

**Understanding and Accounting for Method
Variability in Whole Effluent Toxicity Applications
Under the National Pollutant Discharge Elimination
System Program**

June 30, 2000

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NOTICE AND DISCLAIMER

This document provides guidance to NPDES regulatory authorities and persons interested in whole effluent toxicity testing. This document describes what EPA believes to be sources of variability in the conduct of whole effluent toxicity testing under the Clean Water Act. The document is designed to reflect national policy on these issues. The document does not, however, substitute for the Clean Water Act, an NPDES permit, or EPA or State regulations applicable to permits or whole effluent toxicity testing; nor is this document a permit or a regulation itself. The document does not and cannot impose any legally binding requirements on EPA, States, NPDES permittees, and/or laboratories conducting whole effluent toxicity testing for permittees (or for States in the evaluation of ambient water quality). EPA and State officials retain discretion to adopt approaches on a case-by-case basis that differ from this guidance based on an analysis of site-specific circumstances. This guidance may be revised without public notice to reflect changes in EPA policy.

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State Case Example and Chapter 4

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EXECUTIVE SUMMARY

Background

The Federal Water Pollution Control Act, commonly known as the Clean Water Act, was enacted in 1972 with the objective of “*restoring the chemical, physical, and biological integrity of the Nation’s waters.*” Among the U.S. Environmental Protection Agency’s (EPA’s) efforts toward this objective is the National Pollutant Discharge Elimination System (NPDES) program. This program is designed to control toxic discharges, implement water quality standards, and restore waters to “fishable and swimmable” conditions. Point sources that discharge pollutants must do so under the terms and conditions of an NPDES permit. One approach EPA employs to control toxic pollutants under the NPDES permits program is using whole effluent toxicity (WET) controls.

EPA is issuing this document to both address questions raised on WET test method variability and to satisfy a requirement of a July 1998 settlement agreement with litigants for the Western Coalition of Arid States (WestCAS) and Edison Electric Institute et al. This document was developed by an EPA workgroup consisting of EPA’s Office of Water’s (OW) Headquarters, Office of Enforcement and Compliance Assurance, Office of Research and Development, and Regional staff. The document was externally peer reviewed in accordance with EPA’s peer review guidelines. The document addresses WET test method variability by identifying the potential sources of variance associated with WET testing, discusses how to minimize it and, finally, describes how to address it within the NPDES permitting program. The document cites both Agency and external ongoing research on this topic and scientific findings, particularly technical information that support efforts to minimize WET test result variability.

While the document provides recommendations on how to reduce or minimize WET test variability, the document does not supersede current Agency guidance, policy, or regulation, including EPA’s promulgated test methods (40 CFR Part 136), which remain in effect. EPA expects that implementation of the NPDES program and NPDES permits will continue to comply with regulatory requirements and follow applicable EPA guidance and policy.

Why WET Testing?

Whole effluent toxicity is the aggregate toxic effect of an aqueous sample (e.g., effluent, receiving water) measured directly by an aquatic toxicity test. Aquatic toxicity tests are laboratory experiments that measure the biological effect (e.g., growth, survival, and reproduction) of effluents or receiving waters on aquatic organisms. In aquatic toxicity tests, organisms of a particular species are held in test chambers and exposed to different concentrations of an aqueous sample, for example, a reference toxicant, an effluent, or a receiving water, and observations are made at predetermined exposure periods. At the end of the test, the responses of test organisms are used to estimate the effects of the toxicant or effluent.

Whole effluent toxicity test results are an integral tool in the assessment of water quality. For the protection of aquatic life, the integrated strategy includes the use of three control approaches: the chemical-specific control approach, the WET control approach, and the biological criteria/bioassessment/bioassay approach. The primary advantage of using WET controls over individual, chemical-specific controls is that WET integrates the effects of all chemical(s) in the aqueous sample. Reliance solely on chemical-specific numeric criteria or biological criteria would result in only a partially effective State toxics control program. These toxicity tests therefore must be performed using best laboratory practices, and every effort must be made to enhance repeatability of the test method. This document presents EPA’s approaches to achieve the goals listed below.

Effect of This Guidance

This document clarifies several issues regarding WET variability and reaffirms EPA's guidance in the *Technical Support Document for Water Quality-Based Toxics Control* (TSD, USEPA 1991a). This document provides NPDES regulatory authorities and all stakeholders, including permittees, with guidance and recommendations on how to address WET variability. EPA's recommendations and conclusions are detailed in Chapter 7, and Appendix C provides sample NPDES permit language reflecting these recommendations.

The most significant recommendation is to use and report the values for the percent minimum significant difference (PMSD) with all WET data results. The minimum significant difference (MSD) represents the smallest difference between the control mean and a treatment mean that leads to the statistical rejection of the null hypothesis (i.e., no toxicity) at each concentration of the WET test dilution series. The MSD provides an indication of within-test variability and test method sensitivity. Using this information, the regulatory authority and permittees can better evaluate WET test results.

This document makes several other recommendations, such as continue to use the TSD statistical approach without adjusting for test method variability, obtain sufficient representative effluent samples, verify effluent toxicity data against reference toxicant data, maintain clear communication between the regulatory authority and permittee, and maintain good laboratory checks and certification programs.

Three Goals of This Document

This document describes three goals EPA has defined to address issues surrounding WET variability. In addition, the document is intended to satisfy the requirements of a settlement agreement to resolve litigation over rulemaking to standardize WET testing procedures.

1. Quantify the variability of promulgated test methods and report a coefficient of variation (CV) as a measure of test method variability (see Chapter 3 and Appendix A).
2. Evaluate the statistical methods described in the *Technical Support Document for Water Quality-Based Toxics Control* (TSD) for determining the need for and deriving WET permit conditions (see Chapter 6 and Appendix G).
3. Suggest guidance for regulatory authorities on approaches to address and minimize test method variability (Chapter 6). In addition, the document is intended to provide guidance to regulatory authorities, permittees, and testing laboratories on conducting the biological and statistical methods and evaluating test effect concentrations (Chapter 5).

Data Evaluated

EPA assembled a comprehensive data base to examine variability in the WET test methods from the EPA Regions, several States, and private laboratories, which represent a widespread sampling of typical laboratories and laboratory practices. EPA applied several criteria to the data before they were accepted, including detailed sample information, strict adherence to published EPA WET test methods, and test acceptability criteria (TAC). The resulting data base contains data from 75 laboratories for 23 methods for tests concluded between 1988 and 1999.

Approach Taken To Evaluate Test Method Variability

The variability that EPA is assessing is associated with replicate tests using reference toxicants and WET testing methods within analytical laboratories. The focus of this guidance is *not* to quantify test variability between laboratories or to quantify the total variability of WET tests conducted on effluents. Rather, the purpose is to quantify method variability within laboratories (repeatability) to enable NPDES

programs to distinguish between variability caused by the testing method and variability associated with toxicity of multiple effluent samples taken from the same facility.

To quantify test method variability within and between laboratories using this data base, EPA examined two key parameters: (1) the effect concentrations [effect concentration (EC25), lethal concentration (LC50), no observed effect concentration (NOEC)] estimated by the test, which are used to derive WET permit limits and evaluate self-monitoring data with those limits; and (2) the minimum significant difference (MSD), which summarizes the variability of organism responses at each test concentration within an individual test. The MSD represents the smallest difference that can be distinguished between the response of the control organisms and the response of the organisms exposed to the aqueous sample. The MSD provides an indication of within-test variability and test method sensitivity.

Principal Conclusions

The principal conclusions of this document follow.

Evaluation of Test Method Variability

- Comparisons of WET method precision with method precision for analytes commonly limited in NPDES permits clearly demonstrate that the variability of the promulgated WET methods is within the range of variability experienced in other types of analyses. Several independent researchers and studies also have concluded that method performance improves when prescribed methods are followed closely by experienced analysts (Section 4.3).
- This document provides interim CVs for promulgated WET methods in Appendix A, Tables A-1 (acute methods) and A-2 (chronic methods), pending completion of between-laboratory studies, which may affect these interim CV estimates.

Evaluation of Approach To Incorporate Test Method Variability

- EPA's TSD presents guidance for developing effluent limits that appropriately protect water quality, regarding both effluent variability and analytical variability, provided that the WET criteria and waste load allocation (WLA) are derived correctly (Section 6 and Appendix G).
- EPA's analysis of data gathered in the development of this document indicates that the TSD approach appropriately accounts for both effluent variability and method variability. EPA does not believe a reasonable alternative approach is available to determine a factor that would discount the effects of method variability using the TSD procedures, because the approach would not ensure adequate protection of water quality (Section 6.1.1 and Appendix G).

Development of Guidance to Regulatory Authorities

- EPA recommends that regulatory authorities implement the statistical approach as described in the TSD to evaluate effluent for reasonable potential and to derive WET limits or monitoring triggers (Section 6.1 and Appendix G).
- EPA recommends that regulatory authorities calculate the facility-specific CVs using point estimate techniques to determine the need for and derive a permit limit for WET, even if self-monitoring data are to be determined using hypothesis testing techniques, for example, to determine a "no effect" concentration ("NOEC"). This document describes such facility-specific calculations (Section 3.4.1 and 6.2).

Additional Recommendations and Guidance

This document also provides recommendations and guidance on minimizing variability in three specific areas in order to generate sound WET test results: (1) obtaining a representative effluent sample; (2) conducting the toxicity tests properly to generate the biological endpoints; and (3) conducting the appropriate statistical analysis to obtain defensible effect concentrations (EC25, LC50, NOEC). If these recommendations are addressed, the reliability of the test endpoint values should improve.

- **Regulatory Authorities:** Design a sampling program that collects representative effluent samples to fully characterize effluent variability for a specific facility over time (Sections 6.1.3 and 6.2).
- **Regulatory Authorities:** Ensure proper application of WET statistical procedures and test methods (Sections 5.2 through 5.5).
- **Regulatory Authorities:** Incorporate both the upper and lower bounds using the percent minimum significant difference (PMSD) to control and to minimize within-test method variability and increase test sensitivity. To achieve the PMSD upper bound, either the replication should increase or within-test method variability should decrease, or both (Section 6.4 and Table 3-6).
- **Testing Laboratories:** Encourage WET testing laboratories to maintain control charts for PMSD and the control mean and report the PMSD with all WET test results (Section 5.3.1.1).
- **Regulatory Authorities:** Participate in the National Environment Laboratory Accreditation Program and routine performance audit inspections to evaluate laboratory performance (Section 5.3.1.1).
- **Regulatory Authorities:** Incorporate EPA's guidance on error rate assumption adjustments, concentration-response relationships, confidence intervals, acceptable dilution waters, how to block by parentage for the chronic *Ceriodaphnia dubia* test, and control of pH drift (USEPA 2000a).

LIST OF ACRONYMS AND ABBREVIATIONS¹

ACR	acute-to-chronic ratio
AML	average monthly limit
ANOVA	analysis of variance
APHA-AWWA- WEF	American Public Health Association-American Water Works Association-Water Environment Federation
ASTM	American Society for Testing and Materials
BSAB	Biomonitoring Science Advisory Board
CCC	criteria continuous concentration
CFR	Code of Federal Regulations
CMC	criteria maximum concentration
CV	coefficient of variation
CWA	Clean Water Act
DMR	discharge monitoring report
EMS	error mean square [also referred to as mean square error (MSE)]
EPA	U.S. Environmental Protection Agency (also, the Agency)
FR	<i>Federal Register</i>
IC	inhibition concentration
IWC	instream waste concentration (sometimes referred to as receiving water concentration)
LC50	lethal concentration, 50 percent
LOEC	lowest observed effect concentration
LTA	long-term average (LTAA = acute LTA; LTAc = chronic LTA; LTAA,c = acute-to-chronic LTA)
MDL	maximum daily limit
MSD	minimum significant difference
MSE	mean square error [also referred to as error mean square (EMS)]
MZ	mixing zone
NELAP	National Environment Laboratory Accreditation Program
NOEC	no observed effect concentration
NPDES	National Pollutant Discharge Elimination System
NTRD	National Toxicant Reference Database
PAI	Performance Audit Inspections
PMSD	percent minimum significant difference

¹ Note: These acronyms and abbreviations may have other meanings in other EPA programs or documents.

QA	quality assurance
QC	quality control
rMSE	square root of the mean square error
RP	reasonable potential
RWC	receiving water concentration (sometimes referred to as instream waste concentration)
SCTAG	Southern California Toxicity Assessment Group
SETAC	Society of Environmental Toxicology and Chemistry
TAC	test acceptability criteria
TIE	toxicity identification evaluation
TMDL	total maximum daily load
TRE	toxicity reduction evaluation
TSD	EPA's <i>Technical Support Document for Water Quality-based Toxics Control</i> (March 1991, EPA505/2-90-001)
TU	toxic unit (TU _a = acute toxicity; TU _c = chronic toxicity)
VF	variability factor
WET	whole effluent toxicity
WLA	waste load allocation
WQBEL	water quality based effluent limit

GLOSSARY

Acute Toxicity Test is a test to determine the concentration of effluent or ambient waters that causes an adverse effect (usually death) on a group of test organisms during a short-term exposure (e.g., 24, 48, or 96 hours). Acute toxicity is measured using statistical procedures (e.g., point estimate techniques or a *t*-test).

Acute-to-Chronic Ratio (ACR) is the ratio of the acute toxicity of an effluent or a toxicant to its chronic toxicity. It is used as a factor for estimating chronic toxicity on the basis of acute toxicity data, or for estimating acute toxicity on the basis of chronic toxicity data.

Ambient Toxicity is measured by a toxicity test on a sample collected from a receiving waterbody.

ANOVA is analysis of variance.

Average Monthly Limit (AML) is the calculated average monthly limit of waste load allocation assigned by a State or EPA for a particular facility.

CCC are water quality criteria for chronic exposure (criteria continuous concentrations).

Chronic Toxicity Test is a short-term test in which sublethal effects (e.g., reduced growth or reproduction) are usually measured in addition to lethality. Chronic toxicity is defined as $TUc = 100/NOEC$ or $TUc = 100/ECp$ or ICp .

CMC are water quality criteria for acute exposures (criteria maximum concentration).

Coefficient of Variation (CV) is a standard statistical measure of the relative variation of a distribution or set of data, defined as the standard deviation divided by the mean. It is also called the relative standard deviation (RSD). The CV can be used as a measure of precision within (within-laboratory) and between (between-laboratory) laboratories, or among replicates for each treatment concentration.

Confidence Interval is the numerical interval constructed around a point estimate of a population parameter.

Effect Concentration (EC) is a point estimate of the toxicant concentration that would cause an observable adverse effect (e.g., death, immobilization, or serious incapacitation) in a given percent of the test organisms, calculated from a continuous model (e.g., Probit Model). EC_{25} is a point estimate of the toxicant concentration that would cause an observable adverse effect in 25 percent of the test organisms.

Hypothesis Testing is a statistical technique (e.g., Dunnett's test) for determining whether a tested concentration is statistically different from the control. Endpoints determined from hypothesis testing are $NOEC$ and $LOEC$. The two hypotheses commonly tested in WET are:

Null hypothesis (H_0): The effluent is not toxic.

Alternative hypothesis (H_a): The effluent is toxic.

Inhibition Concentration (IC) is a point estimate of the toxicant concentration that would cause a given percent reduction in a non-lethal biological measurement (e.g., reproduction or growth), calculated from a continuous model (i.e., Interpolation Method). IC_{25} is a point estimate of the toxicant concentration that would cause a 25-percent reduction in a non-lethal biological measurement.

Instream Waste Concentration (IWC) is the concentration of a toxicant in the receiving water after mixing. The IWC is the inverse of the dilution factor. It is sometimes referred to as the receiving water concentration (RWC).

LC50 (lethal concentration, 50 percent) is the toxicant or effluent concentration that would cause death in 50 percent of the test organisms.

Lowest Observed Effect Concentration (LOEC) is the lowest concentration of an effluent or toxicant that results in adverse effects on the test organisms (i.e., where the values for the observed endpoints are statistically different from the control).

Long-term Averages (LTAs) of pollutant concentration or effluent toxicity are calculated from waste load allocations (WLAs), typically assuming that the WLA is a 99th percentile value (or another upper bound value) based on the lognormal distribution. One LTA is calculated for each WLA (typically an acute LTA and a chronic LTA for aquatic life protection). The LTA represents expected long-term average performance from the permitted facility required to achieve the associated WLA.

Maximum Daily Limit (MDL) is the calculated maximum WLA assigned by a State or EPA for a particular facility.

Minimum Significant Difference (MSD) is the magnitude of difference from control where the null hypothesis is rejected in a statistical test comparing a treatment with a control. MSD is based on the number of replicates, control performance, and power of the test.

Mean Square Error (MSE) is the average dispersion of the items around the treatment means. It is an estimate of a common variance, the within variation, or variation among observations treated alike. [Also referred to as error mean square (EMS).]

Mixing Zone is an area where an effluent discharge undergoes initial dilution and is extended to cover the secondary mixing in the ambient waterbody. A mixing zone is an allocated impact zone where water quality criteria can be exceeded as long as acutely toxic conditions are prevented.

No Observed Effect Concentration (NOEC) is the highest tested concentration of an effluent or toxicant that causes no observable adverse effect on the test organisms (i.e., the highest concentration of toxicant at which the values for the observed responses are not statistically different from the controls).

National Pollutant Discharge Elimination System (NPDES) program regulates discharges to the nation's waters. Discharge permits issued under the NPDES program are required by EPA regulation to contain, where necessary, effluent limits based on water quality criteria for the protection of aquatic life and human health.

Power is the probability of correctly detecting an actual toxic effect (i.e., declaring an effluent toxic when, in fact, it is toxic).

Precision is a measure of reproducibility within a data set. Precision can be measured both within a laboratory (within-laboratory) and between laboratories (between-laboratory) using the same test method and toxicant.

Quality Assurance (QA) is a practice in toxicity testing that addresses all activities affecting the quality of the final effluent toxicity data. QA includes practices such as effluent sampling and handling, source and condition of test organisms, equipment condition, test conditions, instrument calibration, replication, use of reference toxicants, recordkeeping, and data evaluation.

Quality Control (QC) is the set of more focused, routine, day-to-day activities carried out as part of the overall QA program.

Reasonable Potential (RP) is where an effluent is projected or calculated to cause an excursion above a water quality standard based on a number of factors.

Reference Toxicant Test is a check of the sensitivity of the test organisms and the suitability of the test methodology. Reference toxicant data are part of a routine QA/QC program to evaluate the performance of laboratory personnel and the robustness and sensitivity of the test organisms.

Significant Difference is defined as a statistically significant difference (e.g., 95 percent confidence level) in the means of two distributions of sampling results.

Statistic is a computed or estimated quantity such as the mean, standard deviation, or coefficient of variation.

Test Acceptability Criteria (TAC) are specific criteria for determining whether toxicity test results are acceptable. The effluent and reference toxicant must meet specific criteria as defined in the test method (e.g., for the *Ceriodaphnia dubia* survival and reproduction test, the criteria are as follows: the test must achieve at least 80 percent survival and an average of 15 young per surviving female in the controls).

Total Maximum Daily Load (TMDL) is a determination of the amount of a pollutant, or property of a pollutant, from point, nonpoint, and natural background sources, including a margin of safety, that may be discharged to a water quality-limited waterbody.

t-Test (formally Student's *t*-Test) is a statistical analysis comparing two sets of replicate observations, in the case of WET, only two test concentrations (e.g., a control and 100 percent effluent). The purpose of this test is to determine if the means of the two sets of observations are different [e.g., if the 100-percent effluent concentration differs from the control (i.e., the test passes or fails)].

Type I Error (alpha) is the rejection of the null hypothesis (H_0) when it is, in fact, true (i.e., determining that the effluent is toxic when the effluent is not toxic).

Type II Error (beta) is the acceptance of the null hypothesis (H_0) when it is not true (i.e., determining that the effluent is not toxic when the effluent is toxic).

Toxicity Test is a procedure to determine the toxicity of a chemical or an effluent using living organisms. A toxicity test measures the degree of effect of a specific chemical or effluent on exposed test organisms.

Toxic Unit-Acute (TUa) is the reciprocal of the effluent concentration (i.e., $TUa = 100/LC50$) that causes 50 percent of the organisms to die by the end of an acute toxicity test.

Toxic Unit-Chronic (TUc) is the reciprocal of the effluent concentration (e.g., $TUc = 100/NOEC$) that causes no observable effect (NOEC) on the test organisms by the end of a chronic toxicity test.

Toxic Unit (TU) is a measure of toxicity in an effluent as determined by the acute toxicity units (TUa) or chronic toxicity units (TUc) measured. Higher TUs indicate greater toxicity.

Toxicity Identification Evaluation (TIE) is a set of procedures used to identify the specific chemicals causing effluent toxicity.

Toxicity Reduction Evaluation (TRE) is a site-specific study conducted in a step-wise process designed to identify the causative agents of effluent toxicity, isolate the source of toxicity, evaluate the effectiveness of toxicity control options, and then confirm the reduction in effluent toxicity.

Variance is a measure of the dispersion in a set of values, defined as the sum of the squared deviations divided by their total number.

Whole Effluent Toxicity (WET) is the total toxic effect of an effluent measured directly with a toxicity test.

Waste Load Allocation (WLA) is the portion of a receiving water's total maximum daily load that is allocated to one of its existing or future point sources of pollution.

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1.0 INTRODUCTION

1.1 Background

The Federal Water Pollution Control Act, commonly known as the Clean Water Act (CWA), was enacted in 1972 with the objective of “*restoring the chemical, physical, and biological integrity of the Nation’s waters.*” Several goals and policies were established in the Act, including the following:

- Eliminating the discharge of pollutants into navigable waters by 1985;
- Wherever attainable, achieving an interim goal of water quality that provides for the protection and propagation of fish, shellfish, and wildlife, and provides for recreation in and on the water by November 1, 1983; and
- Prohibiting the discharge of toxic pollutants in toxic amounts.

In the 28 years since the CWA was enacted, the U.S. Environmental Protection Agency (EPA) and States authorized to administer EPA’s National Pollutant Discharge Elimination System (NPDES) permitting program have made significant progress toward achieving these goals. NPDES is designed to control toxic discharges, implement a water quality standards program, and restore waters to “fishable and swimmable” conditions. A point source that discharges pollutants to waters of the United States must do so under the terms and conditions of an NPDES permit. In setting these terms and conditions, EPA and the States have integrated their control of toxic pollutants through combined use of three approaches [*Technical Support Document for Water Quality-based Toxics Control* (USEPA 1991a, referred to as the TSD)]:

- Chemical-specific controls,
- Whole effluent toxicity (WET) controls, and
- Biological criteria/bioassessments and bioassays.

The WET approach to protection of water quality is the primary subject of this document.

In 1989, EPA defined whole effluent toxicity as “*the aggregate toxic effect of an effluent measured directly by an aquatic toxicity test*” [54 Federal Register (FR) 23868 at 23895, June 2, 1989]. Aquatic toxicity tests are laboratory experiments that measure the biological effect (e.g., growth, survival, and reproduction) of effluents or receiving waters on aquatic organisms. In aquatic toxicity tests, groups of organisms of a particular species are held in test chambers and exposed to different concentrations of an aqueous test sample, for example, a reference toxicant, an effluent, or a receiving water. Observations are made at predetermined exposure periods. At the end of the test, the responses of test organisms are used to estimate the effects of the toxicant or effluent.

In the early 1980s, EPA published methods (USEPA 1985, 1988, 1989) for estimating the short-term acute and chronic toxicity of effluents and receiving waters to freshwater and marine organisms. WET data gathered in the 1980s indicated that approximately 40 percent of NPDES facilities nationwide discharged an effluent with sufficient toxicity to cause water quality problems. Further reductions in the toxicity of wastewater discharges were necessary to achieve compliance with narrative water quality standards expressed as “no toxics in toxic amounts.” In response to these findings, EPA implemented a policy to reduce or eliminate toxic discharges. The *Policy for the Development of Water Quality-based Permit Limitations for Toxic Pollutants* (49 FR 9016, March 9, 1984) introduced EPA’s integrated toxics control program. To support this policy, EPA developed the TSD (USEPA 1991a). The TSD provides guidance to

regulators in implementing WET testing requirements in NPDES permits. In 1989, EPA promulgated regulations specifying procedures for determining when water quality-based effluent limitations are required in NPDES permits [40 CFR, 122.44(d)]. On October 26, 1995, EPA promulgated WET test methods (USEPA 1993, 1994a, and 1994b) and added them to the list of EPA methods approved under Section 304(h) of the CWA (40 CFR, 136) for use in the NPDES program. Although the rulemaking was challenged in court, that challenge has been stayed pending completion of a settlement agreement. The rulemaking remains in force and effect unless and until EPA takes further action.

1.2 Effect of This Guidance

This document attempts to clarify several issues regarding WET variability and reaffirms EPA's earlier guidance and recommendations published in the TSD (USEPA 1991a). This document is intended to provide NPDES regulatory authorities and all stakeholders, including permittees, with guidance and recommendations on how to understand and account for measurement variability in WET testing. The document's recommendations and conclusions are detailed in Section 7. Appendix C provides sample NPDES permit language reflecting these recommendations.

The most significant recommendation is to use and report the values for the percent minimum significant difference (PMSD) with all WET data results. The minimum significant difference (MSD) is the smallest difference that can be distinguished between the response of control organisms and the response of test organisms at each concentration of the WET test dilution series. The MSD provides an indication of the within-test variability and test method sensitivity. Using this information, the regulatory authority and permittees can better evaluate WET test results.

This document also recommends the following:

- Continue to use the EPA TSD statistical approach for NPDES permit limit development (no test method variability adjustments are needed);
- Collect and evaluate a sufficient number of representative effluent samples;
- Verify effluent toxicity data carefully along with reference toxicant data;
- Maintain good communication between the regulatory authority and permittee throughout all phases of the permitting process;
- Implement the PMSD to evaluate both WET and reference toxicant data to minimize within-test method variability and increase test sensitivity;
- Maintain laboratory checks with good laboratory certification programs to encourage experienced laboratories and skilled analysts for the toxicity testing program for individual WET laboratory performance.

1.3 Three Goals of This Document

EPA prepared this document to achieve the following three goals:

1. Quantify the variability of promulgated test methods and report a coefficient of variation (CV) as a measure of test method variability (see Chapter 3 and Appendix A).

2. Evaluate the statistical methods described in the *Technical Support Document for Water Quality-Based Toxics Control* (TSD) for determining the need for and deriving WET permit conditions (see Chapter 6 and Appendix G).
3. Suggest guidance for regulatory authorities on approaches to address and minimize test method variability (Chapter 6). In addition, the document is intended to provide guidance to regulatory authorities, permittees, and testing laboratories on conducting the biological and statistical methods and evaluating test effect concentrations (Chapter 5).

This document does not address effluent variability. It does, however, discuss how handling effluent samples can affect tests. Chapter 2 provides definitions of terms used and discusses the ways in which variability can be quantified. Chapter 3 describes the variability of the effect concentration estimates (EC25, LC50, and NOEC) and the variability of endpoint measurements (survival, growth, and reproduction). Chapter 4 discusses WET variability in the context of chemical-specific method variability. Chapter 5 provides guidance to permittees, testing laboratories, and regulatory authorities to minimize test method variability. Chapter 6 provides guidance to regulatory authorities on how to determine reasonable potential (RP) and derive permit limits or monitoring triggers and evaluate self-monitoring data. Chapter 7 presents EPA's principal conclusions. Chapter 8 is a bibliography containing a list of documents cited herein and additional reading material.

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2.0 DEFINITION AND MEASUREMENT OF METHOD VARIABILITY IN WET TESTING

The terms used to express toxicity test results are defined in this chapter, and methods for quantifying WET test method variability are discussed. Additional terms used throughout this document, along with their definitions, are provided in the Glossary as part of the front matter of this document.

2.1 Terms and Definitions

Biological endpoints are the biological observations recorded when conducting toxicity tests. These observations may include the number of surviving organisms or the number of young produced. There are two basic types of biological endpoints: responses recorded as response/no response (e.g., dead or alive) are quantal data; responses recorded as a measured response (e.g., weight) or as a count (e.g., number of young produced) are considered continuous data. For most WET tests, the observations for each tested concentration are combined and then reported as an average or percentage to represent the biological endpoint. For example, the fathead minnow larval survival and growth chronic test method has two biological endpoints (i.e., percent survival and average dry weight for each test concentration).

Effect concentrations are concentrations of a test material (i.e., effluent, referent toxicant, receiving water) derived from the observed biological endpoints followed by data analysis using either hypothesis testing procedures or point estimate techniques. Effect concentrations derived using point estimation techniques represent the concentration of a test material at which a predetermined level of effect occurs. For example, LC50 is the lethal concentration at which 50 percent of the organisms respond. Effect concentrations commonly estimated for WET methods are LC50, EC50 (effect concentration at which a 50-percent effect occurs), and IC25 (inhibition concentration at which a 25-percent effect occurs). Hypothesis test methods are used to determine the no observed effect concentration (NOEC). The NOEC represents the highest effect concentration in the test concentration response that is not significantly different from the control response. Multiple statistical endpoints can be derived for each WET method. For example, the endpoints for the fathead minnow larval survival and growth chronic test can be reported as an EC25 for growth, an NOEC for growth, an LC50 (or EC50) for survival, and an NOEC for survival.

2.2 Defining WET Test Variability

As with any measurement process, WET tests have a degree of variability associated with the test method performance. Three measures of variability related to WET tests are within-test variability, within-laboratory variability, and between-laboratory variability.

- **Within-test (intra-test) variability** is the variability in test organism response within a concentration averaged across all concentrations of the test material in a single test.
- **Within-laboratory (intra-laboratory) variability** is the variability that is measured when tests are conducted using specific methods under reasonably constant conditions in the same laboratory. Within-laboratory variability, as used in this document, includes within-test variability. The American Society for Testing and Materials (ASTM) uses the term “repeatability” to describe within-laboratory variability. Repeatability is estimated (as a sample variance or standard deviation) by repeating a test method under realistically constant conditions within a single laboratory.
- **Between-laboratory (inter-laboratory) variability** is the variability between laboratories. It is measured by obtaining results from different laboratories using the same test method and the same

test material (e.g., reference toxicant). Between-laboratory variability, as used in this document, does *not* include the within-laboratory component of variance. ASTM uses the term “reproducibility” to describe between-laboratory variability. Reproducibility is estimated by having nearly identical test samples (duplicates or splits) analyzed by multiple laboratories using similar standard methods. Although reproducibility is generally synonymous with between-laboratory variability, estimates of reproducibility may combine within-laboratory and between-laboratory components of variance, making between-laboratory variability numerically larger than within-laboratory variability as defined above.

For purposes of consistency, EPA uses the terms within-laboratory and between-laboratory variability throughout this document.

Numerous factors can affect the variability of any toxicity test method. These factors include the number of test organisms, the number of treatment replicates, randomization techniques, the source and health of the test organisms, the type of food used, laboratory environmental conditions, and dilution water quality. The experience of the analyst performing the test, analyzing the data, and interpreting the results may also affect variability (Grothe et al. 1996, Fulk 1996).

2.3 Quantifying WET Test Variability

Historically, information on the variability of toxicity tests has been developed using effect concentrations, such as the NOEC, EC25, EC50, and LC50 for survival, fecundity, and growth. Variability measures should be quantified based on the end use of the data (i.e., effect concentrations) and be directly related to the WET permit requirement. Typically, the effect concentrations are the endpoints used for evaluating self-monitoring results. The variability of the effect concentrations is quantified by obtaining multiple test results under similar test conditions using the same test material. For example, the sample standard deviation and mean for EC25 obtained from multiple monthly reference toxicant tests for the fathead minnow survival and growth chronic test conducted at one laboratory would quantify “within-laboratory” variability for that laboratory. EPA used this approach to evaluate data for the development of this document (see Chapter 3).

Examining variability for each effect concentration of each biological endpoint for each test method is essential. The biological endpoints may be different for various toxicants and effluents. One biological endpoint, such as reproduction, may be more sensitive to a certain toxicant than another endpoint, such as survival. That sensitivity may be reversed for a different toxicant. Alternatively, an endpoint may be more sensitive to one toxicant than another toxicant.

Three other measures of variability (which are not addressed in this document) that have been applied to WET tests are:

1. Determine the variability of the biological endpoint response. For example, the variance of the biological response (e.g., growth and survival) can be calculated. This approach is useful, but does not quantify variability of the WET test effect concentration, which is important in the context of this document.
2. Quantify the uncertainty of each test point estimate (e.g., the EC50, EC25, or LC50) using confidence intervals, which reflect within-test variability.
3. Use the standard deviation to quantify the uncertainty in the mean of the replicate response at each concentration within a particular test. For example, laboratories can compare the standard deviations of the average weight of fathead minnow larvae in four chronic tests at one test concentration, such as 1 mg/L sodium chloride. These standard deviations may be pooled across

all the concentrations when data have been transformed (if necessary) to give similar variances at each concentration. From the pooled variance, one may calculate a minimum significant difference (MSD) value, which is a useful indication of test sensitivity (see Chapters 3 and 5). In this document, the standard deviation at each concentration was not evaluated as a measure of variability. However, the MSD was considered as a measure of WET test variability.

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3.0 VARIABILITY OF WET TEST METHODS

Chapter 3 describes the variability of effect concentration estimates (EC25, LC50, and NOEC) and endpoint measurements (survival, growth, and reproduction). For definitive studies of the variability of WET methods, readers should also refer to the TSD (USEPA 1991a, Part 1.3.3) and to WET methods manuals (USEPA 1993, 1994a, 1994b). EPA will complete and report on a new between-laboratory study of promulgated methods in 2000 or 2001.

3.1 Acquisition, Selection, and Quality Assurance of Data Presented in This Document

EPA solicited data for reference toxicant tests from laboratories that conduct WET tests and use reference toxicant testing as part of their quality control (QC) program. Reference toxicant testing is required, as specified in EPA toxicity test methods, to document laboratory performance over time for laboratories conducting self-monitoring tests. When laboratories are conducting effluent tests, at least one reference toxicant test must be conducted each month using the same toxicant, test concentrations, dilution water, and data analysis methods. These reference toxicant tests must be conducted using the same test conditions (type of dilution water, temperature, test protocol, and species) that are used for WET tests conducted by the laboratory.

Reference toxicant tests were used to characterize method variability because, in contrast to effluent samples, fixed concentrations of known toxicants are used. Only with this standardization is it possible to conclude that variability of the effect concentration estimates is derived from the sources discussed above, rather than from changes in the toxicant.

EPA received reference toxicant test data from several States, private laboratory sources, and the EPA Regions. Data sources used for these analyses include the EPA National Toxicant Reference Database (NTRD), the EPA Region 9 Toxicity Data Base, and laboratory bench sheets voluntarily submitted by independent sources. Although the data do not represent a random sample of laboratories or tests, they do represent a widespread sampling of typical laboratories and practices.

EPA required that reference toxicant tests included in its data base meet the following four criteria:

1. Test records documented the test method, organism, test date, laboratory, reference toxicant, and individual biological responses in the concentration series.
2. Data for each replicate were provided as required in the published method using the current test method.
3. The test used at least five toxicant concentrations and a control for the most commonly reported chronic toxicity test methods—(1) 1000.0, fathead minnow larval survival and growth; (2) 1002.0, *Ceriodaphnia* survival and reproduction; and (3) 1006.0, inland silverside survival and growth. For other chronic toxicity test methods, the test used at least four toxicant concentrations and a control because the methods permitted, in the recent past, the use of only four concentrations.
4. EPA personnel or an EPA contractor calculated the effect concentration, verified that all test acceptability criteria (TAC) had been met, and verified that the statistical flowchart had been followed correctly. Thus, all summary statistics and estimates were calculated from the replicate data and strictly followed the most current EPA test methods.

Details of data quality assurance and test acceptance are provided in a separate document, available at EPA's Office of Water docket, located in the Office of Science and Technology ["Whole Effluent Toxicity (WET) Data Test Acceptance and Quality Assurance Protocol"]. An attachment to that document provides a laboratory-by-laboratory listing of quality assurance flags, test dates, and toxicant concentrations, as well as summary statistics by laboratory for the NOEC, EC25, and LC50 estimates and test endpoints (survival, growth, reproduction, etc.). Laboratories are not identified by name.

The data set of reference toxicant tests includes information from 75 laboratories for 23 methods for tests conducted between 1988 and 1999. This document addresses, and provides specific guidance on, the variability of methods promulgated by EPA in 40 CFR Part 136 (Table 3-1). The data are also used to develop between-laboratory interim estimates of method variability for the promulgated methods (Appendix A). The Agency identifies these CVs as "interim;" EPA may revise some or all of these estimates based on between-laboratory studies to evaluate some of the promulgated test methods.

The next section presents summary statistics for the promulgated methods. Summary statistics for all methods in the data set appear in Appendix B. For methods represented by a few laboratories, summary statistics should not be considered representative of method performance. For example, EPA's Office of Water usually relies on acceptable data from at least six laboratories (USEPA 1996b) when it conducts a multi-laboratory study to quantify method performance. The data used here have not been obtained under conditions as rigorous as those applied to a between-laboratory study and for that reason, may overestimate variability, particularly for the extremes.

Coefficients of variation are used as descriptive statistics for NOECs in this document. Because NOECs can take on only values that correspond to concentrations tested, the distribution (and CV) of NOECs can be influenced by the selection of experimental concentrations, as well as additional factors (e.g., within-test variability) that affect both NOECs and point estimates. This makes CVs for NOECs more uncertain than the CVs for point estimates, and the direction of this uncertainty is not uniformly toward larger or smaller CVs. Despite these confounding issues, CVs are used herein as the best available means of expressing the variability of interest in this document and for general comparisons among methods. Readers should be cautioned, however, that small differences in CVs between NOECs and point estimates may be artifactual; large differences are more likely to reflect real differences in variability (a definition of what is "small" or "large" would require a detailed statistical analysis and would depend upon the experimental and statistical details surrounding each comparison). NOECs can only be a fixed number of discrete values; the mean, standard deviation, and CV cannot be interpreted and applied as they are for a continuous variable such as the EC25 or EC50. For instance, the typical reference toxicant test might result in only three observed NOEC values, most of them at one or two concentrations. The mean will fall between tested concentrations, as will the stated confidence intervals; thus, these do not actually represent expected outcomes, only approximations of the expected outcome.

As an alternative to CVs, ratios are used to quantify variability of EC25, EC50, and NOEC measurements in Appendix B. Ratios of measurements have been used previously to quantify and compare variability of NOEC and EC50 (Chapman et al. 1996b, Dhaliwal et al. 1997).

3.2 Variability of EC25, LC50, and NOEC

3.2.1 Within-Laboratory Variability of EC25, LC50, and NOEC

This section characterizes the within-test and within-laboratory variability of effect concentration estimates. Tables 3-2 through 3-4 summarize variation across laboratories of the within-laboratory coefficients of variation (CVs), without respect to reference toxicant tested. Tables showing more extensive summaries appear in Appendix B (Tables B-1 through B-3).

Table 3-1. Promulgated WET Methods Included in This Report

Test Method No.	Test Method	EPA Data Base		
		Toxicants	Tests	Labs
Freshwater Methods for Chronic Toxicity^a				
1000.0	<i>Pimephales promelas</i> , Fathead Minnow Larval Survival and Growth Test	Cd, Cr, Cu, KCl, NaCl, NaPCP, SDS	205	19
1000.0	<i>Pimephales promelas</i> , Fathead Minnow Embryo-Larval Survival and Teratogenicity Test		0	0
1002.0	<i>Ceriodaphnia dubia</i> , Water Flea Survival and Reproduction Test	Cd, Cu, KCl, NaCl, NaPCP	393	33
1003.0	<i>Selenastrum capricornutum</i> , ^b Green Alga Growth Test	Cu, NaCl, Zn	85	9
Marine & Estuarine Methods for Chronic Toxicity^c				
1004.0	<i>Cyprinodon variegatus</i> , Sheepshead Minnow Larval Survival and Growth Test	Cd, KCl	57	5
1005.0	<i>Cyprinodon variegatus</i> , Sheepshead Minnow Embryo-larval Survival and Teratogenicity Test		0	0
1006.0	<i>Menidia beryllina</i> , Inland Silverside Larval Survival and Growth Test	Cr, Cu, KCl, SDS	193	16
1007.0	<i>Americamysis (Mysidopsis) bahia</i> , Mysid Survival, Growth, and Fecundity Test	Cr, Cu, KCl	130	10
1008.0	<i>Arbacia punctulata</i> , Sea Urchin Fertilization Test		0	0
1009.0	<i>Champia parvula</i> , Red Macroalga Reproduction Test	Cu, SDS	23	2
Methods for Acute Toxicity^{d,e}				
2000.0	Fathead Minnow Survival Test	Cd, Cu, KCl, NaCl, NaPCP	217	21
2002.0	<i>Ceriodaphnia dubia</i> Survival Test	Cd, Cu, KCl, NaCl, NaPCP	241	23
2004.0	Sheepshead Minnow Survival Test	SDS	65	3
2006.0	Inland Silverside Survival Test	Cd, KCl, SDS	48	5
2007.0	Mysid (<i>A. bahia</i>) Survival Test	Cd, Cu, SDS	32	3
2011.0	Mysid (<i>H. costata</i>) Survival Test	Cd, SDS	14	2
2019.0	Rainbow Trout Survival Test	Cu, Zn	10	1
2021.0	<i>Daphnia magna</i> Survival Test	Cd	48	5
2022.0	<i>Daphnia pulex</i> Survival Test	Cu, NaCl, SDS Cd, Cu, NaCl, NaPCP	57	6

^a See publications EPA/600/4-89-001 (USEPA 1989) and EPA/600/4-91-002 (USEPA 1994b).

^b The genus and species names for *Selenastrum capricornutum* have been changed to *Raphidocelis subcapitata*. In this document, however, *Selenastrum capricornutum* is used to avoid confusion.

^c See publication EPA/600/4-91-003 (USEPA 1994a) and EPA/600/4-87/028 (USEPA 1988).

^d See publications EPA/600/4-85/013 (USEPA 1985) and EPA/600/4-90/027F (USEPA 1993).

^e EPA did not assign method numbers for acute methods in EPA/600/4-90/027F. The numbers assigned here were created for use in this document and in related materials and data bases.

Reference toxicant codes:

Cd	cadmium	NaCl	sodium chloride
Cr	chromium	NaPCP	sodium pentachlorophenate
Cu	copper	SDS	sodium dodecyl sulfate
KCl	potassium chloride	Zn	zinc

Table 3-2. Quartiles (25th and 75th) and Median (50th) of the Within-Laboratory Values of CV for EC25 (Chronic Tests)

Test Method ^a	Test Method No.	Endpoint ^b	No. of Labs	Percentiles of CV		
				25 th	50 th	75 th
Fathead Minnow Larval Survival & Growth	1000.0	G	19	0.21	0.26	0.38
Fathead Minnow Larval Survival & Growth	1000.0	S	16	0.11	0.22	0.32
<i>Ceriodaphnia</i> (Cd) Survival & Reproduction	1002.0	R	33	0.17	0.27	0.45
<i>Ceriodaphnia</i> (Cd) Survival & Reproduction	1002.0	S	25	0.11	0.23	0.41
Green Alga (<i>Selenastrum</i>) Growth	1003.0	G	6	0.25	0.26	0.39
Sheepshead Minnow Larval Survival & Growth	1004.0	G	5	0.09	0.13	0.14
Sheepshead Minnow Larval Survival & Growth	1004.0	S	2	0.15	0.16	0.17
Inland Silverside Larval Survival & Growth	1006.0	G	16	0.18	0.27	0.43
Inland Silverside Larval Survival & Growth	1006.0	S	13	0.22	0.35	0.42
Mysid (Ab) Survival, Growth, & Fecundity	1007.0	F	4	0.30	0.38	0.41
Mysid (Ab) Survival, Growth, & Fecundity	1007.0	G	10	0.24	0.28	0.32
Mysid (Ab) Survival, Growth, & Fecundity	1007.0	S	7	0.17	0.21	0.28
Red Macroalga (<i>Champia parvula</i>) Reproduction	1009.0	R	2	0.58	0.58	0.59

^a Cd = *Ceriodaphnia dubia*, Ab = *Americamysis (Mysidopsis) bahia*

^b G = growth, S = survival, R = reproduction, F = fecundity

Table 3-3. Quartiles (25th and 75th) and Median (50th) of the Within-Laboratory Values of CV for LC50

Test Method ^a	Test Method No.	Endpoint	No. of Labs	Percentiles of CV		
				25 th	50 th	75 th
Freshwater Methods for Chronic Toxicity^c						
Fathead Minnow Larval Survival & Growth	1000.0	S	19	0.15	0.23	0.31
<i>Ceriodaphnia</i> (Cd) Survival & Reproduction	1002.0	S	33	0.10	0.16	0.29
Sheepshead Minnow Larval Survival & Growth	1004.0	S	5	0.07	0.08	0.12
Inland Silverside Larval Survival & Growth	1006.0	S	16	0.16	0.28	0.35
Mysid (Ab) Survival, Growth, & Fecundity	1007.0	S	10	0.16	0.26	0.27
Methods for Acute Toxicity^{d,e}						
Fathead Minnow Larval Survival	2000.0	S	21	0.10	0.16	0.19
<i>Ceriodaphnia</i> (Cd) Survival	2002.0	S	23	0.11	0.19	0.29
Sheepshead Minnow Survival	2004.0	S	5	0.12	0.14	0.21
Inland Silverside Larval Survival	2006.0	S	5	0.15	0.16	0.21
Mysid (Ab) Survival	2007.0	S	3	0.17	0.25	0.26
Mysid (Hc) Survival	2011.0	S	2	0.27	0.30	0.34
Rainbow Trout Survival	2019.0	S	1	0.23	0.23	0.23
<i>Daphnia</i> (Dm) Survival	2021.0	S	5	0.07	0.22	0.24
<i>Daphnia</i> (Dp) Survival	2022.0	S	6	0.19	0.21	0.27

^a Cd = *Ceriodaphnia dubia*, Ab = *Americamysis (Mysidopsis) bahia*, Hc = *Holmesimysis costata*, Dm = *Daphnia magna*, Dp = *Daphnia pulex*

^b S = survival

^c See publications EPA/600/4-89-001 (USEPA 1989) and EPA/600/4-91-002 (USEPA 1994b).

^d See publications EPA/600/4-85-013 (USEPA 1985) and EPA/600/4-90/027F (USEPA 1993).

^e EPA did not assign method numbers for acute methods in EPA/600/4-90/027F. The numbers assigned here were created for use in this document and in related materials and data bases.

Table 3-4. Quartiles (25th and 75th) and Median (50th) of the Within-Laboratory Values of CV for NOEC

Test Method ^a	Test Method No.	Endpoint	No. of Labs	Percentiles of CV		
				25 th	50 th	75 th
Freshwater Methods for Chronic Toxicity^c						
Fathead Minnow Larval Survival & Growth	1000.0	G	19	0.22	0.37	0.53
Fathead Minnow Larval Survival & Growth	1000.0	S	19	0.26	0.39	0.48
<i>Ceriodaphnia</i> (Cd) Survival & Reproduction	1002.0	R	33	0.25	0.33	0.49
<i>Ceriodaphnia</i> (Cd) Survival & Reproduction	1002.0	S	33	0.21	0.30	0.43
Green Alga (<i>Selenastrum</i>) Growth	1003.0	G	9	0.40	0.46	0.56
Marine & Estuarine Methods for Chronic Toxicity^d						
Sheepshead Minnow Larval Survival & Growth	1004.0	G	5	0.34	0.40	0.44
Sheepshead Minnow Larval Survival & Growth	1004.0	S	5	0.14	0.18	0.24
Inland Silverside Larval Survival & Growth	1006.0	G	16	0.31	0.46	0.57
Inland Silverside Larval Survival & Growth	1006.0	S	16	0.30	0.42	0.55
Mysid (Ab) Survival, Growth, & Fecundity	1007.0	F	4	0.17	0.36	0.40
Mysid (Ab) Survival, Growth, & Fecundity	1007.0	G	10	0.35	0.39	0.43
Mysid (Ab) Survival, Growth, & Fecundity	1007.0	S	10	0.28	0.33	0.38
Red Macroalga (<i>Champia parvula</i>) Reprod.	1009.0	R	2	0.85	1.00	1.16
Methods for Acute Toxicity^{e,f}						
Fathead Minnow Larval Survival	2000.0	S	21	0.18	0.22	0.34
<i>Ceriodaphnia</i> (Cd) Survival	2002.0	S	23	0.18	0.35	0.41
Sheepshead Minnow Survival	2004.0	S	3	0	0.31	0.33
Inland Silverside Larval Survival	2006.0	S	5	0	0.33	0.35
Mysid (Ab) Survival	2007.0	S	3	0.29	0.38	0.43
Mysid (Hc) Survival	2011.0	S	2	0.21	0.26	0.31
Rainbow Trout Survival	2019.0	S	1	0.35	0.35	0.35
<i>Daphnia magna</i> (Dm) Survival	2021.0	S	5	0.09	0.36	0.47
<i>Daphnia pulex</i> (Dp) Survival	2022.0	S	6	0.21	0.38	0.61

^a Cd = *Ceriodaphnia dubia*, Ab = *Americamysis (Mysidopsis) bahia*, Hc = *Holmesimysis costata*, Dm = *Daphnia magna*, Dp = *Daphnia pulex*

^b G = growth, S = survival, R = reproduction, F = fecundity

^c See publications EPA/600/4-89-001 (USEPA 1989) and EPA/600/4/4-91-002 (USEPA 1994b).

^d See publication EPA/600/4-91-003 (USEPA 1994a) and EPA/600/4-87/028 (USEPA 1988).

^e See publications EPA/600/4-85/013 (USEPA 1985) and EPA/600/4-90/027F (USEPA 1993).

^f EPA did not assign method numbers for acute methods in EPA/600/4-90/027F. The numbers assigned here were created for use in this document and in related materials and data bases.

Effect concentrations having a p-percent effect are symbolized as EC_p and may be calculated for sublethal and lethal (survival) endpoints (USEPA 1993,1994a,1994b). Effect concentrations commonly estimated for WET methods are LC50, EC50, IC25, and EC25. The symbol EC_p is more general and may be used to represent an LC_p, EC_p, or IC_p endpoint. To simplify presentation of results in this document, the term EC25 is used to represent the concentration at which a 25-percent effect has occurred for either lethal

or sublethal endpoints. The term LC50 is used to represent the concentration at which a 50-percent effect has occurred for lethal endpoints. The EC25 for survival is not routinely used in generating self-monitoring data and is presented here for comparison to the EC25 for sublethal endpoints (i.e., IC25). Estimates of EC25, LC50, and NOEC were calculated for this document as required in the EPA test methods (USEPA 1993, 1994a, 1994b). A CV is reported for NOEC measurements in this document. See Appendix A for further details.

The results in Tables 3-2 through 3-4 were obtained as follows, using as an example the EC25 of the growth endpoint in Method 1000.0 (fathead minnow larval chronic test) on the first row of Table 3-2. The CV of the EC25 estimates was calculated for each laboratory. This calculation resulted in 19 CVs (one per laboratory with each laboratory tested using one toxicant). The sample percentiles were calculated for this set of 19 CVs. In Table 3-2, the column headed "50th" shows the 50th percentile (median value) of CV found across these 19 laboratories; the 50th percentile value is 0.26. In the column headed "75th," the 75th percentile CV is reported as 0.38. When a method is represented by fewer than four laboratories, the minimum and maximum CVs are shown in the columns headed "25th" and "75th," respectively. Note that these CVs represent within-laboratory variability, and that Tables 3-2 through 3-4 show the quartiles and median of the within-laboratory CVs. These tables thus report the typical range of within-laboratory test method variation.

Variation across laboratories in the CV for effect concentration estimates (Tables 3-2 through 3-4) may be summarized as follows, ignoring methods represented by only one or two laboratories. [Refer to the column headed "75th" (the 75th percentile).]

For the EC25 of the growth and reproduction endpoints in chronic toxicity tests, 75 percent of laboratories have a CV no more than 0.14 to 0.45 depending on the method (Table 3-2). For the two most commonly used methods (1000.0, fathead minnow larval chronic test; and 1002.0, *Ceriodaphnia* chronic test), 75 percent of the laboratories have CVs no more than 0.38 and 0.45, respectively.

For the LC50 of the survival endpoint in chronic toxicity tests, 75 percent of laboratories have a CV no more than 0.12 to 0.35, depending on the method. For the two most commonly used methods (1000.0 and 1002.0), 75 percent of laboratories have CVs no more than 0.31 and 0.29, respectively (Table 3-3). For the LC50 in acute toxicity tests, 75 percent of laboratories have a CV no more than 0.19 to 0.29, depending on the method. For the two most commonly used methods (2000.0 and 2002.0), 75 percent of laboratories have CVs no more than 0.19 and 0.29, respectively.

For the NOEC of growth or reproduction endpoints in chronic toxicity tests, 75 percent of laboratories have a CV no more than 0.43 to 0.57, depending on the method. For the two most commonly used methods (1000.0 and 1002.0), 75 percent of laboratories have CVs no more than 0.53 and 0.49, respectively (Table 3-4). For the NOEC of survival in chronic toxicity tests, 75 percent of laboratories have a CV no more than 0.24 to 0.55, depending on the method. For the two most commonly used methods (1000.0 and 1002.0), 75 percent of laboratories have CVs no more than 0.48 and 0.43, respectively. For the NOEC of survival in acute toxicity tests, 75 percent of laboratories have a CV no more than 0.34 to 0.61, depending on the method. For the two most commonly used acute methods (2000.0 and 2002.0), 75 percent of laboratories have CVs no more than 0.34 and 0.41, respectively.

Appendix B discusses the range of toxicant concentrations reported as the NOEC. For chronic toxicity tests, most laboratories report the NOEC to within two to three concentration intervals, and half the laboratories report most NOECs within one to two concentration intervals for reference toxicants. For acute toxicity tests, most laboratories report NOECs at one or two concentrations. This outcome agrees with EPA's expected performance for these methods. The normal variation of the effect concentration estimate in reference toxicant tests has been reported for some EPA WET methods (USEPA 1994a, 1994b) to be plus or minus one dilution concentration for the NOEC and less for LC50.

3.2.2 Between-Laboratory Variability of EC25, LC50, and NOEC

The data set compiled for this document provided reasonable estimates of between-laboratory variability for only a few methods. For many methods and toxicants, there were too few laboratories in the data base. Additional summaries of between-laboratory variability of WET methods are included in the TSD (USEPA 1991a, Part 1.3.3) and the WET methods manuals (USEPA 1994a, 1994b). EPA also intends to provide new data in a forthcoming EPA between-laboratory study of promulgated methods.

Using the data set, credible estimates of between-laboratory variability could be made for a few toxicants and methods having data for six or more laboratories (Table 3-5). The statistical methods are described in Appendix B. Table 3-5 shows values of the square root of within-laboratory and between-laboratory variance components (i.e., standard deviations, σ). The standard deviations and mean are expressed in units of toxicant concentration (e.g., g/L or mg/L). Between-laboratory σ_b estimates the standard deviation for laboratory means of EC25, LC50, and NOEC. The “Mean” column in Table 3-5 shows the mean of the laboratory means, not the mean for all tests. Because the number of tests differed among laboratories, these two means are different. These data suggest that between-laboratory variability (σ_b) is comparable to within-laboratory variability (σ_w) for the methods listed in the table.

In Table 3-5, the ratio of σ_b to the mean is an estimate of the relative variability (CV_b) of laboratory means around their combined mean. The ratio of σ_w to the mean may approach the value of the average within-laboratory CV when the sample of laboratories is large, but to characterize within-laboratory CVs, readers should use Tables 3-2 through 3-4.

Table 3-5. Estimates of Within-Laboratory and Between-Laboratory Components of Variability^a

Test Method ^b	Test EC Estimate	Toxicant	End-Point ^c	Tests	Labs	Within-lab σ_w	Between-lab σ_b	Mean	CV_w	CV_b
1000.0	EC25	NaCl	G	73	6	0.67	0.44	2.63	0.25	0.17
1000.0	LC50	NaCl	S	73	6	1.14	0.45	4.15	0.27	0.11
1000.0	NOEC	N Cl	G	73	6	0.72	0.35	2.18	0.33	0.16
1000.0	NOEC	NaCl	S	73	6	0.96	0.51	2.43	0.40	0.21
1002.0	EC25	NaCl	R	292	23	0.29	0.27	0.92	0.32	0.29
1002.0	LC50	NaCl	S	285	23	0.48	0.24	1.78	0.27	0.13
1002.0	NOEC	NaCl	G	292	23	0.28	0.18	0.74	0.38	0.24
1002.0	NOEC	NaCl	S	292	23	0.47	0.26	1.42	0.33	0.18
1006.0	EC25	Cu	G	130	9	45.1	52.4	97.4	0.46	0.54
1006.0	LC50	Cu	S	130	9	48.4	70.7	127.0	0.38	0.56
1006.0	NOEC	Cu	G	130	9	51.8	44.4	80.1	0.65	0.55
1006.0	NOEC	Cu	S	130	9	34.2	39.5	65.4	0.52	0.60
2000.0	LC50	NaCl	S	154	14	1.05	1.24	7.46	0.14	0.17
2002.0	LC50	NaCl	S	167	15	0.36	0.38	1.97	0.18	0.19

^a σ_w = within-laboratory standard deviation, σ_b = between-laboratory standard deviation

CV_w = within-laboratory coefficient of variation, CV_b = between-laboratory coefficient of variation

^b EPA did not assign method numbers for acute methods in EPA/600/4-90/027F. The numbers assigned here were created for use in this document and in related materials and data bases.

^c G = growth, S = survival, R = reproduction

3.3 Variability of Endpoint Measurements

This section characterizes the within-laboratory precision of endpoint measurements (e.g., growth, reproduction, and survival). Endpoint variability in methods for chronic toxicity is characterized here using sublethal endpoints. The sublethal endpoint was designed to be more sensitive than the survival endpoint, and it incorporates the effect of mortality (i.e., it incorporates biomass). For example, for the chronic survival and growth fathead minnow larval test, the total dry weight at each replicate is divided by the original number of larvae, rather than the surviving number of larvae.

EPA reports measures of test precision based on the control CV [(control standard deviation)/(control mean)] and the “Percent MSD” [$100 \times \text{MSD} / (\text{control mean})$], symbolized as PMSD. Recall that MSD, the “minimum significant difference,” is calculated as $[d \sqrt{\text{EMS}} \sqrt{(2/r)}]$, where “d” is the critical value of Dunnett’s statistic when comparing “k” treatments to a control, EMS is the error mean square from the analysis of variance of the endpoint responses, and “r” is the number of replicates at each concentration (USEPA 1993, 1994a, 1994b). These measures of test precision quantify within-test variability, or the sensitivity of each test to toxic effects on the biological endpoint.

Measures of variability relative to the control mean are used for two reasons. First, a laboratory having consistently large mean endpoint values for the control will also tend to have larger values of MSD and control standard deviation. Second, PMSD is readily interpreted as the minimum percent difference between control and treatment that can be declared statistically significant in a WET test. A significant effect occurs when (control mean - treatment mean) exceeds the MSD. Dividing by the control mean and multiplying by 100 states this relationship in terms of the percent difference between control and treatment.

To characterize the distribution of values of PMSD, values from all laboratories and toxicants for a given method and endpoint were combined, and sample percentiles reported. Percentiles are also reported for the CV of the control, which also indicates variability among replicates under non-toxic conditions and may be a useful indicator of uniformity of the test organisms. The sample percentiles are reported in more detail in Appendix B; the 10th and 90th percentiles are shown in Table 3-6. Method 1009.0 (red macroalga) is omitted from Table 3-6 because it would be inadvisable to characterize method variability using only 23 tests from only two laboratories.

The 90th percentile may be used as an upper PMSD bound (i.e., a limit on the insensitivity of a test). The 10th percentile may be used as a lower PMSD bound for declaring a significant difference or a lower limit to test sensitivity. The 90th percentile has been used in other WET programs (Chapter 5). The 95th percentile is used as a practical upper limit for the variability of analytical results in well-controlled between-laboratory studies that use a standard protocol and specific quality assurance procedures (ASTM 1992, 1998; USEPA 1993, 1996a, 1996b). The tests summarized here have not been subjected to the rigorous standardization and quality assurance of collaborative studies, and the data have not been screened for outliers as specified by ASTM Practices D2777 and E691 (ATSM 1992, 1998). These considerations justify using the sample 90th percentile to set an upper bound. A lower bound is necessary to avoid creating a disincentive for improving test precision and to objectively specify a limit to the test sensitivity achieved in practice. If no more than ten percent of tests are more precise than this lower bound, then in practice, the analytical method rarely detects toxic effects of this small magnitude.

When comparing values in Table 3-6 to a test result, it is important that the test’s MSD be calculated according to procedures described in the EPA method manuals (USEPA 1993, 1994a, 1994b) for Dunnett’s test for multiple comparisons with a control (see Section 6.4.1). An analysis of variance (ANOVA) is conducted using several treatments, including the control. EPA methods require excluding from the ANOVA those concentrations for which no organisms survived in any replicate. For a sublethal endpoint, concentrations are excluded from the analysis if they exceed the NOEC for survival. The MSD is calculated

using the square root of the error mean square (rEMS) from the ANOVA, and using Dunnett's critical value (which depends on the number of replicates and concentrations used in the ANOVA).

Table 3-6. Range of Relative Variability for Endpoints of Promulgated WET Methods, Defined by the 10th and 90th Percentiles from the Data Set of Reference Toxicant Tests^a

Test Method ^b	Endpoint ^c	No. of Labs	No. of Tests	PMSD		Control CV ^d	
				10 th	90 th	10 th	90 th
1000.0 Fathead Minnow	G	19	205	9.4	35	0.035	0.20
1002.0 <i>Ceriodaphnia dubia</i>	R	33	393	11	37	0.089	0.42
1003.0 Green Alga	G	9	85	9.3	23	0.034	0.17
1004.0 Sheepshead Minnow	G	5	57	6.3	23	0.034	0.13
1006.0 Inland Silverside	G	18	193	12	35	0.044	0.18
1007.0 Mysid	G	10	130	12	32	0.088	0.28
2000.0 Fathead Minnow	S	20	217	4.2	30	0	0.074
2002.0 <i>Ceriodaphnia</i>	S	23	241	5.0	21	0	0.11
2004.0 Sheepshead Minnow	S	5	65	0 ^e	55	0	0
2006.0 Inland Silverside	S	5	48	7.0	41	0	0.079
2007.0 Mysid (<i>A. bahia</i>)	S	3	32	5.1	26	0	0.081
2011.0 Mysid (<i>H. costata</i>)	S	2	14	18	47	0	0.074
2021.0 Daphnia (<i>D. magna</i>)	S	5	48	5.3	23	0	0.11
2022.0 Daphnia (<i>D. pulex</i>)	S	6	57	5.8	23	0	0.11

- ^a The precision of the data warrants only three significant figures. When determining agreement with these values, one may round off values to two significant figures (e.g., values >3.45000... and ≤3.5000... are rounded to 3.5). Method 1009.0 (red macroalga) is not reported because it is inadvisable to characterize method variability using only 23 tests from just two laboratories.
- ^b EPA did not assign method numbers for acute methods in EPA/600/4-90/027F. The numbers assigned here were created for use in this document and in related materials and data bases.
- ^c G = growth, R = reproduction, S = survival
- ^d CVs were calculated using untransformed control means for each test.
- ^e An MSD of zero will not occur when the EPA flow chart for statistical analysis is followed. In this report, MSD was calculated for every test, including those for which the flow chart would require a nonparametric hypothesis test. EPA recommends using the value 4.2 (the 10th percentile shown for the fathead minnow acute test) in place of zero as the 10th percentile PMSD (lower PMSD bound) for the sheepshead minnow acute test.

The MSD was calculated for all test results reported here, including those for which non-normality and heterogeneity of variance were indicated. Thus, this document presents MSD as an approximate index of test sensitivity. Estimates of power are also approximate. The MSD generally will be related to test sensitivity, even when the assumptions for ANOVA and Dunnett's test are not strictly satisfied.

Table 3-7 shows the number of laboratories in the WET variability data set having tests exceeding the upper PMSD bound reported in Table 3-6. One-half to two-thirds of the laboratories never or infrequently exceeded the bound, and roughly one in five exceeded it in at least 20 percent of their tests. By definition of the 90th percentile, about 10 percent of all the tests exceeded the bound.

Table 3-7. Number of Laboratories Having a Given Percent of Tests Exceeding the PMSD Upper Bound for the Sublethal Endpoint

Test Method	No. Labs	Endpoints ^a	Number of Labs with Various Percentages of Tests Exceeding the PMSD Upper Bound				
			0%	0%-10%	10%-20%	20%-50%	50%-100%
1000.0 Fathead Minnow	19	G	8	2	7	2	0
1002.0 <i>Ceriodaphnia dubia</i>	33	R	15	7	5	6	0
1003.0 Green Alga	9	G	6	1	0	2	0
1004.0 Sheepshead Minnow	5	G	3	1	0	1	0
1006.0 Inland Silverside	16	G	6	5	1	4	0
1007.0 Mysid (growth)	10	G	5	2	0	3	0

^a G = growth, R = reproduction

3.4 Conclusions about Variability of WET Methods

3.4.1 Variability of EC25, LC50, NOEC

For EC25, the quartiles of the within-laboratory CVs ranged across the promulgated methods from 0.09 to 0.45, and the median CV ranged from 0.13 to 0.38. For LC50, the quartiles of the within-laboratory CVs ranged from 0.07 to 0.35, and the median CV ranged from 0.08 to 0.28. For NOEC, the quartiles of the within-laboratory CVs ranged from 0 to 0.61, and the median CV ranged from 0.18 to 0.46. This summary applies to those methods represented by at least 20 tests and three laboratories.

EPA concludes from Tables 3-2 through 3-4 that point estimates are substantially less variable than the NOEC for the same method and endpoint, and that the LC50 for an acute toxicity test usually is less variable than the LC50 for a chronic toxicity test. The estimated NOEC is more variable than ECp *using current experimental designs* because NOEC can take only those values equal to the concentrations tested, while ECp interpolates between tested concentrations (there may be other, more technical reasons as well). In principle, NOEC could be estimated more accurately and precisely by changing the experimental design to use more concentrations at narrower dilution ratios and by using more replicates. The greater variability of the NOEC underscores the desirability of using point estimates to characterize effluent toxicity.

Tables 3-2 through 3-4 may be used as benchmarks for variability, allowing comparison of one laboratory's CV for reference toxicant testing with CVs reported by experienced laboratories reporting tests that passed the TAC. However, CVs for methods represented by too few laboratories in the table may be atypical.

The CVs in Tables 3-2 through 3-4 may be used as an adjunct to the control chart. If the CV for reference toxicant tests is above the 75th percentile in Tables 3-2 through 3-4, variability likely can be reduced, even if the individual EC25 or LC50 values fall within the control limits. If a control chart is constructed using an unreasonably large standard deviation, the control limits will be unreasonable. If a high CV is not fully explained by an unusually small mean, the standard deviation of EC25 or LC50 should be reduced to bring the CV within the normal range. If the CV exceeds the 90th percentile (Appendix B), there is no question that variability is unacceptably large. Detailed guidance is provided in Chapter 5 (Section 5.3.1.1).

Tables 3-2 through 3-4 indicate the magnitude of the analytical variability that becomes part of the variability of effluent test results under certain conditions. This occurs when effluent test results (NOECs, LC50s, or EC25s) fall between the lowest and highest concentrations tested. Under other conditions, these

CVs may not accurately represent analytical variability. If tests give results consistently near or at the lowest or highest concentrations tested, or if the tests often produce “less than” or “greater than” results, Tables 3-2 through 3-4 will not accurately characterize the analytical CV for such tests. To measure the analytical CV under such conditions, reference toxicant tests would have to be designed to have the effect concentration at or near the lowest or highest concentration. The CV and standard deviation measured under such conditions are unknown, but are likely to differ from those for standard reference toxicant tests.

The data set did not contain information supporting an analysis of the causes of between-laboratory variability. Possible causes may include laboratory differences in concentration series, incorrect or ambiguous calculation or reporting of concentrations (e.g., concentration of the metal ion versus the salt), laboratory differences in dilution water (e.g., water hardness or pH), laboratory differences in foods and feeding regimes, and laboratory differences in cultures (genotypic and phenotypic differences in sensitivity to various toxicants).

The lack of a standard or common reference toxicant creates a problem for permittees and regulatory authorities attempting to evaluate and compare laboratories. Real or apparent differences occur between laboratories in the mean values of EC25, LC50, and NOEC. Some of this difference is random and reflects only the within-laboratory variance; some may be systematic. Systematic, between-laboratory differences can be inferred reliably only when laboratories use the same test method, use the same reference toxicants and dilution series, use similar dilution waters, and report a sufficient number of tests.

3.4.2 Variability of Endpoint Measurements

EPA has selected the PMSD to characterize endpoint variability for WET test methods because it integrates variability from several concentrations (always including the control), and it represents the MSD used in the WET hypothesis test. The control CV, by itself, does not fully represent the variability affecting a WET hypothesis test or point estimate. The PMSD also represents the variability affecting point estimates because it is calculated using the EMS for the endpoint measurement. (However, the standard error of a point estimate of an effect concentration may be a complicated function of the EMS.)

PMSD for sublethal endpoints ranged from 6 to 37 across the promulgated chronic methods. For the fathead minnow chronic method, PMSD ranged from 9 to 35; for the *Ceriodaphnia* chronic method, PMSD ranged from 11 to 37. Thus, most chronic tests were able to distinguish a reduction of 37 percent or smaller in the endpoint. Further analysis in Chapter 5 shows that most tests were unable to distinguish consistently a 25-percent reduction. For the survival endpoint of promulgated acute methods, PMSD ranged from 0 to 55. For the two most commonly used acute methods (fathead minnow and *Ceriodaphnia*), PMSD ranged from 4 to 30 and from 5 to 21, respectively. Thus, PMSD varied markedly for some acute methods and not for others.

As shown by the size of PMSD, test sensitivity to detect substantial toxic effects is occasionally insufficient at some laboratories and routinely insufficient at a few laboratories. Inadequate test sensitivity is not always signaled by control charts of EC25, LC50, and NOEC. Laboratories should consider maintaining control charts for MSD or PMSD, and should report MSD and the control mean with all WET tests.

Some portion of MSDs in the WET variability data set could be considered exceptionally large, if not outliers. This observation underscores the importance of a careful review for each WET test, including an examination of means and standard deviations for endpoint responses at each concentration; the plotting of replicate data (not just concentration means); and, when necessary, a search for possible causes of excessive variability. The tables and plots in the promulgated methods (USEPA 1994a, 1994b) provide good examples.

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4.0 VARIABILITY IN CONTEXT

EPA manages the regulation of WET in the same way it manages the regulation of chemical-specific pollutants in order to determine reasonable potential (RP), derive permit limits, determine data quality control, and evaluate self-monitoring data. Many similarities between chemical-specific toxicant and WET controls can be found in the TSD (USEPA 1991a). Determining RP in both cases uses many of the same strategies. Permit limit derivation makes similar exposure assumptions and relies on nearly identical toxicological data bases.

Considering a value other than the best analytical estimate as a measure for WET or for specific chemical analytes is inappropriate. All analytical results, in either chemical-specific analyses or WET tests, incorporate some estimated range of uncertainty. While infrequently discussed for chemical methods, uncertainty does play a role in the meaning of analytical results. One end of the confidence interval likely will be less protective of aquatic resources than the other. The derived limit and therefore final reported analytical results become the best estimate of the actual ecological need and assessment of the effect.

Significant debate has occurred over assertions that WET data have too much inherent variability for reliable use in the NPDES program. This debate has engendered considerable evaluation of WET precision. Groups of scientists and individual researchers have repeatedly concluded that currently promulgated WET methods are technically sound and that the observed precision is within the range of precision of other analyses frequently required in NPDES permits (Grothe et al. 1996). The findings of some of the significant sources of these conclusions are summarized below.

4.1 Society of Environmental Toxicology and Chemistry Pellston WET Workshop

The 1995 Society of Environmental Toxicology and Chemistry (SETAC) Pellston Workshop on Whole Effluent Toxicity convened 47 experts in the discipline to assess applied methods and their application in the regulatory process. Representation at the workshop was intentionally balanced among government, business, and academic participants. These scientists published consensus conclusions and recommendations, including the following.

4.1.1 General Conclusions and Recommendations

Grothe et al. (1996) state *“Existing WET testing methods (USEPA 1985, USEPA 1988, USEPA 1989) are technically sound, but certain modifications would improve endpoint interpretation. Such changes involve implementing improvements to currently used statistical procedures, establishing acceptable limits for MSD values, and adding confidence limits to WET test endpoints.”*

“A number of problems with WET tests are caused by misapplication of the tests, misinterpretation of the data, lack of competence of the laboratories conducting WET testing, poor condition/health of test organisms, and lack of training of laboratory personnel, regulators, and permittees. More widespread use of WET related guidance provided in USEPA’s TSD (1991a) would help alleviate some of these problems. In addition, an effective QA/QC program will improve data quality and reduce test variability.”

“Increase training opportunities for regulators and permittees to improve the implementation of WET objectives and to promote national consistency in permitting and compliance issues.”

“Implement a broadly based and standardized QA/QC program to improve WET testing performance and data quality.”

“Quantify the ‘confidence’ around test endpoints to improve interpretation of WET test results. Specific statistical methods that could improve precision are presented in Chapter 3 of this document and processes to reduce variability are discussed in Chapter 5. In addition, WET tests should be performed using a dilution series of exposure concentrations to establish a dose-response relationship.”

4.1.2 Conclusions about Data Precision

Ausley (1996) compared CVs of chemical analyses and aquatic toxicity tests conducted by North Carolina NPDES permittees. Ausley found that CVs of reported values for chemical analytes (including metals, organic analytes, and non-metal inorganic analytes) ranged from 11.8 percent to 291.7 percent. Coefficients of variation for toxicity parameters (acute and chronic *Ceriodaphnia dubia*, acute and chronic *Pimephales promelas*, acute *Daphnia pulex*, and acute *Mysidopsis bahia*) ranged from 14.8 percent to 67.6 percent. From this review, he concluded that *“the precision of toxicity analyses is within the range of that being reported for commonly analyzed and regulated chemical parameters.”* Ausley highlighted the difficulty in comparing precision estimates of chemical analytes and WET analyses (particularly NOECs), noting that while chemical precision is often determined well above analytical detection, WET precision is often based on the minimum detection level. An assumption that WET precision will vary among toxicants is also logical. To establish “inherent variability,” considering toxicants that cause minimal variability in the analysis may be appropriate. The high coefficients of variation for some chemical parameters reported by Ausley reflect the fact that, in practice, analytical precision can vary widely in individual studies in which the effects of a single (or a few) poorly operating laboratory can adversely affect precision estimates. In practice, this kind of data must be screened for quality prior to use to evaluate self-monitoring data or estimates of overall method quality.

Ausley’s results closely approximate analytical precision of chemical analytes referenced in the TSD (USEPA 1991a, Chapter 1.2). The CVs for metals (aluminum, cadmium, chromium, copper, iron, lead, manganese, mercury, silver, and zinc) ranged from 18 percent to 129 percent at the low end of the measurement detection range. Between-laboratory CVs for organic analytes ranged from greater than 12 percent to 91 percent. The CVs for non-metal analytes (alkalinity, residual chlorine, ammonia nitrogen, Kjeldahl nitrogen, nitrate nitrogen, total phosphorus, biological oxygen demand, chemical oxygen demand, and total organic carbon) ranged from 4.6 percent to 70 percent in between-laboratory studies of precision.

Burton et al. (1996) concluded that *“USEPA-published methods are functional and appropriate in the context of effluent toxicity control programs.”* They recommended developing limits on within-test variability, a quality assurance and audit program, and guidance for permittee procurement of WET analytical services.

Denton and Norberg-King (1996) cited various studies that favorably compare WET methods with chemical analytical methods (Grothe and Kimerle 1985, Rue et al. 1988, Morrison et al. 1989, Grothe et al. 1990). They proposed that improvements in test result consistency could be accomplished by limiting the range of within-test variability through controls of upper and lower statistical power (e.g., limits on test MSD). Three practices to control within-test variability most effectively are (1) controlling within-test sensitivity, (2) following well-defined test methods, and (3) maintaining communication within the regulatory community. For example, the permittee and regulatory authorities should discuss any facility-specific issues to fully characterize the appropriate permit conditions.

4.2 Water Environment Research Foundation Study

Another publication, “*Whole Effluent Toxicity Testing Program: Evaluation of Practices and Implementation*” (DeGraeve et al. 1998), presents the results of a survey of publicly owned treatment works and State regulatory programs about WET issues. The Water Environment Research Foundation (WERF) sponsored this study. Conclusions by DeGraeve et al. (1998) include the following:

“The project team believes that the results demonstrate that the test methods can be routinely completed successfully by well-trained, competent WET testing laboratories and that the results, considered collectively, suggest that the test methods that are being used to measure WET are technically sound.”

“There is a need for better training/guidance in WET-related issues for both the regulatory staff responsible for implementing WET requirements and for permittees responsible for meeting WET limits.”

DeGraeve et al. (1998) considered the conclusions of the SETAC Pellston WET publication concurring that between-laboratory CV values of toxicity test methods were low, training of regulatory and permittee staff is needed nationally, and strengthened quality assurance (QA)/quality control (QC) practices could improve performance of analyses. Unlike the SETAC Pellston WET conclusions, they found that there are enough laboratories to meet the current market demand for analyses. Like the SETAC effort, DeGraeve et al. (1998) concluded that a national center of expertise on WET issues would be beneficial to provide guidance to regulatory agencies, permittees, and laboratories.

WERF also funded a project entitled “*Whole Effluent Toxicity Testing Methods: Accounting for Variance*” (Warren-Hicks et al. 1999). This study compared within- and between-laboratory results of reference toxicant test variation as measures of reproducibility and comparability, respectively. The authors concluded that some laboratories could consistently reproduce test results, while others could not and inferred that test precision is a factor of laboratory experience and not inherent methodological weakness. The authors recommended that national studies be conducted to evaluate within- and between-laboratory precision of promulgated WET test methods. (EPA has already initiated this study.) They also recommended that additional test acceptability criteria (TAC), such as upper and lower bounds of MSD, be established and incorporated in the NPDES process. The latter recommendation corroborates other researchers’ recommendations discussed above.

4.3 Minimizing Variability by Adhering to WET Toxicity Test Methods

Specific factors that affect variability in WET analyses have been described in several papers (Burton et al. 1996, Ausley 1996, Erickson et al. 1998, Davis et al. 1998). The most important initial consideration in developing precise data is a laboratory’s experience and success in performing a specific analysis. Most critical reviews of WET data precision emphasize this initial consideration. Experienced professionals most likely will be able to develop the most consistent and reliable information and can interpret anomalous conditions in the testing or results.

An additional factor in considering WET test method variability is whether the prescribed methods (e.g., the EPA toxicity test methods promulgated in 40 CFR Part 136) are being followed appropriately (see Chapter 5). If tests are submitted that do not meet specified TAC or are produced when laboratory QA testing indicates analyses are beyond control limits, these results should not be used in the NPDES process. Tests performed on effluent samples that have not met required temperature maxima or holding times should not be considered for regulatory purposes. Rigorous QA practices are critical to the success of any analytical program. Both the regulatory authority and permittee should strive to ensure that such practices are in place

for any program developing WET data, whether by national laboratory accreditation, State regulatory certification, direct permittee oversight, or specific contractual agreement with the laboratory.

Comparisons of WET method precision with analytes commonly limited in NPDES permits clearly demonstrate that the promulgated WET methods are within the range of variability experienced in other analyses. Several researchers also noted clear indications that method performance improves when prescribed methods are followed closely by experienced analysts (Grothe et al. 1996, DeGraeve et al. 1998).

A review of WET test results confirms that imprecise WET data are being reported. As with any analytical technique, inexperienced individuals can perform analyses incorrectly or fail to follow appropriate methods and quality assurance practices. Using the training that is available for these methods and quality assurance techniques referenced by this document will help ensure that data of maximum reliability are used and that sound decisions are made based on those results. The Western Coalition of Arid States conducted a study in 1997 (Moore et al. 2000), which reported the results of 16 tests with a non-toxic test sample using the *Ceriodaphnia dubia* chronic test. These results indicated that 43 percent of the tests showed toxicity. EPA is in the process of reviewing the paper and the raw data.

Persons interested in WET issues may consult another source of information developed by the SETAC Whole Effluent Toxicity Expert Advisory Panels. This group, established under a cooperative agreement with EPA, provides scientific opinion and training on WET technical issues. This information is available on the Internet at the SETAC web site, <http://www.setac.org>. Appendix D contains frequently asked questions with answers prepared by the SETAC WET Expert Advisory Panels. The expert panels have identified and discussed various factors that affect WET variability.

4.4 Conclusion

When the variability of WET analyses is viewed in the context of the NPDES program, these techniques produce data that are as precise as those from chemical analyses. As with any other analytical system, lack of experience in performing the analyses, adherence to prescribed QA practices, or good laboratory practices will reduce the precision of the results. Studies of these factors by independent researchers from both the regulatory and regulated communities support these conclusions. While examples of poor-quality, highly variable results from chemical analyses have also been publicized, these results are frequently influenced by the shortcomings mentioned above. Permittees that must generate and use WET data should become well-educated in data quality interpretation, and permittees should require that QC practices be followed by laboratories generating the data. Various sources of information presented in this chapter should assist permittees, testing laboratories, and regulatory authorities with this education process. Examples of practices that can further reduce the imprecision of analyses are also discussed in Chapters 5 and 6 of this document. Additional refinements of TAC can likewise improve test power to detect effects (or the lack thereof) and increase the statistical confidence in results.

5.0 GUIDANCE TO REGULATORY AUTHORITIES, LABORATORIES AND PERMITTEES: GENERATING AND EVALUATING EFFECT CONCENTRATIONS

5.1 Steps for Minimizing Test Method Variability

This chapter provides the background and recommendations on WET test procedures related to sampling, conducting the toxicity test methods, and conducting the statistical methods. Implementing these recommendations should decrease or minimize WET test method variability, thereby increasing confidence to make regulatory decisions (see Figure 5-1). EPA stands behind the technical soundness of the current WET test methods. The critical steps in minimizing WET test method variability are (1) obtaining a representative effluent sample, (2) conducting the toxicity tests properly to generate the biological endpoints, and (3) conducting the appropriate statistical analysis to obtain powerful and technically defensible effect concentrations. Minimizing variability at each step increases the reliability of the WET test results. For example, factors that affect variability include sampling procedures; sample representativeness; deviations from standardized test conditions (e.g., temperature, test duration, feeding); test organisms; source of dilution water; and analyst experience and technique in conducting the toxicity tests properly (Burton et al. 1996).

5.2 Collecting Representative Effluent Samples

The goal of effluent sampling is to obtain a representative sample that reflects real-world biological responses. Factors affecting the representativeness of effluent samples may include the sampling location, frequency, and type (e.g., composite or grab), and sample volume, container, preservation methods, and holding time. Burton et al. (1996) concluded that the above factors considerably influence test result variability.

Effluent samples must be collected at a location that represents the entire regulated flow or discharge. Typically, the sampling site is designated in the discharge permit. As with sampling for any parameter, effluent samples should be collected from a location where the flow is turbulent and well-mixed. Additionally, effluent samples should be collected at a frequency that enables adequate characterization of the discharge over time (e.g., accounts for daily to seasonal changes and variations in effluent quality). Major facilities should conduct WET testing monthly or quarterly, while minor facilities should conduct WET testing semi-annually or annually.

Appropriate sample types should be collected to represent the effluent fully. When the effluent is variable, collecting composite samples may be necessary. When the effluent is less variable, grab samples may be sufficient (e.g., from long-term retention pond facilities).

Sample containers should be non-reactive so that they do not affect sample characteristics. Table II of 40 CFR Part 136 requires that toxicity test samples be collected in glass or plastic containers, as specified in the methods. Sufficient sample volume should be collected for the type of test being conducted, including the number of test dilutions. When samples are collected in Cubitainers[®], headspace should be minimized.

Samples must be properly preserved. Part 136 of 40 CFR requires that samples for WET testing be cooled to 4°C when shipped off-site and between test sample renewals. Samples must be cooled during all phases of collection, transportation, and storage to minimize physicochemical changes. Samples must be tested within the specified maximum holding times before significant changes occur, such as volatilization or biological or chemical degradation. If samples are not tested within specified maximum holding times, the test is invalid and must be repeated by collecting a new effluent sample and conducting a new toxicity test to comply with the NPDES permit.

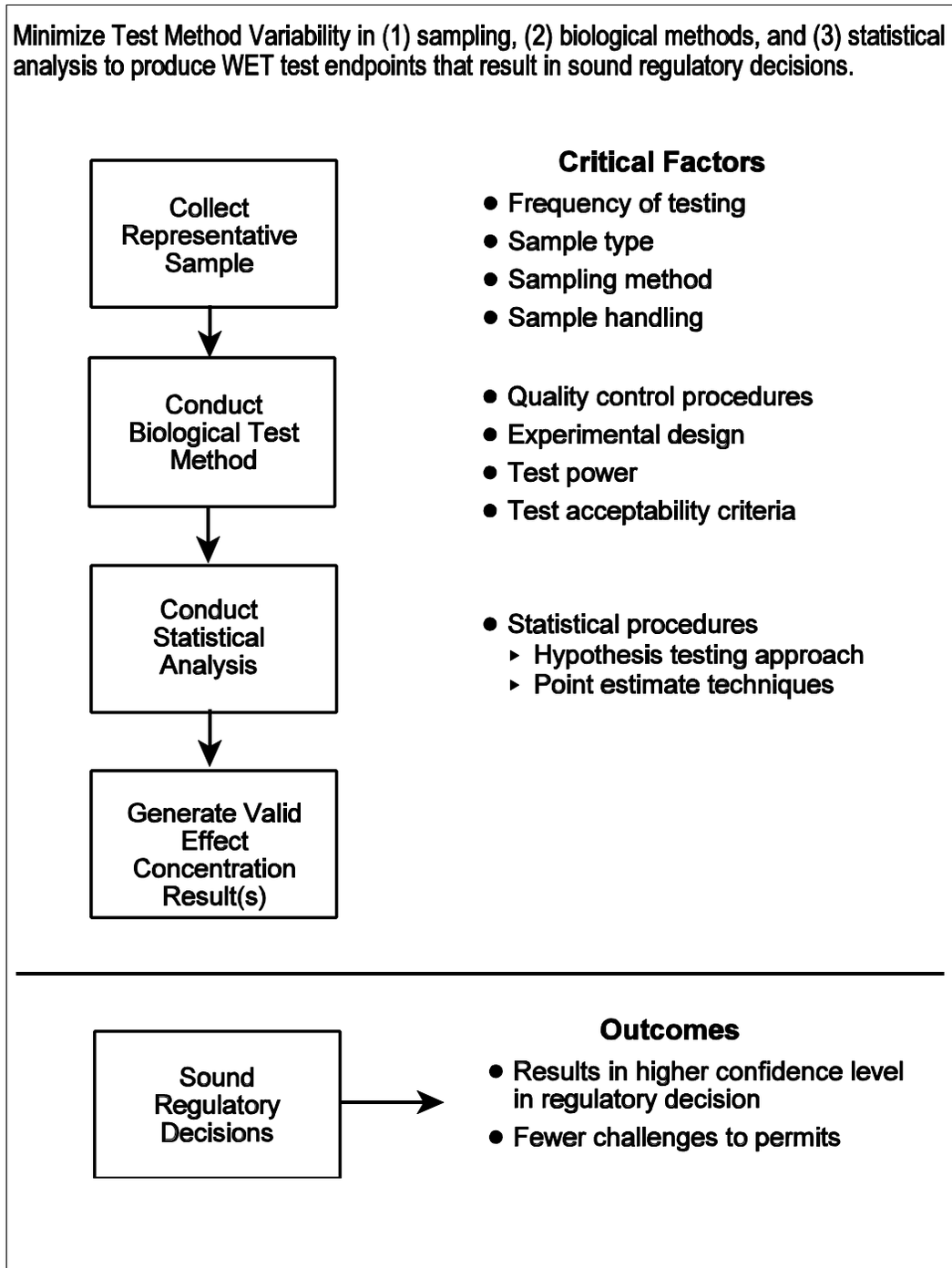


Figure 5-1. Steps to minimize WET test method variability.

5.3 Conducting the Biological Test Methods

Four main components of WET tests afford opportunities to control and minimize variability within tests and within and between laboratories: (1) quality control (QC) procedures; (2) experimental design; (3) test power; and (4) test acceptability criteria (TAC) beyond the minimum requirements specified in EPA’s WET test methods.

5.3.1 Quality Control Procedures

Quality assurance (QA) practices for toxicity tests address all aspects of the tests that affect data quality. These practices include effluent sampling and handling, test organism source and condition, equipment condition, test conditions, instrument calibration, replication, use of reference toxicants, recordkeeping, and data evaluation. The EPA WET toxicity testing manuals specify the minimum requirements for each aspect. Regulatory authorities have the discretion to prepare and implement additional guidance beyond the minimum requirements specified in EPA's WET test methods.

An integral part of the QA program is quality control (QC). The QC procedures are the more focused and routine activities conducted under the overall QA program. An important QC component in WET testing is the requirement to conduct reference toxicant tests with effluent tests. The WET test methods outline when reference toxicant tests are to be conducted. (See sections on quality of test organisms in the manuals.) Reference toxicant testing serves two purposes: (1) determine the sensitivity of the test organisms over time; and (2) assess the comparability of within- and between-laboratory test results. Reference toxicant test results can be used to identify potential sources of variability, such as test organism health, differences among batches of organisms, changes in laboratory water or food quality, and performance by laboratory technicians. In the QA section of each promulgated test method (USEPA 1993, 1994a, 1994b), EPA recommends sodium chloride, potassium chloride, cadmium chloride, copper sulfate, copper chloride, sodium dodecyl sulfate, and potassium dichromate as suitable reference toxicants. The methods do not, however, specify a particular reference toxicant or the specific test concentrations for each test method.

The current characterization of WET test method variability is limited by the ability to quantify sources of within- and between-laboratory variability, because laboratories can use different reference toxicants and test concentrations for a particular method. Future evaluations of method variability would be greatly enhanced by having data to analyze from multiple laboratories for the same reference toxicant, the same dilution water at similar pH and hardness, and the same test concentrations. By standardizing reference toxicants, testing laboratories could compare test results, permittees and regulatory authorities could better compare and evaluate laboratories, and the data could be used to further quantify within- and between-laboratory test precision. Specification of the reference toxicant and test concentrations for a method across laboratories would provide a much larger and consistent data base to assess the comparability of within- and between-laboratory test results.

Standardizing reference toxicants and test concentrations has been discussed in the literature. For example, the chronic methods manual for West Coast species (USEPA 1995) specifies the reference toxicant and test concentrations for each test species. The Southern California Toxicity Assessment Group (SCTAG) is comprised of representatives from permittees, testing laboratories, regulatory authorities, and academic institutions that met to discuss technical aspects of WET testing (e.g., standardization of reference toxicants, control charts). The SCTAG (1996) prepared a report to standardize reference toxicants for the chronic freshwater test methods. This report evaluated an extensive data base of reference toxicant data. The report recommended specific reference toxicants and test concentrations for these methods. The SCTAG (1997) also prepared a QA/QC checklist to help toxicity testing laboratories establish and maintain appropriate data quality measures. Regulatory authorities should review these publications when standardizing reference toxicants.

The selection of reference toxicants and test concentrations should be based on specific criteria. The following criteria, recommended in the SCTAG report, provide an excellent basis for selecting standardized reference toxicants:

1. The toxicant should provide precise and reliable measures of toxicological sensitivity.
2. Toxicant disposal should not be legally or environmentally problematic.

3. The toxicant should produce a concentration-response effect for the test organism.
4. The toxicant should be quantifiable.
5. The toxicant should not pose an unacceptable health hazard to laboratory personnel.
6. The toxicant should be readily available.

Most recently, Warren-Hicks et al. (1999) recommended that national acceptance criteria be specified with upper and lower acceptance limits for reference toxicant test results, which all laboratories would need to achieve to obtain accreditation. Variability could decrease nationally if testing laboratories are provided with more detail on the evaluation and interpretation of reference toxicant control charts (APHA-AWWA-WEF 1998). For example, such guidance could describe how to evaluate test results within the warning limits. Both Environment Canada (1990, 2000) and APHA-AWWA-WEF (1998) have prepared guidance on evaluating control chart data. The Environment Canada (2000) report specifies using zinc as an inorganic reference toxicant and phenol as an organic reference toxicant for many aquatic tests. The report also specifies eight criteria for selecting specific reference toxicants.

1. Previous use
2. Availability in a pure form
3. Solubility
4. Stability in solution
5. Stability during storage
6. Ease of analysis
7. Stable toxicity with normal changes in qualities of laboratory water
8. Ability to detect abnormal organisms

Regulatory authorities may want to evaluate the above reports and the SCTAG reference toxicant recommendations for the chronic freshwater test methods. Regulatory authorities may also want to evaluate and recommend a standard reference toxicant and a specific concentration series for each acute and chronic test method using data from this guidance document.

5.3.1.1 Guidance Related to Quality Control Charts and Laboratory Audits

Ausley (1996) recommends some oversight of data quality, such as evaluating tests in meeting QC criteria, using randomization procedures, and operating in allowed reference toxicant ranges to ensure that QC procedures are properly implemented. Another integral component of QC is the maintenance of control charts for reference toxicants and effluents. Laboratories should provide regular review of control charts. EPA suggests keeping a control chart for each combination of test material, test species, test conditions, and endpoints with a maximum of 20 test results. Modern software makes accumulating data and reviewing key test statistics possible with relatively little effort. Elementary methods can identify problems contributing to variability. Laboratories should practice regular control charting of test PMSDs and control performance for all tests along with control charting of effect concentrations such as NOEC and point estimates for reference toxicants tests. Successive tests should be compared occasionally to detect repeated patterns, such as one replicate's being consistently higher or aberrant, or a trend over time. Time sequence plots of concentration means and standard deviations would be useful in this regard. Occasionally, a set of 5 to 20 tests, in which block positions (see Appendix A in USEPA 1994b) have been recorded, should be subjected to ANOVA for block or position effects. If such effects are significant or large, the laboratory should seek advice on randomizing the replicates and concentrations.

If a laboratory's CV exceeds the 75th percentile CV from Tables 3-2 through 3-4, EPA recommends calculating warning and control limits based on the 75th and 90th percentiles, respectively, of CVs for the method and endpoint (Tables 3-2 and 3-3 and Appendix Tables B-1 and B-2). For example, suppose the mean EC25 for a series of *Ceriodaphnia* chronic tests (Method 1002.0 with reproduction as the endpoint) conducted at one laboratory with reference toxicant is 1.34 g/L NaCl. Also suppose that the standard deviation of the EC25s for these tests is 0.85. The CV for this set of EC25s is thus 0.63. In Table 3-2, the 75th percentile of CVs for this test's reproduction endpoint is 0.45. Calculate the standard deviation corresponding to the 75th percentile CV, $S_{A,75} = 1.34 \times 0.45 = 0.60$. In Appendix Table B-1, the 90th percentile of CVs is 0.62 for this method and endpoint. Calculate $S_{A,90} = 1.34 \times 0.62 = 0.83$. Because the CV for this series of EC25s exceeds the 90th percentile reported in Table B-1, EPA recommends the following:

- Set control limits using $S_{A,90} = 0.83$,
- Set warning limits using $S_{A,75} = 0.60$,
- Promptly take actions to bring results within the control limits, and
- Attempt to bring results within the warning limits in 3-12 months.

If the CV for the set of EC25s is less than the 90th percentile reported in Table B-1, use that CV to set control limits. If the CV for the set of EC25s is less than the 75th percentile in Table 3-2, do not set warning limits using the latter value.

In addition, Burton et al. (1996) encourage regulatory programs to have a laboratory audit component to document the existence and effectiveness of a QA/QC program directed at toxicity testing, including analyst training and experience. Regulatory authorities should use the National Environment Laboratory Accreditation Program (NELAP) (USEPA 1999a) and routine performance audit inspections to evaluate individual laboratory performance. Inspections should evaluate the laboratory's performance with QC control charts based on reference toxicants, examine procedures for conducting the toxicity test procedures, and examine procedures for analyzing test results.

Regulatory authorities should develop a QC checklist to assist in evaluating and interpreting toxicity test results. Appendix E presents examples of State WET implementation procedures related to reviewing reference toxicant data and information on additional QA/QC criteria that have been developed and implemented. Regulatory authorities should also provide additional guidance related to the interpretation of QC control charts. This additional guidance could be that laboratories maintain control charts on within-test variability (e.g., PMSD) and use warning and control limits based on the 75th and 90th percentiles of CVs for the test method and endpoint.

5.3.2 Experimental Design

Experimental design includes randomizing the experimental units (i.e., treatments, organisms, replicates); establishing the statistical significance level (i.e., alpha level); and specifying the minimum numbers of replicates, test organisms, and treatments. Oris and Bailer (1993) recommend that test design and TAC be based, not only on a minimum level of control performance, but also on the ability to detect a particular level of effect (i.e., test power).

A Type I error (i.e., "false positive") results in the false conclusion that an effluent is toxic when it is not toxic. A Type II error (i.e., "false negative") results in the false conclusion that an effluent is not toxic when it actually is toxic. Power (1 - beta) is the probability of correctly detecting a true toxic effect (i.e., declaring an effluent toxic when it is in fact toxic). Acceptable values for alpha range from 0.01 to 0.10 (1 to 10 percent). The current EPA test methods recommend an alpha rate of 0.05 or 5 percent in the toxicity

testing manuals. Currently, EPA is preparing guidance on when an alpha rate of 0.01 or 1 percent would be considered acceptable (USEPA 2000a).

5.3.2.1 False Positives in WET Testing

The hypothesis test procedures prescribed in EPA's WET methods provide adequate protection against incorrectly concluding that an effluent is toxic when it is not. The expected *maximum* rate of such errors is the alpha level used in the hypothesis test. The hypothesis test procedure is designed to provide an error rate *no greater than* alpha when the default assumptions are met. The statistical flow chart provided with each EPA WET method identifies cases when default assumptions are not satisfied and, therefore, when data transformations or alternative statistical methods (e.g., a nonparametric test) should be used.

Alpha and beta are related (i.e., as alpha increases, beta decreases), assuming that the sample size (number of treatments, number of replicates), size of difference to be detected, and variance are held constant. The alpha and beta error rates depend on satisfying the assumptions of the hypothesis test. To ensure that statistical assumptions and methods are properly applied, testing laboratories should review the statistical procedures used to produce WET test results and other factors, such as biological and statistical quality assurance, and verify that test conditions and test acceptability criteria were achieved.

If a test is properly conducted and correctly interpreted, identifying any particular outcome as a "false positive" is impossible. An effluent that is deemed toxic may require that the permittee conduct additional toxicity tests to determine if toxicity is re-occurring. Even if no toxicity is demonstrated in follow-up tests, that does not rule out that the original toxic event was a true toxic spike in the effluent. False negatives, however, impact the environment by allowing the discharge of harmful toxicants without identification. This may occur because the toxic effects are not identified as statistically significant due to lack of test sensitivity (see Sections 5.3.3 and 6.4).

Measurement error should not affect the protection against false positives provided by hypothesis tests and confidence intervals when they are appropriately applied. Measurement error, in the case of WET test treatment mean values, likely consists largely of sampling errors (e.g., variability among organisms or containers), although errors in counting, weighing, and other procedures may also occur. Such sources of imprecision are implicitly accounted for in WET test statistical inferences, because the sample variance among the replicates within each treatment (dilution) is used for inference. The test "size" $1 - \alpha$ will protect adequately against false positives. A larger variance among replicates, however, could make detecting real toxicity (i.e., false negatives) more difficult unless the number of replicates is increased to provide more test sensitivity and power, which will reduce the rate of false negatives.

5.3.2.2 False Negatives in WET Testing

For a given alpha, beta decreases (power increases) as the sample size increases and the variance decreases. Decreasing alpha from 0.05 to 0.01 without otherwise changing the hypothesis test will reduce the ability of the test to detect toxicity, that is, will reduce the power of the test. Thus, as alpha for the hypothesis test is decreased, there is an inevitable trade-off between the rate of false positives when toxicity is not present and the ability to detect toxicity when it is present (i.e., statistical power).

To limit within-test variability and thus increase power, EPA developed a minimum significant difference (MSD) criterion that must be achieved in the chronic West Coast marine test methods (USEPA 1995). The MSD is a measure of the within-test variability and represents differences between treatments and the control that can be detected statistically. Distributions of the MSD values of multiple tests for a specific reference toxicant and test method can be used to determine the level of test sensitivity achievable by a certain percentage of tests. Denton and Norberg-King (1996) analyzed several chronic test methods to quantify the effect size based on the existing toxicity test method experimental design and MSD distributions.

Denton and Norberg-King found when setting the beta error rate at 0.20 (power = 0.80), the effect size detected varies from at least a 15-percent reduction from the control response for the chronic red abalone larval development test to a 40-percent reduction from the control response for the chronic *Ceriodaphnia dubia* test. In this document, EPA has calculated power for each test method (see Section 5.3.3).

5.3.3 Test Power To Detect Toxic Effects

This section describes the statistical power and ability to detect toxic effects achieved by the current WET methods, as inferred from the WET variability data set used to develop this document. These inferences are approximate, because assumptions of normality and homogeneity of variance were not always satisfied.

Power can be characterized only by repeated testing. Power is an attribute not of a single test, but of a sequence of many tests conducted under similar conditions and with the same test design. Therefore, in this document, EPA used the sample averages for each laboratory's data set to characterize each laboratory. The following two parameters were required: (1) the mean endpoint response in the control (growth, reproduction, survival); and (2) the mean value of the error mean square (EMS) for tests.

EPA evaluated the ability to detect toxic effects using three approaches for each test method: (1) number of replicates required to detect a 25-percent difference from the control with power of 0.80; (2) percent difference from the control that can be detected with power of 0.80; and (3) power to detect a 25-percent difference from the control. All calculations are based on a one-sided, two-sample t-test at a level of 0.95 (alpha of 0.05). The power for a multiple comparison (Dunnett's or Steel's test) will be less than the power for this two-sample t-test.

Table 5-1 summarizes the results for this evaluation. Depending on the method, between 30 percent and 80 percent of the laboratories were able to detect a 25-percent effect for the sublethal endpoint consistently. Between 60 percent and 100 percent of the laboratories were able to detect a 33-percent effect.

To examine whether the upper bounds presented in Table 3-6 provide adequate test precision, EPA calculated an estimate of the power to detect a 25-percent effect on a sublethal endpoint when the PMSD equals the upper bound reported in Table 3-6. The upper bounds of the PMSD are shown in Table 3-6 in Chapter 3. At the lower PMSD bound, the power always exceeded 0.98. Tests with PMSD equaling the upper bound are not often able to detect a 25-percent effect. This finding does not mean that the upper bound is ineffective. The PMSD varies between tests, and each laboratory has a distribution of PMSDs. To avoid exceeding this upper bound often, a laboratory would have to achieve substantially lower PMSDs in most tests.

5.3.3.1 Attainment of the PMSD Related to Power

The power of the current experimental design could be reevaluated by comparing it to alternative designs that use increased number of replicates or number of test concentrations (Chapman et al. 1996). In this document, EPA found that about half of the laboratories in the data set were able routinely to detect a 25-percent difference between control and treatment. About two-thirds of the laboratories could routinely detect a 33-percent difference (Table 5-2). For example, mere attainment of the 90th percentile PMSD values shown in Table 3-6 will not ensure the ability to detect a 25-percent effect (Table 5-2). If every acceptable test has a PMSD below that upper bound, however, the average PMSD will be lowered. Based on the within-laboratory variability of PMSD,¹ the average PMSD likely will be substantially lower than the upper bound in Table 3-6, if *most* tests conducted by a laboratory are to have acceptable PMSDs.

¹ The average CV for PMSD is one-third to one-half its mean in commonly used methods.

Table 5-1. Tests for Chronic Toxicity: Power and Ability To Detect a Toxic Effect on the Sublethal Endpoint

Test Method	No. Labs with Power			Power (Range)	No. Labs Having Power at Least 0.8 To Detect Effect of		Effect Detected with Power 0.8 as Percent of Control Mean (Range)
	No. Labs	0.8	0.5		≤ 25%	≤ 33%	
1000.0 Fathead Minnow	19	6	14	0.21 - 1.00	6	13	8.2 - 62
1002.0 <i>Ceriodaphnia</i>	33	10	29	0.38 - 1.00	10	19	14 - 45
1003.0 Green Alga	9	7	8	0.33 - 0.99	7	8	13 - 49
1004.0 Sheepshead Minnow	5	4	5	0.77 - 1.00	4	5	8.6 - 26
1006.0 Inland Silverside	16	7	13	0.23 - 0.97	7	12	17 - 59
1007.0 Mysid (growth)	10	5	8	0.21 - 0.91	5	8	21 - 70

Note: Power was calculated for a two-sample, one-sided t-test at alpha = 0.05, for a 25-percent difference from the control. Effect size detected was calculated for the same test using power 0.80. Calculations used the average EMS from all tests at each laboratory and the minimum number of replicates reported for those tests. Calculations assumed that the parametric mean and variance equal the corresponding sample estimates. They also assumed approximate normality of means and homogeneity of variance. Because these assumptions may be violated, the results here are approximate. By saying “detect a 25-percent difference from control,” this alternative hypothesis is intended: (control mean - treatment mean) > 0.25 × control mean.

Table 5-2. Power To Detect a 25-Percent Difference from the Control at the 90th Percentile PMSD

Chronic Method	Replicates	90 th Percentiles of PMSD	Three Treatments		Four Treatments		Five Treatments	
			alpha = 0.05	alpha = 0.05/3	alpha = 0.05	alpha = 0.05/4	alpha = 0.05	alpha = 0.05/5
1000.0 Fathead Minnow	3	35	0.39	0.25	0.39	0.19	0.39	0.15
	4	35	0.41	0.30	0.42	0.26	0.43	0.23
1002.0 <i>Ceriodaphnia</i>	10	37	0.39	0.31	0.41	0.30	0.43	0.30
1003.0 Green Alga	3	35	0.39	0.25	0.39	0.19	0.39	0.15
	4	35	0.41	0.30	0.42	0.26	0.43	0.23
1004.0 Sheepshead Minnow	3	23	0.72	0.69	0.72	0.62	0.73	0.55
	4	23	0.73	0.71	0.74	0.68	0.75	0.66
1006.0 Inland Silverside	3	23	0.72	0.69	0.72	0.62	0.73	0.55
	4	23	0.73	0.71	0.74	0.68	0.75	0.66
1007.0 Mysid	8	32	0.48	0.41	0.50	0.40	0.52	0.40

Notes: Values are rounded to two significant figures. Number of treatments is the number of concentrations compared with the control in the hypothesis test. The calculations assumed (1) the usual assumptions of the test are satisfied (approximate normality, homogeneity of variances); and (2) equal replication in treatments and control. Because these assumptions may be violated, the results here are approximate. Because the MSD/mean implies a value for $\sqrt{(\text{error mean square})/\text{mean}}$, the latter could be calculated from the MSD, Dunnett’s critical value, and the number of replicates, and then used in a calculation of power. Calculations apply to a one-sided, two-sample t-test of equal means, assuming equal variances and equal replication, with hypotheses $H_0: \{\text{control mean} - \text{treatment mean} = 0\}$ versus $H_a: \{\text{control mean} - \text{treatment mean} > 0.25 \times \text{control mean}\}$. The power achieved by Dunnett’s multiple comparison procedure will lie between the two-sample power at alpha = 0.05 and that for alpha = 0.05/(no. of treatments).

Testing laboratories and permittees can examine the EMS or MSD in Tables B-14 and B-15 (Appendix B) to estimate the ability of a WET test to detect toxic effects. Some regulatory authorities may require a comparison between the control and the receiving water concentration, which requires a two-sample, one-sided t-test. Others may require the multiple comparisons procedure described in the EPA WET methods (Dunnett's or Steel's tests, one-sided, with alpha of 0.05). The power of Dunnett's procedure falls between the power of the one-sided, two-sample t-test with alpha of 0.05 and alpha of 0.01, assuming that no more than five toxicant concentrations are compared to a control. The power of Steel's procedure will be related to, and should usually increase with, the power of Dunnett's procedure and the t-tests. Tables B-14 and B-15 in Appendix B also provide an appropriate guide to achieving power using a nonparametric test.

Recently, the State of Washington (1997) issued guidance specifying an acute and chronic statistical power standard to be achieved for compliance testing. EPA's sediment toxicity testing manuals (USEPA 1994c, USEPA 2000) include power curves for various numbers of experimental units, CV ranges, and associated alpha and beta levels. Sheppard (1999) is a good source to provide a simple explanation of how power helps determine how large a sample should be. Additional information on power may be obtained at: <http://www.psychologie.uni-trier.de:8000/projects/gpower/literature.html>.

EPA recommends that regulatory authorities specify in their State WET implementation procedures that individual test results achieve a level of within-test sensitivity by using the upper and lower PMSD test sensitivity bounds (see Section 6.4). To achieve the test sensitivity bounds, testing laboratories may need to minimize within-test variability (e.g., EMS) or increase the number of replicates tested, or both. If laboratories cannot achieve PMSD values of less than 25 percent for the toxicity test methods that require a minimum of only three replicates (Methods 1000.0, 1004.0, 1006.0), then the numbers of replicates may need to be increased. Appendix B (Section B.4) provides information related to the number of replicates needed and discusses the relationship between test power and effect size achieved. The magnitude of the effect size achieved relates to the test sensitivity.

5.4 Test Acceptability Criteria

EPA test methods have specific TAC that the effluent and reference toxicant tests must meet. A test is considered invalid if the TACs are not met. The recommended test conditions for each test method specify the minimum requirements and the TAC. For example, control survival must be 80 percent or greater and average control reproduction at least 15 young per surviving female in the chronic *Ceriodaphnia dubia* survival and reproduction test.

The chronic West Coast marine methods (USEPA 1995) require additional TAC. For example, to limit the degree of within-test variability, the methods specify a maximum allowable value for PMSD (see Section 5.3.2 on experimental design). Some States have additional TAC in their State WET implementation policies. North Carolina (1998) for example, requires that the chronic *Ceriodaphnia dubia* analyses meet an additional TAC of complete third brood neonate production by at least 80 percent of the control organisms and that the control reproduction CV be less than 40 percent.

Additional TAC might be specified to minimize variability among replicates. Variability of any toxicity test result is influenced by the number of replicates used, number of organisms tested, and variability among replicates at each test concentration and the control. Variability among replicates has been quantified by treatment CV, EMS, or MSD. The application of a maximum acceptable value for CV or MSD helps ensure adequate laboratory QA/QC and increases the reliability of submitted data. One benefit of requiring a maximum allowable within-test variability limit is that laboratories will improve culturing, test handling, and housekeeping, which are usually incorporated into the laboratories' standard operating procedures. For example, the CV requirement might be incorporated directly into the NPDES permit. Sample EPA Region 6 permit language reads:

1. *The coefficient of variation between replicates shall be less than or equal to 40 percent in the control.*
2. *The coefficient of variation between replicates shall be less than or equal to 40 percent at the instream waste concentration (IWC).*
3. *Test failure may not be construed or reported as invalid due to a CV of greater than 40 percent. A repeat test shall be conducted within the required reporting period if any test is determined to be invalid.*

Occasionally, statistical analyses indicate a test failure when as little as 15-percent mortality has occurred in a test dilution. Permit language has been developed to address this occurrence, as in the following example:

If all TAC conditions are met and the percent survival of the test organism is greater than or equal to 80 percent (in a chronic test) or 90 percent (in an acute test) in the critical dilution concentration and all lower dilution concentrations, the test shall be considered to be a valid test, and the PERMITTEES shall report an NOEC of not less than the critical dilution for the discharge monitoring report (DMR) reporting requirements.

Regulatory authorities may consider providing guidance or imposing additional TAC, such as those implemented by EPA Region 6 or like some States have implemented (North Carolina 1998, Washington 1997). Appendix E provides additional examples of States that have implemented further guidance on WET QA/QC procedures and TAC. Warren-Hicks (1999) also recommended that additional national TAC be established for each test method (e.g., upper and lower bounds on the MSD). Therefore, EPA recommends that regulatory authorities require that additional TACs be implemented in permits to minimize within-test variability and increase test sensitivity (see Section 6.4 and Appendix C for sample permit language).

5.5 Conducting the Statistical Analysis To Determine the Effect Concentration

EPA test methods currently recommend two statistical approaches to estimate a chemical or effluent concentration for each biological effect endpoint (e.g., survival, growth, and reproduction). One approach is to derive the NOEC by hypothesis testing, which equates biological significance with statistical significance. The second approach is to estimate an effect concentration that reduces the control response by 25 percent for chronic methods. The expanded use of WET tests in the NPDES program has brought increased attention to the statistical analysis of toxicity test data. A common goal for both regulatory authorities and permittees is to confirm that the effect concentrations were derived correctly using the appropriate analysis approaches. Reliable effect concentrations lead to increased confidence in the data used for making regulatory decisions, such as determining reasonable potential, deriving a permit limit or monitoring trigger, and generating self-monitoring test results.

Another important consideration in conducting statistical analyses is the inconsistent use of statistical programs. The proliferation of statistical packages has been helpful in data analysis; however, these packages also can result in the misapplication of the statistical methods. APCA-AWWA-WEF (1998) cautions the user to confirm the results of each analysis with each package before accepting them. The data user is responsible for evaluating all data submitted to the regulatory authorities.

The 1995 SETAC Pellston Workshop discussed unresolved scientific issues and highlighted significant research needs associated with WET testing. The attendees recommended the following:

Immediately instigate studies to evaluate improvements in the statistical analysis of WET test data. These studies should include, but not necessarily be limited to, the following activities:

(a) investigate the implications of concurrent application of NOEC/MSD, tests of bioequivalence, and EC_p estimators (Chapman et al. 1996a).

In response to this recommendation, EPA began projects to evaluate the bioequivalence approach and additional point estimate models for the WET program. At present, two test methods are being used for this evaluation: (1) the chronic *Ceriodaphnia dubia* survival and reproduction tests and (2) the giant kelp germination and germ-tube length test with reference toxicants.

The bioequivalence approach poses the following question: Do the mean responses of the effluent concentration and the control differ by more than some amount? For example, the control response and the response at the critical effluent concentration (i.e., instream waste concentration) must differ by no more than a fixed value in order to accept the hypothesis of no significant difference (i.e., no toxicity). This approach could address the concern that an imprecise test might not detect toxicity when toxicity is present or that a small but statistically significant effect would detect toxicity that may not be biologically important. Some researchers have suggested that the bioequivalence approach could provide a positive incentive for dischargers to produce test results with lower within-test variability to demonstrate that no toxicity occurs at a level greater than a biologically (bioequivalence approach) significant amount (Shukla et al. 2000, Wang et al. 2000).

Bailer et al. (2000) evaluated the proposed regression-based estimators with the current EPA point estimate models. They found that it appears reasonable to incorporate parametric estimation models in the WET program. Bailer et al. (2000) concluded that these models are appropriate for all response scales (i.e., dichotomous, count, and continuous) and can incorporate monotonicity without bias. However, confidence intervals still need to be developed for these parametric models.

In this document, EPA has not recommended either the bioequivalence or additional point estimate models to supplement the current statistical approaches as described in the testing manuals. An independent, peer-reviewed workshop should be convened to evaluate the benefits of these alternative statistical approaches to enhance the statistical approaches currently applied.

5.6 Chapter Conclusions

In this chapter, EPA provides guidance to permittees and testing laboratories on collecting representative effluent samples, conducting the biological test methods, and evaluating the statistical analyses. EPA recommends that States implement the lower and upper PMSD test sensitivity bounds to achieve an acceptable level of test sensitivity and minimize within-test variability (see Section 6.4). EPA also provides guidance to permittees and testing laboratories on the number of replicates required to achieve the PMSD bounds. Testing laboratories should maintain and evaluate both effluent and reference toxicant data using a measure of within-test variability such as the PMSD.

Permittees and toxicity testing laboratories may need to increase replication in order to reduce PMSD below the upper bound. Table B-15 can be used for initial planning of replication, given knowledge of typical values of the error mean square (EMS) or MSD and the number of concentrations used in the multiple comparison hypothesis test. To ensure that all PMSD values fall below the upper bound in Table 3-6, a laboratory would select the largest EMS value experienced in its past testing.

EPA recommends that testing laboratories require a minimum of four replicates for the fathead minnow, sheepshead minnow, and inland silverside chronic test methods (Methods 1000.0, 1004.0, and 1006.0, respectively). Four replicates are needed to execute the statistical flow chart when a nonparametric test is needed. Three replicates are also sometimes insufficient to keep PMSD below the recommended upper bound. In addition, four replicates are needed to help achieve the upper PMSD bound.

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6.0 GUIDANCE TO REGULATORY AUTHORITIES: DETERMINING REASONABLE POTENTIAL AND DERIVING WET PERMIT CONDITIONS

EPA developed the TSD (USEPA 1991a) to support implementation of national policy to control the discharge of toxic pollutants. The TSD presents a statistical approach for determining the need for and the method of deriving water quality-based effluent limits (WQBELs) based on aquatic life (including WET), human health, and wildlife criteria. This approach accounts for the uncertainty associated with small data sets and data variability by assuming a statistical distribution of effluent data (usually lognormal) and calculating a CV or using a default CV to describe data variability.

6.1 Analytical and Sampling Variability in Calculations for Reasonable Potential and Permit Limits

Section 6.1 discusses use of the CV of sample measurements of toxicity to make a reasonable potential determination and to calculate permit limits. Two points must be understood: (1) this CV is to be calculated using toxic unit (TU) values (USEPA 1991a) (see Section 6.2); and (2) EPA strongly recommends that point estimates (not NOEC or LOEC values) be used to calculate the TU values (USEPA 1994a, 1994b).

Water quality-based effluent limits are required when a discharge causes, has reasonable potential to cause, or contributes to an instream excursion above a water quality standard. Throughout this document, EPA uses the commonly understood, shorthand reference “reasonable potential” to refer to this standard for determining the need for a water quality-based effluent limit.

6.1.1 “Adjusting for Analytical Variability” in Calculations for Reasonable Potential and Permit Limits

Adjustment approaches (see Appendix G.3) have been suggested to “adjust for analytical variability” when deriving permit limits and determining the need for a WET limit in the first place. EPA does not recommend these adjustment approaches (Appendix G.3) and strongly reaffirms the statistical approach and methods for calculating permit limits provided in the TSD (USEPA 1991a). *EPA recommends that regulatory authorities use the statistical approach and calculation methods in the TSD.* The TSD methods were designed to provide a reasonable degree of protection for water quality (i.e., to avoid exceedances of water quality criteria), while providing a reasonable degree of protection from the variability of effluent toxicity and analytical variability. The various “adjustment” approaches would undermine these objectives.

The TSD limit calculation for a point source can be divided into two steps: first, convert the wasteload allocation (WLA) to a long-term average (LTA), and then convert the LTA to effluent limits (maximum daily, average weekly, and average monthly limits). WET limit calculations include an intermediate step in which the acute WLA is converted to a WLA_{a,c}. These calculations employ a facility-specific CV based upon effluent sampling data. The TSD approach uses this CV in both steps.

Adjustment approaches intended to account for analytical variability, discussed in detail in Appendix G, would inappropriately use different CVs in these two steps. The first step would use an estimate of the CV of “true” effluent toxicity, which is smaller than the CV for measured toxicities. This approach would result in a larger calculated LTA. The second step would use the CV for the measured toxicities, which is the same CV used in both steps of the TSD approach.

Use of such adjustment approaches would frequently result in setting an average monthly permit limit (AML) that exceeds the chronic WLA. Appendix G demonstrates that such outcomes (i.e., the AML exceeds

the chronic WLA) generally can be expected to occur when various adjustment approaches are used. Appendix G, Table G-1, presents a numerical example of how an adjustment approach would allow calculation of an AML exceeding the chronic WLA (a four-day average value), even when sampling frequency for the calculation is set at the recommended minimum of four samples per month. [It is acceptable for the maximum daily limit (MDL), which applies to a single sample, to exceed the chronic WLA. It is also acceptable for the AML to exceed the chronic WLA, if the AML calculation is based on fewer than four samples per month. Note, however, that the TSD recommends always assuming at least four samples per month when calculating the AML.]

The TSD reasonable potential calculation first calculates the percentile represented by the maximum observed TU value. For example, the maximum of 10 reported TU values is identified with the 63rd percentile. Then the sample CV is used to project the 95th or 99th percentile TU value, using a table of reasonable potential multiplying factors. This value is combined with the appropriate mixing-zone dilution to project a maximum receiving water toxicity, which is compared with the applicable water-quality criterion. If an adjustment were applied to the reasonable potential calculation, the CV would be adjusted downward and the maximum projected receiving water toxicity would be smaller. This would make a determination of need for a permit limit less likely.

Because of these considerations, EPA strongly recommends that no adjustment be made to the CV or variance of toxicity, either for reasonable potential or permit limit calculations. The TSD statistical approaches already account for analytical variability appropriately. EPA continues to recommend the TSD approach, which ensures that effluent limits and, thereby, *measured* effluent toxicity or pollutant parameter concentrations are consistent with calculated WLAs.

6.1.2 Analytical Variability and Self-monitoring Data

EPA determines compliance with permit limits on the basis of self-monitoring data, and these data include some measure of analytical variability. The influence of analytical variability is accounted for in the TSD statistical procedures used to set water-quality limits and determine the potential for toxicity, as explained in Appendix G.

The permittee is responsible for ensuring that measured discharge toxicity never exceeds the permit limits. No special allowance is made for analytical variability in assessing compliance. The maximum discharge toxicity should incorporate a margin of safety, which will account for sampling and analytical variability. In other words, to avoid exceeding permit limits, the facility's treatment system should be designed so that the maximum toxicity is somewhat lower than its permit limits.

6.1.3 Precision of WET Measurements and Estimates of Effluent CV

Single measurements on effluent involve some uncertainties about the true concentration or toxicity related to representativeness of the sample, including sample holding time and conditions, and the analytical measurement system. Like all analytical measurements, WET measurements (NOEC, EC25, LC50) are inexact. That is, the exact toxicity of an analyte in a sample can be specified only within some range. This imprecision can be reduced by using a suitable number of organisms and replicates for each test (see Section 5.3.2 on experimental design).

The numbers of organisms and replicates required for EPA WET method test acceptability are specified as minimums. Test precision will be approximately proportional to the square root of the number of replicates. Thus, doubling the number of replicates may decrease the MSD to approximately 70 percent of its former value. Increased replication also tightens the confidence interval for a point estimate of the effect concentration (e.g., EC25 and LC50).

EPA strongly recommends that toxicity measurements of an effluent be obtained at least quarterly for three years to provide a good basis for determining the need for limits and for calculating limits. One year should be regarded as the minimum duration needed to characterize effluent variability (due to seasonal, stream flow, or process fluctuations), and ten the minimum number of measurements, unless scientific and technical knowledge supports a shorter period as representative of the distribution of pollutant types and concentrations of toxicity.

Estimates based on multiple measurements involve the same uncertainties that apply to single measurements. They also may involve larger uncertainties related to sampling error, that is, the chance that typical levels of toxicity or concentrations of pollutant may not be encountered during the sampling program. The sampling program may not fully characterize effluent variability if too few samples are taken, the sampling times and dates are not representative, or the duration of the sampling program is not long enough to represent the full range of effluent variability. When determining the need for limits and calculating limits, the variance or the CV of toxicity measurements is key. The larger the number of samples, the more precise is the estimate. Confidence intervals for the variance and CV can be calculated and carried through the calculations for reasonable potential and effluent limits (Appendix G). Even when assumptions are not strictly met, confidence intervals provide a useful perspective on the uncertainty of the results and the need for more samples. The *minimum* number of measurements recommended for calculating estimates of the CV for effluent toxicity is 10.

6.1.4 Between-Laboratory Variability

Between-laboratory variability may increase the CV as discussed in Section 6.1.1, if the toxicity tests were conducted by more than one laboratory for a specific facility. A concern to permittees is that this may increase the likelihood of making a finding of reasonable potential.

Within-laboratory variability is the component of analytical variability that should be reflected in regulatory calculations. If the data used for reasonable potential or permit limit calculations are effluent measurement data reported by at least two laboratories, there are ways to appropriately estimate the variance to be used in TSD statistical calculations.

For example:

- If the same laboratories continue to be used in the same proportion or frequency and the measurements from the individual laboratories represent different sampling dates, the measurement data can be treated as if they were generated by a single laboratory. This approach may increase the estimated variance and the AML, which is not in the interest of the permittee. Selecting one laboratory for future monitoring, based on the variance of its reported reference toxicant test results, should mitigate this problem.
- If only one laboratory has reported data on each sampling date, and the other laboratories report over different time spans or over the same time span on alternating dates, EPA recommends forming a pooled estimate of variance. Calculate the sample variance (S^2) of log(TU) for each laboratory separately, and combine these using the formula:

$$\text{pooled variance of log}(X) = [(N_1 - 1)S_1^2 + (N_2 - 1)S_2^2] / [(N_1 - 1) + (N_2 - 1)]$$

An analogous formula is used for more than two laboratories. The same result can be obtained by conducting a one-way analysis of variance on log(TU) (with laboratories treated as the groups or classes) and using the reported EMS.

Changing a laboratory may change analytical (within-laboratory) variability of measurements and test sensitivity (i.e., PMSD values). That is, the average effect concentration may change (e.g., Warren-Hicks et al. 1999). Ideally, the permittee will anticipate and plan for a change of testing laboratory. Permittees should compare reference toxicant test data for current and candidate replacement laboratories, selecting one with acceptable variability and a similar average effect concentration.

6.2 Determining Reasonable Potential and Establishing Effluent Limits

Effluent characterization is an essential step in determining the need for an NPDES permit limit. NPDES regulations under 40 CFR Part 122.44(d)(1)(ii) specify that reasonable potential include “*whether a discharge causes, has the reasonable potential to cause, or contributes to an instream excursion above a State water quality standard.*” Calculations for reasonable potential determination and for permit limits should follow EPA guidance in the TSD (USEPA 1991a). In particular, the TSD statistical methods should be used. Such calculations should use TUs for WET data, not effect concentrations (percent whole effluent). Toxic units are defined (USEPA 1991a, Chapter 1.3.1, page 6) as the reciprocal of the effect concentration times 100, where the effect concentration is expressed as a percentage of whole effluent, thus $TU_a = 100/LC50$ and $TU_c = 100/EC_p$.

When characterizing an effluent to determine whether a permit limit is necessary, permit writers can use the available effluent WET data and a water-quality model to perform a reasonable potential analysis. The TSD outlines the statistical approach. This approach uses existing effluent data to project a maximum pollutant concentration or a maximum toxicity in the effluent (USEPA 1991a). The projected maximum concentration or toxicity is used as an input in the water quality model to determine whether the effluent has the reasonable potential to cause or contribute to an excursion of ambient water quality criteria. If reasonable potential exists, the permit writer must derive a WET permit limit for that facility.¹

The variability of the existing effluent data, as measured by the CV, has a significant effect on the projected maximum pollutant concentration or toxicity. The higher the CV, the higher the projected maximum, and the more likely that there is reasonable potential and a limit is needed. EPA recommends that regulatory authorities use all valid, relevant, and representative data in making reasonable potential determinations. EPA is developing a national policy clarifying use of the TSD procedures for determining reasonable potential for WET. Important components of this policy include specifying the minimum number of valid WET tests necessary to calculate facility-specific CVs,² as well as recommending a step-wise approach to determining the need for WET permit limits. This approach reflects a strong preference by EPA and its stakeholders to rely on facility-specific WET testing, based on adequate frequency and duration of effluent sampling, for making reasonable potential determinations for toxicity.

EPA recommends that point estimates be used to estimate effluent variability, to determine the need for limits, and to set permit limits. This is recommended whether the self-monitoring test results will be determined using hypothesis tests or point estimates. Point estimates have less analytical variability than NOECs using current experimental designs, as shown in Chapter 3. Point estimates make the best use of the WET test data for purposes of estimating the CV, LTA, and RP factor and calculating the permit limit.

¹ When the State has narrative criteria for toxicity and the TIE/TRE identifies a specific chemical that is the source of toxicity, the permit writer may include a chemical-specific limit for that parameter instead of a WET permit limit in accordance with 40 CFR Part 122.44(d)(v).

² If fewer than ten data points are available, the regulatory authority must use a default CV. As a result, the need for a WET permit limit may be based on a default value rather than actual data.

6.3 Development of a Total Maximum Daily Load for WET

Total maximum daily loads (TMDLs) may be indicated when there is acute or chronic toxicity in a waterbody, leading to the listing of the waterbody as impaired under CWA Section 303(d), and when there are multiple sources of the toxicity. EPA believes that TMDL calculations should be performed on the pollutants causing toxicity whenever possible. In these situations, EPA suggests that the first step of the analysis is to conduct ambient toxicity identification evaluations to identify the pollutant(s) and the source(s) causing the toxicity. Once the pollutant(s) and source(s) causing toxicity have been identified for the waterbody, then a TMDL should be developed for the individual pollutant(s).

6.4 Accounting for and Minimizing Variability In the Regulatory Decision Process

A common goal for the permittee and the regulatory authority is to have confidence in the test results from the biological and statistical procedures. Both permittees and regulatory authorities would then have more confidence in taking regulatory actions, such as evaluating multiple effluent samples to determine reasonable potential and derive permit conditions (e.g., permit limits, monitoring triggers). If steps such as collecting a representative effluent sample to conducting the toxicity tests properly, as discussed in Sections 5.2 through 5.4, and requiring additional TACs (Section 6.4.1) are used to reduce or minimize within-test variability, then the reliability of the WET test results increases.

6.4.1 Recommended Additional TACs: Lower and Upper Bounds for PMSD

Reference toxicant data from a large number of tests and laboratories were used to generate PMSD values; percentiles of these values are reported in Table 3-6. The MSD represents the smallest difference between the control mean and a treatment mean that leads to the statistical rejection of the null hypothesis (i.e., no toxicity) using Dunnett's multiple comparison test. MSD values are divided by the control mean and multiplied by 100 to produce a "percent MSD" (PMSD) value. The PMSD allows comparison of different tests and represents the smallest significant difference from the control as a percentage of the control mean. Thus, it represents the smallest significant value of the relative difference [100 (control mean - treatment mean)/control mean]. The MSD is often expressed as a percentage of the biological endpoint in the control response.

The following formula is used to calculate MSD (as recommended by USEPA 1995):

$$\text{MSD} = d s_w \sqrt{(1/n_1) + (1/n)}$$

where

- d = critical value for the Dunnett's procedure
- s_w = the square root of the error mean square (EMS)
- n₁ = number of experimental units in the control treatment
- n = the number of experimental units per treatment, assuming an equal number at all other treatments

Percent MSD is calculated as follows:

$$\text{PMSD} = \frac{\text{MSD}}{\text{control mean}} \times 100$$

EPA recommends that regulatory authorities implement both the lower and upper PMSD bound approach to minimize within-test variability when using hypothesis testing approaches to report an NOEC. The implementation of the upper PMSD bound should also apply when using point estimate techniques. There are five possible outcomes for regulatory decisions (see Figure 6-1). Two outcomes imply unqualified acceptance of the WET test statistical result:

1. **Unqualified Pass**—The test’s PMSD is within bounds and there is no significant difference between the means for the control and the instream waste concentration (IWC) treatment. The regulatory authority would conclude that there *is no toxicity at the IWC concentration*.
2. **Unqualified Fail**—The test’s PMSD is larger than the lower bound (but not greater than the upper bound) in Table 3-6 and there is a significant difference between the means for the control and the IWC treatment. The regulatory authority would conclude that there *is toxicity at the IWC concentration*.
3. **Lacks Test Sensitivity**—The test’s PMSD exceeds the upper bound in Table 3-6 and there is no significant difference between the means for the control and the IWC treatment. The test is considered invalid. A new effluent sample must be collected and another toxicity test must be conducted.
4. **Lacks Test Sensitivity**—The test’s PMSD exceeds the upper bound in Table 3-6 and there is a significant difference between the means for the control and the IWC treatment. The test is considered valid. The regulatory authority would conclude that there *is toxicity at the IWC concentration*.
5. **Very Small but Significant Difference**—The relative difference (see Section 6.4.2, below) between the means for the control and the IWC treatment is smaller than the lower bound in Table 3-6 and this difference is statistically significant. The test is acceptable. The NOEC is determined as described in Section 6.4.2 below.

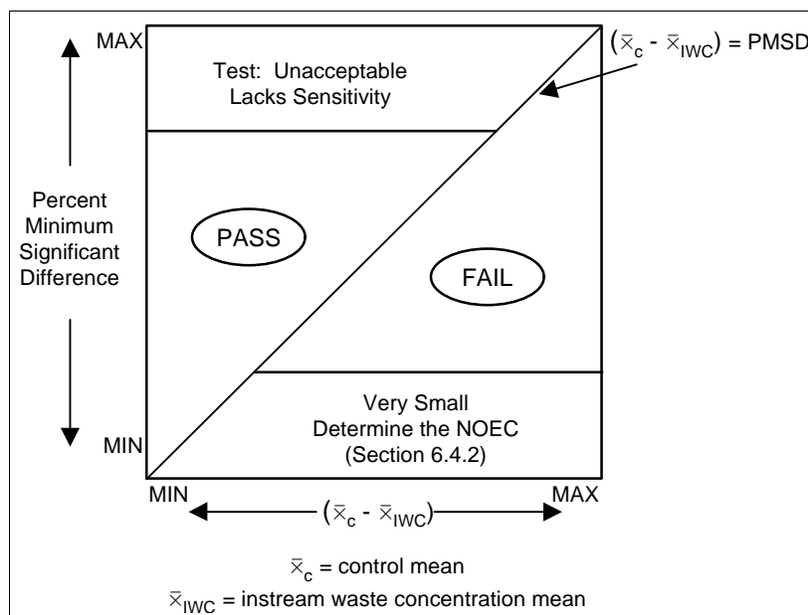


Figure 6-1. Paradigm that incorporates the lower and upper percent minimum significant difference.

Regulatory authorities should examine the sample permit language as provided in Appendix C, for incorporation of the PMSD bound language in a NPDES permit.

Note that “unqualified acceptance” of a WET test result requires that all of the following must be achieved: (1) collect the effluent sample properly; (2) conduct the toxicity test methods as specified in the toxicity manuals; (3) meet the required TACs; (4) meet the proper water quality parameters (e.g., temperature, pH); and (5) conduct the proper statistical calculations. All these conditions must be reviewed and deemed acceptable before a test is evaluated for self-monitoring data and reporting.

Figure 6-2 provides a decision tree that regulatory authorities can use when implementing the lower and upper PMSD bounds.

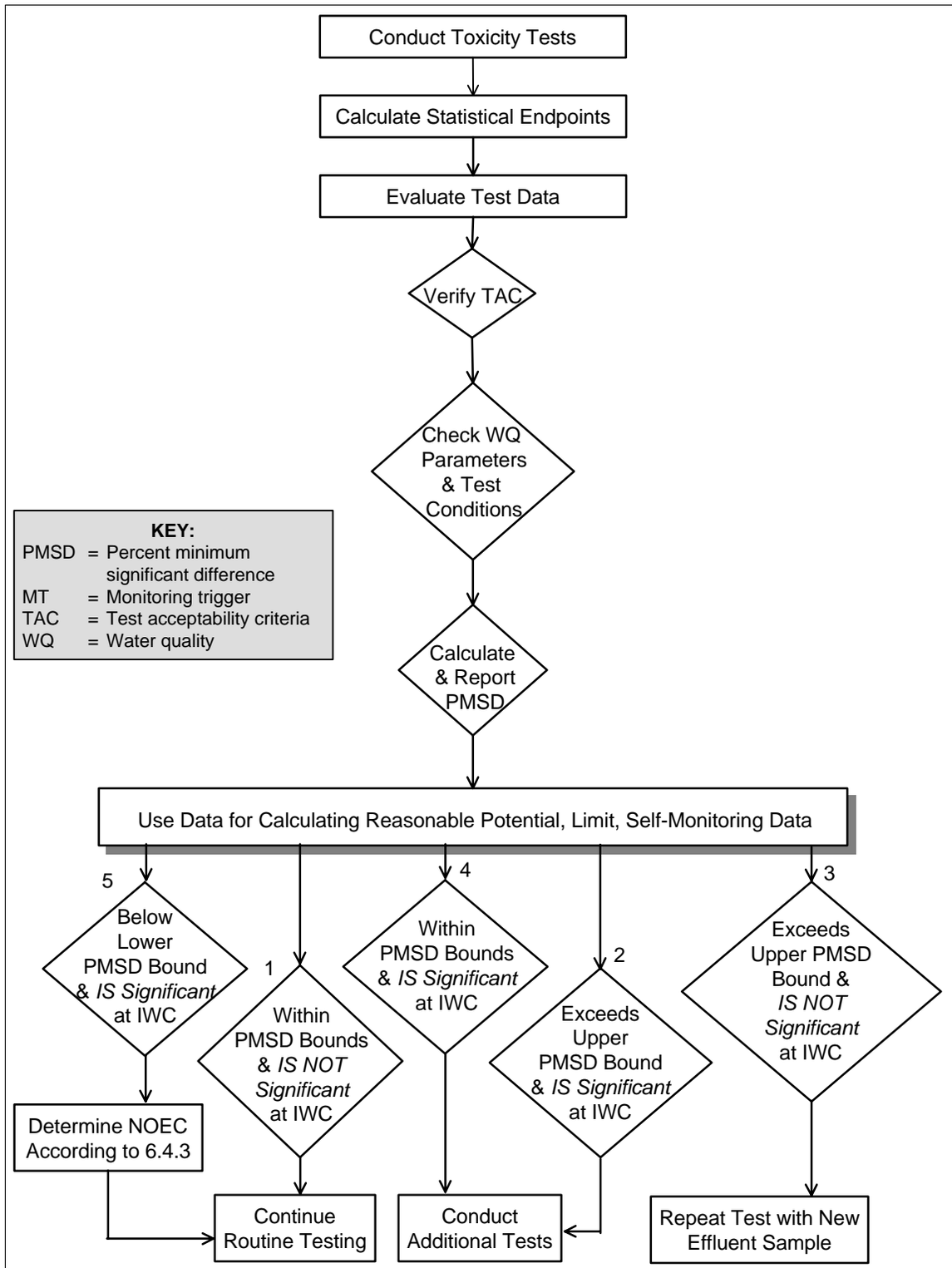


Figure 6-2. Implementing applications of upper and lower PMSD bounds for effluent toxicity testing requirements.

6.4.2 How to Determine the NOEC Using the Lower PMSD Bound

If the permit specifies that self-monitoring data are to be generated using hypothesis testing approaches, then the analyst should report the NOEC as the following. Find the smallest concentration for which (a) the treatment mean differs significantly from the control mean and (b) the relative difference (see example below) is not smaller than the 10th percentile in Table 3-6. Therefore, the NOEC is the next smaller test concentration.

In other words, concentrations having a very small relative difference with control (smaller than the lower PMSD bound) would be treated as if they do not differ significantly from control (even if they do so), for the purpose of determining the NOEC.

Table 6-1 illustrates the application of the lower PMSD bound for the reproduction endpoint of a *Ceriodaphnia* chronic test. In this example, the test's PMSD was 9.9, smaller than the 10th percentile value of 11 found in Table 3-6. The IWC concentration differed significantly from the control. The test falls under outcome number 5, a significant but very small difference at the IWC. The first step is to calculate the relative differences from control (Table 6-1) as [(control mean - treatment mean) divided by (control mean)] × 100. The next step is to determine which relative differences exceed the PMSD lower bound, 11 in this case (see the last column of Table 6-1). Finally, the NOEC is determined as described above. The NOEC is 12.5 percent effluent for this example.

Table 6-1. Example of Applying the Lower PMSD Bound for the Chronic *Ceriodaphnia* Test with the Reproduction Endpoint

Concentration (percent effluent)	Reproduction (mean of ten replicates)	Relative Difference from Control	Does Relative Difference Exceed 11?
100%	5.08 *	82	Yes
50%	12.4 *	56	Yes
25%	23.4 *	17	Yes
IWC = 12.5%	25.3 *	10	No
6.25%	26.1	7.4	No
Control	28.2	0	No

NOTE: The lower PMSD bound for this method and endpoint is 11 (Table 3-6). In this example, the NOEC is 6.25 percent effluent using the test's (very small) PMSD. Therefore, the reported NOEC should be 12.5 percent effluent after applying the lower PMSD bound.

* Differs statistically from the control as determined by MSD = 2.8 neonates. Thus, treatment means that are less than 28.2 - 2.8 = 25.4 would be statistically significant. These correspond to relative differences greater than 100 (2.8 / 28.2) = 9.9 percent.

6.4.3 Justification for Implementing the Test Sensitivity Bounds

A lower bound is needed to avoid penalizing laboratories that achieve unusually high precision. The 10th percentile PMSD represents a practical limit to the sensitivity of the test method because few laboratories are able to achieve such precision on a regular basis and most do not achieve it even occasionally. Several independent researchers have evaluated and provide support for using the MSD approach as additional TAC for the toxicity test methods. Thursby et al. (1997) advocate and provide reasons for using an empirical data base of minimum significant differences to provide TAC using statistical performance assessment. The State of California (Hunt et al. 1996, Starrett et al. 1993) and the West Coast marine toxicity test methods (USEPA 1995) have implemented an upper PMSD bound to minimize insensitive tests. Also the State of North Carolina has implemented additional requirements for the *Ceriodaphnia* chronic tests that reduced method

variability. North Carolina's evaluation of these additional TACs and subsequent improvements in test sensitivity appears in Appendix F.

The North Carolina data base affords the opportunity to evaluate the effectiveness of additional TAC and changes to the toxicity test procedures as they relate to the variability of WET test results (see Appendix F). For example, for PMSD, the median value decreased from 21 percent to 16 percent, while the 90th percentile decreased from 39 percent to 31 percent, indicating an overall increase in test sensitivity. The range in median values across all laboratories before adopting additional TACs was 12 percent to 36 percent. After adopting additional TACs, the range in median values was 10 percent to 27 percent, indicating a decrease in the overall spread between laboratories. The range in control CVs within a laboratory was from 21 percent to 79 percent before adopting TACs, compared to the range in control CVs within a laboratory after adopting TACs, which was narrowed to 17 percent to 36 percent. Overall, laboratories are generating data with more consistency within and between laboratories, after implementation of the additional TACs and additional method guidance provided by the State for the chronic *Ceriodaphnia dubia* test method.

6.4.4 Guidance to Testing Laboratories on How to Achieve the Range of Performance for PMSD

EPA recommends that regulatory authorities use the upper bounds (90th percentiles for PMSD in Table 3-6) to identify tests that are insufficiently sensitive. If PMSD exceeds this upper bound more often than occasionally, the laboratory should thoroughly investigate ways to reduce variability. There are three principal ways to reduce PMSD: (1) decrease within-test variability (that is, decrease the error mean square and therefore the standard deviation at each concentration); (2) increase the control mean; and (3) increase the number of replicates. The number of replicates required could be determined by trial-and-error calculations using the error mean square values obtained from a series of WET tests. At least 20 tests are recommended. The number "n" in the formula for MSD (number of replicates) would be increased and MSD re-calculated for each error mean square value. This approach uses a sample of tests specific to a particular laboratory and reveals the variation among tests. This approach would demonstrate how many replicates would be needed to achieve the upper PMSD bound, as required in Table 3-6.

6.5 Additional Guidance That Regulatory Authorities Should Implement to Further Support the WET Program

As discussed in Section 5.3, regulatory authorities have the discretion to develop and implement additional WET program requirements and guidance to ensure that WET test method variability is reduced by specifying additional guidance beyond the minimum requirements of EPA's WET test method's QA/QC and TACs. Appendix E provides a snapshot of State approaches to implementing NPDES WET programs to minimize WET test variability.

These State approaches include WET information to assist the regulated community with the following:

- Guidance regarding the evaluation of reference toxicant and effluent test results
- Guidance regarding how the State reviews reference toxicant data for laboratory performance
- Guidance regarding additional QA/QC criteria the State has developed and implemented
- Guidance regarding efforts the State has made to minimize test method variability
- Description of how the State reviews or conducts performance laboratory audits
- Description of specific implementation guidance that the State has developed to assist permit writers
- Description of how the State provides or uses toxicity test training

States contemplating such changes should consult with EPA to ensure the changes will be appropriate in the context of the State's overall NPDES WET program. In addition, States should implement a step-wise approach to address toxicity when the permit limit or monitoring trigger is exceeded in their State WET implementation plans.

For example, when an effluent is deemed toxic, then the permittee should take appropriate steps to demonstrate the magnitude, frequency, and potential source(s) of the toxicity. The components of the step-wise approach could include increased frequency of toxicity testing to characterize the magnitude and frequency of toxicity. If continued toxicity is demonstrated, then the permittee could conduct a Toxicity Reduction Evaluation/Toxicity Identification Evaluation (TRE/TIE) with toxic effluent sample(s) (USEPA 1991b, 1992). For example, EPA Regions 9 and 10 have prepared WET implementation guidance to assist their States (Denton and Narvaez 1996). This guidance provides sample permit language for a step-wise approach to address toxic samples (see Appendix C).

6.6 Chapter Conclusions

The TSD statistical approach to reasonable potential determination and permit limit derivation considers combined effluent and analytical variability through the CV of measured effluent values. Because determination of effluent variability is based on empirical measurements, the variability estimated for effluent measurements includes the variability of pollutant levels, sampling variability, and a smaller component owed to method variability. Steps should be taken to reduce these sources of variability. EPA believes that the TSD statistical procedures are appropriately protective in considering both effluent and analytical variability in reasonable potential and effluent limit calculations.

EPA recommends that regulatory authorities use a sampling program that conducts at least ten representative WET tests over a period of three years to represent the full range of effluent variability. Regulatory authorities should use recommended procedures in the TSD to determine when numeric WET limits or WET monitoring triggers are needed. Other permit conditions may include monitoring triggers, such as increased toxicity testing, TREs/TIEs, and follow-up actions initiated because a permit limit is exceeded or a monitoring trigger is not met. Regulatory authorities should implement the additional test sensitivity requirements by requiring that each test result not exceed the upper PMSD bound. In addition, regulatory authorities should determine the appropriate NOEC for test results below the lower PMSD bound as described in Section 6.4.2. These efforts should lead to increased confidence in the effect concentrations that are generated to evaluate self-monitoring data.

7.0 CONCLUSIONS AND GUIDANCE TO LABORATORIES, PERMITTEES, AND REGULATORY AUTHORITIES

This document was prepared to address whole effluent toxicity (WET) test variability. The document has three goals: (1) quantify the variability of promulgated test methods and report a coefficient of variation (CV) as a measure of test method variability; (2) evaluate the statistical methods described in the TSD for determining the need for and deriving WET permit conditions; and (3) suggest guidance for regulatory authorities on approaches to address and minimize test method variability. This document quantified the variability of toxicity test methods based on the end use of the data, that is, the effect concentrations (e.g., NOEC, LC50, EC25). The within-laboratory variability of these effect concentrations was quantified by obtaining multiple test results under similar test conditions using the same reference toxicant. The major conclusions of this document are discussed below.

7.1 General Conclusions

- EPA's *Technical Support Document for Water Quality-based Toxics Control* (referred to as the TSD) presents guidance for developing effluent limits based on three key components: (1) water quality criteria; (2) a calculated dilution factor used to derive a waste load allocation (WLA) from the criteria; and (3) a statistical calculation procedure that uses a CV based on effluent data to calculate effluent limits from the WLA. EPA's TSD statistical approach is appropriately protective, regarding both effluent and analytical variability, provided that the criteria and WLA are derived correctly. It is inappropriate to adjust the TSD statistical methodology for determining when water quality-based effluent limits are needed and for calculating such limits (Section 6 and Appendix G).
- EPA's analysis indicates that the TSD approach appropriately accounts for both effluent variability and method variability. EPA does not believe a reasonable alternative approach is available to determine a factor that would discount the effects of method variability using the TSD procedures (Section 6.1.1 and Appendix G).
- Interim CVs are identified for promulgated WET test methods [Appendix A, Table A-1 (acute methods) and Table A-2 (chronic methods)], pending completion of between-laboratory studies, which may affect these interim CV estimates.
- Comparisons of WET method precision with method precision for analytes commonly limited in the National Pollutant Discharge Elimination System (NPDES) permits clearly demonstrate that the variability of the promulgated WET methods is within the range of variability experienced in other types of analyses. Several researchers also noted that method performance improves when prescribed methods are followed closely by experienced analysts (Section 4.3).
- The hypothesis test procedures prescribed in EPA's WET methods will provide adequate protection against false conclusions that an effluent is toxic. However, the incidence of false negatives can be high because of high within-test variability, making it difficult to detect toxicity when toxicity is truly present. Therefore, evaluating the power of current experimental designs is desirable. EPA expects that regulatory authorities will make prompt and measurable progress toward the goal of requiring all WET tests to detect a toxic effect of 25 percent to 33 percent with power of 0.80 (Section 5.3.3 and Appendix B.4).
- Quality assurance problems became apparent when evaluating the data for this study, especially for the metal reference toxicants and sodium dodecyl sulfate (SDS). Standardizing the choice of reference toxicant and the concentrations to be tested may be appropriate, as well as establishing bounds on the range of acceptable effect concentrations for each test method. As a result,

quantifying between-laboratory variability will be difficult unless these issues can be resolved (Section 5.3.1 and Appendix G.2.5).

- The data analysis did not reveal the potential sources and causes of variability, such as using different sources of test organisms, dilution water, and food. To assess the sources of variability fully, experimenters must carefully design new studies (Section 5.3.1).

7.2 Recommendations for Minimizing Variability and Its Effects

Three critical areas are identified to minimize WET test method variability:

- Obtaining a representative effluent sample,
- Conducting the toxicity tests properly to generate biological endpoints, and
- Calculating the appropriate statistical endpoints to have confidence in the effect concentration.

This document provides guidance to toxicity testing laboratories, permittees, and regulatory authorities in conducting biological and statistical methods and evaluating test effect concentrations. It also develops guidance for regulatory authorities on approaches to address and minimize test method variability. The principal aspects of the guidance are presented in Table 3-6 and re-presented here.

Range of Relative Variability for Endpoints of Promulgated WET Methods, Defined by the 10th and 90th Percentiles from the Data Set of Reference Toxicant Tests^a

Test Method ^b	Endpoint ^c	No. of Labs	No. of Tests	PMSD		Control CV ^d	
				10 th	90 th	10 th	90 th
1000.0 Fathead Minnow	G	19	205	9.4	35	0.035	0.20
1002.0 <i>Ceriodaphnia dubia</i>	R	33	393	11	37	0.089	0.42
1003.0 Green Alga	G	9	85	9.3	23	0.034	0.17
1004.0 Sheepshead Minnow	G	5	57	6.3	23	0.034	0.13
1006.0 Inland Silverside	G	18	193	12	35	0.044	0.18
1007.0 Mysid	G	10	130	12	32	0.088	0.28
2000.0 Fathead Minnow	S	20	217	4.2	30	0	0.074
2002.0 <i>Ceriodaphnia</i>	S	23	241	5.0	21	0	0.11
2004.0 Sheepshead Minnow	S	5	65	0 ^e	55	0	0
2006.0 Inland Silverside	S	5	48	7.0	41	0	0.079
2007.0 Mysid (<i>A. bahia</i>)	S	3	32	5.1	26	0	0.081
2011.0 Mysid (<i>H. costata</i>)	S	2	14	18	47	0	0.074
2021.0 Daphnia (<i>D. magna</i>)	S	5	48	5.3	23	0	0.11
2022.0 Daphnia (<i>D. pulex</i>)	S	6	57	5.8	23	0	0.11

^a The precision of the data warrants only three significant figures. When determining agreement with these values, one may round off values to two significant figures (e.g., values >3.45000... and ≤3.5000... are rounded to 3.5). Method 1009.0 (red macroalga) is not reported because it is inadvisable to characterize method variability using only 23 tests from just two laboratories.

^b EPA did not assign method numbers for acute methods in EPA/600/4-90/027F. The numbers assigned here were created for use in this document and in related materials and data bases.

^c G = growth, R = reproduction, S = survival

^d CVs were calculated using untransformed control means for each test.

^e An MSD of zero will not occur when the EPA flow chart for statistical analysis is followed. In this report, MSD was calculated for every test, including those for which the flow chart would require a nonparametric hypothesis test. EPA recommends using the value 4.2 (the 10th percentile shown for the fathead minnow acute test) in place of zero as the 10th percentile PMSD (lower PMSD bound) for the sheepshead minnow acute test.

7.2.1 Guidance to Toxicity Testing Laboratories

- Testing laboratories should maintain quality assurance/quality control (QA/QC) control charts for percent minimum significant difference (PMSD) along with the statistical endpoints such as NOEC, LC50, and EC25. Testing laboratories should regularly plot the individual raw test data and the average treatment responses to examine possible causes of excessive variability (Section 5.3.1.1).
- The minimum number of replicates for the chronic toxicity tests should be four for the chronic fathead minnow, sheepshead minnow, and inland silverside test methods (Sections 5.3.3.1 and 5.6).
- Testing laboratories should take steps to ensure that the test PMSD does not exceed the upper bound provided in the table above (Sections 3.3, 5.3.3, and 6.4 and Table 3-6). This may require ensuring more uniformity among test organisms and/or using more replicates. Tables are provided to aid in choosing the number of replicates (Tables B-14 and B-15).
- Testing laboratories should examine the power tables to ensure that test results will meet the recommended test sensitivity criteria. These tables can be used to make decisions about replication, given the knowledge of typical values for error mean square (EMS) and number of tested concentrations (Section 5.3.3 and Tables B-9 through B-15).

7.2.2 Guidance to NPDES Permittees

- Permittees should select and conduct all data analyses with one qualified toxicity testing laboratory to determine reasonable potential, derive permit limits, and generate self-monitoring test results. Conducting all effluent testing consistently using one reference toxicant is also prudent (Section 6.1.4 and Appendix G.2.5).
- Permittees should generate WET data ($n = 10$) that have been accumulated over a year or more to fully characterize effluent variability over time. The sampling dates and times should span a sufficient duration to represent the full range of effluent variability (Sections 6.1.3 and 6.2 and Appendix G.2.4).
- Permittees should examine testing laboratories' QA/QC control charts. If the CV for reference toxicant tests is greater than the 75th percentile in Tables 3-2 through 3-4, variability can likely be reduced, even if the individual EC25 and LC50 values fall within the control limits (Section 5.3.1.1).
- Permittees should examine toxicity test data to ensure that data being submitted to regulatory authorities meet specified effluent holding times, temperature, laboratory control limits, and test acceptability criteria, such as requirements for test sensitivity lower and upper PMSD bounds (Sections 5.2 through 5.4).
- Permittees should anticipate and plan for a change if switching to a different testing laboratory. The permittee should compare reference toxicant test data from the current laboratory with data from the candidate replacement laboratory in order to ensure acceptable variability and a similar average effect (Section 6.1.4).

7.3 Guidance to Regulatory Authorities

Guidance to Regulatory Authorities Related to Determining Reasonable Potential and Deriving Permit Limits:

- Regulatory authorities should use EPA's recommended statistical approach in deriving permit limitations. The statistical approach outlined in the TSD represents an effective and appropriately protective approach to effluent limit development (Section 6.1 and Appendix G.1).
- Regulatory authorities should calculate the facility-specific CV using point estimate techniques to determine the need for and derive a permit limit, even if the self-monitoring test results will be determined using hypothesis test procedures (Sections 3.4.1 and 6.2).
- Regulatory authorities that need to cite a characteristic CV for a promulgated method may use Tables A-1 and A-2 in Appendix A, which show the median CV from Tables 3-2 through 3-4, pending completion of between-laboratory studies.
- EPA recommends that regulatory authorities implement a step-wise approach to address toxicity. This approach can determine the magnitude and frequency of toxicity and appropriate follow-up actions for test results that indicate exceedance of a monitoring trigger or a permit limit (Section 6.5).

Guidance to Regulatory Authorities Related to Collecting Effluent Samples, Conducting the Toxicity Test, and Evaluating the Effect Concentrations:

- Regulatory authorities should design a sampling program that collects representative effluent samples to fully characterize effluent variability for a specific facility over time. At least 10 samples are needed to estimate a variance or CV with acceptable precision for a specific facility (Sections 6.1.3 and 6.2).
- Regulatory authorities should ensure that statistical procedures and test methods have been properly applied to produce WET test results. Evaluating other factors and data, such as biological and statistical quality assurance, and ensuring that test conditions and test acceptability criteria (TAC) have been met would be prudent (Sections 5.2 through 5.5).
- Regulatory authorities should apply both the upper and lower bounds using the PMSD as an additional TAC (Section 6.4 and Table 3-6). The State of North Carolina implemented an effective WET program that required additional TAC and guidance for test methods that served to minimize test method variability (Appendix F).
- Regulatory authorities should develop a QC checklist to assist in evaluating and interpreting toxicity test results (Section 5.3.1.1). See Appendix E for examples of State WET implementation procedures.
- Regulatory authorities should consider participation in the National Environment Laboratory Accreditation Program and should conduct routine performance audit inspections to evaluate individual laboratory performance. Inspections should evaluate the laboratory's performance with QC control charts based on reference toxicants, examine procedures for conducting the toxicity test procedures, and examine procedures for analyzing test results (Section 5.3.1.1).
- Regulatory authorities should incorporate revised technical guidance recently published by EPA captioned "Method Guidance and Recommendations for Whole Effluent Toxicity (WET) Testing"

(40 CFR Part 136) (USEPA 2000a). The guidance addresses: (1) error rate assumption adjustments; (2) concentration-response relationships; (3) incorporation of confidence intervals; (4) acceptable dilution waters for testing; (5) guidance on blocking by parentage for the chronic *C. dubia* test method; and (6) procedures for controlling pH drift.

7.4 Future Directions

- An independent peer-reviewed workshop should be convened to evaluate alternatives to the statistical approaches currently used in EPA's WET test methods. Such a workshop might suggest alternatives regarding (1) WET statistical flowcharts, (2) WET statistical methods used to estimate effect concentrations, and (3) test data interpretation and review guidelines (Section 5.5).
- Such a workshop might also evaluate additional QC requirements and recommendations regarding the specification of a reference toxicant and the concentrations to be tested for each test method (Section 5.3.1).

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