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EHD DIV-WIDE PLAN 001 Quality Assurance Manual—General

Environmental Health Division

Wisconsin State Laboratory of Hygiene University of Wisconsin

This manual applies to accredited (1.1) methods within the following analytical sections:

Environmental Toxicology Section
Inorganic Chemistry Section
Trace Element Clean Lab
Metals Section
Organic Chemistry Section
Organic LCMS/PFAS
Radiochemistry Section
Water Microbiology Section

This manual also includes the following support sections:

Glassware/Media, EHD Data Management, Sample Receiving, & Customer Service

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Wisconsin State Laboratory of Hygiene Environmental Health Division

Quality Assurance Manual—General EHD DIV-WIDE PLAN 001 Version 4

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0. 2016 TNI Concordance — Volume 1, Module 2, Quality Systems General Requirements

Standard	TNI	Standard Description	Concordance
1 A Introdu	Page	 cope and Applicability	
1.1	1	Quality system	QAM-General and associated policies and
40 M			SOPs
4.0 Manage		equirements	LIW Office of Local Affairs
4.1.1	8	Legally responsible Meet TNI & other standards and customer	UW Office of Legal Affairs
		needs	QAM-General, section 1.3
4.1.3	8	Management system covers work at all facilities	QAM-General, sections 1.1, 1.17.7
4.1.4	8	Potential conflicts of interest organization-wide	Organizational charts, P-files
4.1.5 a)	8	Employees have resources necessary to carry out their duties	QA Manual, METHOD SOPs
4.1.5 b)	8	Employees are free from undue pressures	EHD DIV-WIDE GENOP 029, "Data
,		that may affect quality of work	Integrity, Ethics, & Data Documentation
			Procedure" (including references)
4.1.5 c)	8	Protection of customers' confidential information	QAM-General, section 1.7
4.1.5 d)	8	Maintain competence, impartiality,	EHD DIV-WIDE GENOP 029 (including
,		judgment, and operational integrity	references)
4.1.5 e)	8	Organization and management structure	Organizational charts, QAM-General, section 1.4
4.1.5 f)	9	Define inter-relationships of personnel	Organizational charts, QAM-General, section 1.4
4.1.5 g)	9	Adequate supervision with assessment of results	QAM-General, sections 1.4 & 1.10
4.1.5 h)	9	Management and resources needed	QAM-General, section 1.4
4.1.5 i)	9	Quality manager with defined	QAM-General, section 1.4
,		responsibility & authority	
4.1.5 j)	9	Appoint deputies	Organizational charts, QAM-General, section 1.4
4.1.5 k)	9	Personnel are aware of importance of their	EHD DIV-WIDE GENOP 029, QAM-
		activities	General
4.1.6	9	Communicate regarding management	EHD DIV-WIDE GENOP 023, "Procedure
		system	for the Management Review of the Quality
			System"
4.1.7.1	9	Duties of quality manager	QAM-General, section 1.4, P-files
4.1.7.2	9	Duties of technical manager	QAM-General, section 1.4, P-files
4.2.1	10	Management system	QA Manual & associated SOPs/documents
4.2.2	10	Quality policy statement	QAM-General, section 1.3
4.2.3	11	Management's commitment to	EHD DIV-WIDE GENOP 023, QAM-

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Standard	TNI Page	Standard Description	Concordance
		implementation and improvement of Quality System	General, section 1.19
4.2.4	11	Meeting customer and regulatory requirements	QAM-General, section 1.3.
4.2.5	11	Structure of documentation	QAM-General, sections 1.11 & 1.26.
4.2.6	11	Responsibilities of QA & technical managers	QAM-General, sections 1.3 & 1.4
4.2.7	11	Maintenance of management system through changes	QAM-General, section 1.4
4.2.8.1	11	Data integrity system	EHD DIV-WIDE GENOP 029 (including references), QAM-General, section 1.10.6
4.2.8.2	11	Quality manager responsible for updating QA Manual	QAM-General, section 1.2.3
4.2.8.3	11	Contents of QA Manual	QA Manual
4.2.8.4	12	Additional requirements to be contained or referenced in QA Manual	QA Manual & referenced documents, Organizational charts, P-files
4.2.8.5	13	SOP requirements	QAM-General, section 1.11, section supplements (sections 3 & 17), SOPs
4.3.1	14	General document control	QAM-General, section 1.11, section supplements (section 3)
4.3.2	14	Document approval & issue	QAM-General, section 1.11, section supplements (section 3), SOPs
4.3.3	14	Document changes	QAM-General, section 1.11, section supplements (section 3), SOPs
4.4	15	Review of requests, tenders and contracts	QAM-General, section 1.15 & section supplements (section 7)
4.5	16	Subcontracting of environmental tests	QAM-General, section 1.23
4.6	16	Procedures for purchasing services & supplies, procedures for receiving, storing, and verifying that supplies & reagents are OK for use.	QAM-General, section 1.18, OnBase BPP, METHOD SOPs
4.7.1	16	Services to clients—clarifying requests, monitoring performance, & sending kits	QAM-General, sections 1.6, 1.15
4.7.2	17	Seeking feedback from clients	QAM-General, section 1.6
4.8	17	Policy & procedure for complaints. Records of complaints & corrective actions	QAM-General, section 1.6. EHD DIV- WIDE GENOP 017, "Managing Customer Feedback"
4.9	17	Policy & procedures for control of nonconforming environmental testing work	QAM-General, section 1.21 & section supplements (section 11), METHOD SOP
4.10	18	Continual improvement of the management system	QAM-General, section 1.3
4.11	18	Policy & procedures for selecting, implementing, & monitoring corrective actions, root cause analysis, additional internal audits, QC responsibilities	QAM-General section 1.21 & section supplements (section 11), METHOD SOPs Non-Conformance Management reports

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Standard	TNI Page	Standard Description	Concordance
4.12	19	Preventive action	QAM-General section 1.21.4, METHOD SOPs
4.13.1.1	19	Procedures for identification, collection, indexing, access, filing, storage, maintenance and disposal of records.	QAM-General, section 1.12, section METHOD and GENOP SOPs, LABWIDE GENOP 1002, "Records Storage & Disposal"
4.13.1.2	19	Records retention times	QAM-General section 1.12, LABWIDE GENOP 1002, https://www.library.wisc.edu/archives/records-management/retention-disposition/general-records-schedules/
4.13.1.3	19	Records held secure & in confidence	LABWIDE GENOP 1002, QAM-General, sections 1.7, 1.12
4.13.1.4	19	Electronic back-up of records	QAM-General, section 1.17.4 OIS procedures
4.13.2.1	19	Retention of technical records	QAM-General section 1.12, https://www.library.wisc.edu/archives/records- management/retention-disposition/general- records-schedules/
4.13.2.2	20	Data & observations recorded immediately	EHD DIV-WIDE GENOP 029, section 9
4.13.2.3	20	Alterations to records	EHD DIV-WIDE GENOP 029, sections 9 & 11.
4.13.3 a)	20	Documentation of history of sample	LIMS, section METHOD and GENOP SOPs, log-books, data packets
4.13.3 b)	20	Record retention minimum of 5 years	QAM-General section 1.12; https://www.library.wisc.edu/archives/records- management/retention-disposition/general- records-schedules/
4.13.3 c)	20	Records available to accreditation body	QAM-General (section 1.7)
4.13.3 d)	20	Electronic records supported by hardware and software necessary for their retrieval	Internal web-site (Admin Services/Info Systems) OIS procedures
4.13.3 e)	20	Access log for archived information	QAM-General section 1.12.3, LABWIDE GENOP 1002
4.13.3 f)	20	Historical reconstruction of data	QA Manual, LIMS, Department-level METHOD and GENOP SOPs, log-books, data packets, PT results, DOC forms
4.13.3 g)	21	Data legibly recorded	EHD DIV-WIDE GENOP 029, "Data Integrity, Ethics, Data Documentation," sections 9 & 11
4.13.3 h)	21	State legal requirements concerning records are followed	QAM-General, section 1.12
4.14	21	Internal audit requirements	QAM-General, section 1.20 & section supplements (section 14), EHD DIV-WIDE QA 120, "Internal Audit Procedures"
4.15	22	Management review requirements	QAM-General, section 1.19, EHD DIV-

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Standard	TNI Page	Standard Description	Concordance
	1 agc		WIDE GENOP 023
4.16	23	Data integrity investigations	EHD DIV-WIDE GENOP 029
5.0 Technic		· · · · · · · · · · · · · · · · · · ·	EIID DIV-WIDE GENOT (2)
5.1.2	23	Take account of factors that affect	QA Manual, METHOD and GENOP SOPs
3.1.2		correctness and reliability of tests	(details below)
5.2.1	23	Ensure competence of analysts	QAM-General, sections 1.4 (education), 1.8
3.2.1		Ensure competence of analysis	(hiring), 1.10 (training)
5.2.2	24	Training program	QAM-General, section 1.10, section
3.2.2	27	Training program	supplements (section 2)
5.2.3	24	Personnel are employed by the lab	HR P-files
5.2.4	24	Maintain current job descriptions for	HR P-files
J.2. T	27	personnel	THE T-MES
5.2.5	24	Authorization of personnel to perform	HR P-files, method training forms,
3.2.3	27	specific tests and duties	certification statements, DOCs
5.2.6.1	24	Technical manager qualifications	HR P-files
5.2.6.2	26	Technical manager qualification	NA NA
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5.2.7	26	Data Integrity Training	EHD DIV-WIDE GENOP 029, personnel
3.2.7	20	Data integrity Training	training files
5.3.1	27	Lab facilities shall facilitate testing,	METHOD SOPs, QAM-General section
5.5.1	2'	document requirements	1.17
5.3.2	27	Monitor, control, and record	METHOD SOPs for requirements, logbook
3.3.2	2'	environmental conditions that affect	for data
		testing. Stop testing when conditions	Tor data
		jeopardize results.	
5.3.3	27	Separation between incompatible	METHOD SOPs, QA Manual section
		activities.	supplements (section 8)
5.3.4	27	Controlled access where necessary for	METHOD SOPs, QA Manual section
		quality of tests	supplements (section 8)
5.3.5	27	Good housekeeping in lab	LABWIDE SAFETY 102, Chemical
			Hygiene Plan, sect. 26
5.4.1	27	Use of appropriate methods	METHOD and GENOP SOPs
5.4.2	28	Use of methods that meet customer needs	METHOD SOPs, QA Manual section
			supplements (sect. 18)
5.4.3	28	Lab-developed methods	METHOD SOPs, QA Manual section
		1	supplements (sect. 18)
5.4.6	30	Procedure for estimating analytical	QAM-General, section 1.14
		uncertainty	
5.4.7.1	31	Appropriate checks of calculations and	METHOD and GENOP SOPs
		data transfers	
5.4.7.2 a)	31	Computer software developed by user is	METHOD and GENOP SOPs and
		documented and validated	associated data
5.4.7.2 b)	31	Procedures for protecting data in	OIS procedures, QAM-General, section
		computers	1.17.4
5.4.7.2 c)	31	Computers maintained to ensure proper	OIS procedures
Ý		functioning	_

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Standard	TNI Page	Standard Description	Concordance
5.5.1	31	Lab must have all equipment needed for test	QAM-General section 1.17, QA Manual section supplements (section 9), METHOD SOPs
5.5.2	31	Equipment must meet specifications	QA Manual section supplements (section 9), METHOD SOPs
5.5.3	32	Equipment shall be operated using up-to- date instructions by authorized personnel	METHOD SOPs, certification statements, DOCs
5.5.4	32	Equipment must be uniquely identified	QA Manual section supplements (section 9), instrument ID lists
5.5.5	32	Equipment records	Sectional instrument ID lists, instrument manuals, calibration data, instrument logbooks, METHOD SOPs, MDL records
5.5.6	32	Proper use, handling, maintenance of equipment	QA Manual section supplements (section 9), METHOD and GENOP SOPs, instrument manuals, and logbooks
5.5.7	32	Equipment not performing correctly must be taken out of service	METHOD and GENOP SOPs, instrument manuals, and logbooks
5.5.8	32	Equipment identified to indicate calibration status (when practicable)	METHOD and GENOP SOPs, instrument logbooks
5.5.9	32	Equipment calibration status checked when returned from outside of lab	QAM-General, section 1.17.1
5.5.10	32	Intermediate checks done according to defined procedure	METHOD SOPs
5.5.11	33	Calibration correction factors used correctly	METHOD SOPs
5.5.12	33	Safeguard equipment from adjustments that would invalidate calibrations	METHOD SOPs
5.5.13.1 a)	33	Support equipment specifications defined	METHOD SOPs & GENOP SOPs for balances, ovens, refrigerators, incubators, measuring devices, etc.
5.5.13.1 b)	33	Support equipment maintained and records kept	Support equipment logs
5.5.13.1 c)	33	Each day of use, balances, ovens, refrigerators, freezers, and water baths must be checked and documented.	METHOD and GENOP SOPs for balances, ovens, refrigerators, incubators, measuring devices, etc., logbooks, data printouts
5.5.13.1 d)	33	Temperature measuring devices calibrated annually	QAM-General, section 1.17.6
5.5.13.1 e)	33	Volumetric dispensing devices (except Class A) must be checked quarterly.	Section SOPs for pipette performance checks, logbooks for quarterly checks
5.5.13.1 f)	34	Support equipment calibrated or verified annually	METHOD and GENOP SOPs for balances, ovens, refrigerators, incubators, measuring devices, etc., logbooks
5.5.13.1 g)	34	Retain raw data for support equipment	Logbooks and data printouts
5.6.2	34	Measurement traceability—calibration of equipment	QA Manual section supplements (sect. 9), METHOD SOPs
5.6.3	36	Measurement traceability—reference	QAM-General section 1.13, & section

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Standard	TNI Page	Standard Description	Concordance
		standards/materials	supplements (sect. 15), METHOD SOPs
5.6.4.1	36	Reference standards/materials—correlation of results (PTs, CRMs)	QAM-General section 1.24, & section supplements (sect. 13), METHOD SOPs
5.6.4.2	37	Documentation & labeling of standards/reagents	QAM-General section 1.18.3, & section supplements (sect. 10, 15), standard & reagent logs
5.7	37	Sampling plan, procedures, & documentation	Test request forms, LIMS
5.8.1	38	Procedures for transportation, receipt, handling, storage, disposal of samples	QAM-General section 1.16, & section supplements (section 5), EHD DIV-WIDE GENOP 033, "Sample Acceptance Policy," section METHOD SOPs, EHD HDM GENOP 116, "Sample Receiving & Login"
5.8.2	38	System for identifying test items	QAM-General, section 1.16.3
5.8.3	38	Record departures from sample receipt protocols, consult with customer.	QAM-General, section 1.16, & section supplements (section 11), EHD DIV-WIDE GENOP 033
5.8.4	38	Safe, secure, & appropriate storage conditions	Section METHOD SOPs for storage conditions, and section GENOPs & logbooks for monitoring and recording environmental conditions
5.8.5	38	System for uniquely identifying samples	QAM-General, section 1.16.3
5.8.6	39	Sample acceptance policy requirements	EHD DIV-WIDE GENOP 033
5.8.7.1	39	Procedure for verifying & documenting preservation	QAM-General, section 1.16, EHD HDM GENOP 103, "Inorganic Sample Check-In," EHD HDM GENOP 116
5.8.7.2	39	What to do if samples do not meet acceptance criteria	EHD DIV-WIDE GENOP 033
5.8.7.3	39	Requirements for sample receipt documentation	HDM dept. SOPs, LIMS
5.8.7.4	40	Retain all documentation sent to the lab with the sample.	EHD DIV-WIDE GENOP 033, QAM-General, sections 1.15.3, 1.16
5.8.7.5	40	Retain COC forms.	EHD DIV-WIDE GENOP 033, QAM- General, section 1.16
5.8.8	40	Legal chain of custody procedures	EHD HDM GENOP 106, "Enforcement Sample Handling," QA Manual section supplements (section 5)
5.8.9 a), b)	40	Sample storage requirements	Section METHOD SOPs
5.8.9 c)	40	SOPs for disposal of samples, digestates, Leachates, extracts, etc.	EHD INORG GENOP 110, EHD ORG CHEM GENOP 0029, EHD RAD GENOP 011, and section METHOD SOPs
5.9.1	41	Quality control procedures and monitoring	QA Manual section supplements (sections 11, 12, 13), EHD DIV-WIDE QA 113, "Horizon QC Limit evaluations & Updates," section METHOD SOPs

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Standard	TNI Page	Standard Description	Concordance
5.9.2	41	Corrective action for QC data outside of pre-defined criteria	QA Manual section supplements (section 11), section METHOD SOPs
5.9.3	41	Monitoring of QC parameters (several required ones are listed), evaluation using established acceptance criteria.	QA Manual section supplements (sections 11, 12), EHD DIV-WIDE QA 113, section METHOD SOPs
5.10.2	42	Test report requirements (list)	QAM-General, section 1.22, WSLH Laboratory Reports
5.10.3.1	43	Additional test report requirements (e.g. flags, comments, uncertainty of measurement, opinions, interpretations)	QAM-General, section 1.22, section METHOD and GENOP SOPs, WSLH Laboratory Reports
5.10.3.2	44	Reporting the results of sampling	WSLH lab reports
5.10.5	44	Reporting opinions and interpretations	QAM-General, section 1.22, WSLH Laboratory Reports
5.10.6	45	Testing results obtained from sub- contractors	QAM-General, section 1.23
5.10.7	45	Reporting requirements met for electronic transmission	WSLH Laboratory Reports
5.10.8	45	Clear, easily understood report format	WSLH Laboratory Reports
5.10.9	45	Requirements for amendments to test reports	WSLH Laboratory Reports
5.10.10	45	Exceptions to reporting requirements	WSLH Laboratory Reports
5.10.11 a)	46	Additional report requirement: report time of analysis if holding time < 72 hrs.	WSLH Laboratory Reports
5.10.11 b)	46	Additional report requirement: report if results are on a basis other than as received (e.g. dry weight)	WSLH Laboratory Reports
5.10.11 c)	46	Non-accredited tests must be clearly identified	WSLH Laboratory Reports, WSLH external web-site
5.10.11 d)	46	Clear ID of numerical results outside the calibration range.	WSLH Laboratory Reports

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1. General Sections

1.1. Applicability

This Quality Assurance Manual—General, along with section supplements are designed to meet requirements of The NELAC Institute (TNI, also referred to as NELAP), Wisconsin Department of Natural Resources (NR 149), and the Environmental Protection Agency (EPA). This applies to lab sections within the Wisconsin State Laboratory of Hygiene (WSLH) Environmental Health Division (EHD). Testing conducted under any of these accreditations, at any WSLH facility (see 1.17.7), must meet the requirements set forth in this manual. Testing conducted under other accreditations or non-accredited testing may use this manual as is to meet requirements. If other accrediting agencies have additional requirements not covered in this manual, those must be met in ancillary documents.

For current scopes of accreditation, see: http://www.slh.wisc.edu/about/compliance/

Note: TNI (The NELAC Institute) was formed in 2006 when NELAC (National Environmental Laboratory Accreditation Conference) combined with INELA (Institute for National Environmental Laboratory Accreditation). NELAP (National Environmental Laboratory Accreditation Program) is one program operated by TNI. For more information, see https://nelac-institute.org/content//programs.php

1.2. Manual Organization and Maintenance

1.2.1. Historical Perspective

The EHD lab sections that are covered under this manual have, over the years, maintained various documents that have served as de facto Quality Assurance Manuals. In November of 1998 an attempt was made to construct a NELAP-compatible Quality Assurance Manual for the Inorganic and Organic Chemistry Departments (Revision 2.0). That manual was revised in May of 1999 (Revision 2.1).

Revision 3.0 was developed during the laboratory's NELAP application process. It was an attempt to cover all of the departments (now known as sections) in the laboratory that would be accredited under NELAP. Those sections are now Inorganic Chemistry, Trace Element Clean Lab, and Metals. Annual reviews resulted in minor revisions (designated as 3.1, 3.2, etc.)

Within revision 4.0, it was necessary to update the NELAC concordance to comply with the new numbering scheme of the 2003 NELAC standards.

Revision 5.0 was completely reorganized so that each section had a separate chapter devoted to it.

Revisions 6.0, 7.0, and 8.0 were reviewed by each section and changes implemented as necessary.

Revision 9.0 was re-written and a new concordance compiled to comply with the new TNI (The NELAC Institute) Standard EL-V1-2009, which is consistent with ISO/IEC 17025:2005 requirements that are

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relevant to the scope of environmental testing services.

Revisions 10.0 and 11 were reviewed by each section and changes implemented as necessary.

Revision 12 was completely reorganized by developing a template with required sections and content for each sectional chapter. The goal was to make the manual more concise, functional, and organized.

Revisions 13, 14, 15, 16, and 17 were reviewed by each section and changes implemented as necessary. Rev. 18 incorporated some updates and notes to correspond with 2016 TNI standards, and the concordance chart was updated for 2016 TNI. Revision 19 was reviewed by each section and changes implemented as necessary.

In Aug. 2022, the QA Manual was updated and then transitioned into the OnBase document management system as Version 1. The structure of the manual was changed as noted below. Subsequent versions of the sectional supplements are managed by each section. Subsequent versions of the General plan are managed by a divisional QA coordinator (or specialist) (see 1.2.3)

1.2.2. Structure

EHD DIV-WIDE PLAN 001, Quality Assurance Manual—General, contains general information that applies to all sections and testing covered by this manual. Each section then has a supplemental plan (e.g. EHD INORG PLAN 001) that contains information specific to that section.

SOP ID	Section	SOP Type	Title
		Type	
EHD DIV-WIDE PLAN 001	DIV-WIDE	PLAN	Quality Assurance ManualGeneral
EHD INORG PLAN 001	INORG CHEM	PLAN	Quality Assurance Manual, Inorg/TECL/Metals*
EHD RAD PLAN 001	RAD CHEM	PLAN	Quality Assurance Manual, Radiochemistry*
EHD ENV TOX PLAN 001	ENV TOX	PLAN	Quality Assurance Manual, Environmental Toxicology*
EHD MICRO PLAN 001	WATER MICRO	PLAN	Quality Assurance Manual, Water Microbiology*
EHD ORG CHEM PLAN 001	ORG CHEM	PLAN	Quality Assurance Manual, Organic Chemistry*
EHD HDM PLAN 001	HDM	PLAN	Quality Assurance Manual, EHD Data Management*
EHD MEDIA PLAN 001	MEDIA	PLAN	Quality Assurance Manual, Glassware/Media*

^{*}Sub-title: (Supplement to EHD DIV-WIDE PLAN 001, QA Manual—General)

1.2.3. Maintenance

At least annually, Quality Assurance Specialists/coordinators will review and update EHD DIV-WIDE PLAN 001, QA Manual—General. In addition, each section of EHD will review and update their QA Manual supplement. Changes to the supplements may occur at any time as needed by the Sections. Changes to the "General" plan will be facilitated by contacting the appropriate QA coordinator if outside

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of the annual renewal timeframe. The electronic version of the manual will reside and be managed in OnBase. Editing within OnBase is controlled through permissions to specific Active Directories. For details, refer to LABWIDE GENOP 700, OnBase User Guide.

1.3. Quality Policy Statement

This quality policy statement describes the overall objectives of the Environmental Health Division's quality system. The complete quality system is documented in the remainder of this Quality Assurance Manual—General, in the section supplements, and in policies and procedures that are referenced in this manual. This quality system is based on the required elements contained in the 2016 TNI (The NELAC Institute) Standard, Volume 1 "Management and Technical Requirements for Laboratories Performing Environmental Analysis." The management of the Environmental Health Division, under the authority of the division leadership, is committed to these quality objectives as a means of maintaining our status as an excellent public and environmental health laboratory. Quality is everyone's responsibility.

Objectives of the Quality System:

High accuracy of work

The EHD's quality system ensures that data is of excellent and documented quality. The management and laboratory professionals are committed to following good professional practices (as defined by our quality system) and to compliance with applicable standards.

Data Integrity

All management and laboratory staff are committed to ethical laboratory practices. All employees are responsible for following the WSLH data integrity, ethics, and data documentation policies.

Continuous quality improvement

Laboratory management and staff are dedicated to continuous quality improvement by means such as corrective and preventive action, root cause analysis, internal audits, and management system reviews. Staff are encouraged to bring suggestions for quality improvement to management. Staff are also empowered to seek out professional development opportunities.

Customer satisfaction

The laboratory's standard of service to our clients and the citizens of the State of Wisconsin includes meeting all quality system objectives, providing timely results, remaining fiscally responsible, and addressing customer questions and concerns. When requested, the EHD also provides outreach services and training if we have sufficient resources to do so. Research and method development work may also be conducted upon request, although this usually would fall outside of any accredited methods.

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Compliance with applicable standards

The EHD conducts testing that may be regulated under one or more of the following agencies: the United States Environmental Protection Agency (EPA), the Wisconsin Department of Natural Resources (DNR), The NELAC Institute (TNI), the Wisconsin Division of Health Services (DHS), Wisconsin Department of Agriculture, Trade and Consumer Protection (DATCP), and local public health agencies. The management and laboratory professionals are committed to upholding the requirements of these standards.

Staff training

Training includes initial and continuing instruction on the quality system documented in this plan and referenced policies and procedures as required for specific job duties. The training ensures that the quality system is communicated, understood, and implemented by the appropriate personnel.

1.4. Organization, Management Structure, and Responsibilities

Organizational charts can be found at O:\Organizational Charts\Current Org Charts.

The Wisconsin State Laboratory of Hygiene (WSLH) is a department of the University of Wisconsin—Madison within the School of Medicine and Public Health. The WSLH was established by state statute in 1903 and is overseen by the WSLH Board. The Board serves to set policy (1.26.3) and direction for the Laboratory, and its members are either designated by state statute or appointed by the Governor of Wisconsin. Operational management of the WSLH is the responsibility of the Laboratory Director.

University of Wisconsin-	WSLH Director	Assoc. Director of
Madison Dept.		Non-Clinical Testing
Wisconsin State Laboratory of	Dr. James Schauer*	Steve Strebel (interim)
Hygiene		

*Address: 465 Henry Mall, Madison, WI 53706

1.4.1. Divisions

The WSLH is divided into several analytical divisions including the Environmental Health Division (EHD). The EHD is further divided into program areas that include Environmental Chemistry, Environmental Biology, Environmental Survey Programs, Forensic Toxicology, and Science & Research. This QA Manual is primarily applicable to some sections within the Environmental Chemistry and Environmental Biology program areas. Lab Directors manage program areas.

Program Area Lab Directors report to the Associate Director of Non-Clinical Testing. In addition, Lab Directors are ultimately responsible for data quality and compliance with applicable standards. They ensure the integrity of the management system when changes are planned and implemented. In the absence of the Lab Director, a temporary appointee will act in his/her place. Note: under TNI, the WSLH has

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named certain staff as technical managers (see 2016 TNI V1M2, 5.2.6.1).

WSLH Division	Applicable Program Area	Lab Director
Environmental Health	Environmental Chemistry	Camille Danielson
Environmental Health	Environmental Biology	Jocelyn Hemming

1.4.2. Offices

The laboratory's analytical divisions are supported by the following: Office of Information Systems, Office of Finance (includes Purchasing, Accounts Receivable, & Accounting departments), Office of Human Resources, Office of the Director. The WSLH support staff offices (e.g., Human Resources) work together with on-campus UW-Madison affiliates to assist in the hiring of staff, approval of contracts, purchases and/or to resolve information system issues. The WSLH Office of Finance works with a wide-variety of contracts and fee-for-service arrangements for private customers and industry, and facilitates complex long-term arrangements with state and federal agencies, other universities and international customers.

1.4.3. Sections

The EHD and its Program Areas are divided into several sections (formerly known as departments): Shipping/Receiving, EHD Data Management/Sample Receiving/Customer Service, Environmental Chemistry (including Organic Chemistry, Air Chemistry, Inorganic Chemistry, Metals, Trace Element Clean Lab, Radiochemistry), Environmental Biology (including Environmental Toxicology, Water Microbiology, Glassware/media, Wastewater Surveillance, and Flow Cytometry), and Forensic Toxicology. Also, within the EHD is the National Atmospheric Deposition Program, PFAS Research Center, and the Soil and Forage Analysis Lab.

All of the sections included in this QA Manual are part of the EHD. The Metals section does work for both EHD and the Occupational Health Division.

Each section has a supervisor who is responsible for day-to-day operation of the laboratory. Supervisors report to the Lab Program Directors, who report to the Associate Director of Non-Clinical Testing. As a group, these supervisors oversee technical operations, sample analysis, quality assurance activities, data entry, report generation, provision of resources, and all other related areas. In addition, they are responsible for employee management and review. Supervisors will appoint a person or persons to cover their duties during an absence. If substitution appointments have not been made in advance, the Lab Directors will make managerial decisions.

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Sections covered by this QA Manual:

Program Area	Section	Supervisor
Environmental Chemistry	Inorganic Chemistry	Graham Anderson
	Metals	Graham Anderson
	Organic Chemistry	Erin Mani
	Organic LCMS/PFAS	Kristen Hannon
	Trace Element Clean Lab	Christa Dahman Zaborske
	Radiochemistry	Jesse Wouters
	EHD Data Management/	Kathleen Dax-Klister
	Customer Service/Sample Recv,	
	Shipping	
Environmental Biology	Water Microbiology/ Glassware/Media	Martin Collins
	Environmental Toxicology	Dawn Perkins

1.4.4. QA Specialists/Coordinators

The EHD QA Manager, and QA Specialists/coordinators are responsible for implementing and maintaining quality assurance procedures throughout the laboratories, and ensuring compliance with applicable standards. They may be QA Specialists who only conduct QA tasks, or they may be staff that have their time split between QA and lab work or management tasks. These coordinators work with the supervisors, lab managers/directors, and division directors to ensure that QA procedures are followed by all staff. Responsibilities of the QA coordinators include oversight of the procedures that generate quality control data. QA coordinators are also responsible for managing internal audits according to a set schedule. The internal audit reports include a listing of deficiencies or action items and a means of monitoring corrective action. The QA coordinators oversee the lab section's certification status and coordinate the various regulatory programs. They also maintain working relationships with regulatory agencies and closely monitor any program or statutory changes. Other duties of QA coordinators include managing performance evaluation (proficiency testing) samples, coordinating the review and writing of SOPs (including the Quality Assurance Manual), and evaluating QC limits.

The QA coordinators report to the Lab Director or Supervisor, and have access to higher levels of management through the Quality Assurance Committee. Their QA duties are independent of any laboratory work that they may perform, or oversight is such that objectivity is maintained.

All QA coordinators have knowledge in the quality system as defined under TNI, DNR, & EPA and experience in the concomitant QA/QC procedures. Documentation of this knowledge and experience includes dated signatures (which may include electronic signatures or acknowledgement within the OnBase document management system) on the QA Manual, SOPs, DOC statements, and other documents that are part of the quality system. QA coordinators also have general knowledge of the analytical test methods performed in their sections.

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If a quality assurance question arises when the QA coordinator for a particular section is absent, the question may be directed to the program area director, section supervisor, or a QA coordinator from another section. In areas where there is not a QA coordinator, the supervisor takes the lead on QA issues with the help of other QA coordinators.

1.4.5. Laboratory Staff

It is the responsibility of the frontline laboratory staff (bench analysts and support/administrative staff) to produce high quality data within the structure of each individual method and within the parameters of the laboratory's quality control guidelines. It is also the responsibility of the staff to identify existing problems or inefficiencies, and to improve the processes of the laboratory whenever possible. Management should be informed by lab staff of any needs or concerns in a timely manner.

1.4.6. Position Descriptions

Specific position descriptions for all personnel are located in the main WSLH Human Resources Office (465 Henry Mall). In addition, each section supervisor can access copies of the position descriptions for their staff in the UW Online Performance Management system. Organizational charts for WSLH and all divisions are located at O:\Organizational Charts\Current Org Charts.

1.4.7. Education and Experience—Division-wide Employees

1.4./.	Education and Experience—Div	ision-wide Employees
Name	Title	Degree
Steve Strebel	Assoc. Director (interim) of Non-Clinical Testing	BS Chemistry
Camille Danielson	Lab Director (Envir. Chemistry)	BA Biology/Chemistry Minor, MS Aquatic Toxicology
Jocelyn Hemming	Lab Director (Envir. Biology)	BA Biology, PhD Environmental Toxicology
Susan Hill	QA Specialist	BS Chemistry, MS Water Chemistry
vacant	QA Manager	

1.5. Security and Access

Access to the WSLH Agriculture Drive location is restricted to authorized individuals to ensure the safety of all staff members and to maintain sample integrity. The exterior doors of the main entrance and the loading dock areas are open to our customers from 7:45 a.m. to 4:30 p.m., Monday through Friday. Saturday hours are 6:30 a.m. – 12:30 p.m. Both of these areas will be secured from the rest of the laboratory by electronic locks. Only staff members, custodial staff, and authorized visitors will have access beyond these two secured areas. All authorized visitors to the labs must be signed in and out of the building and wear a visitor badge. LABWIDE GENOP 1101, "Visitor Access to WSLH Facilities and LABWIDE GENOP 1004, "Building Access "Authorization" detail the specifics of security and access.

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1.6. Service to the Customer & Complaint Resolution

The WSLH will maintain communication with customers to ensure testing is completed as requested by the customer or according to agreement between the customer and the lab (also see section 1.15, Procedures for Accepting New Work/ Review of Requests, Tenders, and Contracts). Any questions, problems, or major delays with a sample will be communicated to the customer as appropriate. Documentation of customer communications will be maintained. When samples and test request/chain of custody forms have already been received at the lab, these communications are documented either on the paperwork (which will then be scanned and saved in the Lab Information System) or in LIMS (go to Clients/Phone Log). E-mail communications with clients will be saved according to the applicable RDA (1.12). Sampling kits or other supplies needed by customers will be sent out in a timely manner according to procedures. Questions will usually be answered by Customer Service representatives. Technical questions regarding specific tests can be passed on to sectional representatives.

Although the Laboratory strives to provide services in a timely and high-quality fashion, it is expected that we will occasionally make mistakes or fail to please a customer. When complaints occur, it is expected that the laboratory staff will handle them in a consistent, courteous, and prompt manner. EHD DIV-WIDE GENOP 017 "Managing Customer Feedback" details how complaints and other feedback should be handled and documented.

The EHD seeks feedback from clients in regular meetings with the Department of Natural Resources and the Department of Health Services. The division has also sent surveys to targeted customer groups.

1.7. Confidentiality Policy

We will handle oral or written requests for sample results according to the following policy: We will release sample results to the person(s) or entity that submitted the sample and/or is paying for the testing. We will also release sample results to the person(s) or agency that was identified as the receiver of the report.

Note: Under Wis. Stat. 280.13(1)(d) and Admin. Rule NR 812 private drinking water test results are released electronically to the WDNR.

If information is requested under the Freedom of Information Act, the request is forwarded to the WSLH Director's Office.

Most test results within the EHD are not considered protected health information under the Health Insurance Portability and Accountability Act (HIPAA).

Laboratory records that support accreditations will be made available to the applicable accreditation

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bodies. These include, but are not limited to quality records, technical records, information necessary for the historical reconstruction of data, administrative records, purchasing records, and personnel records.

Personal information about all staff of WSLH is considered confidential. Conversations regarding staffs' personal and professional information must be conducted in a site where confidentiality can be maintained.

1.8. Hiring Process

The WSLH, as part of the University of Wisconsin-Madison, must conform to the UW's hiring policies. These requirements are designed to ensure that the laboratory's hiring practices are fair and equitable and meet all Federal and State regulations. The WSLH Human Resources and UW Madison Office of Human Resources (OHR) are responsible for developing and maintaining policies, procedures, and documentation related to hiring WSLH personnel. WSLH HR Department is responsible for maintenance and final disposition of personnel records. For details on the hiring process, please see HR Website http://slhicmsprod/administrative-services/human-resources/

1.9. Safety

The "Chemical Hygiene Plan and General Laboratory Safety Plan" (LABWIDE SAFETY 102), contains comprehensive information on general laboratory safety procedures and operations for the WSLH's Agriculture Drive facility. Information includes chemical storage, waste disposal, safety showers, fume hoods, controlling exposure, employee safety training, housekeeping, emergency procedures, personal protective equipment, and more.

For details related to waste management, see EHD DIV-WIDE GENOP 038. This division-level SOP includes a strategy for managing waste streams and promoting techniques such as waste reduction, re-use, recycling, and recovery to protect human health and the environment. It also includes instructions for managing, storing, and disposing of biohazardous, radioactive, and chemical wastes.

At the University level, see UW-Madison policy UW-6066, Chemical Hygiene Plan and Policy: https://policy.wisc.edu/library/UW-6066. In addition, refer to the web-based "Chemical Safety Guide," https://ehs.wisc.edu/labs-research/chemical-safety/chemical-safety-guide/, which is a collection of chemical safety information that can be used to improve laboratory operations, develop standard operating procedures, and inform staff of the latest safety and regulatory requirements.

The WSLH has a Safety committee that meets monthly and conducts safety inspections. Minutes of this committee are available at O:\Teams\Safety\Minutes and Attendance. Membership of this committee consists of a cross section of laboratory personnel.

1.9.1. Note regarding Safety Data Sheets

A Safety Data Sheet (SDS) is a document containing chemical hazard and safe handling information.

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There are 16 sections including: Section 2: Hazard identification, Section 4: First aid measures, Section 7: Handling and storage. For more information about Safety Data Sheets, see the Chemical Hygiene Plan and the Chemical Safety Guide noted above.

1.10. Training

1.10.1. Initial Training of New Employees

Training of employees takes place in a logical progression that meets applicable requirements. The Office of Human Resources has an employee onboarding checklist (http://slhicmsprod/administrative-services/human-resources/) that includes some lab-wide training items (click on the New Employee tab)

WSLH has information of general interest to employees available through the intranet home page: http://slhicmsprod/ (click on the Team Dynamix button, and then click on *Knowledge Base* in the top ribbon.). The Knowledge Base contains informational articles that are organized into categories, which include: hardware, software, media, general, forms, and WSLH-wide information. The WSLH-wide category is further divided into Administrative Policies and Procedures, Human Resources, and Finance Policies and Procedures. Use the "Client Portal" Search bar to find information.

To ensure the safety and well-being of all WSLH personnel, new employees must become familiar with basic safety precautions before working in the laboratory. A key tool in safety training is the Agriculture Drive Employee Safety Checklist (LABWIDE SAFETY 100), which comprehensively lists safety issues such as the location of safety showers and fire extinguishers, evacuation procedures, policies on eating and drinking in the lab, use of potentially dangerous instruments and chemicals, safety apparel use, fume hood use, and much more. The AD Employee Safety Checklist in OnBase also lists external references that contain more information on laboratory safety. Other safety training tools that new employees are required to review are the AG Drive Chemical Hygiene Plan (LABWIDE SAFETY 102) and the Emergency Response Plan for Agriculture Drive (LABWIDE SAFETY 101). All employees are required to take fire extinguisher training offered by the university. Depending on their position, some employees may require more specialized instruction such as blood borne pathogens or radiation safety training offered by the university. All employees have potential exposure to animal or human related samples (i.e., serum, urine, wastewater, biosolids, animal tissue etc.), and therefore must take blood borne pathogen training, which is available through Canvas (Biosafety 102: Bloodborne Pathogens for Laboratory and Research) EHD Data management, Sample Receiving, Customer Service and Shipping staff take Bloodborne Pathogens for Operations in Canvas https://canvas.wisc.edu/enroll/7T9HEF. In addition, they must review LABWIDE SAFETY 302, OSHA Bloodborne Pathogens Exposure Control Plan. EHD Shipping staff also complete Chemical Safety: Hazard Communication - Identifying Chemical Hazards, Chemical Safety: Cryogen Safety Training (only part 1) and Chemical Safety for Non-Lab Personnel all in Canvas.

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New employee training continues with:

- Data Integrity training (EHD DIV-WIDE GENOP 029) and the WSLH Statement of Ethical Expectations (WSLH-WIDE POLICY 001, which is available through employees' My UW accounts in the Canvas app).
- A review of the EHD Quality Assurance Manual (General and applicable section supplements).
- Non-Conformance management training (also see section 1.21.5). The non-conformance management training is available in OnBase as SOPs: LABWIDE GENOP 707 Non-Conforming Event Management System Policy and LABWIDE GENOP 711 Non-Conformance Management Procedure for MediaLab. "Owners" in the system have more advanced training provided in special sessions.
- LABWIDE Safety 307 Personal Electronic Device Usage in Lab Spaces
- HIPAA—Varying levels of HIPAA Privacy Rule training are required depending on an employee's position. For HIPAA training information, consult HR and see the University of Wisconsin—Madison, Office of Compliance, HIPAA training website:

 https://compliance.wisc.edu/hipaa/training/.
- Review of the *QA & You Brown Bag* video. Submit an attestation statement to Human Resources. The video and attestation statement are located on our intranet:

 http://slhicmsprod/regulatory-compliance/quality-management/.
 - Once there, select the *Quality Tools* tab (began 8/2020).
- OnBase document management platform training is available through employees' My UW accounts.
 - o Access enrollment via: https://canvas.wisc.edu/enroll/68JDHH.
 - o Enroll in the Introduction to OnBase module in the Canvas app.
 - o Complete the training and quiz as instructed.
 - Staff that will be involved with editing or approving SOPs will also need to take the more advanced OnBase Workflow training provided within the same Canvas course.
 - For further information go to the WSLH TeamDynamix Service Desk Knowledge Base (access via intranet).
 - Records Training is available through employees' My UW accounts.
 - O Division directors or section supervisors decide which of their staff will take the level 1 training and who will continue with the level 2 training.
 - o Instructions for self-enrollment in the Canvas Course, WSLH Records Training is available through WSLH TeamDynamix Service Desk Knowledge Base (access via intranet).

Each section has a new employee training form or checklist (see section supplements), which ensures that a new employee receives information important to working in that section. Usually, an experienced

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employee guides the new employee through the checklist.

These items complete the initial training of new employees. Next, they will move on to analytical method training if required for their position. Other non-testing training materials may be required by the sections.

1.10.2. Analytical Method Training

Analytical method training is for new employees who have completed the initial training and for any employee who is learning a new procedure. Sections may have method training forms (see section supplements), which help guide a person through the process of learning a new method. Generally, the trainee will review the section's SOP and/or the regulatory method as well as the instrument manual. He/she will observe an experienced analyst prepare samples and operate the instrument. He/she will next work under the direct supervision of the experienced analyst until becoming familiar with the analytical procedures. Training includes sample handling and preparation, safety specific to the method, documentation procedures, calibration procedures, QC requirements, data management, data reporting, and troubleshooting.

If applicable, the trainee will perform an initial Demonstration of Capability (DOC) and document the results on a DOC Certification Statement (EHD DIV-WIDE QA 115, "Initial and Ongoing DOC Procedures"). In addition, the trainee must sign a Certification Statement for Current Technical Methods (Figure 1) or an attestation statement at the end of an SOP, which states that he/she has read, understood, and agreed to perform the most recent version of the test method. Electronic forms and signatures are acceptable. When initial DOC criteria have been satisfied and the experienced analyst and supervisor are confident that the employee is thoroughly familiar with the test, that employee is allowed to work on his/her own with only routine supervision.

1.10.3. Continuing Training

All employees receive continuing training and must demonstrate continued proficiency. Whenever there is a change in instrument type, personnel, or test method a new DOC must be performed. Annually, each analyst must demonstrate continued proficiency on technical methods for which they are responsible (see section supplements for procedures). When a new revision of an SOP is written, analysts who are responsible for that method must sign a new Certification Statement for Current Technical Methods.

All employees must annually review the following documents:

- a. Quality Assurance Manual (General and applicable section supplements)
- b. Chemical Hygiene Plan
- c. Data Integrity, Ethics, and Data Documentation Procedure
- d. Emergency Response Plan
- e. OSHA Bloodborne Pathogens Exposure Control Plan
- f. WSLH Employee Safety Checklist

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1.10.4. Additional Education/Training

- Occasionally, the university and the WSLH offer or require training (e.g. sexual harassment awareness and prevention, HIPAA, disability accommodation, Cybersecurity Awareness, and others). E-mails will detail requirements.
- The laboratory supports continuing education that may include seminars, training offered by vendors, or formal higher education. All employees are encouraged to keep up with changes or advances in analytical methods and instrumentation.

1.10.5. Training Documentation

All training forms, checklists, sign-off sheets, certification statements, and DOC forms related to the above requirements will be signed and dated by the employee and managed by the QA coordinator responsible for that section or documented according to university requirements (e.g., documented within the Canvas learning management system). Electronic forms and signatures may be used see 1.11.3 for signature policy. The QA coordinator will ensure that the documentation is complete and filed appropriately for training within the section. Most training documentation will be filed in the personnel training files in each section or into electronic folders.

1.10.6. Data Integrity Procedures

All employees have access (via OnBase) to the Data Integrity, Ethics, Data Documentation Procedure for the WSLH, Environmental Health Division (EHD DIV-WIDE GENOP 029). The procedure includes our organizational mission relating to data integrity, our steps for data integrity training and training documentation, our methods for monitoring data integrity, and steps for the reporting of data integrity concerns. In addition, it documents our Ethics Policy and our Data Documentation procedure. The Association of Public Health Labs (APHL) has additional ethics training resources that can be accessed online at no cost: https://learn.aphl.org/learn.

1.10.7. Quality Assurance Training

All employees must watch the *QA & You Brown Bag* video once and submit an attestation statement to their supervisor, who in turn will send it to Human Resources. HR will file the statements in employees' P-files. The video and attestation statement are located on our intranet:

http://slhicmsprod/regulatory-compliance/quality-management/.

Once there, select the Quality Tools tab.

1.10.8. Employee Training Summary Tables:

Initial Training of New Employees

Training Document	Link or Reference to	Documentation* of Training
	Document	
Lab-wide new employee checklist	http://slhicmsprod/administrative	Sign checklist & complete
	-services/human-resources/	referenced documentation

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Dept. new employee checklist	See section supplements	Sign checklist
EHD Employee Safety Checklist	Access in OnBase	Sign checklist
(LABWIDE SAFETY 100)		
Chemical Hygiene Plan for Ag Dr	Access in OnBase	My UW account-Canvas app
(LABWIDE SAFETY 102)		or other attestation
Emergency Response Plan for Ag	Access in OnBase	My UW account-Canvas app
Dr (LABWIDE SAFETY 101)		or other attestation
EHD Data Integrity Procedure	Access in OnBase	My UW account-Canvas app
(EHD DIV-WIDE GENOP 029)		or other attestation
Statement of Ethical Expectations	Access in OnBase	My UW account-Canvas app
(WSLH-WIDE POLICY 001)		or other attestation
QA Manual (General/ applicable	Access in OnBase	My UW account-Canvas app
section supplement)		or other attestation
Non-Conformance Management	Access in OnBase	
(LABWIDE GENOPs 707, 711)		
Bloodborne Pathogens	Access in OnBase	My UW account-Canvas app
(LABWIDE SAFETY 302)		or other attestation
HIPAA	https://compliance.wisc.edu/hipa	Through UW website
	<u>a/training/</u>	
QA & You video	http://slhicmsprod/regulatory-	Attestation statement
	compliance/quality-management/	
OnBase Document Management	https://canvas.wisc.edu/enroll/68	My UW account-Canvas app
	<u>JDHH</u>	or other attestation
Biosafety 102: Bloodborne	https://ehs.wisc.edu/training/b	My UW account-Canvas app
Pathogens for Laboratory and	iosafety-102-bloodborne-	or other attestation
Research	pathogens-for-laboratory-and-	
	research/	
WSLH Records Training	https://canvas.wisc.edu/enroll/JY	My UW account-Canvas app
	DJPE	or other attestation

^{*}may be electronic

Analytical Method Training

	Link or Reference to	Documentation of Training
	Document	
Method Training form	See section supplements	Sign form
DOC Certification Statement	EHD DIV-WIDE QA 115	Sign DOC statement
Cert. Statement for Current	QAM-General Figure 1 (Note:	Sign Certification statement
Technical Methods (METHOD	other forms such as SOP Attest	(may be electronic)
SOPs)	Statement are also acceptable)	

Continuing Training

Training Document	Link or Reference to Document	Documentation of Training
DOC or Annual Continued	EHD DIV-WIDE QA 115	Sign DOC statement or other
Proficiency documentation	(OnBase)	doc.
(annually or when there is a		
change)		

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Cert. Statement for Current	QAM-General Figure 1	Sign Certification statement*
Technical Methods (when new SOP	(Note: other forms are also	
revision is written)	acceptable)	
QA Manual, General &	OnBase	My UW account, Canvas app
Supplements (annually)		or other attestation
EHD Chemical Hygiene Plan	LABWIDE_SAFETY 102	My UW account, Canvas app
(annually)	(OnBase)	or other attestation
EHD Data Integrity Procedure	EHD_DIV-WIDE GENOP 029	My UW account, Canvas app
(annually)	(OnBase)	or other attestation
EHD Emergency Response Plan	LABWIDE SAFETY 101	My UW account, Canvas app
(annually)	(OnBase)	or other attestation
Bloodborne Pathogens	LABWIDE SAFETY 302	My UW account-Canvas app
(annually)	(OnBase)	or other attestation
WSLH Employee Safety Checklist	LABWIDE SAFETY 300	My UW account-Canvas app
		or other attestation

^{*}may be electronic

Additional Education & Training

Training Document	Link or Reference to Document	Documentation of Training
Course or class work		Certificate or note about title
		and content of course
HIPAA	https://compliance.wisc.edu/hipa	Through UW website
	<u>a/training/</u>	
QA & You video	http://slhicmsprod/regulatory-	Attestation statement
	compliance/quality-management/	
Sexual Harassment Awareness &	https://compliance.wisc.edu/titlei	Through UW website
Prevention	x/employee-training/	
Cybersecurity Awareness	https://it.wisc.edu/about/division	Completion of two badges on
	-of-information-	training site
	technology/enterprise-	
	information-security-	
	services/office-of-	
	cybersecurity/cybersecurity-	
	<u>awareness-training/</u>	
Disability Accommodation	Reading materials provided by	Quiz provided by HR
Training	HR via e-mail	
Bloodborne Pathogens	https://ehs.wisc.edu/training/bios	My UW account-Canvas app
(Biosafety 102, Bloodborne	afety-102-bloodborne-pathogens-	
Pathogens for Lab & Research	for-laboratory-and-research/	

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Figure 1: Example of a Certification Statement for Current Technical Methods

Analyst Certification Statement For Current Technical Methods Analyst Name: SOP Number: SOP Title: SOP Revision Number: SOP Effective Dates: I, the undersigned, CERTIFY that I have read, understood, and agreed to perform the most recent version of the above test method.	Analyst Cert. Statement Rev. 2 April, 2015 Page 1 of 1	Wisconsin State Laboratory of Hygiene Environmental Health Division
SOP Number: SOP Title: SOP Revision Number: SOP Effective Dates: I, the undersigned, CERTIFY that I have read, understood, and agreed to perform the most recent version of the above test method.		
SOP Number: SOP Title: SOP Revision Number: SOP Effective Dates: I, the undersigned, CERTIFY that I have read, understood, and agreed to perform the most recent version of the above test method.	Analyst Name:	
SOP Revision Number: SOP Effective Dates: I, the undersigned, CERTIFY that I have read, understood, and agreed to perform the most recent version of the above test method.		
I, the undersigned, CERTIFY that I have read, understood, and agreed to perform the most recent version of the above test method.		
I, the undersigned, CERTIFY that I have read, understood, and agreed to perform the most recent version of the above test method.	SOP Revision Number:	
version of the above test method.	SOP Effective Dates:	
Analyst Signature Date	I, the undersigned, CERTIFY that version of the above test method	at I have read, understood, and agreed to perform the most recent
	Analyst Signature	Date

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1.11. Document Control System

1.11.1. Control & Maintenance of Standard Operating Procedures (SOPs)

SOPs that may apply to operations covered by this QA Manual exist at three different levels: Lab-wide, EHD Division-wide, and section-level.

SOPs will be stored, maintained, and accessed through OnBase, an electronic document management platform. OnBase has search and retrieval functions as well as customized workflow for editing and version tracking. Within OnBase, documents are organized by category, or groups of document types that have similar characteristics, purpose, or management requirements. Search and retrieval functions are available.

There are two ways of accessing OnBase: Web Client and Unity Client. Web Client is available via the intranet home page for general access to locate and view approved SOPs (all SOPs are accessible to all staff unless specially restricted). Unity Client must be used for operational, permission-based functions within workflow such as editing or approving documents. Unity Client must be installed on individual workstations, if needed. The history of any particular document within OnBase workflow, including version numbers, effective dates, and who approved each version, is saved within the system.

Within OnBase, approved SOP versions are found under POLICIES-PLANS-PROCEDURES (PPP), SOP-Approved document type.

- Lab-wide: Division=LABWIDE
- EHD Division-wide: Division=EHD, Section=DIVISION-WIDE
- Section-level: search by appropriate division and section

Within OnBase Workflow, various lifecycles (permission-based) exist for updating SOPs:

- SOP Review/Approval LABWIDE
- SOP Review/Approval EHD DIVISION-WIDE
- Each section has a separate lifecycle

For more information regarding the use of OnBase, see LABWIDE GENOP 700, OnBase User Guide. Training is available (see section 1.10 of this document).

Analytical method SOPs will contain all accreditation-required sections and information as applicable.

All SOPs associated with drinking water testing (including administrative as well as technical) must be reviewed for content annually and updated if necessary.

This QA Manual is reviewed at least annually, and it is updated when necessary. OnBase includes a Timers

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and Notifications module, which tracks review periods, due dates, and automatically sends notifications to assigned people.

1.11.2. Components of Document Control

All internally-generated documents (SOPs, policy statements, spreadsheets, forms, bench record cover sheets, certification statements, sign-off sheets, instruction sheets, etc.) must be listed in a table of contents (or equivalent list generated electronically or in OnBase) and have document control [TNI V1M2 4.3.2.3; WI NR 149.39(1)] including:

- a. Issuing authority (e.g., WSLH, division, section)
- b. Unique ID (number or title)
- c. Effective date and/or revision number
- d. Page x of y

1.11.3. Electronic Signature Policy

Electronic signatures through cloud-based services such as DocuSign are acceptable in place of handwritten signatures on hard copy documents. The electronic signatures are unique, traceable, and secure. Within the quality system, staff and supervisor signatures are most often needed on DOCs and other training documents. Documents with electronic signatures must be saved to shared drives (or OnBase) where they are backed up and retained according to the applicable RDA. It is not acceptable to save the electronically signed documents only within the cloud-based signatory service.

1.12. Records Retention, Control, & Storage

1.12.1. Records Disposition Authorization (RDA)

RDAs are also called records schedules. RDAs are key documents for establishing a records management program for organizations within Wisconsin State government. In essence, records schedules describe the organization's information resources, how long they are going to be retained, and what their ultimate disposition will be. They are the policy statements that govern the ultimate disposition of records.

Some EHD records fall under General Records Retention Schedules (GRS), which are approved for UW-Madison campus-wide use. General records schedules codify retention policies for record types that are common to all offices across the UW system. WSLH also has some UDDS/Department-specific RDAs. Refer to https://www.library.wisc.edu/archives/records-management/retention-disposition/

If an RDA needs to be amended or resubmitted upon 10-year sunset, contact:

University Records Officer, University of Wisconsin-Madison

Phone: 608-262-3284

recmgmt@library.wisc.edu

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1.12.2. Laboratory Notebook/Logbook

A laboratory notebook or logbook is used to record testing data, which could include experimental data, standard and reagent logs, instrument logs, etc. Logbooks can be physical or electronic (e.g. LabArchive).

Physical laboratory logbooks are assigned a unique number by QA staff. This number is then used for tracking logbooks in the laboratory. Each logbook will be labeled with the following information: organization name, unique laboratory logbook number, analyst or instrument assigned to, section, logbook's contents, start date and end date.

All logbook information is entered into a database, which also includes information on archival date, storage location, archival box number and disposal date. The database is on drive: R:/EHD/QC/Archived Records.mdb

For more information about the logbook identification labels, see EHD DIV-WIDE GENOP 104, How to Print Laboratory Logbook Labels.

Notebooks and logbooks will be retained according to the applicable RDA.

1.12.3. Storage of Records in Basement room 14

Refer to LABWIDE GENOP 1002, "Records Storage and Disposal," section 4.0 for the procedure for storing documents in record storage boxes. Boxes may be immediately sent to the State Records Center or may be placed in an approved storage area at the WSLH. Approved storage areas are organized with specific locations where new inventory may be added. The approved storage area for Agriculture Drive is room 14 in the basement. Complete records of items placed in storage must be maintained by the divisional or sectional records coordinator. The records coordinator has the responsibility for knowing what is in each box, where each box is located, and the destruction date of each box.

1.12.4. Sending Records to the State Records Center

LABWIDE GENOP 1002, contains information regarding how to use the State Records Center (SRC) Versatile Enterprise Web Module, which is an on-line, web-based process that allows users to submit new inventory (e.g., record storage boxes) to the SRC and to search for and request return of boxes stored there.

1.12.5. Disposal of Records

The SRC can also be used for disposal of records, even when they are not stored at SRC. Refer to LABWIDE GENOP 1002. If the SRC is not used, ensure that documents under RDAs that are confidential are shredded when disposed of.

1.12.6. Other Records Information

LABWIDE GENOP 1002 also details the procedure for electronic records.

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1.13. Traceability of Measurements

All analytical results and measurements are fully traceable to standards, reagents, reference materials, and instrumentation used in deriving the results. Stock and working standards are given codes that are documented in the analytical runs in which they are used. Reagents (including pH paper and other chemical test strips) are also given traceability codes. Standards along with effective and expiration dates are retained in the LIMS (or on the analytical run) and linked to the batches and samples associated with them. Analytical instrumentation and equipment (including filters, pipets, and thermometers) are assigned identification numbers, which are also tracked at the batch level in analytical runs.

Date and time of analysis and analyst initials are documented at the batch and/or sample level with all functions in the LIMS being marked with date/time/user's initials. Test results are entered into LIMS either automatically from the instrument or through manual entry by the analyst. All results are directly linked to each sample.

Individual sections may have more detailed information on how they achieve traceability.

1.14. Policy for Estimating Uncertainty of Measurement

Each section must write a policy or procedure for estimating uncertainty of measurement. The policies will cover all analytical tests performed by the section. The uncertainty needs to be reported only if required by the regulating agency, but it must be available upon request. The policies must attempt to identify all the components of uncertainty, make a reasonable estimation, and ensure that the form of reporting of the result does not give a wrong impression of the uncertainty. Reasonable estimation will be based on knowledge of the performance of the method, on the measurement scope, and will make use of previous experience and validation data.

If the section analyzes samples following a well-recognized test method, which specifies limits to the values of the major sources of uncertainty and also specifies how calculated results will be reported, then the section meets the uncertainty requirements for that method.

1.15. Procedures for Accepting New Work/Review of Requests, Tenders, & Contracts

1.15.1. Requests for service

General water test information for the public can be found on the extranet:

http://www.slh.wisc.edu/environmental/water/

To offer a new test to the public on a fee-for-service basis, consult with the Office of the Director. Contracts and requests for service are reviewed before approval, and if a change in the scope of a contract is requested, a contract modification may be required. The applicable WSLH unit will review the new contract to ensure they have the capability to provide the services. In general, the DocuSign process will

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be used to document this unit review. Copies of WSLH executed contracts are maintained and available by contacting the Contract and Grant Administrator Specialist in the WSLH Office of Finance.

WSLH also offers fee-for-service tests to public water suppliers. Tests requests are made on test request forms that are sent with samples.

1.15.2. Test requests for non-standard analysis:

- Use the template found in the following folder: O:\Quotes\EHD\COC Template
- Update the form for the specific client.
- Update the room number and contact information on the address under the Wisconsin emblem.
- Update the sample description and analysis fields to meet the needs of the client.
- If samples will be logged into the LIMS, include the acode number in the analysis field and identify the LIMS account in the section for notes.
- Delete unnecessary rows for small submissions to limit the number of pages to be printed.
- Save a copy of the form in the file directory of your choice. A naming convention similar to the quotes is advised (Organization Contact Date).
 - 1.15.3. Review of laboratory capability

If new work is to be accepted, a review of laboratory capability and the scope of work is conducted as part of the unit review. The following items may be considered:

- Availability of desired analytical methods and accreditations
- Adequate capacity for facilities, supplies, and instrumentation
- Adequate staffing, training, and experience (both analytical and support staff)
- Capability to meet desired LODs and LOQs
- QA/QC that meets data quality objectives
- Desired turn-around-time
- Desired reporting capabilities and information technology resources
- Appropriate funding to complete the project
- Desired records retention period
 - 1.15.4. Approved Signatory

Contact the Office of Finance for any agreements or purchase orders from an outside entity that require a signature from WSLH. The WSLH Director is authorized by the University of Wisconsin-Madison to sign contracts that create a binding legal or financial obligation upon the university. For more detailed information, please refer to the Signature Authority Memo linked at https://legal.wisc.edu/contract-approval/. Contracts follow UW policies and procedures.

1.15.5. Review of requests for service

Routinely, the lab receives samples or requests from the public and WDNR staff who unsure what they

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want the samples to be tested for. In other cases, the client has asked for tests that may not be ideal for their samples. In these cases, WSLH staff work with the client to determine the best tests that will meet their objectives. When samples and test request/chain of custody forms have already been received at the lab, these communications are documented on the paperwork (which will then be scanned and saved in the LIMS) or in the LIMS. E-mail communications with clients may also be attached to the sample in the LIMS. Each contract (written or oral agreement to provide a customer with testing services) will be acceptable to the lab and the customer; any differences will be resolved prior to work commencing on the samples.

Records will be maintained of reviews of requests, tenders, and pertinent discussions with a customer relating to the customer's requirements, the results of work, any deviation from the accepted contract, and any significant changes to the work being done. If a contract needs to be amended after work begins, the review process noted above will be repeated and all changes communicated to relevant personnel. These records may be in the form of comments appended to test results in the LIMS, DocuSign forms, e-mails, notes from conversations, or written correspondence.

This process also applies to sample testing that may need to be sub-contracted (see section 1.23). The complete history of client and staff agreement regarding sample testing will be traceable through the LIMS and associated documentation. See also section 1.6, Service to the Customer and Complaint Resolution.

Additional details regarding accepting new work may be found in the section supplements of the Quality Assurance Manual.

1.16. Sample Handling and Submission Procedures

1.16.1. Sample Collection

DNR staff who collect samples follow instructions that are posted on their internal website. The lab works with the DNR/Lab Liaison to ensure the instructions are maintained up-to-date. The DNR has training available for staff who collect samples. Sample collection training includes information about the proper supplies to use for collection, how to safely collect the sample, guidance on what parameters should be requested for testing, how to preserve the samples, how to complete and submit required paperwork, and how to ship or deliver samples to the lab.

Special considerations for the collection of enforcement samples include communication and coordination with the lab, chain of custody procedures, and records retention information.

Private homeowners and others who collect samples follow instructions received with their sampling kits.

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1.16.2. Sample Acceptance Policy

Samples arrive at the laboratory in several ways: they may be mailed, shipped by a commercial carrier, brought to the laboratory by client field representatives, or brought to the front desk of the laboratory by the general public. Environmental Toxicology samples may be collected and brought to the lab by WSLH personnel. Each sample received is accompanied by a test request form (TRF). The TRFs, chain of custody forms, and any other documentation received with the sample are retained according to section 1.12. Please see EHD DIV-WIDE GENOP 033, "Sample Acceptance Policy," which describes the policy used by the EHD Receiving Section for accepting samples, into the lab for environmental testing. This SOP also includes links to sectional SOPs that describe specific acceptance and rejection policies for each section.

1.16.3. EHD LIMS

Samples logged into the LIMS receive a unique sample number as described below. Other attributes of the sample documented in the LIMS at the time of sample log-in are preservatives, container type, client/project name, date/time of receipt at the lab, field ID, collection date/time, tests requested, and sample condition at receipt. Durable labels, with unique workorder/letter identifiers and barcodes, are printed for each sample container and test request form. Two labels are generated for each sample container—one is placed on the body of the container, and another one is placed on the lid of the container (if desired for convenience).

Samples are received and noted in the LIMS as being located in the sample receiving section. As samples move through the lab, the location of the sample is changed in the LIMS. Disposal of samples is recorded in the LIMS by the lab section responsible.

1.16.4. Sample ID Generation in the LIMS

Summary

The workorder ID is generated from a database sequencer and is always unique and chronological.

The sample ID is derived from the workorder ID if the sample is part of a workorder, or it is generated from a database sequencer if the sample is an internal QC sample. In both situations the sample ID is unique. The container ID is derived from the sample ID with a letter suffix for each container in the sample, -A, -B, -C, etc. and is always unique. Sub-sample (extracts, aliquots, etc.) IDs are derived from the container ID followed by a number, -1, -2, -3, etc.

Workorder ID Generation

A workorder is a collection of samples that all share various things in common, like customer, due date, collection time, etc. Each workorder is assigned a unique number in the database field PROJECT_SEQ. The system installation setting PSEQASSIGN controls how the PROJECT_SEQ is assigned to the workorder. The laboratory uses the NEXTVAL setting, which assigns each workorder a unique number

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generated by an Oracle database sequencer and is always unique. Each workorder also has a laboratory identifier in the database field LAB_WO_ID. This ID can be the same as the PROJECT_SEQ or formatted differently, and is controlled by the system installation setting named WO_ASSIGN. The laboratory uses the SYSTEM setting, which sets the LAB_WO_ID equal to the PROJECT_SEQ.

Sample ID Generation

Each sample is assigned a unique number in the database field HSN. The system installation setting HSN_ASSIGN controls how the HSN is assigned to a sample. The laboratory uses the NEXTVAL setting, which assigns each sample a unique number generated by an Oracle database sequencer and is always unique. If a sample is part of a workorder (i.e., not an internal QC sample) the HSN number is derived from the PROJECT_SEQ, using three digits to indicate the number of samples within the workorder (i.e. 001, 002, 003, etc.). This creates a unique number for each sample in a workorder. Each sample also has a laboratory identifier in the database field LAB_SAMPLE_ID. This ID can be the same as the HSN or formatted differently, and is controlled by the system installation setting named LAB_ASSIGN. The laboratory uses the SYSTEM setting, which sets the LAB_SAMPLE_ID equal to the HSN.

Container ID Generation

Each container in the LIMS is assigned a unique number in the database field CONTAINER_SEQ. This number is assigned using an Oracle database sequencer, and is always unique. Each container also has a container ID in the database field CONTAINER_ID. The container ID is derived from the LAB_SAMPLE_ID, and is controlled using the system installation setting CONTNAME. The laboratory uses the setting ALPHABETIC, which results in a –A, -B, -C, etc. suffix to the LAB_SAMPLE_ID.

Documentation Access

Requisition forms that are submitted with samples are scanned into the LIMS, associated with the samples via the workorder and are available for viewing. Instrument data that is transferred into the LIMS via the LIMS Data Exchange (the results upload feature) is captured, associated with each sample in the run and available for viewing.

1.16.5. Labeling of Sub-samples

After initial sample log-in, all sample aliquots or sub-samples must be labeled or otherwise identified to ensure that there can be no confusion regarding the identity of such samples at any time.

All digestion or other intermediate processing step tubes, containers, or vessels must have an ID number and associated documentation that will give a one-to-one correspondence between this ID number and the unique sample ID number generated by the LIMS. Examples include etching an ID number onto a digestion tube, creating a LIMS aliquot label to place on a container, and using a permanent marker to

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label a drying pan. There would then be a bench sheet, run log, logbook, or work-list that documents the aliquot ID number and the corresponding LIMS number.

All autosampler vessels will be labeled as above or alternately, the autosampler tray will have ID numbers at each position. These ID numbers will be recorded, and a one-to-one correspondence will be established between this number and the unique sample ID number generated by the LIMS.

1.17. Instrumentation, Equipment, & Facilities

1.17.1. General

The EHD relies heavily on instrumentation. It is imperative that all equipment and instruments are calibrated, verified, operated, and maintained in a proper manner in order to obtain reliable data.

The section supplements of the QA Manual contain lists or references to information about support equipment and analytical instruments used in that section. These lists or references include type of instrument, make and model, type of analysis conducted, where the instrument/equipment is located, and assigned instrument numbers or serial numbers. Calibration, maintenance, and verification procedures are found in the appropriate instrument operation manual, instrument operating procedure (IOP), or standard operating procedure (SOP).

If an instrument fails to operate within defined limits or specifications, the problem is identified, corrective action performed, and samples re-run or qualified as required by the method. It is the responsibility of the analyst to notify the section supervisor if non-routine maintenance is required. Instrument vendor technical staff may be contacted if trouble-shooting is unsuccessful.

Preventive maintenance (general maintenance) should be performed at the frequency recommended by the manufacturer to avoid instrument failure. Most instruments and equipment are on some form of preventive maintenance schedule. Some instruments are covered by maintenance contracts directly with the instrument manufacturer. Preventive maintenance is always recorded in the instrument log (physical logbook or electronic). If there is no recommended preventative maintenance schedule, one should be created.

If equipment is sent out of the lab, its calibration and function will be verified when it is returned to the lab before the equipment is returned to service. The exception to this is when equipment (e.g., motorized pipets) is sent out for calibration. If the equipment is returned to the lab with a certificate of calibration, this certificate will be reviewed, and if acceptable, the instrument will not need independent verification in the lab.

Maintenance and repair (routine or non-routine) performed by other than WSLH staff needs to be summarized in a document provided to WSLH. This document must detail the work performed, which

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should include as found and as left information. If software updates are required, OIS should be notified in advance to determine if a copy of the computer (ghost image) should be taken prior to the new software being installed.

1.17.2. Laboratory Reagent Grade Water

Reverse Osmosis (RO) Water

RO water is plumbed to all the laboratories and is maintained by a service contract with a water treatment vendor. The RO system consists of a pre-filter of graded density non-woven polypropylene for the carbon bed, a filter cartridge of non-woven polypropylene for the 3 - RO/DI 5µm resin traps (replaced every 6 months), a UV Sterilizer in which the Aqua Fine replacement lamp and the Aqua Fine Quartz Sleeve will be replaced annually, a submicron Absolute Rated 0.2 µm bacteria eliminating filter (replaced every 6 months), a recirculating tank vent filter (replaced annually), and cation, anion and mixed bed tanks that will be replaced as needed. The water is ASTM Type II and is used for the glass washing activities, filling water baths, and as a precursor for other "polished" water throughout the building.

ASTM Type I Polisher

These polishers are located throughout the laboratory and provide ASTM Type I water used in the preparation of media, reagents, standards and QC samples (i.e. method blanks) as applicable. There is a service contract with a water treatment vendor to maintain the polishers. The vendor will exchange the carbon filter, the mixed bed cartridges and the organic scavenging Type II ultra-pure anion resin on their contracted schedule. The UV lamp will be changed based on hours of use. Please see EHD DIV-WIDE GENOP 032 "Monitoring and Maintaining Water Purification Systems" for specific water purification system monitoring and maintenance procedures.

1.17.3. Ovens, Incubators, Cold Rooms, Refrigerators, Freezers, and Muffle Furnaces Please see EHD DIV-WIDE GENOP 013 "Responding to Freezer/Refrigerator/Incubator Failures," which describes the procedures that shall be used at the Agriculture Drive facility when constant-temperature storage equipment (e.g., freezers, refrigerators, incubators) fails. The procedure describes the processes that shall be followed to keep samples within the required temperature range after a failure and the steps necessary to obtain service for the repair of constant-temperature storage equipment.

The temperatures of the walk-in refrigerators/incubators are monitored continuously by DOA and a 3rd party vendor, and an alarm will be activated if limits are exceeded. Each section has procedures for monitoring and documenting temperatures and any deviations according to requirements.

1.17.4. Computers

The WSLH Office of Information Systems (OIS) maintains a number of quality objectives to insure the integrity, security and reliability of our systems.

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The OIS quality objectives include:

- A state-of-the-art data center with exceptional security, redundant connectivity and a climate-controlled environment that meets the following certifications:
 - HIPAA/HITECH
 - PCI DSS v3.2.1
 - ISO/IEC 27001:2013
 - EU-U.S. EU Privacy Shield Framework
- Whenever possible, independent test systems are created alongside production to allow a safe environment for training as well as a place where changes can be tested prior to being put into production.
- The use of ITILv4 (Information Technology Infrastructure Library) and risk management frameworks to provide continual process improvement and apply best practices such as Change Management.
- Utilizing a risk-based approach to backup and recovery planning, taking advantage of multiple recovery options when it makes sense to do so (e.g. point-in-time recovery for databases; shadow copies at set intervals during the day) to improve backup and disaster recovery efforts. Backups to tape media utilize a grandfather-father-son rotation scheme and are kept for a corresponding timeframe of weeks, months, or years. These tapes are stored in "media proof safes" that are UL-125 rated (will not exceed 125 degrees Fahrenheit).
- Multiple, redundant network paths for our servers and workstations employing a secure IPSEC tunnel over a fiber-optic ring owned by the "Metropolitan Unified Fiber Network (MUFN) Consortium" https://mufn.org/ (of which we are a voting member) back to the UW-Madison campus and protected behind firewalls. The UW-Madison is our Internet service provider. Secured VPN service with Remote Desktop Connections (RDC) is used for remote access to enterprise resources.
- Policies, standards, and procedures governing user account creation, system access, personal information protection, and the appropriate use of information systems and resources including the use of HIPAA's "minimum necessary standard" whereby users are only granted the minimum access necessary to do their jobs.
- Inventory and asset management including anti-malware/antivirus, vulnerability scans, and routine patching for operating system and 3rd party patches (e.g., all servers are patched within 30 days of critical/security operating system patch releases).
- When computers, drives, and media are no longer useful or usable they are sanitized and/or disposed of in a manner consistent with UW-Madison HIPAA policy.
- See the OIS GENOP 50, "WSLH IT Security Plan" for details and additional items. In OnBase, search SOP-Approved, Division=OIS, Section=Infrastructure.

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Special protections for our Instrument Workstations:

- Instrument workstations typically cannot run the latest operating system patches for fear of breaking the acquisition/analysis software. These are protected by 2 different, segregated VLAN's (virtual networks) with firewall rules:
 - o **Instrument VLAN** contains instrument workstations capable of running antivirus/antimalware software. These PCs are allowed open access to internal servers but Internet traffic is restricted to only a small number of whitelisted sites (e.g. www.PerkinElmer.com).
 - Protected VLAN contains instrument workstations that cannot run antivirus/anti-malware software or those that meet other high-risk criteria. These PCs have very limited access to internal resources and no Internet access.
- When possible, instrument interfaces are established to directly pull data into our LIMS systems (e.g., Epic, WindoPath, Clinysis,) to improve data integrity and increase laboratory productivity.
- We also maintain "bare metal-ghost" backups of our instrument workstations using disk imaging software on an annual basis or whenever laboratory staff notify us of a change to the workstation. It is preferred to take a backup image prior to any vendor changes in case these changes need to be rolled-back. When the work is complete and verified, another backup image should be taken to allow future disaster recovery. It is the responsibility of laboratory staff to notify OIS (via a Team Dynamix ticket) in advance of these changes so the work can be scheduled. These backups are maintained to allow recovery of the PC configuration in case of a catastrophic loss of hardware at the PC level. Status of the ghosting is recorded at O:\OIS\InstrumentBackupSummary.xlsm.
- All EHD sections will assign staff to be responsible for maintaining the above file back-ups and hard drive back-ups.
- All vendor technicians are required to have their removable media scanned before it can be inserted into the instrument PC to prevent the spread of viruses and malware.
 - 1.17.5. Bulk Argon

The Agriculture Drive facility has a bulk liquid argon tank near the loading dock that supplies argon gas throughout the building for instruments that require it. Our supplier of the bulk argon (purity = 99.999%) is Airgas Merchant Gases, Madison, WI.

1.17.6. Thermometers (& other temperature measuring devices)
Thermometers used for NELAC, WDNR, or EPA testing must be calibrated or verified annually. The

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calibration or verification must be traceable to NIST. If the thermometer is used over a range of ≤ 10 °C, a single point verification within that range can be used. If the thermometer is used over a range of > 10 °C, the verification must bracket the range of use. The calibration may be done in-house, by a third-party calibration company, or by the device manufacturer or vendor.

If thermometers are calibrated in-house, the thermometer will be labeled with the date calibrated and the expiration date (same month of next year-e.g., calibrated 01/12/2025, exp. 01/2026). If sections calibrate thermometers in-house, they must have a procedure for doing so. The procedure must include how to do the calibration, how to use correction factors if there is a bias, evaluation criteria, and what to do with thermometers that fail the evaluation criteria.

If thermometers are purchased new annually with a certificate of calibration (NIST-traceable), the thermometer will be labeled with the date it was received and the expiration date (same month of next year-e.g., rec'd 05/10/2025, exp. 05/2026). The person who receives the thermometers must review the calibration certificates to ensure the thermometers have been calibrated within the range of use, and that the expiration date on the certificate extends to cover the section's planned last date of use.

1.17.7. Facilities

The Wisconsin State Laboratory of Hygiene (WSLH) is currently housed in four separate facilities: 465 Henry Mall on the UW Madison Campus, 2811 Agriculture Drive (the State Agriculture Building or SAB), 2601 Agriculture Drive, and 4626 University Ave (Soils and Forage Analysis Lab). The facility at 2601 Agriculture Drive includes the original building built in 1999 and a newer co-located building completed in 2013. The analytical sections covered in this manual are located in the 2601 Agriculture Drive facility.

As part of the University of Wisconsin (UW), the WSLH benefits from services and expertise offered by the University such as the Environment, Health & Safety Department, collaborations with faculty, and access to the library system. Although the EHD laboratories are not physically located on the University campus, the staff are University employees and must follow university policies, procedures, and rules.

The activities of the various EHD lab sections are supported by a fully automated glassware washing room, a media preparation room, shipping & receiving section, and the offices mentioned in section 1.4. Some of these support sections/offices are located at other facilities.

The 2601 Agriculture Drive facility is equipped with an air handling system that is maintained by the Department of Administration. The system consists of three intake fans and three exhaust fans. The air handling system ensures that there is always negative pressure in all fume hoods and snorkels. Except for the trace metal clean lab (rm. 256), there is a 100% exchange of air, (i.e., what comes in, goes out; there is no recirculation) and the volume of air in the building is changed approximately once per minute. In

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addition, the building is designed so that the laboratories will generally be under a negative pressure. The air will flow from the office/cubicle space, into the lab, and then be exhausted.

Section supplements describe facilities that are used within each section.

1.18. Purchasing

1.18.1. Supplies and Services

WSLH Purchasing procures goods and services necessary for operations performed by the WSLH except as noted below. Procedures for purchasing services are the same as for purchasing supplies with the addition of a certificate of insurance, provided by the vendor, prior to coming on-site. Some items are ordered through Acumatica, which is accessed through the WSLH Intranet Home Page. For instructions on submitting purchase requests, see "Purchase Requests" Business Process Procedure in OnBase. Other information regarding purchasing can be found through the intranet Team Dynamix Knowledge Base. Purchasing operates under authority of the UW Madison Purchasing Department and must adhere to University and State of Wisconsin Policies.

In addition to submitting a purchase request through Acumatica, sections have designated staff who are responsible for ordering supplies for their sections from specified vendors. These orders are placed through the University of Wisconsin's procurement tool called ShopUW+ on the Procure-to-Pay (P2P) automated, cloud-based system, https://shopuwplus.wisc.edu/.

A third purchasing option is the use of a Procurement Card (ProCard), which is a State of Wisconsin VISA credit card. A limited number of employees are authorized and trained, following strict guidelines, to make purchases using a ProCard. Please see the WSLH internal website under Administrative Services, Purchasing Department for policies and procedures.

1.18.2. Capital Equipment

Capital equipment purchases (i.e., > \$5,000, with a useful life of more than one year) are generally sent out for bid. Laboratory staff works closely with WSLH Purchasing to assure that all necessary specifications are included in the bid. All items are purchased through Purchasing.

1.18.3. Receipt of Supplies

When standards, reagents, reference materials, and media are received, assign them unique traceability codes (see section 1.13), document their receipt in the appropriate logbook and/or electronic system, and file the certificates of analysis in the appropriate sectional file (either paper or electronic). Certificates of analyses must be maintained at the lab if the manufacturer has them available (in either hard copy or electronic format). It is the responsibility of the person receiving the supplies to seek out the certificates of analysis if they do not arrive with the items. The certificates must be labeled with the unique traceability code. For original containers of standards, reagents, reference materials, chemicals, media, etc., if an

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expiration date is provided by the manufacturer or vendor, it must be recorded on the container.

1.18.4. Verification and Evaluation of Supplies

It is the responsibility of the individual user of supplies, reagents, and consumable materials to verify that the supplies comply with requirements specified in METHOD SOPs. An example would be checking blanks on autosampler tubes that come from a vendor. The records of these verification checks would be maintained along with the normal data output from the instrument that is used. Individual METHOD SOPs also state specific grades of reagents, standards, or chemicals required for the procedure and the reception and storage requirements for reagents, standards, and chemicals.

If goods or supplies are received that do not meet requirements, are damaged, or orders are incomplete, Purchasing must be notified promptly so that they can follow up with the vendor.

1.18.5. Approved Vendor Criteria

The criteria for approval of vendors may include:

- Acceptable historical performance as determined by the verification and evaluation procedure noted above or in specific METHOD SOPs.
- Acceptable reliability and robustness as required by the application are met by instrumentation and equipment.
- Specific applications may require the vendor to hold certification by third parties.
- Acceptable use by other labs with applications similar to ours.
- Acceptability by UW Purchasing and WSLH Purchasing
- All details of vendor acceptability beyond what is noted here will be in section supplements of this QA Manual or in SOPs.

1.18.6. Approved Vendor List

The approved vendor list (O:\SOP\EHD\Division Wide\Draft\In Progress\TNI QA Manual supporting docs for review 03-06-2025\approved vendors list 2025.xlsx) contains vendors commonly used by the sections covered in this QA Manual, and that have met criteria outlined in 1.18.5. Other vendors are used occasionally, and information about them can be found either in the section supplements or in individual SOPs. The approved vendor list is reviewed annually by analytical sections and WSLH Purchasing. Note that UW-Madison Purchasing does not maintain an approved vendor list; rather, they provide instructions on how to check if vendors are not in compliance with Sec. 77.66 Wis. Statutes (certification for collection of sales and use tax). In addition, there is a list of vendors ineligible under the Wisconsin Office of Contract Compliance. If a vendor appears on either of these lists (which are available through doa.wi.gov), we are not allowed to use them. For details and questions, contact Purchasing. Another information source is https://businessservices.wisc.edu/purchasing/how-to-make-a-purchase/how-to-buy-a-product/

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1.19. Management Review of the Quality System

The WSLH Associate Director of Non-Clinical Testing will review the quality system spelled out in this manual annually, and a management review report will be written. The report will include: suitability of our policies and procedures, reports from supervisors, a review of important findings from internal audits, major corrective actions that took place in the past year, a review of any external audit reports, any problems with PT samples that need to be addressed, changes in the volume or type of work performed by the sections, a review of client feedback and complaints, and other relevant factors. For details, please see EHD DIV-WIDE GENOP 023 "Procedure for the Management Review of the Quality System." A schedule for the Management Reviews can be found at O:\Teams\EHD QC Team\Schedules

1.20. Internal Audits & Data Review

1.20.1. Internal Audits:

Internal audits address all elements of the quality system along with environmental testing activities as related to each specific section. Internal audit findings are summarized in the annual management review document.

Internal audits are conducted according to EHD DIV-WIDE QA 120, "Internal Audit Procedures." The purpose of internal audits is to help meet quality goals such as:

- Ensure that procedures specified in SOPs and regulatory methods are being followed.
- Promote consistent practices across all areas of the section, and increase awareness of QA practices.
- Ensure that the section is meeting requirements of regulating agencies (e.g. TNI, USEPA, and Wis. DNR).

Internal Audits will be conducted by the Quality Assurance (QA) coordinators or designees. They will be done according to set schedules or when corrective or preventative actions reveal a need for one. Internal audits consist of two parts: a system internal audit and a group/series of method internal audits. All accredited methods will be audited. Non-accredited methods may also be audited.

- System internal audits are a general review of each section's quality system.
- Method internal audits look at one method or a group of methods in greater detail.

Internal audit reports, including findings/deficiencies, will be generated. There are templates for the audits in O:\SOP\EHD\Division Wide\Final. Corrective actions in response to the internal audit report will be coordinated by the supervisor and reported back to the QA coordinator. In some cases a Non-Conforming Event Management (NCEM) report will need to be filed. If any of the findings of the internal audit casts doubt on the validity of client results, the client will be notified, in writing, within one month of the findings.

A schedule for internal audits can be found at O:\Teams\EHD QC Team\Schedules

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1.20.2. Data Review:

Some data calculations are housed within the LIMS and are performed when results are entered. All data is reviewed by a second person before it is released. Review of data is documented in the LIMS with date/time and reviewer's initials. Note that for results calculated by spreadsheets or other software, the software calculations must be verified as giving accurate results before initial use of the software. This verification must be documented and kept on file. Once verified, the calculations must be secured to prevent unauthorized amendment (see EHD DIV-WIDE GENOP 040, "Protecting Excel and Access Data.") If a calculation cannot be secured (for example, by locking a cell), then some other means of preventing unauthorized amendment must be documented. Some areas use a macro-enabled spreadsheet that protects filled cells. Software used to record data must have some means of maintaining data integrity and preventing unauthorized amendment of those records. This could include printing out spreadsheets or using a track changes feature to document that data was not changed at a later date. Details on review of data for specific analytical tests can be found at the sectional level.

Quality control samples that are analyzed and entered in the LIMS are given a unique sample number. Those QC samples are related to all samples in the batch in LIMS. The LIMS does an evaluation of the QC sample results based on limits entered into the LIMS and displays pass or fail and the samples related to those QC samples.

1.21. Corrective and Preventive Action

1.21.1. Corrective Action for Non-Conforming Work

Generally, when any aspect of sample testing does not conform to our standard operating procedures (nonconforming work), including quality control procedures, corrective action will be promptly performed, starting with a correction of the issue, followed by root cause analysis, and documentation of the non-conformance. Section-specific details of corrective action documentation within analytical runs can be found in the chapter of this manual dedicated to that section and in individual METHOD SOPs. Corrective actions will be performed by the analyst in consultation with the peer review auditor, QA coordinator and their supervisor. The lab supervisor is responsible for the management and evaluation of non-conforming work in consultation with the QA coordinator. The supervisor is responsible for halting work or withholding test reports if deemed necessary and will authorize resumption of work. The supervisor may consult with the client (data user) to determine the usefulness of any qualified data and to determine a subsequent course of action. In some cases, more samples may need to be collected. If a data report has already been released when an error is discovered, the client will be notified as soon as possible and an amended report will be released once the issue has been resolved. All communications with clients will be documented and archived using e-mail, telephone logbooks, written records, or Non-Conforming Event Management System. If the non-conformance of work casts doubts on the laboratory's compliance

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with its own policies and procedures or compliance with TNI standards, an internal audit will be performed.

1.21.2. Corrective action for departures from policies, procedures & quality control Generally, when departures from documented policies, procedures and quality assurance occur corrective action will be promptly performed and documented. Section-specific details of corrective action documentation can be found in the supplemental plan of this manual dedicated to that section and in individual METHOD SOPs. Section-specific corrective action procedures identify the person responsible for assessing each QC data type and initiating and recommending corrective action. The treatment of a data set, the reportability of test sample results, and the use of appropriate laboratory-defined data qualifiers is outlined in section-specific methods. Departures from policies and procedures and out-of-control situations and their subsequent corrective actions are documented through the use of Non-Conforming Event Management (NCEM)reports. If the departures from documented policies, procedures and quality control cast doubt on the laboratory's compliance with its own policies and procedures or compliance with NELAC standards, an internal audit will be performed.

1.21.3. Permitting Departures from Documented Policies and Procedures

Invariably there will be exceptions to the policies and procedures documented and referenced in this QA Manual where corrective action as noted above does not resolve the problem. If an analyst is unsure about how to proceed, they should ask their supervisor or quality assurance coordinator. Any decision to proceed with work that does not conform to standard procedures must be approved by a supervisor or quality assurance coordinator. The supervisor may consult with the client (data user) or scientific experts to determine the usefulness of any qualified data and to determine a subsequent course of action. Decisions must be documented.

1.21.4. Preventive Action

Preventive action is essential in providing accurate, reproducible, reliable data. Preventive action is necessary to ensure that equipment and all quality systems are functioning properly and to prevent potential non-conforming events. If needed improvements or nonconformities arise, actions are taken to ensure that future non-conformances are prevented. Use of the NCEM system will allow for the identification and implementation of further preventive actions. Preventative action is routinely implemented by the laboratory staff. Preventative action includes (but is not limited to):

- Reviewing operational procedures
- Reviewing non-conforming event management (NCEM) reports
- Discussing NCEMs and trends with staff
- Reviewing QC data for trends and outliers
- Conducting periodic instrument maintenance

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- Reviewing instrument logs for problems
- Reviewing customer (internal & external) comments and complaints
- Reviewing PT results
- Reviewing staffing and training needs
- Performing DOCs
 - 1.21.5. Non-Conforming Event Management System

The WSLH utilizes software for Non-Conforming Event Management. Guidelines for use of NCEM for documentation of corrective action include when there is a non-routine QC failure that cannot be resolved within the analytical batch or a systemic problem that needs to be tracked over time. Examples of non-conformances entered are PT failures, customer complaints, reporting errors, instrument/equipment failure, HIPAA violations, etc. Our internal web site includes a link to the software to be used for NCEM.

There are two lab-wide SOPs that document non-conforming event management:

LABWIDE GENOP 711, Non-Conformance Management Procedure for MediaLab.

LABWIDE GENOP 707, Non-Conforming Event Management System Policy

The procedure for review, monitoring, and close-out of Non-Conforming Event Management forms is detailed in LABWIDE GENOP 707. For testing under the quality system as documented in this QA Manual (1.1), open Non-Conforming Events are discussed at section staff meetings or at the EHD QA team meeting. This allows dissemination of information and encourages progression of corrective actions and follow-up activities.

For non-conformances associated with TNI-accredited testing, a root cause analysis must be conducted and documented. Where applicable, follow-up monitoring must be conducted and documented. Whenever possible, link or upload all supporting documentation to the NCEM system.

Documentation of non-conformances must be made by the person discovering the problem, their supervisor, or their QA representative. The documentation must be made as soon as possible after discovery, and at most within a week. Follow-up action taken, monitoring results, and root cause determinations must be documented. Documentation of these processes must also be completed in a timely manner, and at most within a week of the action taken.

NCEM training: Training consists of reading LABWIDE GENOPs 707 and 711. Selected employees of EHD have more advanced training as "owners" to use the non-conformance management software in MediaLab. Also, see section 1.10 of this QA Manual.

Any questions regarding the use of the NCEM system can be directed to any quality assurance coordinator.

1.21.6. Quality, Safety, & Improvement Awards

The WSLH offers Quality, Safety, and Improvement (QSI) Awards to recognize and celebrate staff efforts

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to find improvements to the work environment, processes, data quality, or the delivery of services to our customers. Award criteria include outstanding performance, beyond normal job expectations for an individual or team. Criteria are based on being an advocate, improving a process, innovating, or completing a project in the areas of quality, safety, or improvement. Staff can nominate co-workers at any time through the internal website. Awards are presented twice per year.

1.22. Reporting Analytical Results

Analytical documentation

Reporting of data from the LIMS can be accomplished through either electronic transfer, printed, faxed or emailed version of the result report.

WSLH Laboratory Reports generated by the LIMS for the EHD contain:

- Address and contact information for the Agriculture Drive laboratory site and the responsible managers
- A title, unique report ID, and pagination
- Customer name and address
- Sample identification and information related to sampling if applicable
- Analytical & preparation methods used and date of analysis
- Test results along with appropriate units, LODs, LOQs, comments, qualifiers, test conditions, and additional information as necessary
- Statements indicating compliance with applicable standards, applicability of test results, and instructions for reproducibility of reports.
- Pending tests, if results are reported prior to the completion of all requested analysis

1.22.1. Amendments to Test Reports

- If a report is amended after issue, the amended report will meet all of the requirements listed above.
- The amended report will refer to the original report ID.
- The amended report will include a note stating why it was amended.

Drinking Water Requirement for Chemical Testing

If an MCL (Maximum Contaminant Level) for an analyte regulated under ch. NR 809 has been exceeded for a PWS (Public Water Supply) sample, the water supply facility must be notified within 48 hours of completing the analyses [Wis. Admin. Code (06/29/2021) NR 149.47(1)(f)]. The analysis is considered complete when a batch is finalized in the LIMS after the peer review audit. Applicable MCLs will be listed in SOPs. The EHD LIMS automatically flags results above the applicable MCL for internal use as well as on the test report.

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1.23. Subcontracting of Environmental Tests

If there is an emergency such as an instrument break-down where accredited work would need to be subcontracted, there are several considerations and requirements:

- DNR work certified under NR 149 must only be subcontracted to another lab certified under the same code for the specific field of certification.
- Drinking water compliance samples must only be analyzed by laboratories certified by the EPA or via reciprocity by an authorized state.
- TNI-accredited work must only be subcontracted to another lab that complies with TNI standards for the specific field of accreditation.
- The WSLH must advise the data users of the subcontract arrangement. This must be documented.
- The WSLH must request proof from the subcontractor of compliance with applicable standards (e.g., a certificate and scope of accreditation). These documents must be kept on file.
- The final WSLH report must include the name of the sub-contracted laboratory (and Wisconsin Facility Identification Number—FID, if applicable) performing any subcontracted work. The subcontractor's full report must be made available to the data user if requested.
 - 1.23.1. Subcontract Lab Contingency Lists for Emergencies
- Labs certified under Wisconsin Administrative Code NR 149 (lists include scopes of accreditation): https://dnr.wisconsin.gov/topic/labCert/certified-lab-lists
- Labs accredited under TNI (searchable by location and scope): https://lams.nelac-institute.org/Search
- Labs certified by EPA for drinking water: https://www.epa.gov/dwlabcert/contact-information-certification-programs-and-certified-laboratories-drinking-water

1.24. Proficiency Testing Sample Procedures

All proficiency testing samples are analyzed in the same manner as used for routine environmental samples, including:

- same staff
- Same methods (ensure method codes match TNI scope of accreditation, where applicable)
- Same procedures
- Same equipment and facilities

The lab independently analyzes proficiency testing samples and reports the results according to the following rules:

- Does not sub-contract the analysis nor analyze another lab's PT sample
- Does not communicate with another lab about a PT sample prior to the close of the study
- Does not attempt to obtain the PT results from the proficiency testing provider prior to close of the

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study

Each section is responsible for ordering, analyzing, and reporting PT samples and results that they need to maintain accreditation. See section supplements and associated SOPs for details.

Each section is responsible for performing appropriate corrective action for PT failures and documenting the corrective action in the NCEM system.

Each section must notify the non-clinical QA coordinator of any PT failures.

Each section is responsible for organizing and maintaining PT records including bench sheets, instrument printouts, data calculations, data summary reports, and PT study report forms according to the applicable RDA.

1.25. Method References

1.25.1. Policy

Approved editions of reference methods will be used when required by State of Wisconsin administrative codes or federal rules.

1.26. General References

1.26.1. WSLH Lab-wide Procedures and Policies

Lab-wide policies and references to University of Wisconsin and State of Wisconsin policies and procedures that apply to the WSLH are found in the Knowledge Base of Team Dynamix (see intranet home page). In addition, lab-wide approved SOPs are stored in the OnBase (Web-client) document management platform (see 1.11.1).

1.26.2. EHD Division-wide Procedures and Policies

Division-wide approved SOPs are stored at in the OnBase (Web-client) document management platform.

1.26.3. WSLH Board Policies and Procedures

Policies and Procedures of the WSLH Board, along with meeting minutes, are stored on the extranet at: http://www.slh.wisc.edu/about/board/

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1.27. Major Changes

1.27.1. Ver.2 (OnBase)

Section	Change
	Began version tracking table
	Updated header, footer
	Updated signature page
1.2.1	Added info re. updates of subsequent versions of the QA Plans after 08/2022
1.3	Added "Quality is everyone's responsibility" as per APHL
1.4	Updated the organization of EHD and staff names
1.4	Deleted specific info re. TNI technical managers (left reference)
1.6	Added cross-ref. to RDA (section 1.12), deleted outdated references
1.10.1,	Added WSLH Statement of Ethical Expectations (WSLH-WIDE POLICY 001)
1.10.8	
1.10.5	Updated documentation of training to Canvas app
1.11	Updated for OnBase Timers & Notifications module
1.12	Updated RDA info and hyperlink
1.12.3	Deleted, "for up to three years."—No longer policy
1.15.1	Added Copies of WSLH executed contracts are maintained and available by
	contacting the Contract and Grant Administrator Specialist in the WSLH Office of
	Finance (deleted the previous link to a shared drive).
1.15.3	Added item to be considered for new work: desired records retention period
1.15.4	Changed link to access the actual memo rather than the webpage (suggestion by
	Kevin Karbowski)
1.17.4	Updates to Computers section as directed by Allen Benson
1.18	Updated Purchasing section on the advice of Mark Conklin and Tina Mathew,
	WSLH Purchasing
1.19	Updated position name
1.20	Added link for internal audit templates
1.21.6	Added new section for opportunities to improve processes (verbal suggestion by
	TNI auditor Bill Hall, May, 2023).

1.27.2. Ver. 3

Section	Change
	Updated signature page
1.2.1	Updatd EHD sections accredited under TNI
1.4.2	Added info re. support offices
1.4.4	Changed "annually" to "a set schedule," added "or action items"
1.8	Updated hiring practices wording
1.9	Updated UW-Madison safety information
1.10	Added info throughout section for Bloodborne Pathogen training
1.10.1	Updated for new non-conformance procedures
1.10.6	Added APHL ethics training link
1.10.8	Added other documentation options for those who do not use Canvas
1.12.2	Added electronic logbooks (e.g. LabArchives)

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1.12.5	Added disposal of records through SRC option
1.17.4	Added file back-up locations, hard drive ghosting schedule spreadsheet location,
	and EHD section staff responsibilities concerning back-ups and ghosting.
1.17.7	Updated facility locations
1.18.6	Added info about unapproved vendors through UW-Madison Purchasing and DOA
1.20.1	For internal audits, changed "annual" to "according to set schedule"
1.21	Updated "occurrence" to "non-conformance." Or nonconforming event Updated
	Footprints to MediaLab

1.27.3. Ver. 4

1.2/.	5. Vel. 4
Section	Change
1.4.4	Added place-holder for EHD QA Manager (more details to be added next version update)
1.10	Added records training information
1.10.1	Updated Employee Handbook to the Knowledge Base; updated to make bloodborne pathogen training required for all employees.
1.10.3	Added WSLH Employee Safety Checklist to annual review items.
1.10.8	Added Employee Safety Checklist to Continuing Training Summary table.
1.12.3	Deleted info about access log ESS801 because log was rarely used, and we now define "archived" as those records sent to the SRC (TNI V1M2 1.13.3 e). The basement room 14 is an on-site storage location.
1.15	Minor wording changes suggested by K. Karbowski; addition of DocuSign processes
1.16.1	Added information regarding sampling procedures
1.18.1	For purchasing info, added Team Dynamix Knowledge Base
1.20.1	Added "and increase awareness of QA practices"
1.20.2	Some areas use a macro-enabled spreadsheet that protects filled cells.
1.21.6	Updated the QSI Award info
1.22	Added preparation methods to list of items in reports
1.26.1	Updated location of lab-wide policies from the Employee Handbook to the Knowledge
	Base module of Team Dynamix.
1.26.3	Added, "along with meeting minutes"
	In multiple areas removed Horizon and changed to generic LIMS because Horizon will be renamed Clinisys in 2025