

Guidelines and Standard Procedures for Continuous Water-Quality Monitors: Site Selection, Field Operation, Calibration, Record Computation, and Reporting

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ABSTRACT

The U.S. Geological Survey uses continuous water-quality monitors to assess variations in the quality of the Nation's surface water. A common system configuration for data collection is the four-parameter water-quality monitoring system, which collects temperature, specific conductance, dissolved oxygen, and pH data, although systems can be configured to measure other properties such as turbidity or chlorophyll. The sensors that are used to measure these water properties require careful field observation, cleaning, and calibration procedures, as well as thorough procedures for the computation and publication of final records.

Data from sensors can be used in conjunction with collected samples and chemical analyses to estimate chemical loads. This report provides guidelines for site-selection considerations, sensor test methods, field procedures, error correction, data computation, and review and publication processes. These procedures have evolved over the past three decades, and the process continues to evolve with newer technologies.

INTRODUCTION

Water quality changes over time, necessitating repeated measurements to characterize variations in

quality adequately. When the time interval between repeated measurements is sufficiently small, the resulting record of water quality can be considered continuous. A device that automatically measures water quality in this way is called a continuous water-quality monitor. These monitors have sensors and recording systems to measure physical and chemical water-quality properties at discrete time intervals at point locations. Operation of a water-quality monitoring station provides a nearly continuous record of water quality that can be processed and published or distributed directly by telemetry or on the World Wide Web. The water-quality record provides a nearly complete record of changes in water quality as well as a basis for computation of constituent loads at a station. Data from the sensors also can be used as surrogates for the measurement of other constituents by using regression analyses to provide estimates of instantaneous chemical loads. Emerging sensor technology is broadening the variety of measurable chemical constituents and is reducing limits of detection. As it has become possible to make near real-time water-quality monitoring data available on the World Wide Web, continual progress is being made to improve applications and refine quality-control procedures.

Purpose and Scope

This report provides basic guidelines and procedures for U.S. Geological Survey (USGS) site selection and operation of continuous water-quality monitors, field data evaluation, and subsequent record

computations. The basic guidelines are minimal requirements that may need to be modified to meet local environmental conditions. This report provides examples of the application of scientific judgment in the evaluation of data records. Field training with the monitoring equipment and first-hand knowledge of the watershed form the core of the data-evaluation process. Record-computation procedures provide a uniform set of minimum requirements for computing records. Adoption of these water-quality record procedures will ensure that published data are properly documented.

Representative illustrations of the application of scientific judgment are, by necessity, site specific. Other specific examples are used to demonstrate the range of environmental conditions that affect the evaluation process. The final evaluation to determine portions of the record that are published requires careful review and verification of the data, but much is left to the professional judgment and observations of the hydrographer operating the station. A set of considerations that represent a sound basis for the operation of a water-quality station, data evaluation, and publication criteria is presented in this report.

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Internal (unpublished) continuous water-quality monitoring procedures documented by the Colorado, Maine, Michigan, North Carolina, Ohio, Oregon, Texas, and Washington Districts significantly contributed to the development of these guidelines as well as previously published guidelines. Special thanks go to Brett A. Smith who co-authored an earlier version of this report and provided valuable insights to the record computation process. The encouragement of Brett, Rebecca Deckard, and Gary Pederson is gratefully acknowledged.

WATER-QUALITY MONITORING SITE OPERATION

Physical and chemical measurements in streams can be made at discrete intervals to provide a nearly continuous record of stream water quality. Major factors in the operation of a continuous water-quality site include selection of sensors and types of monitors, the type of monitor configuration, site selection, location of the sensors in the stream cross section, the use and calibration of field meters, and the actual operation of the continuous water-quality monitors. Sensor and site selection are guided by the purpose of monitoring and the data objectives. The main objective in the placement of the sensors is the selection of a stable, secure location that is representative of the stream.

Monitor Selection

The selection of a water-quality monitor involves four major interrelated elements—(1) the purpose of the data collection, (2) the type of installation, (3) the type of sensor deployed at the installation, and (4) the specific sensors needed to satisfy the accuracy and precision requirements of the data-quality objectives.

The most widely used water-quality sensors in monitoring installations are temperature, specific conductance, dissolved oxygen (DO), pH, and turbidity. Sensors also are available to measure oxidation-reduction potential, water level, depth, salinity, ammonia, nitrate, chloride, and chlorophyll; but the focus of this report is on temperature, specific conductance, DO, pH, and turbidity. The sensors needed to measure these properties are available as single instruments or in various combinations. For clarity in this report, a sensor is referred to as a particular lone sensor or a sensor that is detachable from a combination. A group of sensors configured together commonly is referred to as a sonde, which typically has a single recording unit or electronic data logger to record the output from the multiple sensors.

Types of Monitor Configuration

In general, three types of configurations are used for water-quality monitors. The flow-through monitoring system generally has a pump that delivers water to the sensor(s) mounted in a shelter; the second configuration is one in which the sensors are placed in

situ (immersing a field measurement sensor directly into the water); and the third configuration is a self-contained sensor and recording system that requires no external power and is placed in situ. Each configuration has advantages and disadvantages in relation to site location and data-quality objectives.

The flow-through monitoring system has a pump to convey water from the stream to a tank inside a

shelter that contains the monitoring sensor or sonde (fig. 1). Typical pumps require 120-volt alternating current (AC) and deliver about 10 gallons of water per minute. If access to power is not a problem, then other site considerations become important (table 1); the advantages and disadvantages of the flow-through monitoring system must be compared to the data objectives.

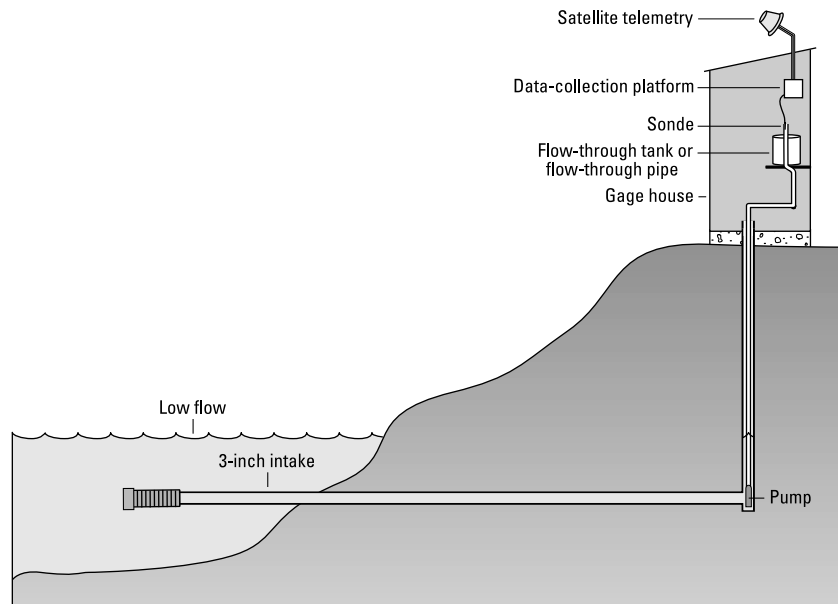
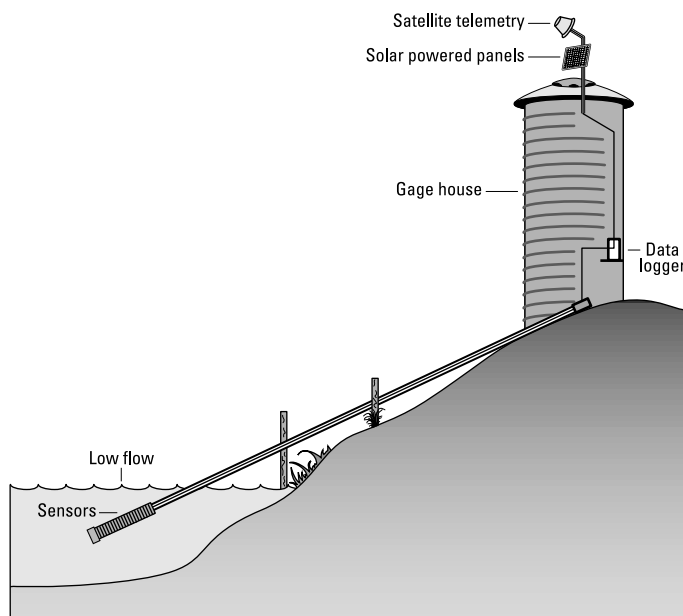


Figure 1. Flow-through water-quality monitoring station. (PHOTOGRAPH: RAMAPO RIVER AT POMPTON LAKES, NEW JERSEY, BY MICHAEL J. DELUCA)

Table 1. Principal advantages and disadvantages of flow-through monitoring systems

Advantages	Disadvantages
Unit can be coupled with chlorinators to reduce membrane fouling.	120-volt AC power source is needed.
Expensive sensor systems can be secured in vandal-proof shelters.	Higher installation costs are incurred.
Calibration can be performed in the shelter.	Pumps tend to clog in streams with algal fouling or high sediment loads.
	Electrical shock protection is required.
	Pumps may be damaged by corrosive waters.
	Pump maintenance is required.
	Pumping may cause changes in water quality.



The sensors in the in-situ monitoring system are placed at the measuring point in the stream cross section (fig. 2). Cables run from the sensors to the recording equipment that is housed in a shelter. The primary advantage of this configuration is that no power is needed to pump water (table 2). Direct current, 12-volt batteries easily meet the power requirements of the sensors and recording equipment. In-situ water-quality monitoring systems can be installed at remote locations where AC power is not available, but the advantages and disadvantages of the in-situ monitoring system also must be considered.

Table 2. Principal advantages and disadvantages of in-situ monitoring systems.

Advantages	Disadvantages
No power is needed to pump water.	Sensors are susceptible to vandalism.
Remote locations are possible.	The water flow cannot be treated to reduce fouling.
Smaller shelters can be used.	In shallow bank installations, the proper location of sensors in the cross section is difficult.
Pumping maintenance is not required.	Servicing sensors during flooding can be difficult.
Freeze protection is provided to the sensors.	Sensors are susceptible to debris or high flow.
Electrical hazards are reduced.	Shifting channels may cause movement of the equipment.



Figure 2. In-situ water-quality monitoring station. (PHOTOGRAPH: SPRING BROOK CREEK AT ORILLIA, WASHINGTON, BY DAVID K. MULLIS)

The third water-quality monitoring system is a combined sensor and recording sonde that is self-contained, requires no external power, and reduces exposure to vandalism. Power is supplied by conventional batteries located in a sealed compartment, and sensor data are stored within the sonde on nonvolatile, flash-memory, recording devices (fig. 3). The advantages and disadvantages of the self-contained sensor and recording system must be considered (table 3).

Table 3. Principal advantages and disadvantages of the self-contained monitoring system.

Advantages	Disadvantages
Location options are flexible.	Data are available only during site visits.
The monitor is protected from vandalism.	Equipment status checks are required.
Electrical hazards are nonexistent.	Servicing sensors and recovering data can be difficult.
	Sensors are susceptible to debris or high flow.
	Shifting channels may cause movement of the equipment.
	Status of the equipment can be checked only while servicing.
	If batteries die, data are lost.

Types of Sensors

Sensors are available for continuous measurement of many physical properties and chemical constituents, but five of the most commonly used sensors are for the measurement of temperature, specific conductance, DO, pH, and turbidity.

Temperature

Temperature has an important influence on the density of water, the solubility of constituents in water, pH, specific conductance, the rate of chemical reactions, and biological activity in water (Radtke, Kurklin, and Wilde, 1998). The preferred water-temperature scale for most scientific work is the Celsius scale. Continuous water-quality sensors usually measure temperature with a thermistor, which is a semiconductor with resistance that changes with temperature. Thermistors are reliable, accurate, and durable sensors that require little maintenance and are relatively inexpensive. Modern thermistors can easily measure temperature to plus or minus (\pm) 0.1 degree

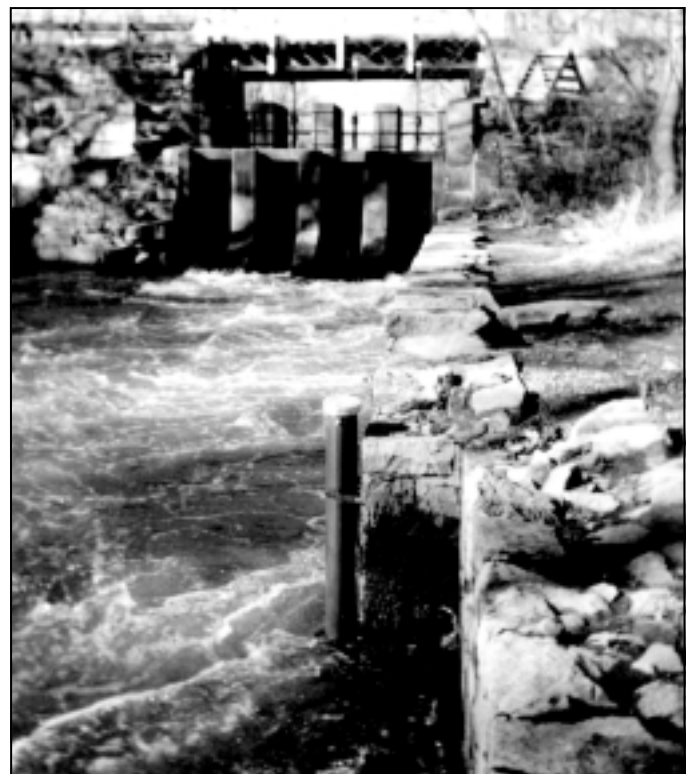
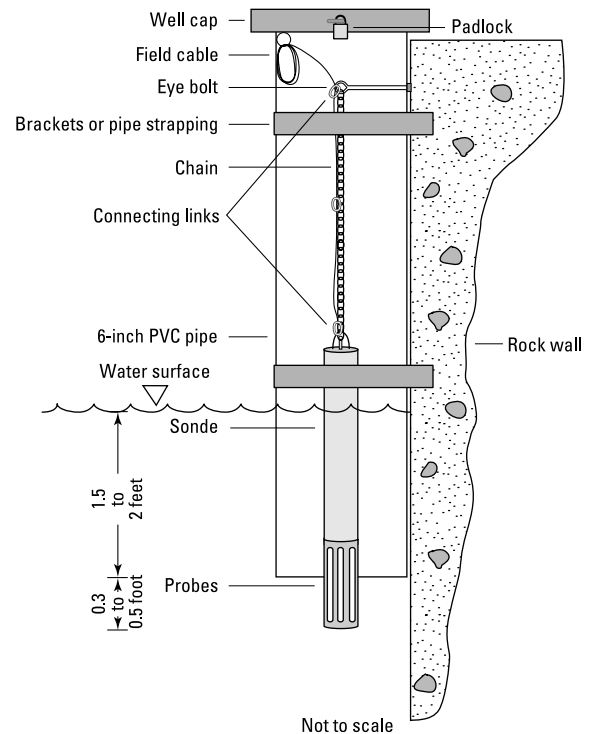


Figure 3. Self-contained water-quality monitoring sensor and recording system. (PHOTOGRAPH: DELAWARE AND RARITAN CANAL FEEDER AT RAVEN ROCK, NEW JERSEY, BY BONNIE GRAY)

Celsius (°C), but the user must verify the accuracy claimed by the manufacturer for the range of application. Typically, temperatures are reported to the nearest 0.5 °C because the thermistor is located at one point, which may not be representative of the entire body of water being measured.

Specific Conductance

Specific conductance is a measure of the capacity of water to conduct an electrical current and is a function of the types and quantities of dissolved substances in water (Radtke, Davis, and Wilde, 1998). As concentrations of dissolved ions increase, specific conductance of the water increases. Specific conductance measurements are a good indication of total dissolved solids and total ion concentrations, but there is no universal linear relation between total dissolved solids and specific conductance. A continuous record of specific conductance can be used in conjunction with chemical analyses and continuous discharge records to estimate constituent loads (Clifton and Gilliom, 1989; Hill and Gilliom, 1993; Christensen and others, 2000). The USGS measures specific conductance at or compensated to 25 °C and reports specific conductance in microsiemens per centimeter at 25 °C ($\mu\text{S}/\text{cm}$ at 25 °C).

Specific conductance sensors are one of two types—contact sensors with electrodes and sensors without electrodes. Continuous water-quality sensors generally have electrodes that require the user to choose a cell constant for the expected range of specific conductance. Multiparameter monitoring systems usually contain automatic temperature compensation circuits; thus, specific conductance usually is compensated to 25 °C, but this should be verified by checking the manufacturer's instruction manual. All modern monitoring systems are designed for medium-strength specific conductance waters (100 to 2,000 $\mu\text{S}/\text{cm}$ at 25 °C) or higher. In general, specific conductance electrodes are reliable, accurate, and durable but are susceptible to fouling from aquatic organisms and sediment.

Dissolved Oxygen

The DO concentration in surface water is related primarily to atmospheric reaeration and photosynthetic activity of aquatic plants (Radtke, White, and others, 1998). The range of observed DO in surface waters typically is from 2 to 10 milligrams per liter (mg/L) at

20 °C. The value for 100-percent saturation of DO decreases with increased temperature and increases with increased atmospheric pressure. Occasions of excess oxygen (supersaturation) often are related to extreme photosynthetic production of oxygen by aquatic plants as a result of nutrient (nitrogen and phosphorus) enrichment, sunlight, and low-flow conditions. Occasions of saturated oxygen commonly are related to cascading flow conditions, both natural and artificial. DO may be depleted by inorganic oxidation reactions or by biological and chemical processes that consume dissolved, suspended, or precipitated organic matter (Hem, 1989). DO is a significant factor in chemical reactions in water and the survival of aquatic organisms.

The most commonly used technique for measuring DO concentrations with continuous water-quality sensors is the amperometric method, which measures DO with a temperature-compensated polarographic membrane-type sensor. While polarographic membrane-type sensors generally provide accurate results, they commonly are sensitive to temperature and water velocity and are prone to fouling. Because the permeability of the membrane and solubility of oxygen in water change as functions of temperature, it is critical that the DO sensors be temperature compensated.

The measuring process consumes DO; therefore, water flow past the sensor is critical. If water velocity at the point of measurement is less than 1 foot per second (ft/s), an automatic or manual stirring mechanism or a different measurement technique is required. DO sensors are prone to inaccuracies from algal fouling, sedimentation, low velocity, and very high velocities. A complete discussion of DO calibration, measurement, and limitations can be found in Radtke, White, and others (1998).

pH

The pH of an aqueous solution is controlled by interrelated chemical reactions that produce or consume hydrogen ions (Hem, 1989). The pH of a solution is a measure of the effective hydrogen-ion concentration (activity; Radtke, Busenberg, and others, 1998). More specifically, pH is a measure that represents the negative base-10 logarithm of hydrogen-ion activity of a solution, in moles per liter. Solutions having a pH below 7 are described as acidic; solutions with a pH greater than 7 are described as basic or

alkaline. Dissolved gases, such as carbon dioxide, hydrogen sulfide, and ammonia, appreciably affect pH. Degasification (for example, loss of carbon dioxide) or precipitation of a solid phase (for example, calcium carbonate) and other chemical, physical, and biological reactions may cause the pH of a water sample to change appreciably soon after sample collection (Radtke, Busenberg, and others, 1998).

The electrometric pH measurement method, using a hydrogen-ion electrode, commonly is used for continuous water-quality pH sensors. Sensors used in submersible monitors typically are combination electrodes in which a proton (H⁺)-selective glass-bulb reservoir is filled with an approximate pH-7 buffer. A silver wire coated with silver chloride is immersed in the internal reference electrode buffer reservoir. Protons on both sides of the glass pH electrode (media and buffer reservoir) selectively interact with the glass, setting up an external potential gradient across the outer glass membrane. Because the hydrogen-ion concentration in the internal buffer solution is constant, this external potential difference across the outer glass membrane, which is determined relative to the internal silver/silver-chloride reference electrode, is proportional to the pH of the medium. A correctly calibrated pH sensor can accurately measure pH to ± 0.2 pH unit; however, the sensor can be scratched, broken, or fouled easily. Detailed instructions for the calibration and measurement of pH are described by Radtke, Busenberg, and others (1998) or by the instrument manufacturer.

Turbidity

Turbidity sensors operate differently from those for temperature, specific conductance, DO, and pH, which convert electrical potentials into the measurement of the constituent of interest. A turbidity sensor operates by directing a light beam from a light-emitting diode into the water sample and measuring the light that scatters off the suspended particles present in the water. The nephelometric measurement method is used in most commercially available sensors with a sensor range of 0–1,000 nephelometric turbidity units (NTU) and an accuracy of ± 5 percent or 2 NTU, whichever is greater. However, some sensors can report values reliably up to about 1,500 NTU.

Turbidity readings are affected by suspended-sediment particle size, entrained air bubbles, floating debris, and other particles in the water that may collect on or near the optic sensor during data collection. The

effect of temperature on turbidity sensors is minimal, and the software for modern sensors provides temperature compensation. Calibration and measurement of turbidity by using a submersible sensor are discussed by Wilde and Gibs (1998), but manufacturer's instructions and recommendations must be followed. Sensors that are maintained and calibrated routinely will be relatively error free and will provide valid data at all times.

Instrument Acceptance Criteria

Independent testing to ensure accuracy and reliability is an important part of any quality-assurance program for hydrologic field instrumentation. As stated in the Water-Resources Division (WRD) Hydrologic Field Instrumentation and Equipment Policy and Guidelines (U.S. Geological Survey WRD Memorandum 95-35, 1995), any USGS District, program, or project that procures an instrument is responsible for ensuring that adequate testing is carried out and that the documented results fully characterize the performance and capabilities of the instrument. One of the primary responsibilities of the USGS Hydrologic Instrumentation Facility (HIF) is the testing, evaluation, and documentation of instrument performance. Districts are encouraged to purchase instruments through the HIF when possible or perform the necessary steps of independent testing to ensure accuracy and reliability as stated by the instrument manufacturer. District personnel are encouraged to work with the HIF to evaluate new instrumentation and actively participate in the web-based instrument quality-assurance data base managed by the HIF by entering, reviewing, and overseeing District data in the data base and by taking corrective actions when necessary.

Site Selection

The location of a water-quality monitoring site is directly related to the purpose of monitoring and the data-quality objectives. Stream characteristics, location of the site, and other data-quality objectives determine whether a data sonde will be placed in situ or whether a flow-through receptacle with a pumping sampler will be a better choice. More site-specific considerations in monitor placement include site-design considerations, monitor installation, physical constraints, and service requirements (table 4).

Table 4. Factors for consideration in the placement and installation of continuous water-quality monitoring systems

Site-design considerations
Representative of cross-sectional variability
Constraints of channel configuration
Range of streamflow (from low flow to flood)
Velocity of streamflow
Turbulence
Avoidance of high-water debris damage
Range of values for water-quality physical properties
Protection from vandalism
Safety hazards
Monitor installation
Permits for installation
Type of installation
Difficulty and cost of installation
Ability to install monitor in representative location
Logistics (service requirements)
Accessibility of site
Frequency of service interval to meet data-quality objectives
Rate of fouling
Proximity to cross-section measuring location
Event related (for example, flooding event)
Proximity to electrical power or telephone service
Need for real-time reporting

Balancing the numerous site considerations for placement of a continuous water-quality monitoring system is difficult. The optimum site consideration for achieving the data-quality objectives is placing the pump intake, sensor package, or sonde in a location that best represents the section of the water body being measured. Lateral mixing in large rivers, however, often is not complete for tens of miles downstream from a tributary or outfall. A location near the streambank may be more representative of local runoff or affected by point-source discharges upstream, whereas a location in center channel may be more representative of areas farther upstream in the drainage basin. Turbulent streamflow may aid in mixing, but turbulence can create problems for some monitored parameters such as DO or turbidity. For a medium to small stream with alternating pools and riffles, the best flow and mixing occurs in the riffle portion of the stream; however, flooding may change the locations of shoals upstream from the monitoring site, and the measurement point may no longer represent the overall water-quality characteristics of the water body.

Sufficient cross-section measurements must be made at the site to determine if a prospective site is sufficiently well mixed and to ensure that the site will not be subject to significant difference in a cross section. If significant horizontal or vertical variability is determined, consideration must be given to choosing another site or using a different approach to meet the data-quality objectives (see **Placement of Sensors in the Cross Section**).

Large streams and rivers may be monitored best from the downstream side of bridge abutments, assuming that safety hazards and other difficulties can be reduced or overcome. The measurement point in the vertical dimension for larger flow systems also needs to be appropriate for the primary purpose of the monitoring installation. The vertical measurement point can be chosen for low-, medium-, or high-flow conditions.

Selection of a water-quality monitoring site is determined by the data-quality objectives, and the best location for a site is often one that is best for measuring surface-water discharge. Although hydraulic factors in site location must be considered, it is more important to consider factors that affect the water-quality data. It is important to select a site that is representative of the stream, and the cross-section characteristics of specific conductance, turbidity, or pH may be helpful in selecting the best sites. Assessment of a site may be dependent on fouling potential, high-tension power lines, or radio towers that may interfere with data collection or telemetry. The same hydraulic factors that are used in selecting a specific site for gaging discharge in a channel also can be used in selecting a water-quality monitoring location. Both purposes seek a representative site that approaches uniform conditions across the entire width of the stream. Rantz and others (1982) identified nine hydraulic conditions of the ideal gage site, and these also must be considered in site selection for water-quality sites (table 5).

The location of water-quality monitoring sensors in cold regions requires additional considerations in order to obtain data during periods of ice formation. White (1999) discusses environmental factors in the site selection of an automated water-quality station in British Columbia, Canada, but also generalizes morphological stream factors and the importance of selecting a site that has a minimal chance of damage or destruction from natural forces and a minimal source of bubbles. White also

Table 5. Hydraulic conditions of the ideal gage site (after Rantz and others, 1982)

<ol style="list-style-type: none">1. The general course of the stream is straight for about 300 feet upstream and downstream from the gage site.2. The total flow is confined to one channel at all stages, and no flow bypasses the site as subsurface flow.3. The streambed is not subject to scour and fill and is free of aquatic growth.4. Banks are permanent, high enough to contain floods, and free of brush.5. Unchanging natural controls are present in the form of a bedrock outcrop or other stable riffle for low flow and a channel constriction for high flow, or a falls or cascade that is unsubmerged at all stages.6. A pool is present upstream from the control at extremely low stages to ensure a recording stage at extremely low flow, and to avoid high velocities at the streamward end of gaging-station intakes during periods of high flow.7. The gaging station is far enough upstream from a confluence with another stream or from tidal effect to avoid any variable influence from the other stream or tide on the stage at the gage site.8. A satisfactory reach for measuring discharge at all stages is available within reasonable proximity of the gage site. (It is not necessary that low and high flows be measured at the same stream cross section.)9. The site is readily accessible for ease of installation and operation of the gaging station.10. The site is not susceptible to manmade disturbances, nearby tributaries, or point-source discharges.

emphasizes that a site should (1) meet minimum stream-depth requirements for instruments, (2) be safe and accessible under all conditions, and (3) be located to avoid the danger of vandalism.

Placement of Sensors in the Cross Section

The location of a water-quality monitoring site is dependent on the purpose of monitoring and the data-quality objectives. In some cases, such as the location of a monitoring site in a lake, bay, or estuary, the data-quality objectives may require only point measurements; whereas, data-quality objectives for the measurement of load or flux in a large river generally require placement of a water-quality monitoring site at the most representative measurement point in the cross section of the stream.

Determinations of cross-sectional variability are required before a monitoring site is installed to properly locate the most representative measurement point in the stream cross section and to determine if a cross-section correction is necessary. A minimum of two cross-section measurements per year is required to verify that no significant changes have occurred in the distribution of constituents across the stream. Documentation of vertical mixing is required at least annually at a minimum of two depths for all cross-section measurements. For monitoring sites that are operated more than 1 year, a minimum of six cross-section measurements representing different flow

conditions is required to determine if discharge or seasonal changes are affecting significantly the distribution of constituent values in the cross section. By choosing a monitoring site with well-mixed streamflow, cross-section corrections may not be needed.

Standard USGS procedure is to use either discharge- or area-weighted methods for selecting locations of the most representative measurement points in a cross section to ensure that chemical loads can be calculated correctly (Wilde and Radtke, 1998). The measurement point in a cross section is established by using either method to obtain data by using calibrated field instruments. Generally, a minimum of five increments across a large stream or river is sufficient to provide a discharge-weighted mean and establish the horizontal cross-sectional variability of the measured physical property. Alternatively, an area-weighted value can be determined by measuring the centroid of equal-width increments with calibrated field instruments. Generally, a minimum of 10 and a maximum of 20 measurement points across a large stream or river is sufficient to establish the horizontal cross-sectional variability of a physical property. Computations of mean physical property values for a large number of cells will produce a cross-section correction for any measurement location with good flow; however, as the number of cells exceed 40 or more, the computational precision exceeds the measuring precision. Examples of both area- and discharge-weighted field measurements are demonstrated by Wilde and Radtke (1998).

The number of vertical measurements needed for deeper flow regimes is dependent on the degree of vertical mixing. If physical or chemical stratification is observed, the number of vertical measurements may need to be increased from mid-depth to two measurements (0.2 and 0.8 depth) or more. The vertical stratification may be sharply defined, which requires the measurements across the transition zone to be more closely spaced to represent the position and degree of stratification adequately.

The most efficient means of obtaining physical property multiparameter measurements is with a multisensor monitor that can measure more than one physical property at each measurement point. For locations with higher stream velocities, point sampling with a heavy sampler may be necessary. Measurements of discrete samples from the cross section should be made quickly before measurable changes occur in some physical properties. Discrete samples should not be composited for measurement of cross-section corrections.

In the case of pH, a median value is used for determining the measuring-point correction. Alternatively, pH values can be converted to hydrogen ion concentrations (antilogarithms of negative pH) for computational purposes (see Appendix 1). Computed mean pH values tend to be more acidic than median values; for large flow systems with 40 or more pH measurements, the median and the properly computed mean pH values should be very close.

Specific information about a field site (station) is summarized in a station description (see Appendix 2). The station description, which contains specific information about a field site location, history, and operation, should be placed in an appropriate field folder and carried into the field. Cross-section measurements and extreme values must be updated annually in the station analysis (see Appendix 3) as part of the data-evaluation process. For many stations with critical data-quality objectives, service intervals may be a week or less. The service interval must be included in the station description as well as in the report manuscript station description, as described by Novak (1985).

Field Meter Calibration

Measurements that compare field meter physical property values with the continuous monitor readings should be made before, during, and after servicing the

monitor to document any environmental changes during the service interval. Measurements are made at the monitoring site by using calibrated field instruments as close to the sensor as possible and within 5-minute intervals. Before site visits, all support field meters should be checked for operation and accuracy. Minimum calibration frequency is detailed by Wilde and Radtke (1998) for each type of meter, and all calibrations are recorded in the corresponding instrument log books. All information related to pre- and post-trip, National Field Quality Assurance, and other periodic calibrations is recorded in the instrument log books.

Temperature

The proper calibration and documentation for thermometers and thermistor thermometers are detailed by Radtke, Kurklin, and Wilde (1998). The USGS procedures specify that thermometers be calibrated or checked against a National Institute of Standards and Technology (NIST)-certified thermometer, and thermistors should be accurate within ± 0.2 °C. For both thermistors and liquid-in-glass thermometers, an annual 5-point calibration over the temperature range of 0–40 °C is required. The annual 5-point check requires laboratory equipment. In addition, 2-point calibration checks over the maximum and minimum expected annual temperature range should be made three or more times per year for thermistors and two or more times per year for liquid-in-glass thermometers. Calibrated thermometers and thermistors must be marked with the date of calibration.

Specific Conductance

The proper calibration and documentation for specific conductance meters are detailed by Radtke, Davis, and Wilde (1998). Standards bracketing the expected full range of anticipated values are used to calibrate the specific conductance meter to the appropriate units for particular field conditions. Calibration is performed at the field site with standards that have been allowed to equilibrate to the temperature of the water being monitored. The accuracy of the meter should be within 5 percent for specific conductance values less than or equal to 100 $\mu\text{S}/\text{cm}$, or within 3 percent for specific conductance values more than 100 $\mu\text{S}/\text{cm}$. Specific conductance standards are

available from the USGS Ocala Quality Water Service Unit (QWSU). Standards must be discarded after use.

Dissolved Oxygen

The proper calibration and documentation for DO meters are detailed by Radtke, White, and others (1998). DO sensors measure the partial pressure of DO by the flow of oxygen through a porous membrane and oxygen consumption at a cathode. The calibrated accuracy of DO meters should be within the lesser of 5 percent or ± 0.3 mg/L. The accuracy of a DO meter at 0.0 mg/L is verified by measuring the DO of a sodium sulfite solution, as described by Radtke, White, and others (1998). Calibration of a DO meter at 100-percent oxygen saturation is made by adjusting the meter reading for air saturated with water vapor to a value obtained from a solubility table generated from the equations of Weiss (1970) and listed in Radtke, White, and others (1998). The 100-percent saturation value is based on the water temperature and the uncorrected barometric pressure. A reliable pocket altimeter can be used to measure uncorrected (true) atmospheric pressure to the nearest 1 millimeter (mm) of mercury. An additional correction for salinity, based on temperature and specific conductance, should be calculated for waters that have specific conductance values greater than 2,000 $\mu\text{S}/\text{cm}$. Calibration and operation procedures differ among instrument types and makes, and manufacturer's instructions must be followed closely. All calibration information must be recorded in the instrument log book.

pH

A detailed 10-step description of the calibration of pH meters and that covers a wide range of available equipment is described by Radtke, Busenberg, and others (1998). Calibration and adjustments for multiparameter sensor systems are found in the manufacturer's service manuals. Accuracy of field pH meters should be at least ± 0.1 pH unit. Two standard buffers bracketing the expected range of environmental values are used to calibrate a pH electrode, and a third is used as a check for linearity. The pH-7 buffer is used to establish the null point, and a pH-4 or pH-10 buffer is used to establish the slope of the calibration line at the temperature of the solution. The slope of a pH electrode is temperature sensitive, but modern sensors can adjust the pH slope to the observed temperatures through manual or automatic temperature

compensation. It is important, however, that the temperatures of the buffers be as close as possible to the samples being measured. Immersing the pH buffer bottles in the surface water for about 15 minutes allows the stream or river temperature and the buffer temperature to equilibrate. Standard buffers of pH 4, 7, and 10 are readily available from the QWSU. All calibration information is to be recorded in the instrument log and on the field notes. Proper calibration of pH sensors for low specific conductance waters, less than 100 $\mu\text{S}/\text{cm}$, is described by Busenberg and Plummer (1987) and by Radtke, Busenberg, and others (1998).

Turbidity

The proper calibration and documentation for turbidity meters are described by Wilde and Gibs (1998). Calibration of a turbidity instrument by using Formazin or another primary standard usually is done in the laboratory, with instrument checks being performed in the field. Most sensor manufacturers recommend either Formazin or other approved primary standards, such as styrene divinylbenzene (SDVB) polymer standards, for calibrating turbidity sensors. Formazin-based standards can be diluted by using a dilution formula; however, errors may be introduced during the dilution process, thus reducing the accuracy of the standard solution. Formazin-based standards also are temperature dependent, and accurate readings may be difficult to obtain during field conditions. Wilde and Gibs (1998) suggest that the effect of thermal fluctuations can be minimized by calibrating the instrument at room temperature in an office laboratory using a Formazin-based standard and secondary standard. Instrument calibration can then be checked at the field site by using a secondary standard. Turbidity standards for various ranges are available commercially. Before placing the sensor in a standard, the sensor must be cleaned, rinsed three times with turbidity-free water, and carefully dried. Turbidity-free water is prepared as described by Wilde and Gibs (1998).

Monitor Operation

The operation of a water-quality monitoring station is intended to produce the greatest amount of correctable field record that can be verified. The general operational categories include maintenance of

the station and equipment; inspection and recording of sensor readings; cleaning, calibration, and troubleshooting of sensors and recording equipment; cross-section measurements; and accurate record keeping.

Maintenance

Maintenance frequency generally is governed by the fouling rate of the sensors, and this rate varies by sensor type, hydrologic environment, and season. The performance of temperature and specific conductance sensors tends to be less affected by fouling, whereas the dissolved oxygen, pH, and turbidity sensors are more prone to fouling. The use of wiper or shutter mechanisms on modern turbidity instruments has decreased the fouling problem significantly. For stations with critical data-quality objectives, service intervals may be weekly or more often. Monitoring sites with nutrient-enriched waters and moderate to high temperatures may require service intervals as frequently as every third day. In cases of severe environmental fouling or remote locations, the use of an observer for servicing the water-quality monitor should be considered. In addition to fouling problems, physical disruptions (such as pump failure, recording equipment malfunction, sedimentation, electrical disruption, debris, ice, or vandalism) also may require additional site visits. The service needs of water-quality monitoring stations equipped with telemetry can be recognized quickly, and the use of satellite telemetry to verify proper equipment operation is recommended.

The general maintenance functions at a water-quality monitoring station include:

- Daily review of sensor function for sites equipped with telemetry
- Inspection of the site for signs of physical disruption
- Inspection of sensor(s) for fouling, corrosion, or damage
- Battery (or power) check
- Time check
- Routine sensor cleaning and servicing
- Calibration (if needed)
- Downloading of data

The specific maintenance requirements depend on the site configuration and equipment. A useful discussion of the maintenance requirements for the flow-through and USGS minimonitor installations can

be found in Gordon and Katzenbach (1983), but nearly all operational requirements are fulfilled by the completion of a water-quality continuous monitor field-inspection form (Appendix 4). The manufacturer's instructions must be followed for other types of equipment.

Sensor Inspection

The purposes of the sensor inspection are to provide an ending point for the interval of water-quality record since the last service visit, a beginning point for the next interval of water-quality record, and verification that the sensor is working properly. This is accomplished by recording the initial sensor readings, servicing the sensors, recording the cleaned sensor readings, performing a calibration check of sensors by using appropriate standards, and if the readings of the monitoring sensor are outside the range of acceptable differences (see next section, **Monitor Calibration Criteria**), recalibrating the sensor. A final environmental sensor reading is required after the calibration check or after recalibration. The difference between the initial sensor reading and the cleaned sensor reading is the sensor error as a result of fouling; the difference between the calibration-check reading and calibrated-sensor reading, if necessary, is a result of drift. All information related to the sensor inspection must be recorded on a field form or in a field notebook. The sensor readings in the field notes become the basis for corrections (shifts) during the record-processing stage; thus, a complete and thorough documentation of the sensor inspection is important.

The initial sensor readings (before cleaning) of the monitoring equipment are compared to a calibrated field meter before removing the monitor sonde for servicing. The comparison ideally is made at the measuring point in the stream. This initial sensor reading becomes the ending point of the data record since the last servicing, and the field meter reading provides a sense of the reasonableness of the monitor readings and an indication of potential drift and fouling. However, except for temperature sensors, the field meter readings are not used directly in record computation. The field meter is used as a tool to assess cross-sectional variability and environmental changes that may occur while the monitor is being serviced.

Upon removal from the water, the monitoring sensors are inspected for signs of chemical precipitates, stains, siltation, or aquatic growth. These observations are recorded in the field notes before cleaning, and the

individual sensors must be cleaned according to the manufacturer’s specifications. The cleaned sonde or sensor is then returned to the stream and checked against the field meter. Both instrument readings are recorded in the field notes. The observed difference between the initial sensor reading and the cleaned sensor reading is a result of fouling (chemical precipitates, stains, siltation, or aquatic growth).

Monitor Calibration Criteria

The concept of not calibrating sensors for small changes from a standardized value was defined by Gordon and Katzenbach (1983). This idea recognizes limits in recording electronic data under field conditions that are close to the operating accuracy of the equipment. Adjustments within these acceptable limits may not improve the overall data accuracy. With improvements in sensor technology, this acceptable-limit concept has been refined to calibration criteria for recording field sensors based on the stabilization criteria defined by Wilde and Radtke (1998) and also takes into consideration the lesser accuracy of some continuous water-quality sensors. A calibration check using known standards is performed on cleaned monitoring sensors; if the monitor sensors are outside the range of acceptable differences, the sensor must be recalibrated. If the calibration-check sensor readings for the monitor are within the calibration criteria (table 6), the monitoring sensors are considered checked and no further adjustments are required.

Table 6. Calibration criteria for continuous water-quality monitors

[±, plus or minus value shown; °C, degree Celsius; μS/cm, microsiemens per centimeter at 25 °C; mg/L, milligram per liter; pH unit, standard pH unit; NTU, nephelometric turbidity unit]

Measured physical property	Calibration criteria for measurements (variability should be within the value shown)
Temperature	± 0.2 °C
Specific conductance	The greater of ± 5 μS/cm or ± 3 percent of the measured value
Dissolved oxygen	± 0.3 mg/L
pH	± 0.2 pH unit
Turbidity	The greater of ± 2 NTU or ± 5 percent of the measured value

All sensor readings are recorded in the field notes, and all calibration information is recorded in the monitor instrument log. The calibrated monitoring sensor is returned to the stream and allowed to

equilibrate to the stream temperature. The observed difference between the cleaned sensor calibration-check reading and the calibrated sensor reading (if necessary) is a result of drift. The calibrated sensor stream reading is the beginning observation of the new water-quality record interval. If the calibrated monitoring sensor fails to agree with the calibrated field meter within the calibration criteria, the faulty sensor must be repaired or replaced after verifying that the readings of the field meter are not in error (see **Troubleshooting Procedures**). The alternative is to replace the monitoring sonde or sensor with a calibrated backup unit and repair the malfunctioning monitor in the laboratory or return it to the manufacturer for repair.

Field Cleaning of Sensors

Most commercially available temperature sensors can be cleaned with a detergent solution and a soft-bristle brush. Film on the sensor that resists removal usually can be removed by soaking the sensor in a detergent and water solution (Ficken and Scott, 1989), but the manufacturer’s recommended cleaning procedures should be followed carefully for multiparameter sensor systems.

Radtke, Davis, and Wilde (1998) recommend cleaning the specific conductance sensors thoroughly with de-ionized water before and after making a measurement. Oily residue or other chemical residues (salts) can be removed by using a detergent solution. Specific conductance sensors can soak in detergent solution for many hours without damage. Oil or other residues can be removed by dipping the sensor in a solvent or diluted hydrochloric acid solution (5 percent), but the manufacturer’s recommendations should be checked before using acid solution or solvents on sensors. The sensor should never be in contact with acid solution for more than a few minutes. Carbon and stainless steel sensors can be cleaned with a soft brush, but platinum-coated sensors should never be cleaned with a brush.

Routine cleaning of DO sensors includes the use of a soft-bristle brush to remove silt from the outside of the sensor, wiping the membrane with a damp towel, and then rinsing with de-ionized water. The sensor usually is covered with a permeable membrane and filled with a potassium chloride solution. The membrane is easily fouled and will need to be replaced at least every 30 days, or as needed. When the membrane is replaced, the potassium chloride solution

should be rinsed out of the sensor with de-ionized water, followed by several rinses with potassium chloride solution before the sensor is refilled. The membrane must be replaced with care so that the surface of the membrane is not damaged or contaminated with grease and no bubbles are trapped beneath the membrane. The surface of the membrane should be smooth, and the membrane should be secured tightly with the retaining ring. The sensor should be stored in water overnight to relax the membrane before installation and calibration. This procedure normally requires either replacing the DO sensor or revisiting the site for calibration at a later date.

Inspection of the DO sensor must be noted in detail on the field sheet. The gold cathode of the DO sensor can be fouled with silver over an extended period of time, and a special abrasive tool usually is required to recondition the sensor. A fouled anode, usually indicated by the white silver electrode turning grey or black, can prevent successful calibration. As with the cathode, the sensor usually can be reconditioned as directed by the manufacturer. Following reconditioning, the sensor cup must be refilled with potassium chloride solution and a new membrane installed.

The pH electrode must be kept clean and the liquid junction (if applicable) free flowing in order to produce accurate pH values (Radtke, Busenberg, and others, 1998). The body of the electrode should be thoroughly rinsed with de-ionized water before and after use. In general, this is the only routine cleaning needed for pH electrodes; however, for specific cases of fouling or contamination, the manufacturer's cleaning instructions must be followed. All cleaning and maintenance on the pH sensor are recorded in the instrument log book.

Optical sensors are extremely sensitive to fouling; thus, frequent maintenance trips may be necessary to prevent fouling of the turbidity sensor in an aquatic environment high in sediment build up, algae accumulation, larvae growth, or other biological or chemical debris. Mechanical cleaning devices or shutters that remove or prevent this build up are available for modern sensors. If the turbidity sensor is not equipped with a mechanical cleaning device that removes solids accumulation or a shutter that prevents build up on the lens before readings are recorded, reliable data collection will be more difficult.

Field Calibration of Sensors

A water-quality monitoring sensor or sonde should be calibrated in the laboratory before installation at a field location. Calibration in the laboratory or the field is done only by using standards of known quality. Field calibration is performed if the cleaned sensor readings obtained during the calibration check differ by more than the calibration criteria (table 6). Spare monitoring sondes or sensors are used to replace water-quality monitors that fail calibration after troubleshooting steps have been applied (see **Troubleshooting Procedures**). All calibration equipment must be kept clean, stored in protective cases during transportation, and protected from extreme temperatures.

Temperature Sensors

Modern water-quality temperature sensors are quite sturdy and accurate, and the manufacturers commonly make no provisions for field calibration of the temperature sensor. Monitoring temperature sensors generally are not calibrated, but comparisons of temperature readings are made by using District-certified thermometers or thermistors as described by Radtke, Kurklin, and Wilde (1998). Temperature accuracy is especially important because of the effect of temperature on the performance of other sensors. Before the field trip, the "District Certification" labels on thermometers or thermistors must be checked for the last date of certification and whether the instruments are appropriate for use in extreme field conditions.

The water-quality monitoring temperature sensor and the calibrated field thermistor are placed adjacent to one another, preferably in flowing water. If a liquid-in-glass thermometer is used, it must be the total-immersion type. Sufficient time for temperature equilibration must elapse before a reading is made. The two temperature sensors must be read and the temperatures recorded within 2 minutes. If the monitoring temperature sensor fails to agree within ± 0.2 °C, troubleshooting steps must be taken; if calibration fails, the sensor should be replaced. The faulty sensor or sonde should be returned to the manufacturer for proper calibration or repair.

Specific Conductance Sensors

If sensor inspection and cleaning processes fail to bring a specific conductance sensor within the

calibration criteria (table 6), the sensor must be calibrated. Field calibration differs among the types of available instruments but generally relies on calibration with two standard solutions, bracketing the expected range of environmental conditions, and a third standard solution near the ambient specific conductance value of the water. All calibration information is recorded on the field sheet and in the monitoring instrument log.

Sensor inspection and field cleaning procedures must be followed to confirm that the monitor thermistor and the field thermistor or thermometer agree within ± 0.2 °C. Expiration dates for the standard solutions should be checked, and the standard solution bottles must be allowed to equilibrate with the stream temperature (by soaking 15 to 30 minutes). The sensor, thermistor or thermometer, and measuring container are rinsed three times with a standard solution that approximates ambient conditions. Fresh standard solution is poured into the measurement container; the temperature setting is adjusted if necessary; the specific conductance, the standard values, and the temperature are read and recorded in the field notes and the monitoring instrument log; and the monitor reading is adjusted to the standard solution value. A temperature correction may be necessary if the monitor does not have automatic temperature correction (Radtke, Davis, and Wilde, 1998).

The used standard solution is discarded into a waste container, and the procedure is repeated using the second and third standard solutions. The monitor readings for the standards are read and recorded in the field notes and the instrument log. If the monitor sensor readings differ from the standards by more than 5 $\mu\text{S}/\text{cm}$ or 3 percent, whichever is greater, the calibration sequence must be repeated. If the second calibration sequence still differs by more than the calibration criteria, troubleshooting techniques will need to be attempted (see **Troubleshooting Procedures**). If these steps fail, the sonde or monitoring sensor must be replaced and the backup instrument calibrated.

Dissolved Oxygen Sensors

Dissolved oxygen in water is related to water temperature, atmospheric pressure, and salinity. Radtke, White, and others (1998) provide detailed calibration steps for four different procedures: (1) air-calibration chamber in water, (2) air-saturated water, (3) air-calibration chamber in air, and (4) iodometric (Winkler) titration. The first three procedures determine DO by an amperometric method that

depends on diffusion of oxygen through a Teflon membrane. The iodometric titration measures DO directly by a dye color change upon reduction of available oxygen. The appropriate procedure depends on the type of monitoring equipment. Manufacturer's calibration procedures should be followed closely to achieve a calibrated accuracy of ± 0.3 mg/L DO. In general, calibration of DO sensors involves measuring DO at 100-percent saturation and measuring a zero-DO solution. For measured temperature and atmospheric pressure, the saturated DO is obtained from a table developed by Weiss (1970) and reproduced in Radtke, White, and others (1998). Ambient atmospheric pressure should be determined with a calibrated pocket barometer to the nearest 1 mm of mercury.

In sea water or brackish water, a salinity-correction factor must be applied to the measured DO concentration. Manufacturer's instructions should be referred to before applying a correction for saline waters (waters with specific conductance values greater than 2,000 $\mu\text{S}/\text{cm}$ or chloride concentrations greater than 1,000 mg/L); but generally, specific conductance values less than 8,000 $\mu\text{S}/\text{cm}$ have correction factors less than the calibration criteria of the monitoring sensor for DO. For specific conductance values greater than 8,000 $\mu\text{S}/\text{cm}$, refer to salinity correction tables developed by Weiss (1970) and reproduced in Radtke, White, and others (1998).

A fresh zero-DO standard solution should be prepared before each field trip, as described by Radtke, White, and others (1998). Calibration at 100-percent saturation in the field presents a problem because replacement of the Teflon membrane frequently is required, and the replaced membrane must be allowed to "relax" in water for 24 hours before calibration. One solution to this problem is to carry clean and serviced spare DO sensors, stored in water, to the field. The replacement DO sensors can then be calibrated in the field, and an interruption in the record and a next-day return site visit can be avoided. Direct measurement of ambient DO with the iodometric procedure requires preventing exposure of the sample to atmospheric oxygen, and this often makes accurate and precise field determinations difficult. The accuracy of this procedure also depends on the experience and technique of the data collector (Radtke, White, and others, 1998).

pH Sensors

Field calibration of pH sensors is often a time-consuming process that requires patience. Before going to the field, the field meter pH response and thermistor should be checked, as described by Radtke, Busenberg, and others (1998). Expiration dates for the pH 4, 7, and 10 buffers should be checked, and spare monitoring pH sensors or backup sondes will need to be prepared in case replacement of field sensors is required.

Upon arrival at the field site, the tightly capped buffer solutions are placed in the stream to allow time for temperature equilibration, usually 15 to 30 minutes. The pH sensor is checked for physical damage and replaced if necessary. Sensor inspection and field-cleaning procedures are followed as described by the manufacturer or Radtke, Busenberg, and others (1998). If the pH monitoring sensor reading exceeds the calibration criteria (table 6), the monitoring sensor should be calibrated as described by the manufacturer's instructions.

Temperature effects on pH buffer solutions vary with individual solutions, and the temperature-correction factor should be checked with the manufacturer. See Appendix 5 for examples of common pH buffer solution variances with temperature. The pH sensor reading should be standardized to the temperature-corrected pH value. The pH sensor, thermistor or thermometer, and measurement container are rinsed with pH-7 buffer solution, which is then discarded along with all subsequent rinsates in a waste container. Fresh pH-7 buffer solution is poured into the rinsed measurement container, and the instruments are allowed to equilibrate for 1 minute before the buffer solution is discarded and fresh pH-7 buffer solution is poured into the measurement container. The solution is slowly stirred with the pH sensor or sonde. The temperature, pH, and associated millivolt reading are measured and recorded. This standardization process is repeated with fresh pH-7 buffer solutions until two successive values of the temperature-adjusted pH-7 readings are obtained.

The pH sensor, thermistor or thermometer, and measurement container are rinsed with de-ionized water, and the standardization process is repeated with a pH-4 or pH-10 buffer solution to establish the response slope of the pH sensor by choosing the buffer that brackets the expected range of pH values. The second temperature-corrected pH value, temperature, and millivolt readings are recorded, and the pH sensor,

thermistor or thermometer, and measurement container are rinsed with de-ionized water. The pH-7 buffer solution is then used to rinse, fill, and check the pH-7 calibration measurement. If the pH sensor reading is 7 ± 0.1 pH units, the slope adjustment has not affected the calibration. If the accuracy standard is not met, the calibration and slope adjustment steps must be repeated. If the repeated calibration and troubleshooting steps fail, the pH sensor or monitoring sonde must be replaced.

Once the slope adjustment step is completed satisfactorily, the third buffer solution can be used as a calibration range and linearity check. The temperature and pH values are read and recorded, but the ± 0.1 accuracy should not be expected to be achieved over the full range from pH 4 to pH 10 for a monitoring sensor.

Waters with specific conductance values less than $100 \mu\text{S}/\text{cm}$ require special low-ionic strength buffers and pH sensors. The extra preparations, precautions, and troubleshooting steps for measuring low-ionic strength waters are described by Busenberg and Plummer (1987).

Turbidity Sensors

Field inspection or calibration of the turbidity sensor is made by using Formazin or other approved primary standards and following the manufacturer's calibration instructions as described by Wilde and Gibs (1998). Turbidity standards with various ranges are commercially available, and most sensor manufacturers recommend either Formazin or SDVB polymer standards for calibrating turbidity sensors. Formazin-based standards can be diluted by using a dilution formula; however, errors may be introduced during the dilution process, thus reducing the accuracy of the standard solution. Turbidity-free water, used in the preparation of standards, dilution, and rinsing, is prepared as described by Wilde and Gibs (1998).

Sensors should first be inspected for damage, ensuring that the optical surfaces of the probe are in good condition. Before placing the turbidity sensor in standards, the sensor should be cleaned, rinsed three times with turbidity-free water, and carefully dried. If the readings are unusually high or erratic during the sensor inspection, entrained air bubbles may be present on the optic sensor and need to be removed. If the sensor readings exceed the calibration criteria (the greater of ± 5 percent or 2 NTU) during the inspection process, the sensor must be calibrated by following the

manufacturer's instructions. A three-point calibration process is recommended, covering the expected range of values, although some instruments may be limited to calibration with only one or two standards. If the instrument allows only a two-step process, then two primary standards covering the expected range should be used for calibration and a third mid-point standard used to check for linearity. Similarly, if the instrument requires calibration only with turbidity-free water and one standard, another mid-point standard should be used to check for linearity.

Cross-Section Measurements

The data objectives for most continuous water-quality monitoring sites require that the sensors be installed in locations that are most representative of stream cross sections. A minimum of two cross-section measurements per year is required to verify that no significant changes in the distribution of physical properties have occurred. Variability in some streams may require more frequent measurements. For sites that are operated more than 1 year, a minimum of six cross-section measurements representing different seasons and flow conditions is required to determine if discharge or seasonal changes have significantly affected the distribution of constituent values in the cross section (see **Placement of Sensors in the Cross Section** for details).

Troubleshooting Procedures

When a parameter cannot be calibrated with standard solutions, the hydrographer must determine if the problem resides with the monitoring sensor or with the monitor itself and make necessary corrections to ensure that the monitor is operational. The hydrographer should carry spare sensors and sondes so that troubleshooting, if necessary, can be accomplished at the time of the service visit. Troubleshooting in the field can prevent the need for extra trips and greatly reduce lost record and the amount of time spent in processing the records in the office later. A successful service trip results in a properly calibrated and operating monitor. Some of the more common problems that are likely to be encountered in the field when servicing monitors are listed in Appendix 6 to assist in the troubleshooting process.

Field Notes and Instrument Logs

Field notes and instrument logs are the basis for the accurate and verifiable computation of water-quality monitoring records. Minimum requirements in the field notes for water-quality monitors include the following items:

- Station number and name
- Name(s) of data collector(s)
- Date and times of each set of measurements
- Field meter and monitor serial or "W" numbers
- Purpose of the site visit
- Horizontal and vertical locations of sensors in the cross section (if applicable)
- Recorded monitor values and corresponding field values (initial, after cleaning, and final instream readings)
- Cross-section measurement data (locations of verticals, constituent values, and measurement time)
- Cross-section measurement summaries and corrections
- Pertinent gage-height data
- Remarks that describe channel conditions, condition of the sensors, and so forth
- Battery voltage of monitor at arrival and departure (replaced?)
- Were sensors changed or replaced? (other remarks or observations that may aid in further processing of the record)

Forms including these items encourage consistency and help to avoid the costly omission of critical information. A field form that constitutes a comprehensive checklist for data collection at many water-quality monitoring sites is shown in Appendix 4. Alternative forms should be approved by the USGS District water-quality specialist.

Each field meter and water-quality monitor has an instrument log book, and all pertinent information regarding the monitor is recorded in the instrument log book. One of the most significant pieces of recorded information is the instrument calibration—both field and laboratory. Calibration information can be recorded initially on field forms or field notebooks, but the information then must be copied into the instrument log book. The instrument log book should contain a complete record of all maintenance in the field, the laboratory, or by the manufacturer. Permanent instrument logs contain critical calibration and

maintenance information that document instrument performance throughout the useful service life of the instrument. Calibration log information that is important from a record-processing standpoint includes:

- Calibration dates, times, and temperatures
- Calibration standard values and lot numbers
- Initial and final monitor calibration data
- Field meter calibration values

Field notes and calibration log information should be clearly written, and all notations should be self explanatory. The goal is to have sufficient information for another individual to complete the record processing steps, if necessary. Clear notes simplify the record check and review processes.

RECORD COMPUTATIONS

The record-computation process is accomplished to verify the data and document its quality. Accurate field notes and calibration logs are essential in processing the record. The hydrographer who services the water-quality monitoring installation typically computes the water-quality data record; writes the manuscript that describes the data; updates the station description and station analysis; and prepares the data record review package.

The primary steps in processing the record are an initial data evaluation, application of corrections and shifts, application of cross-section corrections, and a final data evaluation. The initial data evaluation should begin immediately upon completion of the field trip to ensure that all necessary information is available and to check for possible instrument malfunctions that may not have been observed in the field. Record checks and review of the record confirm that the data are accurate and ready for publication.

Data-Processing Procedures

The processing of water-quality monitoring records should be completed in a timely manner according to water-quality-assurance plan policies. Complete and accurate field notes are an important part of the data processing and reduce the amount of time required to process the data. Corrections to data should not be made unless the causes of errors can be validated or explained by information or observations in the field

notes or by comparison to information from adjacent stations (Ritz and others, U.S. Geological Survey, written commun., 1998). Ritz and others summarize data processing as a process that can be classified in six major categories: (1) initial data evaluation, (2) application of corrections and shifts, (3) application and evaluation of cross-section corrections, (4) final data evaluation, (5) record checking, and (6) record review. The first four categories are discussed in this section, and the latter two categories are combined in the section **Preparation of the Review Package**.

Initial Data Evaluation

The initial data evaluation checks the success of the transfer of raw field data (instrument readings) to the office data base and provides the opportunity for initial checks to evaluate and correct erroneous data (daily, if telemetry is available). Raw field data may be stored in a variety of formats, depending on the recording equipment and the means of downloading data from the recording equipment. The conversion of raw data from a variety of recording devices into a standard entry format to the District data base, or Automated Data-Processing System (ADAPS), generally is accomplished with the use of an on-line computer program, or Device Conversion and Delivery System (DECODES). After entry of data into ADAPS, primary data tables and plots can be produced for review. See Appendix 7 for an example of a primary data table.

Sensors, recorders, transmitters, receivers, relays, transmission systems, or unforeseen events can, at times, produce unexplainable data. Data should be reviewed daily, if possible, to edit obviously erroneous data caused by data transmission problems. Data should be processed immediately after the service visit and viewed graphically. In addition to confirming the accurate transfer of data, this may aid in detecting instrument or sensor errors. Data that are missing (for example, because of instrument or transmission problems) should not be estimated, but missing data should be documented and no statistics should be calculated that involve missing data. An example in figure 4 of data record for a continuous specific conductance monitor shows truncated record at the default maximum setting of the data logger. Exceedance of the default maximum was not noticed during routine site visits, and because the record was not computed and viewed graphically after service visits, the truncated maximum values were not

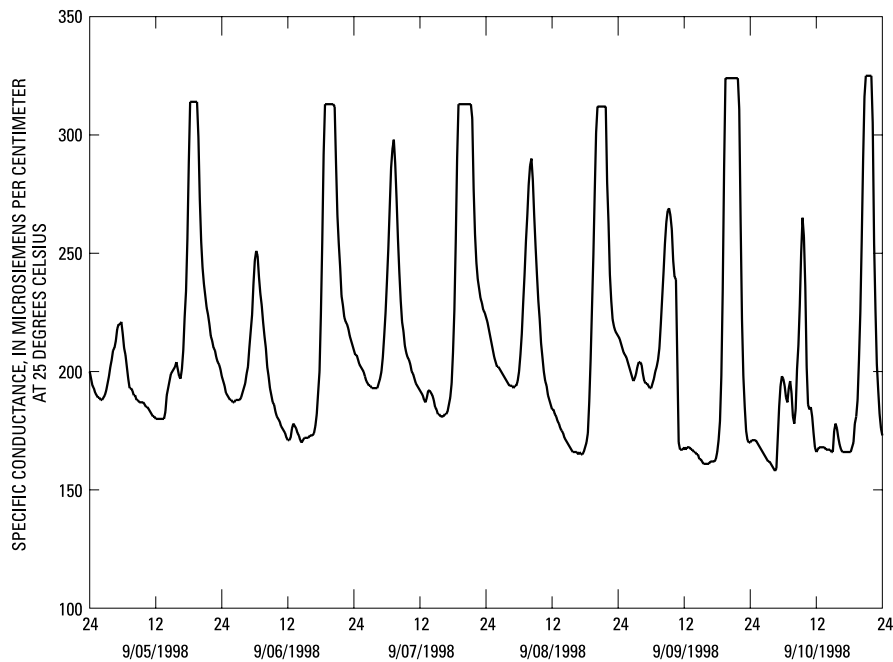


Figure 4. Specific conductance at Duwamish River at Tukwila, Washington, September 5–10, 1998.

observed until the end of the year. Only minimum values were published for the period of record with truncated values, and the manuscript station description noted that maximum values may have been exceeded during periods of missing record (see **Data Qualification Statements** for details).

Great emphasis has been placed on the relation between variations of the five water-quality properties (temperature, specific conductance, DO, pH, and turbidity) and discharge variations, but other event-related changes are equally important and can be factored into the relation only through field experience and first-hand observation at a site. Some examples include changes in air temperature, periods of sustained cloud cover, chemical spills, increased photosynthesis (influenced by a variety of factors), increased wind conditions, combined sewer overflows, forest fires in the watershed, road construction, and ice formation.

As mentioned previously, temperature, specific conductance, DO, pH, and turbidity have complex interdependencies. Understanding these relations is a necessary component for accurate water-quality record computation. Periodically visiting the upstream portions of the drainage basin and keeping field notes on evidence of changes in land-use activities are helpful in explaining observed changes in water quality.

Application of Corrections and Shifts

The application of corrections and shifts allows data to be adjusted to compensate for errors that occurred during the service interval as a result of environmental or instrumentation effects. Three types of measurement-error corrections are described here—fouling, drift, and cross-section correction. Corrections only should be made to measurements when the type and degree of correction is known. Because of the complex interdependencies of temperature and the other measured physical properties, a malfunctioning thermistor can invalidate some or all of the other measured physical properties. The permeability of a DO membrane, for example, changes the DO concentration 3 percent for each temperature change of 1 °C (Wilde and Radtke, 1998). Specific conductance also can vary as much as 3 percent per 1 °C.

The sequence for determining the type and degree of measurement error in the field and the application of corrections in ADAPS generally is fouling, drift, and cross-section correction. If the deviation between actual value and sensor reading exceed the criterion for water-quality data shifts (table 7), a correction is required. The correction is a linear interpolation over the time between sensor inspections.

Table 7. Criteria for water-quality data shifts

[±, plus or minus value shown; °C, degree Celsius; µS/cm, microsiemens per centimeter at 25 °C; mg/L, milligram per liter; pH unit, standard pH unit; NTU, nephelometric turbidity unit]

Measured physical property	Shift criteria (apply shift when deviation exceeds this value)
Temperature (may affect the other physical properties)	± 0.2 °C
Specific conductance	The greater of ± 5 µS/cm or ± 3 percent of the measured value
Dissolved oxygen	± 0.3 mg/L
pH	± 0.2 pH unit
Turbidity	The greater of ± 2 NTU or ± 5 percent of the measured value

The same measurement criteria used to determine the need for calibration (table 6) are used to determine the need for water-quality data shifts. In general, shifts are required when the deviation between actual and recorded data exceed the shift criteria listed in table 7.

The allowable limits shown in table 7 are minimal requirements. More stringent requirements may be required to meet the data objectives of the project or particular hydrologic and chemical environments and are determined at the discretion of the project chief or District water-quality specialist.

The example of graphic corrections in Appendix 8 shows two standards that exceed the shift criteria. Deviation from the value of the third standard does not exceed the shift criteria, but the shifted values also are shown if shifts were made for the third standard. The quality of data is excellent whether or not the shift is made (see **Publication Criteria** section); thus, shifting the data does not improve the data. Decisions for data shifts and corrections must be resolved by the hydrographer and members of the review team.

Decisions about the costs of more frequent service visits compared to the decrease in accuracy and precision in the record must be considered carefully for every water-quality monitoring site for various seasons and flows. It generally is understood that the more publishable record there is for a site, the more consistently the data objectives can be met for accuracy and precision. So it follows that it may take some time to finalize the appropriate service intervals at new installations.

Fouling may result from several sources and also may be event-related. Identification of electronic drift or loss of sensor sensitivity should be distinguished from fouling drift, if at all possible. The degree of fouling is determined from the difference between sensor measurements before and after the sensors are cleaned.

Modern temperature sensors are sturdy, dependable, and resistant to fouling. Fouling has the least effect on temperature sensors, which is fortunate because modern sensor packages use internal thermistors to provide continuous temperature corrections to specific conductance, DO, pH, and turbidity values, all of which are temperature dependent.

Specific conductance sensors can be affected by fouling problems when mineralization, severe sedimentation (for example, during the recession period of a flood), or aquatic growth occurs. In general, the sensors are sturdy, easy to clean, and the calibrated field meters are reliable. The specific conductance correction for fouling generally is a datum correction.

Dissolved oxygen sensors are susceptible to fouling of the Teflon membrane, loss of elasticity of the membrane, and chemical alteration of the cathode–anode measuring electrodes. Fouling of the membrane can include coating from organic or oily substances, siltation, attachment of aquatic organisms (for example, barnacles in estuarine locations), growth of algae, or deposition of other materials. Replacement of the sensor membrane is simple but requires calibration and eliminates the ability to distinguish drift from fouling. The reconstruction of reliable record ranges from straight forward (for example, onset of a flood) to complex (for example, gradual growth of an algal coating). The DO error correction for the appropriate part of the service interval generally is by use of a datum correction in ADAPS.

Chemical alteration of the DO electrodes can be caused by either a strong oxidizing or reducing chemical agent, such as in a spill situation, a metal-rich drainage basin, or organic-rich waters, such as a wetland. A more common alteration is sulfide poisoning over a longer period of time. Poisoning would be corroborated by chronically low dissolved-oxygen readings, even after sensor membrane replacement. For many investigations, sulfide poisoning usually is evident as a grey or black color on the silver anode.

Continuous records of DO and turbidity are often the most difficult to maintain within acceptable limits for accuracy and precision. Because of this, the time period between service visits may need to be shortened to maintain the quantity and quality of the DO or turbidity record. Sensor fouling, as a result of biological or chemical particles accumulating on the sensors, can occur on any monitor left in the stream for a period of time. Although fouling frequently begins as soon as the monitor is placed in the stream for deployment, a rise in gage height generally will contribute more to fouling overall. If a turbidity probe is equipped with wiper or shutter technology and is properly maintained, the need for a fouling correction will be minimized. The optic mechanism on the turbidity sensor is extremely sensitive to build up; if the probe is not equipped with wiper or shutter technology, accuracy of turbidity data will be impaired significantly. A fouling shift generally is applied as a datum correction from either the last time the sensor was cleaned, from the last rise in gage height, or from a significant event noted in the field notes.

Fouling of a pH sensor can be similar to the fouling of a DO membrane. The pH sensor must be cleaned delicately in order to avoid scratching the glass surface, which would render it useless. The before- and after-cleaning measurements of pH are critical because the chance of sensor drift is higher for pH than for the other four sensors. Distinction between drift and fouling in the field may be difficult; since replacement of a pH sensor is often required as a troubleshooting step, it may not be possible to distinguish drift from

fouling. Accurate instrument log information on pH monitors is vital in recognizing sensor as opposed to monitor malfunctions. The consequence in ADAPS is a single datum correction for both drift and fouling. The preferred method of correction in ADAPS is the variable shift program.

Drift

A calibration drift is an electronic drift in the equipment from the last time it was calibrated and is determined by the difference between readings of a cleaned sensor in standards or buffers and a calibrated sensor. If, after checking, the deviation from calibrations is within the calibration criteria of the sensor (table 6), then no sensor drift is present. Drift is assumed to occur at a constant rate across the service interval. If the sensor readings exceed the shift criteria (table 7), then the correction is a linear interpolation over the time between calibration checks. This is called a prorated shift in ADAPS (Bartholoma, 1997) and is illustrated for temperature in figure 5.

The Automated Data-Processing System (ADAPS) does not have an option for recording the percentage of shift by recorded value, but it does have a provision that allows three standards to be entered as value adjustments for a measured parameter. Corrections to the record can thus be applied as a 3-point variable shift in ADAPS. This value-dependent adjustment is called a variable shift and is applied in situations where standards at the beginning and ending of the service interval have different amounts of deviation (variable shifts). The equivalent situation

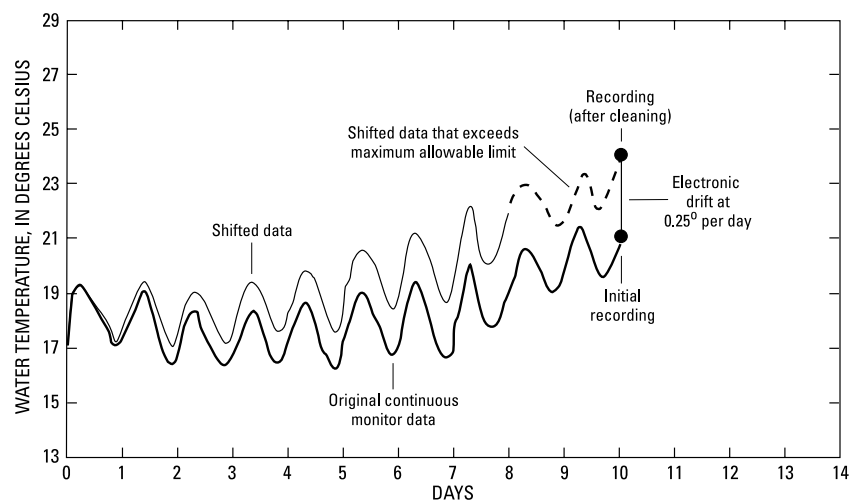


Figure 5. Generalized graph of corrections to continuous temperature record.

for computation of streamflow in ADAPS is called a V-shift (Bartholoma, 1997). A graphic representation of a variable shift for specific conductance is shown in Appendix 8. Variable shifts can be accomplished for any parameter, and this is the recommended technique for corrections related to instrument drift. Because three standards frequently are used for monitor calibration, the variable shift is the preferred method of correction for drift (see Appendix 8).

The treatment for suspected sensitivity loss is sensor reconditioning or replacement. The pH sensors are particularly prone to sensitivity loss (Busenberg and Plummer, 1987). Sensitivity loss in the pH sensor may result from a partially clogged reference electrode junction or a change in the concentration of the potassium chloride filling solution. Distinguishing between pH sensor drift and electronic drift is possible by determining the sensor slope; however, the correction is the same. Improvements in electronic devices in the past two decades have greatly reduced the amount of true electronic drift.

Evaluation and Application of Cross-Section Corrections

The purpose of cross-section corrections is to adjust the measurements of the monitoring equipment to reflect conditions more accurately in the entire stream cross section, from bank to bank and surface to bottom. The principal value of such adjusted measurements is realized when concurrent discharge measurements permit the computations of constituent loads transported past the station. Primary tables and plots from ADAPS that represent drift and fouling corrections should be made before the application of corrections to the cross section. The application of cross-section corrections is intended to improve the accuracy and representativeness of monitoring measurements; however, cross-section corrections should be made only if the variability in the cross section exceeds the shift criteria.

Corrections to the cross section are based on field measurements taken both horizontally and vertically in the stream cross section. Shared characteristics of the specific corrections for temperature, specific conductance, DO, pH, and turbidity are that they may differ from each other at each site, and they all are site dependent and may be discharge and seasonally dependent as well. Several considerations govern the use of corrections. Cross-section measurements must be made under a variety of

flow conditions (minimum of six), all seasonal conditions, and at least twice annually to confirm that the correction being applied to the water-quality measurements truly represents the median or mean cross-section values. Corrections should not be applied to water-quality measurements beyond the range of discharge measurements nor during periods of unsteady flow. If the correction for a measured physical property is consistent across a range of discharges and seasons, then correction of the measurement values by simple adjustment is warranted. Seasonal changes in water quality may be representative of the hydrologic system, but they also may be a result of local conditions.

Often, an increase in specific conductance may accompany low-flow conditions in basins where ground water has higher specific conductance than surface water. It is important to consider, however, that during summer low-flow conditions, extremes in temperature, pH, and DO may be representative of local biological activity rather than seasonal or discharge-related cross-sectional variability. Professional judgment and observations in the field logs will assist in evaluating the need for the application of cross-section corrections. Continuous water-quality monitoring has great value in recognizing the interdependence of water-quality physical properties under varying flow regimes. Understanding these relations is essential in the appropriate application of discharge-dependent corrections.

Final Data Evaluation

Final data evaluation consists of reviewing the data record, checking shifts, and making any needed final corrections. When completed, the data are verified for publication and then rated for quality. Data that cannot be verified or are rated as unacceptable are retained for record-checking and review purposes but are not published or stored in ADAPS. Unacceptable or unverified data, however, should be archived following established District policies.

Maximum Allowable Limits for Reporting Continuous Data

Systematic adoption of a standardized final data-evaluation process, including maximum allowable limits and publication criteria, are important components in finalizing District water-quality records. Many USGS District offices have established quality-control limits for shifting data. These

commonly are referred to as “maximum allowable limits.” The concept is simple—if the recorded values differ from the field-measured values by more than the maximum allowable limits, the data are not published. For the purpose of consistency within the USGS and to inform data users of the maximum allowable limits, the limits are established at 10 times the calibration criteria, except for more stringent requirements for DO and turbidity, for all standard continuous-monitoring, data-gathering activities (table 8).

Table 8. Maximum allowable limits for continuous water-quality monitoring sensors

[±, plus or minus value shown; °C, degree Celsius; mg/L, milligram per liter; pH unit, standard pH unit]

Measured physical property	Maximum allowable limits for water-quality sensor values
Temperature	± 2.0 °C
Specific conductance	± 30 percent
Dissolved oxygen	The greater of ± 2.0 mg/L or 20 percent
pH	± 2 pH units
Turbidity	± 30 percent

The maximum allowable limits shown in table 8 are considered minimum standards for quality, and Districts are encouraged to establish stricter requirements. Even with the establishment of maximum allowable limits, professional judgment by the hydrographer still is needed in record processing. For example, a decision must be made regarding at what point in the data-collection interval the physical property values exceeded the maximum allowable limit. In the cases of instrument drift, gradual fouling, or event-related fouling, reasonable judgment can be

made as to the time in the record interval when the maximum allowable limits were exceeded. The hydrographer computing the record and the record reviewer must agree on this determination and note the time when the maximum allowable limits were exceeded on the station analysis sheet.

Publication Criteria

An assessment of accuracy is required for the publication of all continuous water-quality records. After the record has been evaluated for maximum allowable limits, one of four accuracy classifications is applied to each measured physical property on a scale ranging from poor to excellent. The accuracy rating is based on data values recorded before any shifts or corrections are made (table 9).

Additional consideration must be given to the amount of publishable record and to the amount of data that have been corrected or shifted. Missing data or data that do not meet the criteria for maximum allowable limits can have much to little or no effect on the accuracy of the published data, depending on the situation; the decision to publish the data is left to the professional judgment of the hydrographer. However, to publish a daily value for any physical property, the following minimum requirements apply:

1. On days when less than 100 percent of the data collected at a discrete fixed or varying time interval (unit values) are available, in order to publish a daily mean value, both the expected daily maximum and daily minimum values must be present.

Table 9. Rating continuous water-quality records

[≤, less than or equal to; ±, plus or minus value shown; °C, degree Celsius; >, greater than; %, percent; mg/L, milligram per liter; pH unit, standard pH unit]

Measured physical property	Ratings			
	Excellent	Good	Fair	Poor
Water temperature	≤ ± 0.2 °C	> ± 0.2 to 0.5 °C	> ± 0.5 to 0.8 °C	> ± 0.8 °C
Specific conductance	≤ ± 3%	> ± 3 to 10%	> ± 10 to 15%	> ± 15 %
Dissolved oxygen	≤ ± 0.3 mg/L	> ± 0.3 to 0.5 mg/L	> ± 0.5 to 0.8 mg/L	> ± 0.8 mg/L
pH	≤ ± 0.2 unit	> ± 0.2 to 0.5 unit	> ± 0.5 to 0.8 unit	> ± 0.8 unit
Turbidity	≤ ± 5%	> ± 5 to 10%	> ± 10 to 15%	> ± 15 %

2. Daily maximum and minimum values may be published independently if they occur during the expected times and when the following criteria are met:

Water temperature—A minimum of 25 percent of the unit values is present, and flow is gradually changing or at a steady state.

Specific conductance—A minimum of 50 percent of the unit values is present, and flow is gradually changing or at a steady state. Consider the relation of specific conductance to streamflow.

Dissolved oxygen—A minimum of 25 percent of the unit values is present, and flow is gradually changing or at a steady state. Consider the temperature and pH relation.

pH—A minimum of 25 percent of the unit values is present, and flow is gradually changing or at a steady state.

Turbidity—A minimum of 50 percent of the unit values is present, and flow is gradually changing or at a steady state. Consider specific conductance/turbidity correlation. Reporting daily mean values when less than 100 percent of the unit values is available is a judgment decision and generally determined by data trends. If more than 50 percent of the recorded data is missing, then the mean value should be reported as estimated.

The current station description format for the USGS annual data report requires a description of the specific service interval ratings (Novak, 1985). The record-rating system described in this report must be systematically applied to all continuous water-quality monitoring data published in USGS reports. Accuracy ratings are listed in the REMARKS section of the station description for USGS annual reports. The appropriate rating and accuracy of each water-quality monitoring record also must be included in the station analysis form (see **Preparation of the Review Package**).

The calculation of daily mean pH values in ADAPS should be discontinued in District water-quality data-collection programs and replaced by the calculation of median daily pH values or a more accurate computational approach for the calculation of mean pH values. The correct calculation involves the conversion of pH units to hydrogen-ion concentrations, calculation of a mean concentration value, and conversion back into pH units (as shown in Appendix 1). This additional step is mathematically correct but tends to bias the results toward lower pH values than the simpler median value.

Preparation of the Review Package

Typically, the same individual who services the water-quality monitoring site also computes the water-quality monitoring record, writes the water-quality station analysis that describes annual operation of the site, makes necessary changes to the station description, and prepares a package of auxiliary information to aid in the review of the record.

Record Checking

All data used in producing the final water-quality record should be checked thoroughly for completeness and accuracy before final review and publication. The hydrographer who is responsible for computation of the water-quality record should review the record, followed by a second check for completeness and accuracy by an experienced hydrographer before final inspection by the USGS District water-quality specialist or District-designated reviewer. All field data should be verified for accuracy of transcription from field sheets; all shifts should be checked to assure that the correct values were used for a shift; and all dates and numbers in the station manuscript should be checked for accuracy.

Station Description

All stations are required to have a station description that identifies the location, history, and operation of the station. A complete description is prepared for each new station and is revised only upon changes in location or operation. The contents of a continuous water-quality station description vary, but in addition to the minimum requirements of location, history, and operation of the station, the station description includes information on instrumentation and the location and placement of water-quality sensors (see Appendix 2).

Station Analysis

An individual station analysis is required annually and includes an analysis of each constituent monitored at a site. The purpose of the station analysis is to compile information annually from the field notes, instrument log, and station description that bear directly on the accuracy and completeness of the record. At a minimum, the station analysis must include the instrumentation, period of record, sampling procedures, special conditions, and any variations from

normal procedure (see Appendix 3). In addition, the station analysis includes information relating to the accuracy of the instrumentation and the record produced for the year, dates that probes were changed or replaced, procedures and methods used in the calibration process, a statement of how corrections or shifts were applied to the data, and any unusual events that may have affected the record. Reasons are given for how data were validated or why anomalous record was removed or retained. Details are recorded relating to how and when cross-section measurements were made and variations throughout the cross section.

Contents of the Review Package

The record review package includes the following figures, forms, or tables (examples of the figures, forms, or tables are included in the Appendix section of this report):

1. Station description form (Appendix 2)
2. Station analysis form (Appendix 3)
3. USGS Water-Quality Monitor Field-Inspection Form (Appendix 4)
4. Primary computations table from ADAPS (Appendix 7)
5. Annual data report draft manuscript (Appendix 9)
6. Annual field measurement summary form (Appendix 10)
7. Water-quality variable shift correction tables from ADAPS (Appendix 11)
8. Datum correction value table from ADAPS (Appendix 12)
9. Computed (type 3) daily-values tables (Appendix 13)
10. Discharge hydrograph (Appendix 14)
11. Graph of individual water-quality physical property for review (Appendix 15)

Documentation

The methods for collecting, correcting, and processing continuous water-quality monitoring data are defined and publication standards are established. Systematic application of monitoring site operation, the record rating system, and a standard record-review process are part of the necessary quality assurance in producing and documenting complete and accurate water-quality monitoring records. The use of

DECODES for data conversion and ADAPS for data processing ensures national consistency and a valid data trail. Additionally, participation in the National Field Quality-Assurance Program (Stanley and others, 1998) ensures the proficiency of the analyst and the instruments used in making measurements.

Level of Review Required

Another individual (a senior colleague or section supervisor) examines the record-review package for completeness and accuracy. If areas of interpretation exist (for example, when to start or terminate a shift), a discussion between these two individuals is held. When all issues are resolved, the decisions are documented on the appropriate form. The completed record-review package is then inspected by the District water-quality specialist or District-designated reviewer for completeness and accuracy.

PUBLICATION

It is the policy of the USGS that, with only a few exceptions, all data collected by the USGS are published or made publicly available. Although paper reports are the traditional means of publication, data currently (2000) are being made available in other electronic formats, such as computer disks, compact disks-read only to memory (CD-ROMs), or the World Wide Web. The same requirements for checking for completeness and accuracy of the record apply to all formats of publication.

Data Reports

All nonproprietary water-quality data collected during the water year are published in the USGS annual data report, "Water Resources Data, [State name], Water Year __," or in individual project data reports. Approval of hydrologic data reports is in accordance with applicable USGS policies (Alt and Iseri, 1986). USGS policy (U.S. Department of the Interior, 1992) states that data and information are to be published; however, publication is not limited to paper media (U.S. Geological Survey, 1990; U.S. Department of the Interior, 1993). Electronic formats for public distribution include computer storage media, such as CD-ROM and the World Wide Web.

Data Qualification Statements

The accuracy of published water-quality monitoring records is described by the rating of the record and in the **Publication Criteria** section (table 9), and the accuracy attributed to the water-quality records is indicated in the REMARKS paragraph of the manuscript station description (Appendix 9). Different accuracies may be attributed to different parts of a given record.

ARCHIVING OF RECORDS

According to USGS policy, all original data that are published or support published scientific analyses are to be archived (Hubbard, 1992; U.S. Geological Survey, 1992). Original data—from automated data-collection sites, laboratories, outside sources, and nonautomated field observations—are unmodified data as collected or received and in conventional units (engineering units, generally with a decimal). Original data should be preserved in this form, no matter how they may be modified later (Hubbard, 1992). Recent USGS policy (U.S. Geological Survey, 1999) reaffirms the requirement of preserving original unaltered field data, and further requires that the data be preserved in computer-readable (electronic) digital format. In addition, paper copies of digital field-recorded time-series data are no longer required. All personnel responsible for the collection, analysis, manipulation, and storage of water-quality monitoring data should ensure that the specified requirements of archiving original electronic data are implemented.

In addition to electronic field data, original water-quality monitoring data on paper may include field notes, field measurements, calibration notes, analytical service requests, and water-quality analytical printouts. These data are archived when the project is completed or terminated, or if data are more than 7 years old. It is the responsibility of the District Chief to ensure that project files are entered into the District archive and are organized and complete. The District archive is well documented and maintained by specified personnel in the District.

SUMMARY

Recording systems that measure physical and chemical water-quality properties at discrete time

intervals provide a nearly continuous record of water quality. A common configuration is the four-property water-quality monitoring system that collects temperature, specific conductance, dissolved oxygen, and pH data, although systems can be configured to measure other properties such as turbidity or chlorophyll. The sensors that are used to measure these water properties require careful field observation, cleaning, and calibration. Producing an accurate final record requires thorough procedures for the computation, publication, and archiving of the data.

This report provides guidelines for U.S. Geological Survey site-selection considerations, sensor test methods, field procedures, error correction, data computation, and review and publication processes. Emerging sensor technology is broadening the variety of measurable chemical constituents and will continue to lower the limits of detection. Recent improvements have been made in our ability to make near real-time water-quality monitoring data available through the World Wide Web. This progress will improve applications and allow quality-control procedures to be refined.

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