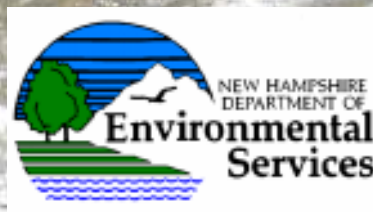


NH Volunteer River Assessment Program

Water Quality Monitoring Field Sampling Protocols for Volunteer Monitors

**LaMotte 2020 Turbidity Meter, Orion 210A pH Meter,
YSI 95 Dissolved Oxygen Meter, YSI 30 Conductivity Meter**



New Hampshire Department of Environmental Services

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Concord, NH 03301

<http://des.nh.gov/organization/divisions/water/wmb/vrap/index.htm>

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Sampling Reminders: Remember...

- Calibrate the pH and DO meters before each measurement!
- Do not turn off the DO meter until the end of the day!
- Run a replicate sample once a day!
- Test the pH 6.0 Buffer, Turbidity DI Blank, and Zero Oxygen once a day!
- Check the Conductivity meter with the known standard at the beginning & end of the day!
- Calibrate the Turbidity meter with the 1.0 NTU standard at the beginning of the day and check the Turbidity meter with the 1.0 NTU standard at the end of the day.
- Rinse everything with DI water - a lot!

Introduction: What is VRAP?

In 1998, the New Hampshire Department of Environmental Services (NHDES) established the New Hampshire Volunteer River Assessment Program (VRAP) as a means of expanding public education of water resources in New Hampshire. VRAP promotes awareness and education of the importance of maintaining water quality in rivers and streams. VRAP was created in the wake of the success of the existing New Hampshire Volunteer Lake Assessment Program (VLAP), which provides educational and stewardship opportunities pertaining to lakes and ponds to New Hampshire's residents.



Today, VRAP continues to serve the public by providing water quality monitoring equipment, technical support, and educational programs to volunteer groups on numerous rivers and watersheds throughout the state. These groups conduct water quality monitoring on an ongoing basis and increase the amount of river water quality information available to local, state and federal governments, which allows for effective financial resource allocation and watershed planning.

This manual is meant to be used as a guide for VRAP monitors. Take this manual with you in the field as a reminder of the proper sampling procedures. Each meter has a very important calibration procedure, which must be followed to ensure the sampling results are as accurate as possible. If you encounter problems during calibration, refer to the manufacturer's operation manuals or contact the VRAP staff.

This manual is designed to compliment the annual VRAP volunteer training workshop and is not a replacement for attending a training workshop. VRAP staff are also available to visit with each group in the field. Please contact us to schedule a visit.

Informational Resources

- **Water Quality Monitoring Field Sampling Protocols:**
http://des.nh.gov/organization/divisions/water/wmb/vrap/documents/protocols_old_meters.pdf
- **VRAP Field Data Sheet:**
http://des.nh.gov/organization/divisions/water/wmb/vrap/documents/field_data_sheet.pdf
- **A Quick-Reference Guide to Water Quality Standards**
http://des.nh.gov/organization/divisions/water/wmb/vrap/documents/wq_standards.pdf
- **Interpreting VRAP Water Quality Parameters:**
http://des.nh.gov/organization/commissioner/pip/publications/wd/documents/vrap_parameters.pdf
- **Troubleshooting Guide to VRAP Water Quality Meters:**
http://des.nh.gov/organization/divisions/water/wmb/vrap/documents/troubleshooting_meters.pdf

For More Information

- **Jen Drociak, VRAP Coordinator**
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Equipment & Supplies

Please ensure that you have all necessary equipment and supplies before each sampling event. An **Equipment & Supply Checklist** can be found in each VRAP kit and the VRAP website at <http://des.nh.gov/organization/divisions/water/wmb/vrap/documents/checklist.pdf>.

Necessary Equipment & Supplies

Turbidity

- 1.0 NTU Standard
- 10.0 NTU Standard
- DI Blank (0.0 NTU Standard)
- Sample Vial
- Turbidity Meter

pH

- pH Electrode Storage Solution
- pH Electrode Filling Solution
- 4.0 pH Buffer
- 6.0 pH Buffer
- 7.0 pH Buffer
- pH Meter & Electrode Probe

Dissolved Oxygen/Temperature

- Zero DO H₂O Dissolved Oxygen Standard
- Extra DO Membranes and Membrane Solution
- DO/Temperature Meter

Specific Conductance

- Conductivity Standard
- Conductivity Meter

Other Supplies

- Sample Container (Plastic)
- Deionized (DI) Water Bottle
- Kimwipes
- Cooler & Ice Packs (*If sampling for laboratory parameters*)
- Plastic Sample/Laboratory Bottles (*If sampling for laboratory parameters*)
- Clipboard
- Pencils & Pens (*Pencils will write in wet weather*)
- Permanent Marker
- Extra Batteries (AA and 9V)
- Extra Field Data Sheets
- Bucket & Rope Coil

Safety in the Field

It is important to ensure that adequate safety precautions are taken in the field. The most common hazards associated with conducting VRAP sampling are listed below along with recommended safety precautions. This is not a comprehensive safety protocol, but rather a guideline to ensure safe field conditions.

General

- Always monitor with at least one person.
- Listen to weather reports. Caution should be exercised if severe weather is forecasted.
- Have a first-aid kit accessible.
- If at any time you feel uncomfortable about the condition of the stream or your surroundings, monitoring efforts should be terminated.

Stream Safety

- Do not enter the water if the stream is at flood stage.
- If you must cross the stream, use a walking stick to steady yourself and to probe for deep water and unstable terrain.
- Streambeds composed of coarse substrates and/or bedrock can be slippery and have deep pools.
- Streambeds composed of finer substrates can prove treacherous in areas where mud, silt, or sand creates unstable terrain.

Poison Ivy

- Watch for poison ivy, poison oak, sumac, and other types of vegetation in your area that can cause rashes and irritation.

Mosquitoes

Mosquito-borne illnesses such as West Nile virus and Eastern Equine Encephalitis (EEE) have become more prevalent. The following steps can reduce the incidence of mosquito bites.

- Apply an effective repellent to exposed skin and clothing.
- Wear long-sleeves, long pants, and socks when outdoors.

Ticks

Ticks, which can carry the Lyme disease bacterium, prefer wooded and bushy areas with high grass and abundant leaf litter. Extra precaution should be taken in May, June, and July, when ticks that transmit Lyme disease are most active.

- Wear long pants, long sleeves, and long socks to keep ticks off your skin. Light-colored clothing will help you spot ticks more easily. Tucking pant legs into socks or boots and tucking shirts into pants help keep ticks on the outside of clothing.
- Perform daily tick checks after being outdoors. Inspect all parts of your body carefully. Remove ticks immediately using fine-tipped tweezers.

Sun/Heat

- Dress for the heat. Wear a hat and lightweight, light-colored clothing. Light colors will reflect away some of the sun's energy.
- Drink water.
- If you recognize that you or someone else is showing the signals of a heat-related illness, stop activity and find a cool place.

Quality Assurance & Quality Control

Collecting Data

In order for VRAP data to be used in the assessment of New Hampshire's surface waters, the data must meet quality control guidelines as outlined in the VRAP Quality Assurance Project Plan (QAPP). The VRAP QAPP was approved by NHDES and reviewed by EPA in the summer of 2003. The QAPP is reviewed annually and is officially updated and approved every five years. The VRAP Quality Assurance/Quality Control (QA/QC) measures include a six-step approach to ensuring the accuracy of the equipment and consistency in sampling efforts.

- **Calibration:** Prior to each measurement, the pH and DO meters must be calibrated. Conductivity and turbidity meters are checked against a known standard before the first measurement and after the last measurement of the day.
- **Replicate Analysis:** A second measurement by each meter is taken from the original sample at one of the stations during the sampling day. If the same sampling schedule is used throughout the monitoring season, the replicate analysis should be conducted at different stations. Replicates should be measured within 15 minutes of the original measurements. If more than one team is out sampling each team should do a replicate analysis. Record the measurements on the bottom front page of the VRAP Field Data Sheet.
- **6.0 pH Standard:** A reading of the pH 6.0 buffer is recorded at one of the stations during the sampling day. If the same sampling schedule is used throughout the monitoring season, the 6.0 pH standard check should be conducted at different stations. Record the reading, station, and time on the bottom front page of VRAP Field Data Sheet. Another calibration does not need to be done before this measurement as it is intended to detect drift in the meter
- **Zero Oxygen Solution:** A reading of a zero oxygen solution is recorded at one of the stations during the sampling day. If the same sampling schedule is used throughout the monitoring season, the zero oxygen standard check should be conducted at different stations. Record the reading, station, and time on the bottom front page of the VRAP Field Data Sheet. The value should be below 1.0 mg/L.
- **DI (De-Ionized) Turbidity Blank:** A reading of the DI turbidity blank (0 NTU) is recorded at one of the stations during the sampling day. If the same sampling schedule is used throughout the monitoring season, the blank check should be conducted at different stations. Record the reading, station, and time on the bottom front page of the VRAP Field Data Sheet.
- **End of the Day Conductivity and Turbidity Meter Check:** At the conclusion of each sampling day, the conductivity and turbidity meters are re-checked against a known standard. Record the readings on the bottom front page of the VRAP Field Data Sheet.

Completing the VRAP Field Data Sheet

One person should perform all of the data entry on the VRAP Field Data Sheet for that particular sampling date so that entries are consistent.

When completing the VRAP Field Data Sheet, please:

1. Enter data neatly, legibly, and thoroughly so that there is not any missing, incomplete, or incorrect information.
2. Ensure that all sections of the VRAP Field Data Sheet are completed before moving to the next sampling station.
3. If a parameter cannot or was not measured (i.e., instrument failure), record the reason for the missing data under the “Comments” section on the back of the VRAP Field Data Sheet.
4. When the observations have been completed for a station, review the VRAP Field Data Sheet to ensure that all the necessary data are accurately recorded and that everything is legible.
5. If an error is made in recording data, do not erase. Draw a single pencil line through the incorrect values and enter the correct values.
6. Lastly, remember to legibly print the first and last name of each volunteer for that particular sampling date on the VRAP Field Data Sheet, so that we can give all volunteers proper credit!

Collecting Samples for Field & Laboratory Analysis

Order of Field Tests

- Pour and/or collect samples for laboratory analysis
- Turbidity (*LaMotte 2020*)
- pH (*Orion*)
- Water Temperature, Dissolved Oxygen (*YSI 95*)
- Specific Conductance (*YSI 30*)

Collecting Samples for Field Analysis

Please label all bottles *prior to filling them* with the date and time of collection, NHDES Station Name/ID, collector’s initials and (when necessary) the test(s) requested. It is important to use assigned NHDES Station IDs.

When sampling, begin with the most downstream sampling station so that sampling activities do not affect water quality at downstream stations. It is also important to pour off water for laboratory test(s) before sampling field water quality.

Method 1: Bridge Sampling

1. Lower the bucket from the **upstream** side of the bridge into the river and fill the bucket $\frac{1}{4}$ full of water.
2. Pull the bucket up, swish the water around in the bucket to rinse, and dump the water off the downstream/opposite side of the bridge. Repeat this process two more times.
3. Return the bucket to the river on the upstream side of the bridge and fill the bucket $\frac{1}{2}$ - $\frac{3}{4}$ full of water as slowly as possible (you may wish to weight one side of the bucket).

4. Pull the bucket up and carry to a safe location (away from the road!) for analyses. Ensure not to bump the bucket against the bridge for this may introduce additional oxygen and sediment.

Method 2: Offshore Sampling

1. Carefully wade out into the river as close as possible to the center. Do not enter water above your waist, and do not enter the water if there is any concern for your safety. Be sure to have someone on shore that knows where you are. NHDES highly recommends that volunteers wear an appropriate personal floatation device when working in or near the water.
2. Position yourself facing upstream and rinse the bucket in the river three times. **Do not collect the water that is running over your legs/boots.** Discard the rinse water behind yourself.
3. With the bucket facing upstream and held in front of your body, slowly dip the lip of the bucket into the flowing water and allow the bucket to fill. Rivers receive oxygen from the atmosphere through mixing. Just as riffles and rapids increase the oxygen in a river or stream, rushing water over the side of the bucket will add oxygen to the sample and yield inaccurate readings.
4. Carefully return to shore with the bucket $\frac{1}{2}$ - $\frac{3}{4}$ full and place it on the bank for immediate analysis. Do not place the bucket on the hot tar/asphalt or in direct sunlight.

Collecting Samples for Laboratory Analysis

If you are collecting *E. coli* samples for laboratory analysis the bottles may be filled using either of the following two methods:

Method 1: Bridge Sampling

1. Fill a sterilized *E.coli* bottle using the water in the bucket. This should be done immediately after the bucket sample is collected. **Avoid touching the neck, inside the bottle or cap to prevent contamination.**
2. Replace the cap on the bottle. **Please leave some air at the top of the bottle to allow sample to be shaken by lab staff.**

Method 2: Offshore Sampling

1. Wade out into the river and sample about knee deep or midway between the top and bottom of a flowing stream.
2. Remove the cap of a sterilized *E.coli* bottle when you are ready to collect the sample. **Avoid touching the neck, inside the bottle or cap to prevent contamination.**
3. Point the mouth of the bottle down towards the water's surface.
4. Using a continuous "U"-shaped motion, thrust the bottle under the water's surface and fill in one continuous upstream motion (*away from you*), turning the bottle right side-up at the bottom of the "U". In this fashion, the water will flow into the bottle, then over your hand. If sampled in a downstream fashion, the water would flow over your hand then into the bottle causing contamination from the sampler.
5. Replace the cap on the bottle and carry the sample to shore. **Please leave some air at the top of the bottle to allow sample to be shaken by lab staff.**

This technique should not be used for bottles which have acid preservatives in them such as **total phosphorous, total Kjeldahl nitrogen** and **metals**. The preservatives are very strong acids which will burn the skin on contact. These bottles should only be filled by from another bottle or a bucket. Preserve the bottles properly (in a cooler on ice), and submit the samples to the NHDES Laboratory Services within the sample holding time appropriate to each test. Always return samples to the NHDES Laboratory Services before 3 pm (Monday – Thursday) or before 12 pm on Friday.

Completing the NHDES Laboratory Services Login & Custody Sheet

VRAP staff are available to assist you with the login procedure for laboratory samples. If you know you will be relinquishing laboratory samples, please contact Jen Drociak. However, during the sampling season VRAP staff are frequently out of the office. The following is a guideline for how to login in samples that must be logged in immediately due holding time requirements. For samples with longer hold times (chloride, TP, TKN) you can ask the laboratory staff to put them in the refrigerator and ask a VRAP staff person to log them in.

- **Client ID:** Please leave blank.
- **Lab Account (Billing):** Please write in either your group's laboratory account number or the VRAP lab account number which is 05-0022518
- **One Stop Project:** Please write in "VRAP".
- **NHDES Site Number:** Please leave blank.
- **Description:** Please write in your river or watershed name.
- **Town:** Please leave blank.
- **Collected By:** Please write in the name and phone number of the person who should be contacted if there are any questions about the samples.
- **Contact & Phone Number:** Please write in "Jen Drociak, 271-0699"
- **Sample Location/Station ID:** Please use NHDES VRAP Station IDs. This is important because if the ID is not exactly the same as the one in our database it holds up the reporting of the results.
- Common mistakes include not including a zero where it is needed (1-BKB instead of 01-BKB), adding text to the end of the station ID (00M-BKB Front Street instead of just 00M-BKB), and forgetting to include the dash between the number and Station ID or REP for replicates (01BKB instead of 01-BKB or 01-BKBREP instead of 01-BKB-REP).
- It is important to remember that samples with mistakes in the Station ID hold up the reporting of the entire sampling batch.
- **Date/Time Sampled:** Please write the date and time of each sample collected.
- **# of Containers:** Please list the number of sample bottles per station.
- **Matrix:** Please write "AQ" for aqueous.
- In the columns to the right of the "Matrix" column, please write in the parameter to be analyzed. (For example TP, TKN, *E.coli*, chloride, nitrate, etc.)
- **Sampler Comments:** Please leave blank.
- **Lab Login #.** Please leave blank.
- **Relinquished By:** Please sign your name on the top line and leave the bottom line blank.
- **Date & Time:** Please write the date and time you relinquished the samples on the top line and leave the bottom line blank.
- **Received By:** Please leave blank. This will be completed by the NHDES Laboratory Services personnel.
- Please fill in the number of pages (Example: Page 1 of 1) at the bottom of the sheet.

Turbidity: *LaMotte 2020 Turbidity Meter*

I. Before Sampling:

Perform & Record the Initial Turbidity Meter Check Value

1. To turn the meter on, press the **READ** button. A triangle should be displayed in the upper left corner of the display screen.

Note: If the triangle is not displayed, turn the meter off by holding the **READ** button down until the screen reads **OFF**. Press the **CAL** button while pressing the **READ** button to turn the meter on. If the triangle does not appear, gently repeat this step until it does. This step places the meter into “EPA mode”, which means the meter will automatically round readings to US Environmental Protection Agency standards for uniform data reporting.



2. From the turbidimeter case (black, separate from the VRAP Kit) remove the standard vial marked “**1.0 NTU**” and carefully wipe off any water, dust and/or fingerprints from the vial with a Kimwipe only.
3. Open the lid of the turbidimeter and align the etched arrow on the “1.0 NTU” vial with the arrow under the meter lid. Insert the vial into the chamber and close the lid.
4. Press the **READ** button.
5. If the displayed value is the same as the 1.0 NTU Standard, calibration is unnecessary at this time. Record 1.0 on the top left of the VRAP Field Data Sheet as the “**Initial Turbidity Meter Check Value**”.

II. Before Sampling: Calibrate the Meter

Note: The turbidimeter needs to be calibrated once prior to the first measurement and checked once after the last measurement at the end of the day. Please turn the meter off when not in use to conserve battery power.

1. If the displayed value differs from the standard value (1.0 NTU), record the value on the top left of the VRAP Field Data Sheet as the “**Initial Turbidity Meter Check Value**”, and push the **CAL** button until **CAL** is displayed. This should take around 5 seconds. Release the button. The display will flash.
2. Adjust the value with the up and down buttons until the value of the standard is displayed.
3. Push the **CAL** button again to complete calibration.

III. Measuring Turbidity

1. **Rinse** the small plastic sample container with DI water. Then rinse the same container twice with a small amount of river water from the bucket.

2. Slowly pour sample water from the bucket into the plastic sample container (1/2 full) to avoid adding bubbles to the sample.
3. From the turbidimeter case remove the sample vial labeled “Sample” or “S”.
4. **Rinse** the sample vial once with DI water and then twice with river water from the plastic sample container.
5. Fill the sample vial with river water by carefully and slowly pouring the water down the side of the sample vial to avoid introducing any bubbles.
6. Wipe any water, dust and/or fingerprints off the sample vial with a Kimwipe. **Note: Any residue on the vials will interfere with an accurate turbidity reading. Anything other than Kimwipes may scratch the vials, causing inaccurate readings.**
7. If the meter is off, turn it on by pressing the **READ** button.
8. Open the lid of the turbidimeter and align the etched arrow on the cleaned sample vial with the arrow under the turbidimeter lid, and 8) Close the lid. Press READ.
9. **Record** the displayed turbidity reading on the VRAP Field Data Sheet.
10. If the turbidity value is great than 10 NTU you should recalibrate the meter with the 10 NTU standard and take another reading. This will give a more accurate measurement of how high the turbidity level is. If you do recalibrate with the 10 NTU standard, be sure to indicate this under the “Comments” section on the back the VRAP Field Data Sheet. **Recalibrate with the 1.0 NTU at the next station to prevent the readings from being artificially elevated.**
11. Turn the meter **OFF** by holding the **READ** button down until the screen reads **OFF**.

IV. QA/QC Meter Check

1. At one of the stations during the sampling day read the DI Turbidity Blank (0.0 NTU) standard. If the same sampling schedule is used throughout the monitoring season, the DI turbidity blank check should be conducted at different stations.
2. **Record** the value, station, and time on the VRAP Field Data Sheet.

V. End of the Day Meter Check

1. At the end of the day, read the 1.0 standard.
2. **Record** the value under the “**End of Day Meter Check**” on the VRAP Field Data Sheet.
3. Turn the meter off.
4. **Rinse** the sample vial with DI water and fill the vial with DI water for storage.

pH: Orion 210A pH Meter

I. Before Sampling: Calibrate the Meter

Note: The pH meter must be calibrated prior to each measurement (at each station) including a replicate.

There are two different types of pH probes that can be used with this meter. One has a blue plug on the side which covers a small hole. This probe is refillable and must be checked to be sure that the reference solution inside is at the proper level. The second type of probe has no hole or blue plug and is not refillable.



1. Ensure electrode connections are properly fastened in the appropriate ports.
2. Unscrew the cap on the electrode solution storage container and remove the pH probe (the screw cap can remain on the electrode). Clean any salty deposits off by rinsing the probe with DI water. Blot dry with a Kimwipe. **CAUTION:** Be sure to never touch the glass bulb on the bottom; even with a Kimwipe.
3. If necessary, remove the blue plug from the hole in the side of the probe and refill the electrode with pH electrode filling solution (it may have spilled out). Fill to just below the hole. Remove the plug during calibration and sampling. Return the blue plug for storage and travel between stations.
4. Press the **POWER** key to turn the meter on. All the features of the display will light up. Then the model number, "210", will be displayed. Once all power up procedures are complete the meter advances to **MEASURE** mode.
5. Select calibration mode by pressing the **MODE** key until **CALIBRATE** is displayed.

First Standard to Test (7.0 pH Buffer):

6. The last calibration standards or buffers used will be displayed (7 and 4). Press **YES** to accept this setting. "**P1**" will be displayed in the lower display field and the standard measurement will be displayed in the main display field. A black arrow will be displayed on the bottom of the screen pointing to 7 indicating that the meter is ready to measure the 7.0 buffer.
7. **Rinse** the electrode with DI water and blot dry with a Kimwipe.
8. **If necessary, remove the blue plug** from the side of the electrode and immerse the probe into the 7.0 buffer (yellow solution). Allow at least one inch of the pH electrode filling solution volume inside the probe to remain above the standard level during measurement/calibration. The end of the probe must be completely immersed into the sample.
9. When **READY** is displayed the electrode is stabilized. Press **YES** while **READY** is displayed. The **READY** may flash on and off at first but will soon stabilize.

Second Standard to Test (4.0 pH Buffer):

Note: "**P2**" will be displayed in the lower display field indicating the meter is ready for the second standard. Make sure "**P2**" appears before continuing. If it does not appear, keep the electrode in the 7.0 buffer until **READY** appears again and press **YES**. A black arrow will be displayed on the bottom of the screen pointing to 4 indicating that the meter is ready to measure the 4.0 buffer.

10. Remove the electrode from the 7.0 buffer, **rinse** it with DI water and blot dry with a Kimwipe.

11. Place the electrode in the second (4.0) buffer (red solution). When **READY** is displayed press **YES. WATCH!**
12. **SLP** (Slope Value) will appear in the lower display field and the current electrode slope will be displayed in the main field. Record the slope on the VRAP Field Data Sheet as the pH Calibration Slope. An acceptable range for the slope is 92-102%. If you get a slope outside of this range repeat the calibration procedure and check the batteries. If the slope is still outside of the range continue to sample but notify VRAP staff immediately. If you miss the slope you must go back and repeat the calibration procedure. pH data without calibration slopes will not pass our quality control process.
13. The meter will proceed to the measure mode. **MEASURE** is displayed above the main display field. Remove the electrode from the 4.0 buffer, rinse with DI water and blot dry. The meter is now ready for use.

II. Measuring pH

Note: The pH meter must be calibrated prior to each measurement (at each station) including a replicate.

1. Remove the probe, **rinse** with DI water and blot the plastic areas dry with a Kimwipe. **CAUTION:** Be sure to never touch the glass bulb/measurement end; even with a Kimwipe.
2. If necessary, remove the blue plug from the probe and ensure it is clean.
3. Immerse the pH probe into the small plastic sample container. The meter should be in **MEASURE** mode.
4. Submerge the bottom two inches of the electrode and agitate by slowly moving the electrode back and forth in the sample until the pH value stabilizes.
5. Wait for the **READY** indicator to be displayed and record the value on the VRAP Field Data Sheet. The READY indicator may blink on and off. **It is important to wait until drifting of the pH value has stopped before recording measurement.**
6. **Rinse** the probe with DI water and return it to the electrode solution storage container. Ensure the pH electrode storage container is filled about halfway with pH storage solution. Be careful not to push the electrode against the bottom of the container as this could damage the electrode. **Never store pH probe in DI water!**
7. Turn the meter **OFF**. Place the electrode **VERTICALLY** in the storage solution container, being careful not to hit the bottom of the container with the probe and screw the cap on the container. If necessary, ensure the blue plug is secured in the probe, even with a little masking tape or a rubber band, and set the meter in the kit until you are ready to take a reading.

III. QA/QC Meter Check

1. At one of the stations during the sampling day read the 6.0 pH buffer. If the same sampling schedule is used throughout the monitoring season, the blank check should be conducted at different stations.
2. **Record** the value, station, and time on the VRAP Field Data Sheet.

IV. End of the Day

1. Turn the meter off. **Rinse** the probe with DI water and blot dry with a Kimwipe.
2. Return the probe to the storage solution container. Store probe upright.

Dissolved Oxygen & Water Temperature: YSI 95 Dissolved Oxygen Meter

I. Before the Meter is Turned On:

Check the Dissolved Oxygen Membrane & Calibration Chamber

To ensure the probe remains moist inside the meter calibration/storage chamber, pull the probe out of the chamber and add a few drops of deionized (DI) water to the sponge at the bottom of the calibration/storage chamber. Turn the meter on its side to allow any excess water to drain out of the chamber. *This step will only be necessary once per sampling day, but be sure the sponge in the calibration/storage chamber is moist before storage.* Be careful not to over-wet the sponge. Calibrate the meter after 15 minutes and after any excess “puddled” water is drained from the chamber.

Note: The wet sponge creates a 100% saturated air environment within the chamber for ideal calibration conditions. Ensure that the sensor does not contact the wet sponge by inserting the probe only until the rubber seal is flush with the outer edge of the chamber.



II. If Bubbles **ARE** Detected: Change the Membrane

Membrane life depends on usage. Membranes will last a long time if installed properly and treated with care. Erratic readings are a result of loose, wrinkled, damaged, or fouled membranes, or from large bubbles in the electrolyte reservoir. If erratic readings or evidence of membrane damage occurs, you should replace the membrane and the solution, following Step A through Step G. (Record the performed maintenance under the “Comments” section on the back of the VRAP Field Data Sheet). To keep the membrane from drying out, store the probe in the calibration chamber with the damp sponge.

- A. Unscrew and remove the black protective cage.
- B. Unscrew and remove the old membrane cap.
- C. Thoroughly rinse the sensor tip (gold and silver areas) with DI water.
- D. Hold the membrane cap upside down and add enough drops of the membrane probe solution to form a meniscus in the probe membrane cap.
- E. Tap the bottom of the cap with your finger a few times to remove any trapped air bubbles.
CAUTION: Do not touch the membrane surface.
- F. Screw the membrane cap onto the probe tightly by hand (to prevent leakage of probe solution). A small amount of probe solution should overflow.
- G. Shake off any excess probe solution and rinse the sensor thoroughly with DI water to prevent corrosion.

III. Turn the Meter ON and Wait 15 Minutes

1. Turn the meter on by pressing the **ON/OFF** button. **Ensure the meter has been turned ON with the probe in its chamber for at least 15 minutes before calibrating.**
2. Record the time the dissolved oxygen meter was turned on - on the upper right front page of the VRAP Field Data Sheet.

IV. Before Sampling: Calibrate the Meter for Dissolved Oxygen

Note: The Dissolved Oxygen/Temperature meter must be calibrated prior to each dissolved oxygen measurement (at each station) including a replicate.

1. **Record** the time of the first dissolved oxygen calibration on the upper right front page of the VRAP Field Data Sheet.
3. Press the **MODE** button until dissolved oxygen is displayed in % saturation. The **UP** button on the YSI 95 can also be used to toggle back and forth between % saturation and mg/L.
4. Press and release both the **DOWN** and **UP** arrow buttons (DOWN slightly prior to UP) to enter the DO/Temperature meter calibration menu. You will see **CAL** in the lower left hand corner when you have successfully entered calibration mode.
5. The screen will prompt you to enter the local altitude in hundreds of feet. Use the **UP** and **DOWN** arrows to adjust the value appropriately (For example, entering a 12 indicates 1200 feet above sea level) and press **ENTER**.
6. The screen will then prompt you to enter the salinity of the sample you will be measuring. Be sure the screen reads zero and press **ENTER**.
7. **Record** the calibration value (displayed on the bottom right-corner of the LCD screen) on the VRAP Field Data Sheet. The calibration value will vary with altitude and thus may be different at each station if the altitude varies.
8. Press **ENTER** again. The display should read **SAVE** and then return to normal measurement mode.
9. Wait approximately one minute for dissolved oxygen % saturation to stabilize. Once it has stabilized, record the dissolved oxygen % saturation (chamber reading) on the VRAP Field Data Sheet. If drift occurs (goes up or down by more than 5%) ensure you have waited long enough for the reading to stabilize. If drift still occurs, recalibrate.

Note: The Dissolved Oxygen/Temperature meter should remain on until the last station has been sampled. If the meter is turned off prior to the end of the sampling day, the meter must be turned on and allowed a 15-minute warm-up period, with the probe in its chamber, prior to calibration and additional sampling.

V. Measuring Water Temperature & Dissolved Oxygen

Note: The Dissolved Oxygen/Temperature meter should remain on until the last station has been sampled. If the meter is turned off prior to the end of the sampling day, the meter must be turned on and allowed a 15-minute warm-up period, with the probe in its chamber, prior to calibration and additional sampling. Remember, the dissolved oxygen/temperature meter must be calibrated prior to each dissolved oxygen measurement.

1. Remove the probe from the calibration chamber and **rinse** the probe and cable (approximately 6 inches) with DI water. If necessary, press the **UP** button until dissolved oxygen is displayed in % saturation.
2. Submerge the probe into the bucket and agitate by slowly moving the probe back and forth in the sample until the water temperature stabilizes. **Record** the water temperature (°C) on the VRAP Field Data Sheet.
3. After the water temperature has stabilized, wait for the dissolved oxygen (% saturation) to stabilize. Once it is stable, record the value on the VRAP Field Data Sheet.
4. Immediately press the **DOWN** button and immediately record the value for dissolved oxygen concentration (mg/L) on the VRAP Field Data Sheet.

VI. QA/QC Meter Check

1. At one of the stations during the sampling day read the Zero Dissolved Oxygen Standard (% saturation and mg/L). If the same sampling schedule is used throughout the monitoring season, the blank check should be conducted at different stations.
2. **Record** the value, station, and time on the VRAP Field Data Sheet.
3. **Rinse** the probe with DI water and return it to the storage chamber.

VII. End of the Day

1. **Rinse** the probe with DI water.
2. Return the probe to the chamber with a wet sponge. Drain any water from the chamber.
3. Turn the meter off.

Specific Conductance: YSI 30 Conductivity Meter

I. Before Sampling:

Perform & Record the Initial Conductivity Meter Check Value

1. Turn the meter on by pressing the **ON/OFF** button. The meter will activate all segments of the display screen for a few seconds, followed by a self-test. If the meter is not functioning properly, a continuous error message will be displayed.
2. Ensure the meter is in the temperature-compensated specific conductance mode by pressing the **UP** or **DOWN** buttons until the °C is flashing to indicate this mode.
3. **Rinse** the probe with DI water and blot dry with a Kimwipe. Gently shake the probe to remove water from the oval upper conductivity opening.
4. Submerge the entire probe in the **conductivity standard solution**, and allow to stabilize. Ensure there is enough solution to cover the top opening of the probe.
5. Record the **“Initial Conductivity Meter Check Value”** on the top left of the VRAP Field Data Sheet. A 20% error regardless of the standard used (1,600 – 2,400 μ S for 2,000 μ S standard, 160–240 μ S for 200 μ S standard or 80 μ S - 120 μ S for 100 μ S standard) is acceptable. If the reading is outside of this range, please check again with new standard if available. If new standard is unavailable please sample anyway and contact VRAP staff as soon as possible.



II. Measuring Specific Conductance

1. Immerse the probe in the sample and ensure it is deep enough to cover the entire probe. Do not allow the probe to touch any solid object or the bottom of the bucket while you are taking readings. It is also important that there are no air bubbles on/in the electrode. To dislodge any bubbles, gently move the electrode through the water before recording the measurement.
2. Agitate by slowly moving the probe back and forth in the sample until the specific conductance value stabilizes. **Record** the specific conductance value on the VRAP Field Data Sheet.

Note: Please keep the reference temperature of the meter at 25 °C at all times of the year, instead of changing it to 20 °C during the winter months and 25 C after April 1st. If you receive an error message of “LErr”, switch modes to conductivity (the °C will not flash on and off in this mode), and take that measurement instead. Cross out the word “Specific” on the VRAP Field Data Sheet and write “Actual”.

3. Rinse the probe and return it to the storage chamber between measurements. Please turn off when not in use to conserve battery power.

III. End of the Day Meter Check

1. Ensure the meter is in the temperature compensated specific conductance mode by pressing the **UP or DOWN** button until the °C is flashing to indicate this mode.
2. **Rinse** the probe with DI water and blot dry with a Kimwipe. Gently shake the probe to remove water from the oval upper conductivity opening.
3. Submerge the entire probe in the **conductivity standard solution**, and allow to stabilize. Ensure there is enough solution to cover the top opening of the probe.
4. **Record** the “**End of the Day Meter Check**” value on the VRAP Field Data Sheet. A 20% error regardless of the standard used (1,600 – 2,400 μ S for 2,000 μ S standard, 160-240 μ S for 200 μ S standard or 80 μ S - 120 μ S for 100 μ S standard) is acceptable. If the reading is outside of this range, please check again with new standard if available. If new standard is unavailable please contact VRAP staff as soon as possible.
5. Rinse the probe with DI water and return it to the storage chamber.
6. Turn the meter off.