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## Final Butte Reduction Works (BRW) Phase I QAPP

Pioneer Technical Services, Inc.

Nikia Greene Environmental Protection Agency

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## **EPA REGION 8 QA DOCUMENT REVIEW CROSSWALK**

					A DOCUMENT RE		
QAPP/FSP/S			Entity (gra	ntee, contract, EPA AO	, EPA Program, Other)	Regulatory	2 CFR 1500 for Grantee/Cooperative
(check approprie						Authority	Agreements
GR	ANTEE		Atlantic Ric	chfield (PRP)			48 CFR 46 for Contracts
CO	NTRACTOR					and/or	Interagency Agreement (FFA, USGS)
EPA	1						EPA/Court Order
Oth	er					Funding	EPA Program Funding
	••					Mechanism	EPA Program Regulation
							EPA CIO 2105
Document T	itle l be repeated in Hea	udar l	Final Butte Reduction Works (BRW) Phase I QAPP				
	AP Preparer	uerj	Dionoor Too	chnical Services, Inc. for	r Atlantia Diabfield		
QALITISITS	Al l'Ieparei		Company (		I Atlantic Richneid		
Period of Pe			2022			Date Submitted	7/20/2022
(of QAPP/FSP/S						for Review	
EPA Project Officer						PO Phone #	
	EPA Project Manager		Nikia Greene			PM Phone #	(406) 457-5019
	QA Program Reviewer or					Date of Review	
Approving (			Nikia Green		<u> </u>		
<b>Document</b>	s Submitted f	or QAI	PP Review	(QA Reviewer must	Notes for Document		
complete):					<b>1.</b> A QAPP written by a Grantee, EPA, or Federal Partner <u>must include</u> for review:		
1. QA Docu	ment(s) submit	ted for <b>r</b>	review:		<ul> <li>Work Plan (WP) / Statement of Work (SOW) / Program Plan (PP) / Research Proposal (RP) and funding mechanism</li> <li>2. A QAPP written by Contractor must include for review:</li> </ul>		
QA	Document	Docur	nent	Document with			
Document	Date	Stand	-alone	QAPP		Order Work Assignn	
QAPP		Yes / I	No				opy of the contractor's approved QMP
FSP		Yes / I	No	Yes / No		act SOW if no QMP	
SAP		Yes / I	No	Yes / No		Court Order, if appli	
SOP(s)				Yes / No	e) The QA Review must determine (with the EPA CO or PO) if a QARF was completed		
	//TO/PP/RP Da				for the environmental data activity described in the QAPP.		
	//TO/RP Perfor				3. a. Field Sampling Plan (FSP) and/or Sampling & Analyses Plan (SAP) must include the		
3. QA document consistent with the:			Project QAPP or must be a stand-alone QA document that contain all QAPP required				
WP/SOW/PP for grants? <u>Yes / No / NA</u>			elements (Project Management, Data Generation/Acquisition, Assessment and				
SOW/TO for contracts? <u>Yes / No / NA</u> 4. QARF signed by R8 QAM <u>Yes / No / NA</u>			Oversight, and Data Validation and Usability).				
				ibmitted with a QA	document that contains all QAPP required		
Funding Mechanism <u>IA / contract / grant / NA</u> Amount			elements.				
Summary of	Comments (hi)	ehlieht s	ignificant co	oncerns/issues):			
<b>,</b>	(	5	0	···· ··· · · · · · · · · · · · · · · ·			

Element	Acceptable Yes/No/NA	Page/ Section	Comments
A. Project Management		•	
A1. Title and Approval Sheet			
a. Contains project title		Title Page	
b. Date and revision number line (for when needed)		Approval Page	
c. Indicates organization's name		Title Page	
d. Date and signature line for organization's project manager		Approval Page	
e. Date and signature line for organization's QA manager		Approval Page	
f. Other date and signatures lines, as needed		Approval Page	
A2. Table of Contents			
a. Lists QA Project Plan information sections		Pages i to iii	
b. Document control information indicated		Footer	
A3. Distribution List			
Includes all individuals who are to receive a copy of the QA Project Plan and identifies their organization		Distribution List	
A4. Project/Task Organization	•		
a. Identifies key individuals involved in all major aspects of the project, including contractors		Section 6.0	
b. Discusses their responsibilities		Section 6.0	
c. Project QA Manager position indicates independence from unit generating data		Section 6.0	
d. Identifies individual responsible for maintaining the official, approved QA Project Plan		Section 6.0	
e. Organizational chart shows lines of authority and reporting responsibilities		Figure 25	
A5. Problem Definition/Background	-	•	
a. States decision(s) to be made, actions to be taken, or outcomes expected from the information to be obtained		Section 1.2	
b. Clearly explains the reason (site background or historical context) for initiating this project		Section 1.1, Section 2.0	
c. Identifies regulatory information, applicable criteria, action limits, etc. necessary to the project		Section 2.4	

A6. Project/Task Description		
a. Summarizes work to be performed, for example, measurements to be made, data files to be obtained, etc., that support the project's goals	Section 1.2	
b. Provides work schedule indicating critical project points, e.g., start and completion dates for activities such as sampling, analysis, data or file reviews, and assessments	Section 2.5	
c. Details geographical locations to be studied, including maps where possible	Section 2.5, Figure 5, Figure 8, Figure 10, Figure 12	
d. Discusses resource and time constraints, if applicable	Section 2.5	
A7. Quality Objectives and Criteria	· · ·	
<ul> <li>a. Identifies</li> <li>performance/measurement criteria for all information to be collected and acceptance criteria for information obtained from previous studies,</li> <li>including project action limits and laboratory detection limits and</li> <li>range of anticipated concentrations of each parameter of interest</li> </ul>	Section 2.5	
b. Discusses precision	Section 2.5	
c. Addresses bias	Section 2.5	
d. Discusses representativeness	Section 2.5	
e. Identifies the need for completeness	Section 2.5	
f. Describes the need for comparability	Section 2.5	
g. Discusses desired method sensitivity	Section 2.5	
A8. Special Training/Certifications		
a. Identifies any project personnel specialized training or certifications	Section 3.1	
b. Discusses how this training will be provided	Section 3.1	
c. Indicates personnel responsible for assuring training/certifications are satisfied	Section 3.1	
d. identifies where this information is documented	Section 3.1	
A9. Documentation and Records		

a. Identifies report format and summarizes all data report package information	Section 4.6, Section 5.4
b. Lists all other project documents, records, and electronic files that will be produced	Section 4.6
c. Identifies where project information should be kept and for how long	Section 4.6
d. Discusses back up plans for records stored electronically	Section 4.6
e. States how individuals identified in A3 will receive the most current copy of the approved QA Project Plan, identifying the individual responsible for this	Section 3.1, Section 6.0
<b>B.</b> Data Generation/Acquisition	
<b>B1.</b> Sampling Process Design (Experimental Design)	
a. Describes and justifies design strategy, indicating size of the area, volume, or time period to be represented by a sample	Section 2.5
b. Details the type and total number of sample types/matrix or test runs/trials expected and needed	Section 2.5
c. Indicates where samples should be taken, how sites will be identified/located	Section 2.5
d. Discusses what to do if sampling sites become inaccessible	Section 2.5
e. Identifies project activity schedules such as each sampling event, times samples should be sent to the laboratory, etc.	Section 2.5
f. Specifies what information is critical and what is for informational purposes only	Section 2.5
g. Identifies sources of variability and how this variability should be reconciled with project information	Section 2.5
B2. Sampling Methods	
a. Identifies all sampling SOPs by number, date, and regulatory citation, indicating sampling options or modifications to be taken	Section 2.5, Section 3.0, Appendix 2
b. Indicates how each sample/matrix type should be collected	Table 3,       Appendix 2

c. If in situ monitoring, indicates how instruments should be deployed and operated to avoid contamination and ensure maintenance of proper data	Section 4.4	
d. If continuous monitoring, indicates averaging time and how instruments should store and maintain raw data, or data averages	Section 4.4	
e. Indicates how samples are to be homogenized, composited, split, or filtered, if needed	Section 3.0, Appendix 2	
f. Indicates what sample containers and sample volumes should be used	Table 3	
g. Identifies whether samples should be preserved and indicates methods that should be followed	Table 3	
h. Indicates whether sampling equipment and samplers should be cleaned and/or decontaminated, identifying how this should be done and by-products disposed of	Appendix 2	
i. Identifies any equipment and support facilities needed	Section 3.9, Section 4.4	
j. Addresses actions to be taken when problems occur, identifying individual(s) responsible for corrective action and how this should be documented	Section 5.0	
B3. Sample Handling and Custody		
a. States maximum holding times allowed from sample collection to extraction and/or analysis for each sample type and, for in-situ or continuous monitoring, the maximum time before retrieval of information	Table 3	
b. Identifies how samples or information should be physically handled, transported, and then received and held in the laboratory or office (including temperature upon receipt)	Section 3.8	
c. Indicates how sample or information handling and custody information should be documented, such as in field notebooks and forms, identifying individual responsible	Section 3.7, Section 3.8	
d. Discusses system for identifying samples, for example, numbering system, sample tags and labels, and attaches forms to the plan	Section 3.6	
e. Identifies chain-of-custody procedures and includes form to track custody	Section 3.8	
B4. Analytical Methods		

a. Identifies all analytical SOPs (field, laboratory and/or office) that should be followed by number, date, and regulatory citation, indicating options or modifications to be taken, such as sub-sampling and extraction procedures	Table 3,Appendix 1,Appendix 2,Appendix 3,Appendix 7,Appendix 8
b. Identifies equipment or instrumentation needed	Section 4.4
c. Specifies any specific method performance criteria	Section 2.5, Table 3
d. Identifies procedures to follow when failures occur, identifying individual responsible for corrective action and appropriate documentation	Section 5.0
e. Identifies sample disposal procedures	Appendix 1,Appendix 2,Appendix 8,Lab SOPs
f. Specifies laboratory turnaround times needed	Section 3.10
g. Provides method validation information and SOPs for nonstandard methods	Section 9.0
B5. Quality Control	
a. For each type of sampling, analysis, or measurement technique, identifies QC activities which should be used, for example, blanks, spikes, duplicates, etc., and at what frequency	Section 4.0
b. Details what should be done when control limits are exceeded, and how effectiveness of control actions will be determined and documented	Section 5.0
c. Identifies procedures and formulas for calculating applicable QC statistics, for example, for precision, bias, outliers and missing data	Section 2.5
B6. Instrument/Equipment Testing, Inspection, and Mainte	nance
a. Identifies field and laboratory equipment needing periodic maintenance, and the schedule for this	Section 4.4
b. Identifies testing criteria	Section 4.4
c. Notes availability and location of spare parts	Section 4.4
d. Indicates procedures in place for inspecting equipment before usage	Section 4.4, Appendix 2

Section 6.0

inspection and maintenance

e. Identifies individual(s) responsible for testing,

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f. Indicates how deficiencies found should be resolved,	Section 4.4
re-inspections performed, and effectiveness of	and Section
corrective action determined and documented	5.0
<b>B7.</b> Instrument/Equipment Calibration and Frequency	
a. Identifies equipment, tools, and instruments that	Section 3.9,
should be calibrated and the frequency for this	Section 4.0,
calibration	Appendix 2
b. Describes how calibrations should be performed and	Section 3.9,
documented, indicating test criteria and standards or	Section 4.0,
certified equipment	Appendix 2
c. Identifies how deficiencies should be resolved and	Section 3.9,
documented	Section 4.0,
	Appendix 2
<b>B8.</b> Inspection/Acceptance for Supplies and Consumables	
a. Identifies critical supplies and consumables for field	Section 4.5
and laboratory, noting supply source, acceptance	
criteria, and procedures for tracking, storing and	
retrieving these materials	
b. Identifies the individual(s) responsible for this	Section 4.5
<b>B9.</b> Use of Existing Data (Non-direct Measurements)	
a. Identifies data sources, for example, computer	Section 2.2
databases or literature files, or models that should be	
accessed and used	
b. Describes the intended use of this information and the	Section 2.2
rationale for their selection, i.e., its relevance to project	
c. Indicates the acceptance criteria for these data	Section 2.2,
sources and/or models	Section 2.5
d. Identifies key resources/support facilities needed	Section 2.2,
	Section 2.5
e. Describes how limits to validity and operating	Section 2.2,
conditions should be determined, for example, internal	Section 2.5
checks of the program and Beta testing	
B10. Data Management	
a. Describes data management scheme from field to	Section 3.7,
final use and storage	Section 4.6
h Discusses standard record-keeping and tracking	Section 4.6
b. Discusses standard record-keeping and tracking practices, and the document control system or cites	Section 4.6

c. Identifies data handling equipment/procedures that should be used to process, compile, analyze, and	Section 4.6
transmit data reliably and accurately	
d. Identifies individual(s) responsible for this	Section 4.6
e. Describes the process for data archival and retrieval	Section 4.6
f. Describes procedures to demonstrate acceptability of hardware and software configurations	Section 4.6
g. Attaches checklists and forms that should be used	Section 4.6; Appendix 10
C. Assessment and Oversight	
C1. Assessments and Response Actions	
a. Lists the number, frequency, and type of assessment activities that should be conducted, with the approximate dates	Section 5.0
b. Identifies individual(s) responsible for conducting assessments, indicating their authority to issue stop work orders, and any other possible participants in the assessment process	Section 5.0
c. Describes how and to whom assessment information should be reported	Section 5.0
d. Identifies how corrective actions should be addressed and by whom, and how they should be verified and documented	Section 5.0
C2. Reports to Management	
a. Identifies what project QA status reports are needed and how frequently	Section 5.0
b. Identifies who should write these reports and who should receive this information	Section 5.0
D. Data Validation and Usability	
D1. Data Review, Verification, and Validation	
Describes criteria that should be used for accepting, rejecting, or qualifying project data	Section 9.0
D2. Verification and Validation Methods	
a. Describes process for data verification and validation, providing SOPs and indicating what data validation software should be used, if any	Section 9.0

b. Identifies who is responsible for verifying and validating different components of the project data/information, for example, chain-of-custody forms, receipt logs, calibration information, etc.	Section 9.0	
c. Identifies issue resolution process, and method and individual responsible for conveying these results to data users	Section 9.0	
d. Attaches checklists, forms, and calculations	Section 9.0; Appendix 10	
D3. Reconciliation with User Requirements		
a. Describes procedures to evaluate the uncertainty of the validated data	Section 9.0	
b. Describes how limitations on data use should be reported to the data users	Section 9.0	