QUANTITY | CONTENTS | CODE |
|----------|----------------------------------|-----------|
| 15 mL | Chlorine Inhibitor | 3990-E |
| 250 mL | Ozone Buffer | 3991-K |
| 30 mL | Indigo Blue Stock Solution | 3989-G |
| 1 | Sampling Apparatus | 0681 |
| 1 | Pipet, transfer, 1.0 mL | 2-2170 |
| 1 | Pipet, transfer, 5 mL | 0329 |
| 1 | Pump, 10 mL | 30527 |
| 1 | Bottle, HR Reagent, amber glass | 3988-MT-G |
| 1 | Graduated Cylinder, 50 mL, glass | 0418 |

***WARNING:** Reagents marked with an * are considered to be potential health hazards. To view or print a Safety Data Sheet [SDS] for these reagents go to www.lamotte.com. Search for the four digit reagent code number listed on the reagent label, in the contents list or in the test procedures. Omit any letter that follows or precedes the four digit code number. For example, if the code is 4450WT-H, search 4450. To obtain a printed copy, contact LaMotte by email, phone or fax.

Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

Ozone is sometimes used in place of, or in conjunction with, chlorine or other halogens for disinfection of pool, spa, or drinking waters. Recently, large aquatic facilities have begun using ozone as a disinfectant in many artificial habitats.

| | |
|---------------------------------|---|
| APPLICATION: | Drinking, pool and aquatic waters. |
| RANGE: | 0.00–0.40 ppm Ozone, Low Range 0.00–2.50 ppm Ozone, High Range |
| MDL: | 0.01 ppm Ozone, Low Range 0.05 ppm Ozone, High Range |
| METHOD: | Ozone rapidly and stoichiometrically decolorizes Indigo Trisulfonate under acidic conditions. |
| SAMPLE HANDLING & PRESERVATION: | Ozone is extremely unstable in aqueous solutions. Test must be performed immediately and the sample must not be agitated. |
| INTERFERENCES: | Manganese at any level interferes. |

PROCEDURE—LOW RANGE

A. PREPARATION OF HR REAGENT

NOTE: The quantity of Indigo Blue Stock Solution [3989] supplied will prepare one batch of HR Reagent for the High Range Ozone procedure or five batches of HR Reagent for the Low Range Ozone procedure.

1. Use the 50 mL graduated cylinder to carefully add 45 mL of *Ozone Buffer [3991] to amber glass bottle marked HR Reagent [3988-MT-G].
2. Use the 5 mL transfer pipet [0329] and pump [30527] to add 5 mL of Indigo Blue Stock Solution [3989] to the amber glass bottle. Cap and mix.

B. DETERMINATION OF OZONE

3. Use the 1.0 mL transfer pipet [2-2170] and pump [30527] to add 1.0 mL of HR Reagent [3988] to each of 2 clean tubes [0290].
4. If chlorine is present add 3 drops Chlorine Inhibitor [3990] to each tube. Cap tubes.
5. Take one of the prepared tubes [0290] and sampling apparatus [0681] to sampling site.
6. Lower end of tubing of sampling apparatus to desired depth. Slowly withdraw and depress plunger several times to purge syringe and tubing. Slowly withdraw plunger to fill purged syringe.
7. Remove plastic tubing from syringe. Remove cap from the prepared tube. Place tip of syringe against inside of the prepared tube. Slowly depress plunger and fill to the 10 mL line and cap. This is the Sample Tube.
NOTE: DO NOT SHAKE OR INVERT THE SAMPLE.
8. Fill the second prepared tube [0290] to the 10 mL line with ozone free water. This is the Reagent Blank.
9. Press and hold  until colorimeter turns on.
10. Press  to select **Testing Menu**.
11. Select **All Tests** [or another sequence containing **071 Ozone IB LR**] from **Testing Menu**.
12. Scroll to and select **071 Ozone IB LR** from menu.
13. Insert the Reagent Blank tube into chamber, close lid and select **Scan Blank**.
14. Insert reacted Sample Tube into chamber, close lid and select **Scan Sample**. Record result.
15. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTE: HR Reagent must be made fresh each week. If reagent is refrigerated, it may be kept up to 3 weeks.

PROCEDURE-HIGH RANGE

A. PREPARATION OF HR REAGENT

NOTE: The quantity of Indigo Blue Stock Solution [3989] supplied will prepare one batch of HR Reagent for the High Range Ozone procedure or five batches of HR Reagent for the Low Range Ozone procedure.

1. Use the 50 mL graduated cylinder to carefully add 25 mL of *Ozone Buffer [3991] to amber glass bottle marked HR Reagent [3988-MT-G].
2. Use the 50 mL graduated cylinder to carefully add 25 mL of Indigo Blue Stock Solution [3989] to the amber glass bottle. Cap and mix.

B. DETERMINATION OF OZONE

3. Use the 1.0 mL transfer pipet [2-2170] and pump [30527] to add 1.0 mL of HR Reagent [3988] to each of 2 clean tubes [0290].
4. If chlorine is present add 3 drops Chlorine Inhibitor [3990] to each tube. Cap tubes.
5. Take one of the prepared tubes [0290] and sampling apparatus [0681] to sampling site.
6. Lower end of tubing of sampling apparatus to desired depth. Slowly withdraw and depress plunger several times to purge syringe and tubing. Slowly withdraw plunger to fill purged syringe.
7. Remove plastic tubing from syringe. Remove cap from the prepared tube. Place tip of syringe against inside of the prepared tube. Slowly depress plunger and fill to the 10 mL line and cap. This is the Sample Tube.
NOTE: DO NOT SHAKE OR INVERT THE SAMPLE.
8. Fill the second prepared tube [0290] to the 10 mL line with ozone free water. This is the Reagent Blank.
9. Press and hold  until colorimeter turns on.
10. Press  to select **Testing Menu**.
11. Select **All Tests** (or another sequence containing **072 Ozone IB HR**) from **Testing Menu**.
12. Scroll to and select **072 Ozone IB HR** from menu.
13. Insert the Reagent Blank tube into chamber, close lid and select **Scan Blank**.
14. Insert reacted Sample Tube into chamber, close lid and select **Scan Sample**. Record result.

15. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTE: HR Reagent must be made fresh each week. If reagent is refrigerated, it may be kept up to 3 weeks.

PHOSPHATE – LOW RANGE

ASCORBIC ACID REDUCTION METHOD · CODE 3653-SC

| QUANTITY | CONTENTS | CODE |
|----------|----------------------------|-----------|
| 60 mL | *Phosphate Acid Reagent | *V-6282-H |
| 5 g | Phosphate Reducing Reagent | V-6283-C |
| 1 | Pipet, 1 mL, plastic | 0354 |
| 1 | Spoon, 0.1 g, plastic | 0699 |

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Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

Phosphorus is an important nutrient for aquatic plants. The amount found in water is generally not more than 0.1 ppm unless the water has become polluted from waste water sources or excessive drainage from agricultural areas. When phosphorus is present in excess of the concentrations required for normal aquatic plant growth, a process called eutrophication takes place. This creates a favorable environment for the increase in algae and weeds. When algae cells die, oxygen is used in the decomposition and fish kills often result. Rapid decomposition of dense algae scums with associated organisms give rise to foul odors and hydrogen sulfide gas.

| | |
|---------------------------------|--|
| APPLICATION: | Drinking, surface and saline waters; domestic and industrial wastes [Method based on reactions that are specific for orthophosphate]. |
| RANGE: | 0.00–3.00 ppm Orthophosphate |
| MDL: | 0.05 ppm |
| METHOD: | Ammonium molybdate and antimony potassium tartrate react in a filtered acid medium with dilute solution of PO_4^{-3} to form an antimony-phosphomolybdate complex. This complex is reduced to an intense blue colored complex by ascorbic acid. The color is proportional to the amount of phosphate present. [Only orthophosphate forms a blue color in this test.] Polyphosphates [and some organic phosphorus compounds] may be converted to the orthophosphate form by sulfuric acid digestion. Organic phosphorus compounds may be converted to the orthophosphate form by persulfate digestion. |
| SAMPLE HANDLING & PRESERVATION: | If benthic deposits are present in the area being sampled, great care should be taken not to include these deposits. If the analysis cannot be performed the same day of collection, the sample should be preserved by the addition of 2 mL of concentrated sulfuric acid or 40 mg mercuric chloride per liter and refrigerated at 4°C. |
| INTERFERENCES: | <p>a. No interference from copper, iron, or silicate at concentrations many times the concentration of sea water. However, high iron concentrations can cause precipitation and subsequent loss of phosphorus.</p> <p>b. Salt error for samples ranging from 5% to 20% salt content was found to be less than 1%.</p> <p>c. Mercuric chloride, HgCl_2, when used as the preservative, interferes when the chloride levels are low [less than 50 mg/L]. This interference is overcome by spiking samples with a minimum of 50 mg/L of sodium chloride.</p> |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** (or another sequence containing **081 Phosphate LR**) from **Testing Menu**.
4. Scroll to and select **081 Phosphate LR** from menu.
5. Rinse a clean tube [0290] with sample water. Fill to the 10 mL line with sample.
6. Insert tube into chamber, close lid and select **Scan Blank**.
7. Remove tube from colorimeter. Use 1.0 mL pipet [0354] to add 1.0 mL of *Phosphate Acid Reagent [V-6282]. Cap and mix.
8. Use the 0.1 g spoon [0699] to add one measure of Phosphate Reducing Reagent [V-6283]. Cap and mx until powder dissolves. Wait 5 minutes for full color development. Solution will turn blue if phosphates are present.
9. At end of 5 minute waiting period, mix, insert tube into chamber, close lid and select **Scan Sample**. Record result.
10. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

PHOSPHATE – HIGH RANGE

VANADOMOLYBDOPHOSPHORIC ACID METHOD · CODE 3655-SC

| QUANTITY | CONTENTS | CODE |
|-----------|------------------------|---------|
| 4 x 30 mL | *VM Phosphate Reagent | *4410-G |
| 1 | Pipet, 1.0 mL, plastic | 0354 |

***WARNING:** Reagents marked with an * are considered to be potential health hazards. To view or print a Safety Data Sheet (SDS) for these reagents go to www.lamotte.com. Search for the four digit reagent code number listed on the reagent label, in the contents list or in the test procedures. Omit any letter that follows or precedes the four digit code number. For example, if the code is 4450WT-H, search 4450. To obtain a printed copy, contact LaMotte by email, phone or fax.

Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

Phosphate treatments in boiler and cooling water and other industrial water systems are run at levels up to 100 ppm orthophosphate. These high levels permit the use of a simpler, high range test.

| | |
|---------------------------------|---|
| APPLICATION: | Boiler, cooling, and industrial water. |
| RANGE: | 0.0–70.0 ppm Orthophosphate |
| MDL: | 0.5 ppm |
| METHOD: | Orthophosphate reacts in acid conditions with ammonium vanadomolybdate to form vanadomolybdophosphoric acid. This yellow color is proportional to the concentration of orthophosphate and is read colorimetrically. |
| SAMPLE HANDLING & PRESERVATION: | If the analysis cannot be performed the same day of collection, the sample should be preserved by the addition of 2 mL of concentrated sulfuric acid or 40 mg mercuric chloride per liter and refrigerated at 4°C. |
| INTERFERENCES: | Silica interferes only if the sample is heated. Arsenate, fluoride, thorium, bismuth, sulfide, thiosulfate, and thiocyanate cause negative interference. |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** (or another sequence containing **080 Phosphate HR**) from Testing Menu.
4. Scroll to and select **080 Phosphate HR** from menu.
5. Rinse a clean tube (0290) with sample water. Fill to the 10 mL line with sample.
6. Insert tube into chamber, close lid and select **Scan Blank**.
7. Remove tube from colorimeter. Use the 1.0 mL pipet (0354) to add 2.0 mL of *VM Phosphate Reagent (4410). Cap and mix. Wait 5 minutes for full color development.
8. After 5 minute waiting period, mix, insert tube into chamber, close lid and select **Scan Sample**. Record result.
9. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

POTASSIUM

TETRAPHENYLBORON METHOD · CODE 3639-SC

| QUANTITY | CONTENTS | CODE |
|----------|--------------------------|-----------|
| 30 mL | *Sodium Hydroxide, 1.0N | *4004WT-G |
| 5 g | *Tetraphenylboron Powder | *6364-C |
| 1 | Spoon, 0.05 g, plastic | 0696 |

***WARNING:** Reagents marked with an * are considered to be potential health hazards. To view or print a Safety Data Sheet (SDS) for these reagents go to www.lamotte.com. Search for the four digit reagent code number listed on the reagent label, in the contents list or in the test procedures. Omit any letter that follows or precedes the four digit code number. For example, if the code is 4450WT-H, search 4450. To obtain a printed copy, contact LaMotte by email, phone or fax.

Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

Potassium, as the seventh most common element on the Earth, may be found in minor quantities in most water supplies. It seldom exceeds 10 ppm in drinking water and usually is less than 2 ppm. In some brine or runoff in agricultural areas the potassium concentration may reach 100 ppm.

| | |
|---------------------------------|---|
| APPLICATION: | Drinking, surface, and saline water. |
| RANGE: | 0.0–10.0 ppm Potassium |
| MDL: | 0.8 ppm |
| METHOD: | Potassium reacts with sodium tetraphenylborate to form a colloidal white precipitate in quantities proportional to the potassium concentration. |
| SAMPLE HANDLING & PRESERVATION: | Store samples in polyethylene bottles, not in soft glass where leaching of potassium from the glass may occur. Samples may be acidified to pH 2 with nitric acid, but should be neutralized before analyzing. |
| INTERFERENCE: | Calcium and magnesium interfere at very high concentrations. Check for stray light interference [see p. 69]. |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** (or another sequence containing **085 Potassium**) from **Testing Menu**.
4. Scroll to and select **085 Potassium** from menu.
5. Rinse a clean tube [0290] with sample water. Fill to the 10 mL line with sample.
6. Insert tube into chamber, close lid and select **Scan Blank**.
7. Remove tube from colorimeter. Add 4 drops of *Sodium Hydroxide, 1.0N [4004WT]. Cap and mix.
8. Use the 0.05 g spoon [0696] to add one measure of *Tetraphenylboron Powder [6364]. Cap and shake vigorously for 20 seconds until all of the powder has dissolved. Wait 5 minutes.
9. At end of 5 minute waiting period, mix tube again to suspend any settled precipitate. Insert tube into chamber, close lid and select **Scan Sample**. Record result.
10. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTES: For best possible results, a reagent blank should be determined to account for any contribution to the test result by the reagent system. To determine the reagent blank, follow the above test procedure to scan a distilled or deionized water blank. Then follow the above procedure to perform the test on a distilled or deionized water sample. This test result is the reagent blank. Subtract the reagent blank from all subsequent test results of unknown samples. It is necessary to determine the reagent blank only when a new lot number of reagents are obtained.

For the most accurate results, the sample and reagents should be at 25±4°C.

SILICA – LOW RANGE

HETEROPOLY BLUE METHOD · CODE 3664-SC

| QUANTITY | CONTENTS | CODE |
|----------|-----------------------|-----------|
| 30 mL | *Silica Reagent #1 | *V-4466-G |
| 30 mL | *Silica Reagent #2 | *V-4467-G |
| 30 mL | *Silica Reagent #3 | *V-4468-G |
| 10 g | Silica Reagent #4 | V-6284-D |
| 1 | Spoon, 0.1 g, plastic | 0699 |

***WARNING:** Reagents marked with an * are considered to be potential health hazards. To view or print a Safety Data Sheet [SDS] for these reagents go to www.lamotte.com. Search for the four digit reagent code number listed on the reagent label, in the contents list or in the test procedures. Omit any letter that follows or precedes the four digit code number. For example, if the code is 4450WT-H, search 4450. To obtain a printed copy, contact LaMotte by email, phone or fax.

Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

Silicon dioxide, SiO₂, commonly known as silica, occurs in all natural water. Silica may be present as suspended, insoluble particles in a colloidal or polymeric state. It may also be present in a reactive form as silicic acid or silicate ions. Silica is a major nutrient for diatoms. A silica cycle occurs in many bodies of water containing organisms, such as diatoms, that use silica in their skeletal structure. The silica removed from the water may be slowly returned to solution by the decomposition of the dead organisms. The major source of silica in natural water is from the decomposition of silicate minerals in the drainage basin from which the waters flow.

The presence of silica is particularly objectionable in water used for boiler feed water purposes, as it may cause the formation of a hard, dense scale which has unusually high resistance to heat transfer. Serious loss of turbine efficiency results from insoluble silica turbine blade deposits caused by vaporization of silica from boiler water.

| | |
|---------------------------------|---|
| APPLICATION: | Drinking, surface and saline waters; domestic and industrial wastes. |
| RANGE: | 0.0–4.0 ppm Silica |
| MDL: | 0.05 ppm |
| METHOD: | Reactive silica forms a complex with ammonium molybdate in an acidic solution to produce a yellow-green color in proportion to the amount of silica present. Phosphate also reacts with molybdate but the addition of oxalic acid eliminates the molybdophosphoric acid complex. This silica molybdate complex is then reduced by ascorbic acid to produce an intense blue color. |
| SAMPLE HANDLING & PRESERVATION: | Silica samples may be preserved by refrigeration at 4°C in plastic containers up to one week without any change in silica concentration. |
| INTERFERENCES: | Sulfides and large amounts of iron interfere. Color and turbidity may be removed by standardizing the instrument with the original water sample. Since silica is a component of glass waste and a common contaminant, it is suggested to run a reagent blank using silica-free water. The blank value is subtracted from the sample concentrations. |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** [or another sequence containing **087 Silica LR**] from **Testing Menu**.
4. Scroll to and select **087 Silica LR** from menu.
5. Rinse a clean tube [0290] with sample water. Fill to the 10 mL line with sample.
6. Insert tube into chamber, close lid and select **Scan Blank**. [See Note]
7. Remove tube from colorimeter. Add 6 drops *Silica Reagent #1 [V-4466]. Cap and invert to mix.
8. Add 12 drops of *Silica Reagent #2 [V-4467]. Cap and mix. Wait 5 minutes.
9. Add 8 drops of *Silica Reagent #3 [V-4468]. Cap and mix. Wait 2 minutes.
10. Use the 0.1 g spoon [0699] to add one measure of Silica Reagent #4 [V-6284]. Cap and mix gently until powder has dissolved. Wait 5 minutes for full color development.
11. At end of 5 minute waiting period, mix, insert tube into chamber, close lid and select **Scan Sample**. Record result.
12. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTE: For best possible results, a reagent blank should be determined to account for any contribution to the test result by the reagent system. To determine the reagent blank, follow the above test procedure to scan a distilled or deionized water blank. Then follow the above procedure to perform the test on a distilled or deionized water sample. This test result is the reagent blank. Subtract the reagent blank from all subsequent test results of unknown samples. It is necessary to determine the reagent blank only when a new lot number of reagents are obtained.

SILICA – HIGH RANGE

SILICOMOLYBDATE METHOD · CODE 3687-SC

| QUANTITY | CONTENTS | CODE |
|----------|--------------------|-----------|
| 30 mL | *Silica Reagent #1 | *V-4466-G |
| 30 mL | *Silica Reagent #2 | *V-4467-G |
| 15 mL | *Silica Reagent #3 | *V-4468-G |

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Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

Silicon dioxide, SiO_2 , commonly known as silica, occurs in all natural water. Silica may be present as suspended, insoluble particles in a colloidal or polymeric state. It may also be present in a reactive form as silicic acid or silicate ions. Silica is a major nutrient for diatoms. A silica cycle occurs in many bodies of water containing organisms, such as diatoms, that use silica in their skeletal structure. The silica removed from the water may be slowly returned to solution by the decomposition of the dead organisms. The major source of silica in natural water is from the decomposition of silicate minerals in the drainage basin from which the waters flow.

The presence of silica is particularly objectionable in water used for boiler feed water purposes, as it may cause the formation of a hard, dense scale which has unusually high resistance to heat transfer. Serious loss of turbine efficiency results from insoluble silica turbine blade deposits caused by vaporization of silica from boiler water.

| | |
|---------------------------------|---|
| APPLICATION: | Boilers and cooling towers; domestic and industrial wastes. |
| RANGE: | 0–75 ppm Silica |
| MDL: | 1 ppm |
| METHOD: | Silica forms a complex with ammonium molybdate in an acidic solution to produce a yellow color in proportion to the amount of silica present. Phosphate also reacts with molybdate but the addition of oxalic acid eliminates the molybdophosphoric acid complex. |
| SAMPLE HANDLING & PRESERVATION: | Silica samples may be preserved by refrigeration at 4°C in plastic containers up to one week without any change in silica concentration. |
| INTERFERENCES: | Sulfides and large amounts of iron interfere. Color and turbidity may be removed by standardizing the instrument with the original water sample. |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** [or another sequence containing **086 Silica HR**] from **Testing Menu**.
4. Scroll to and select **086 Silica HR** from menu.
5. Rinse a clean tube [0290] with sample water. Fill to the 10 mL line with sample.
6. Insert tube into chamber, close lid and select **Scan Blank**.
7. Remove tube from colorimeter. Add 6 drops *Silica Reagent #1 [V-4466]. Cap and invert to mix.
8. Add 12 drops of *Silica Reagent #2 [V-4467]. Cap and mix. Wait 5 minutes.
9. At end of 5 minute waiting period, add 8 drops of *Silica Reagent #3 [V-4468]. Cap and mix.
10. Insert tube into chamber, close lid and select **Scan Sample**. Record result.
11. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTE: To extend the range to 100 ppm, perform a 2:1 dilution of water sample, with silica-free water. Perform test and multiply result by 2.

SULFATE – HIGH RANGE

BARIUM CHLORIDE METHOD · CODE 3665-SC

| QUANTITY | CONTENTS | CODE |
|----------|-----------------------|-----------|
| 10 g | *Sulfate Reagent | *V-6277-D |
| 1 | Spoon, 0.1 g, plastic | 0699 |

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Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

The most common mineral forms of sulfur are iron sulfide, lead sulfide, zinc sulfide and as calcium sulfate and magnesium sulfate. In most fresh waters the sulfate ion is the second or third most abundant anion, being exceeded only by bicarbonate and, in some cases, silicate. Sulfur, in the form of sulfate, is considered an important nutrient element. Mineral springs are rich in sulfate and feed appreciable quantities of this compound to the watershed. Acid mine water drainage is a form of pollution which may contribute extremely large amounts of sulfate content to natural waters. Other sources of sulfate include waste material from pulp mills, steel mills, food processing operations and municipal wastes. Many bacteria obtain sulfur from sulfate for the synthesis of amino acids. In lakes and streams low in oxygen, this process of sulfate reduction causes the production of hydrogen sulfide, with its characteristic offensive odor. Calcium sulfate and magnesium sulfate contribute significantly to the hardness of water. Under natural conditions, the quantities ordinarily to be expected in lakes are between 3 and 30 parts per million.

| | |
|---------------------------------|---|
| APPLICATION: | Drinking and surface waters, domestic and industrial wastes. |
| RANGE: | 0–100 ppm Sulfate |
| MDL: | 3 ppm |
| METHOD: | Sulfate ion is precipitated in an acid medium with barium chloride to form a barium sulfate suspension in proportion to the amount of sulfate present. |
| SAMPLE HANDLING & PRESERVATION: | Sulfate samples may be preserved by refrigeration at 4°C up to 7 days in glass or plastic containers without any change in concentration. |
| INTERFERENCE: | Suspended matter and color interference may be removed by a filtration step. Silica in excess of 500 mg/L will interfere. Check for stray light interference [see page 69]. |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** (or another sequence containing **089 Sulfate HR**) from **Testing Menu**.
4. Scroll to and select **089 Sulfate HR** from menu.
5. Rinse a clean tube (0290) with sample water. Fill to the 10 mL line with sample.
6. Insert tube into chamber, close lid and select **Scan Blank**.
7. Remove tube from colorimeter. Use the 0.1 g spoon (0699) to add one measure of *Sulfate Reagent (V-6277). Cap and shake until powder dissolves. A white precipitate will develop if sulfates are present. Wait 5 minutes.
8. Mix tube again. Insert tube into chamber, close lid and select **Scan Sample**. Record result.
9. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTE: If the sulfate concentration of the test sample is greater than 100 ppm, it is recommended that a dilution be made with deionized water and the results multiplied by the dilution factor.

A white film is deposited on the inside of test tubes as a result of the sulfate test. Thoroughly clean and rinse test tubes after each test.

For the most accurate results, samples and reactions should be at 25±4°C.

SULFIDE – LOW RANGE

METHYLENE BLUE METHOD · CODE 3654-02-SC

| QUANTITY | CONTENTS | CODE |
|-----------|-------------------------|-----------|
| 2 x 30 | *Sulfide Reagent A | *V-4458-G |
| 15 mL | *Sulfide Reagent B | *V-4459-E |
| 2 x 60 mL | Sulfide Reagent C | 4460-H |
| 2 | Pipets, 1.0 mL, plastic | 0354 |

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Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

Sulfide occurs in many well water supplies and sometimes is formed in lakes or surface waters. In distribution systems, it may be formed as a result of bacterial action on organic matter under anaerobic conditions. It may also be found in waters receiving sewage or industrial wastes. Lake muds rich in sulfates produce hydrogen sulfide during periods of very low oxygen levels that result from stagnation. Concentrations of a few hundredths of a part per million [or milligram per liter] cause a noticeable odor. At low concentrations, this odor is described as “musty”; at high concentration, as “rotten eggs.” Removal of sulfide odor is accomplished by aeration or chlorination. Hydrogen sulfide, a toxic substance, acts as a respiratory depressant in both humans and fish.

| | |
|---------------------------------|--|
| APPLICATION: | Drinking, surface and saline waters; domestic and industrial wastes. |
| RANGE: | 0.00–1.50 ppm Sulfide |
| MDL: | 0.06 ppm |
| METHOD: | Under suitable conditions the sulfide ion reacts with p-aminodimethylaniline and ferric chloride to produce methylene blue in proportion to the sulfide concentration. Ammonium phosphate is added to remove the color due to the ferric iron. |
| SAMPLE HANDLING & PRESERVATION: | Samples must be taken with a minimum of aeration since sulfide is volatilized by aeration and any oxygen which is taken up will destroy sulfides by chemical action. Samples that are used for total sulfide concentrations may be preserved by adding 2M zinc acetate solution at a dosage of 2 mL per liter of sample. This precipitates sulfide as inert zinc sulfide. Determination of dissolved sulfides in samples not preserved with zinc acetate must be started within 3 minutes of sampling. |
| INTERFERENCES: | Strong reducing agents such as sulfite, thiosulfate, and hydrosulfite prevent the formation of the color or diminish its intensity. High concentrations of sulfide will inhibit the reaction, but dilution of the sample prior to analysis eliminates this problem. |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** [or another sequence containing **090 Sulfide LR**] from **Testing Menu**.
4. Scroll to and select **090 Sulfide LR** from menu.
5. Rinse a clean tube [0290] with sample water. Fill to the 10 mL line with sample.
6. Insert tube into chamber, close lid and select **Scan Blank**.
7. Remove tube from colorimeter. Use the 1.0 mL pipet [0354] to add 1.0 mL of *Sulfide Reagent A [V-4458]. Cap and mix.
8. Add 6 drops of Sulfide Reagent B [V-4459]. Cap and mix. Wait 1 minute. Solution will turn blue if sulfides are present.
9. Use the 1.0 mL pipet [0354] to add 2.0 mL of Sulfide Reagent C [4460]. Cap and mix. Color development is immediate and stable.
10. Insert tube into chamber, close lid and select **Scan Sample**. Record result.
11. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

TANNIN

TUNGSTO-MOLYBDOPHOSPHORIC ACID METHOD CODE 3666-01-SC

| QUANTITY | CONTENTS | CODE |
|-----------|------------------------|---------|
| 30 mL | *Tannin Reagent #1 | *7833-G |
| 2 x 60 mL | *Tannin Reagent #2 | *7834-H |
| 1 | Pipet, plain, plastic | 0352 |
| 1 | Pipet, 1.0 mL, plastic | 0354 |

***WARNING:** Reagents marked with an * are considered to be potential health hazards. To view or print a Safety Data Sheet (SDS) for these reagents go to www.lamotte.com. Search for the four digit reagent code number listed on the reagent label, in the contents list or in the test procedures. Omit any letter that follows or precedes the four digit code number. For example, if the code is 4450WT-H, search 4450. To obtain a printed copy, contact LaMotte by email, phone or fax.

Emergency information for all LaMotte reagents is available from Chem-Tel: [US, 1-800-255-3924] [International, call collect, 813-248-0585].

Tannin and lignin are examples of hydroxylated aromatic compounds found in discharge wastewater from paper mills, in some boiler water treatment, in natural brackish water, and in wastewater from leather tanning plants. The taste and odor of these compounds is generally offensive so that their control is important in many areas.

| | |
|---------------------------------|---|
| APPLICATION: | Industrial wastewater, boiler water, and natural water. |
| RANGE: | 0.0-10.0 ppm Tannic Acid |
| MDL: | 0.1 ppm |
| METHOD: | The hydroxylated aromatic compounds will reduce a mixture of tungstophosphoric and molybdophosphoric acids to form a blue color in proportion to the concentration of aromatic hydroxyl groups. |
| SAMPLE HANDLING & PRESERVATION: | Sample should be analyzed as soon as possible after collection. |
| INTERFERENCES: | Other reducing compounds such as ferrous iron and sulfites. Results may be expressed as tannin like compounds, or aromatic hydroxy compounds. |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** (or another sequence containing **093 Tannin**) from **Testing Menu**.
4. Scroll to and select **093 Tannin** from menu.
5. Rinse a clean tube (0290) with sample water. Fill to the 10 mL line with sample.
6. Insert tube into chamber, close lid and select **Scan Blank**.
7. Remove tube from colorimeter. Use the plain pipet (0352) to add 4 drops of *Tannin Reagent #1 (7833). Cap and mix.
8. Use the 1.0 mL pipet (0354) to add 2.0 mL of *Tannin Reagent #2 (7834). Cap and mix. Wait 30 minutes for full color development.
9. At end of 30 minute waiting period, mix, insert tube into chamber, close lid and select **Scan Sample**. Record result.
10. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTES: For best possible results, a reagent blank should be determined to account for any contribution to the test result by the reagent system. To determine the reagent blank, follow the above test procedure to scan a distilled or deionized water blank. Then follow the above procedure to perform the test on a distilled or deionized water sample. This test result is the reagent blank. Subtract the reagent blank from all subsequent test results of unknown samples. It is necessary to determine the reagent blank only when a new lot number of reagents is obtained.

For the most accurate results, the sample and reagents should be at $20 \pm 2^{\circ}\text{C}$.

TURBIDITY

ABSORPTION METHOD · NO REAGENTS REQUIRED

Turbidity is a measure of water clarity and is independent of color. Turbidity is caused by undissolved and suspended solids. Mud, silt, algae, and microorganisms can all cause turbidity. Turbidity is a gross measurement of water quality.

| | |
|---------------------------------|---|
| APPLICATION: | Surface and industrial water for non-compliance monitoring. [For compliance monitoring at low turbidity levels, use a commercial nephelometer.] |
| RANGE: | 0–500 FAU [Formazin Attenuation Units] |
| MDL: | 3 FAU |
| METHOD: | Absorptimetric, 180° detector |
| SAMPLE HANDLING & PRESERVATION: | Measure sample as soon as possible after collection. |
| INTERFERENCES: | Check for stray light interference [see page 69]. |

PROCEDURE

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** [or another sequence containing **095 Turbidity**] from **Testing Menu**.
4. Scroll to and select **095 Turbidity** from menu.
5. Rinse a clean tube [0290] with deionized water [turbidity free]. Fill to the 10 mL line with deionized water.
6. Insert tube into chamber, close lid and select **Scan Blank**.
7. Rinse a second clean tube [0290] with sample water. Fill to the 10 mL line with sample. Cap tube. Wipe off excess water and fingerprints. Shake to resuspend particulate matter. Remove all bubbles before measurement.
8. Insert tube into chamber, close lid and select **Scan Sample**. Record result. Turbidity measurements should be taken as soon as possible after sample has been collected.
9. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTE: For the most accurate results, the sample should be at $25 \pm 4^{\circ}\text{C}$.

FORMAZIN STOCK SOLUTIONS

The turbidity calibration curve for this instrument was prepared by using formazin solutions as a reference. A 4000 FTU standard solution is available [Order Code 6195-H, 60 mL] that can be diluted with low turbidity water to prepare solutions within the test range. Dilutions from this stock solution should be prepared fresh daily with low turbidity water.

Alternatively, a stock turbidity solution of 400 NTU can be prepared by observing safety precautions and carefully following the procedure below.

Preparation of Formazin Stock Solution

1. Dissolve 1.000 g of Hydrazine Sulfate in deionized water and dilute to the mark in a 100 mL volumetric flask.
2. Dissolve 10.00 g of hexamethylenetetramine in deionized water and dilute to the mark in a 100 mL volumetric flask.
3. Mix 5 mL of each solution in a 100 mL volumetric flask and allow to sit undisturbed for 24 hours at 25 ± 3 °C.
4. At the end of the waiting period, dilute to the mark with deionized water and mix. Store in amber glass.
5. The concentration of this stock solution is 400 FTU. This stock solution is stable for one month. Dilutions from this stock solution should be prepared fresh daily with low turbidity water.

ZINC – LOW RANGE

ZINCON METHOD · CODE 3667-01-SC

| QUANTITY | CONTENTS | CODE |
|----------|---|-----------|
| 30 mL | *Zinc Indicator Solution | *6314-G |
| 120 mL | *Methyl Alcohol | *6319-J |
| 10 g | Sodium Ascorbate Powder | 6316-D |
| 25 g | *Zinc Buffer Powder | *6315-G |
| 15 mL | *Sodium Cyanide, 10% | *6565-E |
| 30 mL | *Formaldehyde Solution, 37% | *5128-G |
| 1 | Dilute Zinc Indicator Solution” Bottle, w/1 pipet assembly | 6321-MT-G |
| 1 | Graduated Cylinder, 10 mL, glass | 0416 |
| 1 | Spoon, 0.5 g, plastic | 0698 |
| 2 | Pipets, plain, plastic | 0352 |
| 1 | Spoon, 0.1 g, plastic | 0699 |

***WARNING:** Reagents marked with an * are considered to be potential health hazards. To view or print a Safety Data Sheet (SDS) for these reagents go to www.lamotte.com. Search for the four digit reagent code number listed on the reagent label, in the contents list or in the test procedures. Omit any letter that follows or precedes the four digit code number. For example, if the code is 4450WT-H, search 4450. To obtain a printed copy, contact LaMotte by email, phone or fax.

Emergency information for all LaMotte reagents is available from Chem-Tel: (US, 1-800-255-3924) (International, call collect, 813-248-0585).

Zinc enters the domestic water supply from the deterioration of galvanized iron and brass pipes, and from industrial wastes. Zinc is an essential element for body growth and development and is an important plant nutrient. Concentrations of zinc above 5.0 mg/L in drinking water can cause a bitter astringent taste. In the U.S., zinc concentrations may vary between 0.06 to 7.0 mg/L, with an average value of 1.33 mg/L.

APPLICATION: Drinking and surface waters, domestic and industrial waste water.

RANGE: 0.00–3.00 ppm Zinc

MDL: 0.05 ppm

METHOD: Zinc forms a blue colored complex with Zincon in a solution buffered at pH 9.0. Other heavy metals are complexed by cyanide and the zinc cyanide complex is released by the addition of formaldehyde before the other metal cyanide complexes are destroyed. Sodium ascorbate is added to reduce the interference of manganese.

SAMPLE HANDLING & PRESERVATION: Sample should be analyzed within 6 hours after collection. The addition of hydrochloric acid will help preserve the metal ion content, however the acid should be neutralized before analysis.

INTERFERENCES: The following ions interfere in concentrations greater than those listed.

| Ion | mg/L | Ion | mg/L |
|----------|------|-----------------------|------|
| Cd(II) | 1 | Cr(III) | 10 |
| Al (III) | 5 | Ni(II) | 20 |
| Mn (II) | 5 | Co (II) | 30 |
| Fe (III) | 7 | CrO ₄ (II) | 50 |
| Fe (II) | 9 | | |

PROCEDURE

A. PREPARATION OF DILUTE ZINC INDICATOR SOLUTION

1. Use a pipet [0352] to add exactly 5.0 mL of *Zinc Indicator Solution [6314] to a 10 mL graduated cylinder [0416]. The bottom of the curved surface [the meniscus] of liquid should be at 5.0 mL mark. Pour this into the bottle labeled *Dilute Zinc Indicator Solution" [6321-MT-G].
2. Use unrinsed graduated cylinder to add 10.0 mL and then 7.8 mL [total of 17.8 mL] of *Methyl Alcohol [6319] to bottle labeled *Dilute Zinc Indicator Solution [6321]. Cap and mix ingredients in this bottle. Do not leave this bottle uncapped.

B. DETERMINATION OF ZINC

1. Press and hold  until colorimeter turns on.
2. Press  to select **Testing Menu**.
3. Select **All Tests** [or another sequence containing **097 Zinc LR**] from **Testing Menu**.
4. Scroll to and select **097 Zinc LR** from menu.
5. Rinse a clean tube [0290] with sample water. Fill to the 10 mL line with sample.
6. Insert tube into chamber, close lid and select **Scan Blank**. [See Note]
7. Remove tube from colorimeter. Use 0.1 g spoon [0699] to add one measure of Sodium Ascorbate Powder [6316]. Use 0.5 g spoon [0698] to add one measure of *Zinc Buffer Powder [6315]. Cap and shake vigorously for 1 minute. Some undissolved buffer may remain in the bottom of the tube.
8. Add 3 drops of *Sodium Cyanide, 10% [6565]. Cap and mix.
9. Use the 1 mL pipet assembly to add 1 mL of *Dilute Zinc Indicator Solution [6321]. Cap and mix.
10. Use a second plain pipet [0352] to add 4 drops of *Formaldehyde Solution, 37% [5128]. Cap and mix by inverting 15 times.
11. Insert tube into chamber, close lid and select **Scan Sample**. Record result.
12. Press  to turn colorimeter off or press  to exit to a previous menu or make another menu selection.

NOTE: For best possible results, a reagent blank should be determined to account for any contribution to the test result by the reagent system. To determine the reagent blank, follow the above test procedure to scan a distilled or deionized water blank. Then follow the above procedure to perform the test on a distilled or deionized water sample. This test result is the reagent blank. Subtract the reagent blank from all subsequent test results of unknown samples. It is necessary to determine the reagent blank only when a new lot number of reagents is obtained.

APPENDIX

Ammonia in water occurs in two forms: toxic unionized ammonia (NH_3) and the relatively non-toxic ionized form, ammonium ion (NH_4^+). This test method measures both forms as ammonia-nitrogen ($\text{NH}_3\text{-N}$) to give the total ammonia-nitrogen concentration in water. The actual proportion of each compound depends on temperature, salinity, and pH. A greater concentration of unionized ammonia is present when the pH value and salinity increase.

1. Consult the table below to find the percentage that corresponds to the temperature, pH, and salinity of the sample.
2. To express the test result as ppm Unionized Ammonia Nitrogen ($\text{NH}_3\text{-N}$), multiply the total ammonia-nitrogen test result by the percentage from the table.
3. To express the test result as ppm Ammonia Nitrogen ($\text{NH}_3\text{-N}$), subtract the unionized ammonia-nitrogen determined in step 2 from the total ammonia-nitrogen.

| pH | 10°C | | 15°C | | 20°C | | 25°C | |
|-----|------|-------|------|-------|-------|-------|-------|-------|
| | FW1 | SW2 | FW | SW | FW | SW | FW | SW |
| 7.0 | 0.19 | — | 0.27 | — | 0.40 | — | 0.55 | — |
| 7.1 | 0.23 | — | 0.34 | — | 0.50 | — | 0.70 | — |
| 7.2 | 0.29 | — | 0.43 | — | 0.63 | — | 0.88 | — |
| 7.3 | 0.37 | — | 0.54 | — | 0.79 | — | 1.10 | — |
| 7.4 | 0.47 | — | 0.68 | — | 0.99 | — | 1.38 | — |
| 7.5 | 0.59 | 0.459 | 0.85 | 0.665 | 1.24 | 0.963 | 1.73 | 1.39 |
| 7.6 | 0.74 | 0.577 | 1.07 | 0.836 | 1.56 | 1.21 | 2.17 | 1.75 |
| 7.7 | 0.92 | 0.726 | 1.35 | 1.05 | 1.96 | 1.52 | 2.72 | 2.19 |
| 7.8 | 1.16 | 0.912 | 1.69 | 1.32 | 2.45 | 1.90 | 3.39 | 2.74 |
| 7.9 | 1.46 | 1.15 | 2.12 | 1.66 | 3.06 | 2.39 | 4.24 | 3.43 |
| 8.0 | 1.83 | 1.44 | 2.65 | 2.07 | 3.83 | 2.98 | 5.28 | 4.28 |
| 8.1 | 2.29 | 1.80 | 3.32 | 2.60 | 4.77 | 3.73 | 6.55 | 5.32 |
| 8.2 | 2.86 | 2.26 | 4.14 | 3.25 | 5.94 | 4.65 | 8.11 | 6.61 |
| 8.3 | 3.58 | 2.83 | 5.16 | 4.06 | 7.36 | 5.78 | 10.00 | 8.18 |
| 8.4 | 4.46 | 3.54 | 6.41 | 5.05 | 9.09 | 7.17 | 12.27 | 10.10 |
| 8.5 | 5.55 | 4.41 | 7.98 | 6.28 | 11.18 | 8.87 | 14.97 | 12.40 |

¹ Freshwater data from Trussel [1972].

² Seawater values from Bower and Bidwell [1978].

Salinity for Seawater values = 34% at an ionic strength of 0.701m.

FOR EXAMPLE:

If a fresh water sample at 20°C has a pH of 8.5 and the test result is 1.0 ppm as Total Ammonia-Nitrogen:

1. The percentage from the table is 11.18% [or 0.1118].
2. 1 ppm Total Ammonia-Nitrogen x 0.1118 = 0.1118 ppm Unionized Ammonia-Nitrogen.

| | | |
|-----------------------------------|---|-------------------|
| 3. Total Ammonia-Nitrogen | | 1.0000 ppm |
| <u>Unionized Ammonia-Nitrogen</u> | - | <u>0.1118 ppm</u> |
| Ionized Ammonia-Nitrogen | = | 0.8882 ppm |