Quality Assurance Project Plan

Statewide Survey of Per- and Poly-fluoroalkyl Substances in Washington State Rivers and Lakes

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Per- and Poly-fluoroalkyl Substances
in Washington State Rivers and Lakes

April 2016

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Signatures are not available on the Internet version.
EAP:  Environmental Assessment Program
HWTR: Hazardous Waste and Toxics Reduction
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2.0 Abstract

Per- and poly-fluoroalkyl substances (PFASs) are a large group of chemicals used in many industrial and consumer applications, such as water-, stain-, and oil-repelling coatings and firefighting foams. Some of the chemicals in this group – such as perfluorooctane sulfonate (PFOS) and perfluorooctanoic acid (PFOA) – have been identified as persistent, bioaccumulative, and toxic chemicals (PBTs). PFOS, PFOA, and their known precursors were largely phased out in the United States in the mid-2000s and early 2010s, yet it is not known whether this has resulted in lower environmental levels of PFASs in Washington State freshwater systems.

Washington State Departments of Ecology (Ecology) and Health (DOH) are currently in the process of developing a chemical action plan for PFASs to identify the steps the state may take to reduce the threat of PFASs in Washington’s environment. In 2008, Ecology carried out a statewide survey measuring PFASs in a variety of environmental media to evaluate their presence in the state. The study found widespread presence of PFASs in surface waters, WWTP effluent, fish tissue, and osprey eggs in Washington State at levels consistent with other non-point source waterbodies in North America.

In 2016, Ecology will conduct a follow-up study to the 2008 statewide survey to characterize the current level of these contaminants and to determine whether the concentrations and/or compound make up has changed. This data will also provide an up-to-date dataset to support the PFAS chemical action plan actions and recommendations.
3.0 Background

Per- and poly-fluoroalkyl substances (PFASs) are a large group of chemicals used in many industrial and consumer applications, such as water-, stain-, and oil-repelling coatings and fire-fighting foams. PFASs have been manufactured since the 1950s, but manufacturers began phasing out certain compounds – perfluorooctane sulfonates (PFOS), perfluorooctanoic acid (PFOA) and their known precursors – in the 2000s due to the concern over their toxicity and persistence in humans and the environment. The primary manufacturer of PFOS phased out production in 2002, and eight major companies joined EPA’s PFOA Stewardship Program to work toward eliminating PFOA and other long-chained PFASs by 2015.

The Washington State Departments of Ecology (Ecology) and Health (DOH) are currently in the process of developing a chemical action plan for PFASs to identify the steps the state may take to reduce the threat of PFASs, such as PFOS, in the environment. The PFAS chemical action plan is expected to be finalized by the end of 2016.

In 2008, the Washington State Department of Ecology (Ecology) carried out a statewide survey measuring PFASs in a variety of environmental media to determine their occurrence in the state’s freshwater systems (Furl and Meredith, 2010). This study found widespread presence of PFASs in surface waters, wastewater treatment plant (WWTP) effluent, fish tissue, and osprey eggs in Washington State at levels consistent with other non-point source waterbodies in North America. The following year, Ecology and Herrera (2010) analyzed PFASs in effluent of ten Puget Sound area wastewater treatment plants (WWTPs), and reported higher loading estimates for total (T-) PFASs than loading estimates for T-polychlorinated biphenyls, T-polybrominated diphenyls, and T-polycyclic aromatic hydrocarbons. Since then, Ecology has also found PFASs in marine sediments (Dutch et al., 2014) and reported rising PFAS concentrations in freshwater sediment cores (Mathieu, 2013).

In 2016, Ecology will conduct a follow-up study to the 2008 statewide survey to characterize the current level of these contaminants and to qualitatively assess whether the concentrations and/or compound makeup has changed over time. The 2016 study will analyze PFASs in surface water, WWTP effluent, freshwater fish, and osprey eggs.

3.1 Study area and surroundings

Ecology will collect surface water, freshwater fish, and osprey eggs from lakes and rivers distributed throughout the state (Figure 1). The waterbodies sampled in the 2008 survey will be targeted for 2016 sampling, with the addition of three sites to include potential PFAS sources that weren’t captured in the 2008 study. Angle Lake in western Washington and Moses Lake in eastern Washington are being added to the study locations for their proximity to use of aqueous film-forming foams (AFFFs). Lake Meridian is being added for additional data on PFAS contamination in urban lakes. Four WWTPs from the 2008 study and one additional WWTP will be targeted for effluent collection. Osprey eggs will be collected from several sites along the lower Columbia River (to match the 2008 study), as well as near Lake Washington and West Medical Lake.
3.1.1 Logistical problems

Ecology will work with a contracted wildlife biologist consultant to obtain osprey eggs. The consultant will help work out the logistical problems anticipated in egg collection, such as obtaining necessary wildlife collection permits, determination of nest occupancy, securing safe access to nests, and collection of the egg samples with minimal disturbance to the nesting ospreys.

Other logistical problems associated with access will be alleviated by desk reconnaissance of boat launches and access points. All waterbody sites in this study have been targeted by Ecology field crews before and access has been verified.

3.1.2 History of study area

Furl and Meredith (2010) were the first to report a broad survey of PFAS concentrations in Washington State freshwater systems. Out of fourteen waterbodies sampled in 2008 for PFASs in surface water, West Medical Lake – which is impacted by wastewater treatment plant effluent – contained the highest concentrations, followed by South Fork Palouse River (also impacted by WWTP effluent), and Lake Washington (urban waterbody). Surface water concentrations of PFOA and PFOS were generally lower than those reported in other parts of the United States.
Effluent samples from four WWTPs were also analyzed for this study, with total PFASs in the order of Spokane (Riverside Park) > West Medical Lake > Sumner > Marine Park. Concentrations of PFOA and PFOS were within the range of values reported in other regions of the U.S., with greater median values of PFOA and lower medians of PFOS in the Washington samples.

The 2008 survey also analyzed fish tissue collected from seven waterbodies throughout the state (Furl and Meredith, 2010). Fish from the lower Columbia River and Lake Washington contained the highest T-PFAS concentrations. These two sites are both within industrial or urban areas. West Medical Lake and F.D.R. Lake also had detected PFAS concentrations in fish samples. PFASs were not detected in fish samples from the two reference sites – Entiat and Quinault Rivers. A largemouth bass sample from the lower Columbia River and a peamouth sample from Lake Washington were above the Minnesota Department of Health fish consumption advisory level of 40 ng/g in fillets (MDH, 2008). No fish consumption advisory level currently exists for Washington State.

Osprey eggs were collected from the lower Columbia River in 2008, upstream and downstream of the Willamette River confluence (Furl and Meredith, 2010). PFOS concentrations in the eggs were similar to values recorded at Delaware Bay and lower than those recorded from Chesapeake, Casco, and Penobscot Bays. However, the highest PFOS concentration in the eggs (downstream of the Willamette River confluence) was the second highest value of recorded osprey egg concentrations in the U.S at the time.

### 3.1.3 Parameters of interest

This study will analyze four environmentally relevant groups of PFAS analytes, for which analytical methods have been established. Table 9 shows the analyte groups and individual chemicals in each group.

Perfluoroalkyl acids (PFAAs) will be analyzed in all samples. The PFAA group includes perfluorooalkyl sulfonates (PFSAs) and perfluorooalkyl carboxylates (PFCAs) with carbon chain lengths between 4 and 12. PFOS is a perfluorooalkyl sulfonate with 8 carbons and PFOA is the 8 chain perfluorooalkyl carboxylate. Long-chain PFAAs (those with a carbon chain length of 8 or greater) are widespread in the environment, highly persistent, bioaccumulative and have shown toxicity in studies of animals (EPA, 2009). A panel of scientists identified a probable link between PFOA exposure and several health concerns in humans, including thyroid disease, testicular and kidney cancer, and pregnancy-induced hypertension, high cholesterol, and ulcerative colitis (C8 Science Panel, 2013).

In addition to PFAA contamination in the environment through direct releases, other PFAS chemicals can break down through biotic and abiotic pathways into PFAAs as the terminal end product (Butt et al., 2014). These chemicals are referred to as precursors. This study will analyze known and potential PFAA precursors in surface water and WWTP effluent samples. The known precursor analyte group includes polyfluorinated sulfonamides and fluorotelomer carboxylates. Potential PFAA precursors include polyfluorooalkyl phosphates (PAPs) and fluorotelomer sulfonates (FTSs). PFAA precursors are most likely to be present in WWTP effluent and in surface water of rivers downstream of WWTP effluent discharge.
Polyfluoroalkyl phosphates (PAPs) are phosphate-based acid esters possessing at least one polyfluoroalkyl group. PAPs are potentially biotransformed to PFCAs (Lee et al., 2010; D’Eon et al., 2009), and may be an important source of PFAS exposure to humans (Eriksson and Karrman, 2015). PAPs are used in products such as food packaging, paints, sealers, and personal care products and are ubiquitous in indoor dust (Eriksson and Karrman, 2015; De Silva et al., 2012). Only WWTP effluent will be analyzed for PAPs, as the compounds are unlikely to be above quantitation limits in surface water. Fluorotelomer sulfonates (FTSs) are a group of fluorotelomers used in AFFFs (Herzke et al., 2012; Schultz et al., 2004) and in metal plating applications (OECD, 2013).

3.1.4 Results of previous studies

Table 1 presents summary statistics of Washington State T-PFAS concentrations previously reported for the target media of this study (surface water, WWTP effluent, freshwater fish tissue, and osprey eggs).

Table 1. Total PFAS (T-PFAS) Concentrations Reported in Previous Washington State Studies.

<table>
<thead>
<tr>
<th>Sample Matrix</th>
<th>Sample Type</th>
<th>Collection Year</th>
<th>Units</th>
<th>T-PFAS</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surfacewater (fresh)</td>
<td>Spring</td>
<td>2008</td>
<td>ng/L</td>
<td>1.11 185 21.9 7.47</td>
<td>(1)</td>
</tr>
<tr>
<td>Surfacewater (fresh)</td>
<td>Fall</td>
<td>2008</td>
<td>ng/L</td>
<td>&lt; 0.9 170 21.1 3.60</td>
<td>(1)</td>
</tr>
<tr>
<td>Surfacewater (fresh and marine)</td>
<td>Spring/summer/fall/winter mean</td>
<td>2009-2010</td>
<td>ng/L</td>
<td>1.5 41 --- ---</td>
<td>(2)</td>
</tr>
<tr>
<td>WWTP effluent</td>
<td>Spring</td>
<td>2008</td>
<td>ng/L</td>
<td>61.0 418 97.0 218</td>
<td>(1)</td>
</tr>
<tr>
<td>WWTP effluent</td>
<td>Fall</td>
<td>2008</td>
<td>ng/L</td>
<td>73.3 188 148 140</td>
<td>(1)</td>
</tr>
<tr>
<td>WWTP effluent</td>
<td>Summer</td>
<td>2009</td>
<td>ng/L</td>
<td>46.3 146 100 93.4</td>
<td>(3)</td>
</tr>
<tr>
<td>WWTP effluent</td>
<td>Winter</td>
<td>2009</td>
<td>ng/L</td>
<td>35.3 194 92.0 73.5</td>
<td>(3)</td>
</tr>
<tr>
<td>Freshwater fish</td>
<td>Fillet</td>
<td>2008</td>
<td>ng/g ww</td>
<td>&lt; 10 76 --- &lt; 10</td>
<td>(1)</td>
</tr>
<tr>
<td>Freshwater fish</td>
<td>Liver</td>
<td>2008</td>
<td>ng/g ww</td>
<td>&lt; 25 527 --- 47.6</td>
<td>(1)</td>
</tr>
<tr>
<td>Freshwater fish</td>
<td>Fillet</td>
<td>2011</td>
<td>ng/g ww</td>
<td>2.13 21.5 12.1 12.3</td>
<td>(4)</td>
</tr>
<tr>
<td>Freshwater fish</td>
<td>Whole body</td>
<td>2011</td>
<td>ng/g ww</td>
<td>3.27 91.9 35.1 22.7</td>
<td>(4)</td>
</tr>
<tr>
<td>Osprey</td>
<td>Egg</td>
<td>2008</td>
<td>ng/g ww</td>
<td>37.5 910 194 90.7</td>
<td>(1)</td>
</tr>
</tbody>
</table>

T-PFAS: sum of detected PFAS chemicals (13 PFAAs)
ww: wet weight
(1) Furl and Meredith, 2010
(2) Dinglasan-Panlilio et al., 2014
(3) Ecology and Herrera, 2010
(4) Johnson and Friese, 2012
3.1.5 Regulatory criteria or standards

No environmental regulatory criteria or standards exist for PFASs in Washington State. The EPA has issued a provisional health advisory for PFOS in drinking water of 200 ng/L (ppt). The Minnesota Department of Health issued PFOS screening levels for fish tissue at which fish consumption advisories are issued: 40 ppb (1 meal/week), >40 – 200 ppb (1 meal/month), and >800 ppb (do not eat) (MDH, 2008). Environment Canada drafted Federal Environmental Quality Guidelines (FEQGs) to assess PFOS levels in the environment (Environment Canada, 2013). PFOS levels above the FEQG indicate an increased likelihood that adverse effects may occur to fish at these levels: water (6,000 ng/L) and fish tissue (8,300 ng/g ww). The FEQG thresholds for fish tissue that indicate a risk to wildlife predators are much lower: 4.6 ng/g ww (mammalian) and 8.2 ng/g ww (avian). The Minnesota and Canadian values have no regulatory relevance to Washington State fish, but will be used when evaluating data to help provide context.

4.0 Project Description

4.1 Project goals

This project is being carried out with the following goals:

- To characterize current levels of PFASs in selected Washington State freshwater systems.
- To qualitatively assess whether concentrations and/or compound makeup has changed since the last statewide survey in 2008.
- To provide data to support PFAS chemical action plan actions and recommendations.

4.2 Project objectives

The following objectives will be carried out in 2016 to meet project goals:

- Ecology will collect surface water samples from 15 waterbodies during the spring and fall for analysis of 25 PFAS chemicals.
- Ecology will collect effluent during the spring and fall from 5 WWTPs for analysis of 35 PFAS chemicals.
- Ecology will collect osprey eggs at a subset of 3 waterbodies in the spring for analysis of 13 PFAS chemicals.
- Ecology will collect freshwater fish from 11 waterbodies in the fall for analysis of 13 PFAS chemicals in liver and fillets.
4.3 Information needed and sources

This project will generate new environmental data. Results from the 2008 PFAS survey will be used for comparisons with the 2016 data.

4.4 Target population

The target populations include surface water, osprey (*Pandion haliaetus*) eggs, and freshwater fish of selected rivers and lakes in Washington State, and final effluent from selected WWTPs in the state. Fish collections will target species obtained in the 2008 survey or those that are available at added sites: cutthroat trout, largemouth bass, largescale sucker, mountain whitefish, peammoth, pumpkinseed, rainbow trout, smallmouth bass, tench, walleye, and yellow perch. A complete list of target fish species and their scientific and family names is located in Appendix A. Table 2 displays the fish species collected in 2008, as well as those targeted in 2016. Efforts will be made to obtain two species at each waterbody covering two trophic levels (bottom feeder and predator). However, the first priority is to match species collected in 2008.

Table 2. Target Freshwater Fish Species by Location.

<table>
<thead>
<tr>
<th>Study Location</th>
<th>Species Collected in 2008</th>
<th>Species Targeted in 2016</th>
</tr>
</thead>
<tbody>
<tr>
<td>Angle Lake</td>
<td>---</td>
<td>LMB, YP</td>
</tr>
<tr>
<td>F.D.R. Lake</td>
<td>SMB, WAL</td>
<td>SMB, WAL</td>
</tr>
<tr>
<td>Lake Washington</td>
<td>LMB, LSS, PEA, YP</td>
<td>LMB, LSS, PEA, YP</td>
</tr>
<tr>
<td>Lower Columbia River</td>
<td>LMB, LSS</td>
<td>LMB, LSS</td>
</tr>
<tr>
<td>Meridian Lake</td>
<td>---</td>
<td>LMB</td>
</tr>
<tr>
<td>Mid-Columbia River (McNary)</td>
<td>---</td>
<td>LMB, LSS</td>
</tr>
<tr>
<td>Moses Lake</td>
<td>---</td>
<td>LMB, LSS</td>
</tr>
<tr>
<td>Quinault River</td>
<td>CTT</td>
<td>CTT</td>
</tr>
<tr>
<td>Snohomish River</td>
<td>---</td>
<td>MWF, PEA</td>
</tr>
<tr>
<td>Spokane River</td>
<td>LSS</td>
<td>LSS</td>
</tr>
<tr>
<td>West Medical Lake</td>
<td>PS, RBT, TCH</td>
<td>PS, RBT, TCH</td>
</tr>
</tbody>
</table>

*See Appendix A for explanation of abbreviations.

4.5 Study boundaries

Figure 1 displays the study locations for this project. All samples will be collected as close to the sampling geographic coordinates of the 2008 study as possible. At individual study locations, fish will be collected from the entire lake, or within a two river mile stretch of river. Fish collections will target areas with habitat that is most likely to contain the species of interest. Osprey egg collection sites will depend on successful nesting osprey sites identified during the nesting surveys. Water resource inventory areas and hydrologic unit codes for the waterbodies are listed in Table 3.
Table 3. Water Resource Inventory Area (WRIA) and 8-digit Hydrologic Unit Code (HUC) Numbers for the Study Area.

<table>
<thead>
<tr>
<th>Study Location</th>
<th>WRIA</th>
<th>HUC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Angle Lake</td>
<td>9</td>
<td>17110013</td>
</tr>
<tr>
<td>Lake Washington</td>
<td>8</td>
<td>17110012</td>
</tr>
<tr>
<td>Lower Columbia River</td>
<td>28</td>
<td>17080003</td>
</tr>
<tr>
<td>Snohomish River</td>
<td>7</td>
<td>17110011</td>
</tr>
<tr>
<td>South Fork Palouse River</td>
<td>34</td>
<td>17060108</td>
</tr>
<tr>
<td>West Medical Lake</td>
<td>43</td>
<td>17020013</td>
</tr>
<tr>
<td>Mid-Columbia River (McNary Dam)</td>
<td>31</td>
<td>17070101</td>
</tr>
<tr>
<td>Meridian Lake</td>
<td>9</td>
<td>17110013</td>
</tr>
<tr>
<td>Moses Lake</td>
<td>41</td>
<td>17020015</td>
</tr>
<tr>
<td>Nooksack River</td>
<td>1</td>
<td>17110004</td>
</tr>
<tr>
<td>Puyallup River</td>
<td>10</td>
<td>17110014</td>
</tr>
<tr>
<td>Spokane River</td>
<td>54</td>
<td>17010307</td>
</tr>
<tr>
<td>Upper Columbia River</td>
<td>61</td>
<td>17020001</td>
</tr>
<tr>
<td>Franklin D. Roosevelt Lake</td>
<td>53</td>
<td>17020001</td>
</tr>
<tr>
<td>Quinault River</td>
<td>21</td>
<td>17100102</td>
</tr>
</tbody>
</table>

4.6 Tasks required

The following tasks will be carried out for this project:

- Conduct desktop reconnaissance of study locations.
- Work with MEL’s QA officer to secure a contract laboratory for analysis of PFASs.
- Develop service contract for collection of osprey eggs by wildlife biologist.
- Assist consulting wildlife biologist with nesting survey and collection of osprey eggs in the spring on the lower Columbia River. Consultant will deliver osprey eggs from other two locations to Ecology HQ.
- Process osprey eggs samples and send to contract laboratory.
- Collect spring surface water and WWTP effluent samples and send to contract laboratory.
- Collect fall surface water and WWTP effluent samples and send to contract laboratory.
- Collect target fish species in the fall, process fish samples, and send to contract laboratory.
- Review data quality of laboratory results and work with MEL’s QA officer to resolve any issues.
- Write draft report summarizing results, route the draft following EAP review procedures, and publish final report.
- Load data into Ecology’s EIM database.

4.7 Practical constraints

See Section 3.1.1.
4.8 Systematic planning process

This Quality Assurance Project Plan addresses the elements of the systematic planning process.

5.0 Organization and Schedule

5.1 Key individuals and their responsibilities

Table 4. Organization of Project Staff and Responsibilities.

<table>
<thead>
<tr>
<th>Staff (all are EAP except client)</th>
<th>Title</th>
<th>Responsibilities</th>
</tr>
</thead>
<tbody>
<tr>
<td>Holly Davies HWTR program Phone: 360-407-7398</td>
<td>EAP Client</td>
<td>Clarifies scope of the project. Provides internal review of the QAPP and approves the final QAPP.</td>
</tr>
<tr>
<td>Callie Mathieu Toxics Studies Unit, SCS Phone: 360-407-6965</td>
<td>Project Manager and Principal Investigator</td>
<td>Writes the QAPP. Oversees field sampling and transportation of samples to the laboratory. Conducts QA review of data, analyzes, and interprets data. Writes the draft report and final report.</td>
</tr>
<tr>
<td>Dave Serdar/Christopher Clinton Toxics Studies Unit, SCS Phone: 360-407-6060</td>
<td>Field Lead</td>
<td>Leads field collections, records field information, and enters data into EIM.</td>
</tr>
<tr>
<td>Brandee Era-Miller Toxics Studies Unit, SCS Phone: 360-407-6771</td>
<td>Acting Unit Supervisor for the Project Manager</td>
<td>Provides internal review of the QAPP, approves the budget, and approves the final QAPP.</td>
</tr>
<tr>
<td>Jessica Archer SCS Phone: 360-407-6698</td>
<td>Section Manager for the Project Manager</td>
<td>Reviews the project scope and budget, tracks progress, reviews the draft QAPP, and approves the final QAPP.</td>
</tr>
<tr>
<td>Joel Bird MEL Phone: 360-871-8801</td>
<td>Director</td>
<td>Reviews and approves the final QAPP.</td>
</tr>
<tr>
<td>Karin Feddersen MEL Phone: 360-871-8829</td>
<td>MEL Quality Assurance Coordinator</td>
<td>Reviews draft QAPP, oversees laboratory contract, and coordinates with contract lab.</td>
</tr>
<tr>
<td>William R. Kammin Phone: 360-407-6964</td>
<td>Ecology Quality Assurance Officer</td>
<td>Reviews and approves the draft QAPP and the final QAPP.</td>
</tr>
<tr>
<td>James L. Kaiser Osprey Solutions, LLC Phone: 206-938-1600</td>
<td>Consulting Raptor Biologist</td>
<td>Coordinates logistics surrounding osprey egg collection, conducts nest surveys, and collects sample eggs.</td>
</tr>
</tbody>
</table>

EAP: Environmental Assessment Program  
EIM: Environmental Information Management database  
HWTR: Hazardous Waste and Toxics Reduction  
MEL: Manchester Environmental Laboratory  
QAPP: Quality Assurance Project Plan  
SCS: Statewide Coordination Section
5.2 Special training and certifications

Osprey eggs will be collected by the consulting wildlife biologist who has specialized experience in collecting eggs from nesting osprey in the Pacific Northwest. All Ecology field crew carrying out fish collections have specialized training in electro-shocking techniques for fish collections. Field crew conducting field measurements of temperature, pH, and conductivity must meet the personnel qualifications listed in the corresponding SOPs listed in 8.1.

5.3 Organization chart

See Table 4.

5.4 Project schedule

Table 5. Proposed Schedule for Completing Field And Laboratory Work, Data Entry into EIM, and Reports.

<table>
<thead>
<tr>
<th>Field and laboratory work</th>
<th>Due date</th>
<th>Lead staff</th>
</tr>
</thead>
<tbody>
<tr>
<td>Field work completed</td>
<td>12/2016</td>
<td>Christopher Clinton</td>
</tr>
<tr>
<td>Laboratory analyses completed</td>
<td>03/2017</td>
<td></td>
</tr>
</tbody>
</table>

Environmental Information System (EIM) database

<table>
<thead>
<tr>
<th>Product</th>
<th>Due date</th>
<th>Lead staff</th>
</tr>
</thead>
<tbody>
<tr>
<td>EIM data loaded</td>
<td>08/2016</td>
<td>Christopher Clinton</td>
</tr>
<tr>
<td>EIM data entry review</td>
<td>09/2017</td>
<td>Melissa McCall</td>
</tr>
<tr>
<td>EIM complete</td>
<td>10/2017</td>
<td>Christopher Clinton</td>
</tr>
</tbody>
</table>

Final report

<table>
<thead>
<tr>
<th>Author lead / Support staff</th>
<th>Callie Mathieu / Christopher Clinton</th>
</tr>
</thead>
<tbody>
<tr>
<td>Schedule</td>
<td></td>
</tr>
<tr>
<td>Draft due to supervisor</td>
<td>07/2017</td>
</tr>
<tr>
<td>Draft due to client/peer reviewer</td>
<td>08/2017</td>
</tr>
<tr>
<td>Final (all reviews done) due to publications coordinator</td>
<td>09/2017</td>
</tr>
<tr>
<td>Final report due on web</td>
<td>10/2017</td>
</tr>
</tbody>
</table>

5.5 Limitations on schedule

Osprey egg collection is dependent on nesting success and timing. The consulting wildlife biologist will conduct nesting surveys in mid-April 2016 to determine schedule of egg collections. The final collection date is not likely to affect the overall schedule of the project.
5.6 Budget and funding

The total budget for laboratory and contracting services for this project is $130,100. Table 6 presents the estimated costs of PFAS laboratory analyses, MEL contracting services, and contracting costs for osprey egg collection. The number of quality control (QC) samples includes only those tests that are not included in the cost of analysis. Field QC samples – replicates and equipment blanks – are included in the number of QC samples.

Table 6. Project Budget.

<table>
<thead>
<tr>
<th>Analyte Group</th>
<th>Matrix</th>
<th>Number of Samples</th>
<th>Number of QC Samples</th>
<th>Cost per Sample</th>
<th>Total Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>PFAAs</td>
<td>Water/Effluent</td>
<td>40</td>
<td>8</td>
<td>$410</td>
<td>$19,680</td>
</tr>
<tr>
<td>PFAAs</td>
<td>Fish Tissue</td>
<td>44</td>
<td>---</td>
<td>$475</td>
<td>$20,900</td>
</tr>
<tr>
<td>PFAAs</td>
<td>Osprey Eggs</td>
<td>12</td>
<td>---</td>
<td>$475</td>
<td>$5,700</td>
</tr>
<tr>
<td>FTS</td>
<td>Water/Effluent</td>
<td>40</td>
<td>8</td>
<td>$425</td>
<td>$20,400</td>
</tr>
<tr>
<td>PAPs</td>
<td>Water/Effluent</td>
<td>10</td>
<td>4</td>
<td>$500</td>
<td>$7,000</td>
</tr>
<tr>
<td>precursors</td>
<td>Water/Effluent</td>
<td>40</td>
<td>8</td>
<td>$500</td>
<td>$24,000</td>
</tr>
</tbody>
</table>

Lab subtotal: $97,680
MEL contracting costs: $24,420
Contracting costs for osprey egg collection: $8,000

Total Lab and Contracting Costs: $130,100
### 6.0 Quality Objectives

#### 6.1 Decision Quality Objectives (DQOs)

This study will not require decision quality objectives.

#### 6.2 Measurement Quality Objectives (MQOs)

Table 7. Measurement Quality Objectives.

<table>
<thead>
<tr>
<th>Analyte</th>
<th>Matrix</th>
<th>Precision</th>
<th>Bias</th>
<th>Instrument performance</th>
<th>Sensitivity</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Lab Duplicates (RPD)*</td>
<td>LCS (% recov.)</td>
<td>Method Blanks</td>
<td>Surrogate Standards (% recov.)</td>
</tr>
<tr>
<td>PFAAs</td>
<td>Tissue</td>
<td>&lt; 40%</td>
<td>70 - 130%</td>
<td>&lt; 0.5 ng/g¹</td>
<td>40 - 150%²</td>
</tr>
<tr>
<td>PFAAs</td>
<td>Water/ effluent</td>
<td>&lt; 40%</td>
<td>PFSAs: 70 - 130% PFCAs: 80 - 120%</td>
<td>&lt; 0.25 ng/sample</td>
<td>40 - 150%³</td>
</tr>
<tr>
<td>PAPs</td>
<td>Water/ effluent</td>
<td>&lt; 40%</td>
<td>40 - 160%</td>
<td>&lt; LOQ</td>
<td>n/a</td>
</tr>
<tr>
<td>FTS</td>
<td>Water/ effluent</td>
<td>&lt; 40%</td>
<td>50 - 150%</td>
<td>&lt; 1 ng/sample</td>
<td>40 - 150%</td>
</tr>
<tr>
<td>Precursors</td>
<td>Water/ effluent</td>
<td>&lt; 40%</td>
<td>70 - 150%⁴</td>
<td>&lt; 0.13 to &lt;1.0 ng/sample</td>
<td>50 - 150%⁵</td>
</tr>
</tbody>
</table>

* RPD for concentrations > 5 times reporting limit.
¹ < 1 ng/g for PFBS, PFHxS, and PFOS.
² 20 - 150% for 13C₄-PFBA; 20 - 130% 13C₈-PFOSA
³ 20 - 150% for 13C₄-PFBA
⁴ Method acceptance limits are not established. Interim limits range from 10-140% to 50-350%.
⁵ 50 - 200% for FOSAA.

RPD: relative percent difference. LCS: laboratory control sample. LOQ: limit of quantitation.

#### 6.2.1 Targets for precision, bias, and sensitivity

**6.2.1.1 Precision**

Precision is a measure of the variability in the results of replicate measurements due to random error. Laboratory analysis precision will be assessed through laboratory duplicate samples for all matrices and analyses. Table 7 shows MQOs for laboratory duplicate samples.

One field replicate per batch of surface water and WWTP effluent samples will be collected and analyzed alongside the field samples. A field replicate sample will be collected immediately
after the field sample using the same sampling technique. Field replicate relative percent difference (RPD) should be < 40% for concentrations greater than 5 times the reporting limit.

6.2.1.2 Bias

Bias is the difference between the population mean and the true value. Laboratory analysis bias will be assessed through laboratory control samples. MQOs for laboratory control sample recoveries are included in Table 7.

6.2.1.3 Sensitivity

Sensitivity is a measure of the capability of a method to detect a substance. Laboratory analysis sensitivity is defined here as the quantitation limit. See Table 7 for quantitation limits.

6.2.2 Targets for comparability, representativeness, and completeness

6.2.2.1 Comparability

Section 8.1 lists the standard operating procedures (SOPs) to be followed for field sampling. Use of SOPs for field sampling will help ensure comparability between results from this study and the 2008 study. Samples collected in 2016 will be collected as close to spring and fall sampling dates as those used in 2008 as possible; however, differences in weather and/or flow may affect comparability. Comparability of laboratory analyses for PFAAs may be affected by using a different laboratory in 2016. Slight differences exist in methods, such as different solvents used for extraction. Both studies are utilizing the same instrument (high resolution LC-MS/MS) and general analytical approach.

6.2.2.2 Representativeness

Surface water sampling is being conducted during May and October to capture spring run-off and summer low-flow conditions in rivers. Sampling at the WWTPs will occur concurrently with surface water sampling. The selected WWTPs discharge into surface waters being sampled for this study and represent a range of flow capacities and sources (domestic and domestic/industrial). Fish samples will be analyzed as three to five fish composites in order to integrate variability within a waterbody and provide a representative sample. Only one osprey egg will be collected per nest as per wildlife permits to reduce effects on nest productivity.

The study locations were chosen to represent various levels of contamination potential and to characterize PFAS levels in different watershed types.

6.2.2.3 Completeness

The project manager will consider the study to have achieved completeness if 95% of the samples are analyzed acceptably.
7.0 Sampling Process Design (Experimental Design)

7.1 Study Design

Ecology will collect samples of fresh surface water, WWTP effluent, freshwater fish tissue, and osprey eggs in 2016 throughout Washington State for analysis of PFASs. Table 8 displays the overall study design, including study locations and sample types. This study is a follow-up to Ecology’s 2008 PFAS survey and is based on locations that were sampled previously. However, the study is being expanded to include additional sites and a higher number of biota samples to capture potential contamination from sources that weren’t characterized in 2008 (such as AFFFs) and to characterize contaminant burdens in biota from urban sites, which was limited in 2008. This study will also be expanded to include additional PFAS chemicals in surface water and WWTP that were not analyzed in 2008.

Table 8. Study Locations and Sample Types.

<table>
<thead>
<tr>
<th>Study Location</th>
<th>Water Samples</th>
<th>Fish Samples</th>
<th>Osprey Eggs (# of samples)</th>
<th>Waterbody Type</th>
<th>Contamination Potential</th>
<th>Potential Sources/Pathways of Interest</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Surface Waters</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Angle Lake*</td>
<td>SP, F</td>
<td>F</td>
<td></td>
<td>Lake</td>
<td>High</td>
<td>Stormwater, AFFF</td>
</tr>
<tr>
<td>Lake Washington</td>
<td>SP, F</td>
<td>F</td>
<td>SP* (8)</td>
<td>Lake</td>
<td>High</td>
<td>Stormwater</td>
</tr>
<tr>
<td>Lower Columbia River</td>
<td>SP, F</td>
<td>F</td>
<td>SP (2)</td>
<td>River</td>
<td>High</td>
<td>WWTP, Stormwater</td>
</tr>
<tr>
<td>Snohomish River</td>
<td>SP, F</td>
<td>F*</td>
<td>---</td>
<td>River</td>
<td>High</td>
<td>WWTP, Stormwater</td>
</tr>
<tr>
<td>South Fork Palouse River</td>
<td>SP, F</td>
<td>---</td>
<td>---</td>
<td>River</td>
<td>High</td>
<td>WWTP</td>
</tr>
<tr>
<td>West Medical Lake</td>
<td>SP, F</td>
<td>F</td>
<td>SP* (2)</td>
<td>Lake</td>
<td>High</td>
<td>WWTP</td>
</tr>
<tr>
<td>Mid-Columbia River (McNary Dam)</td>
<td>SP, F</td>
<td>F*</td>
<td>---</td>
<td>Impoundment</td>
<td>Medium</td>
<td>WWTP</td>
</tr>
<tr>
<td>Meridian Lake*</td>
<td>SP, F</td>
<td>F</td>
<td>---</td>
<td>Lake</td>
<td>Medium</td>
<td>Stormwater</td>
</tr>
<tr>
<td>Moses Lake*</td>
<td>SP, F</td>
<td>F</td>
<td>---</td>
<td>Lake</td>
<td>Medium</td>
<td>AFFF</td>
</tr>
<tr>
<td>Nooksack River</td>
<td>SP, F</td>
<td>---</td>
<td>---</td>
<td>River</td>
<td>Medium</td>
<td>Atmospheric Dep.</td>
</tr>
<tr>
<td>Puyallup River</td>
<td>SP, F</td>
<td>---</td>
<td>---</td>
<td>River</td>
<td>Medium</td>
<td>WWTP</td>
</tr>
<tr>
<td>Spokane River</td>
<td>SP, F</td>
<td>F</td>
<td>---</td>
<td>River</td>
<td>Medium</td>
<td>WWTP</td>
</tr>
<tr>
<td>Upper Columbia River</td>
<td>SP, F</td>
<td>---</td>
<td>---</td>
<td>River</td>
<td>Medium</td>
<td>Atmospheric Dep.</td>
</tr>
<tr>
<td>Franklin D. Roosevelt Lake</td>
<td>SP, F</td>
<td>F</td>
<td>---</td>
<td>Impoundment</td>
<td>Low</td>
<td>Atmospheric Dep.</td>
</tr>
<tr>
<td>Quinault River</td>
<td>SP, F</td>
<td>F</td>
<td>---</td>
<td>River</td>
<td>Low</td>
<td>Atmospheric Dep.</td>
</tr>
<tr>
<td><strong>Wastewater Treatment Plants (Receiving Waters)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Marine Park (Lower Columbia River)</td>
<td>SP, F</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>Domestic/Industrial</td>
</tr>
<tr>
<td>Puyallup (Puyallup River)</td>
<td>SP, F</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>Domestic</td>
</tr>
<tr>
<td>Spokane (Spokane River)</td>
<td>SP, F</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>Domestic</td>
</tr>
<tr>
<td>West Medical Lake (West Medical Lake)</td>
<td>SP, F</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>Domestic</td>
</tr>
<tr>
<td>Pullman (South Fork Palouse River)*</td>
<td>SP, F</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>---</td>
<td>Domestic</td>
</tr>
</tbody>
</table>

*Indicates study location or sample type that was not sampled in 2008.

SP: spring
F: fall
AFFF: aqueous film-forming foam
WWTP: wastewater treatment plant

Ecology will collect surface water samples from 15 waterbodies in the spring (May) and fall (September) to assess concentrations of 25 PFASs during spring runoff and early fall low-flow.
conditions. Surface water sampling locations include rivers, lakes, and impoundments of varying watershed size and contamination potential (Table 8).

Final effluent from 5 WWTPs will be collected concurrently with surface water samples, for analysis of 35 PFAS chemicals. The WWTPs are located upstream of surface water collection locations and represent a range of flow capacities and sources (domestic and domestic/industrial). The City of Pullman WWTP was added to the sampling plan in 2016 because PFAS concentrations in surface water samples from its receiving water – South Fork Palouse River – were the second highest of all study locations in 2008. WWTP discharges from upstream municipalities, including Pullman, potentially account for the majority of total river flow during low-flow periods (Pelletier, 1993).

Ecology will collect two species of freshwater fish from 11 of the surface water sampling locations in the fall (September - November). One composite sample of liver and one composite sample of fillet tissue from each species will be analyzed for 13 PFASs. PFASs are not lipophilic and preferentially accumulate in blood, liver, and kidneys of fish (Martin et al., 2003). Ecology crew will target the same species and similar size classes as those obtained in 2008. Where possible, the species will be from different trophic levels (bottom feeder and predator). Table 2 displays target species at each location. Composite samples will consist of 3-5 individual fish. The subset of lakes targeted for freshwater fish sampling also cover a range of waterbody type, watershed size, and contamination potential.

In May, Ecology and a consulting wildlife biologist will collect osprey eggs for analysis of 13 PFASs from a subset of three study locations. One viable egg will be collected from eight individual nests along the lower Columbia River, two nests near Lake Washington, and two nests near West Medical Lake. Osprey are useful biomonitoring species, as they feed almost exclusively on fish near their nests. A total of 11 osprey eggs were collected from the lower Columbia River in the 2008 study. In 2016, two additional sites have been added to gain more information on PFAS levels at the top of the trophic chain. All three osprey egg collection waterbodies have high potential of PFAS contamination. Candidate sites for egg collection at other study locations were limited.

### 7.1.1 Field measurements

Conductivity, pH, and temperature measurements will be recorded in the field at all study locations. The latitude and longitude of sampling locations will be located by GPS and recorded. Fish total length (mm) and weight (g) will be measured and recorded in the field at time of collection. All field measurements will be recorded on project field logs.

### 7.1.2 Sampling location and frequency

Table 8 outlines the locations and timing of sample collections. Geographic coordinates for surface water, fish tissue, and WWTP effluent sampling locations are included in Appendix B. Precise sampling locations for osprey egg collection is dependent upon nesting activity identified by nesting surveys. The sampling strategy is based on the 2008 statewide survey, with additional sites and sample types for inclusion of other potential contamination sources. One sampling
event will occur per season for surface water (spring and fall), WWTP effluent (spring and fall), fish (fall), and osprey eggs (spring). All samples will be collected in 2016.

### 7.1.3 Parameters to be determined

Table 9 lists the parameters to be analyzed during this project, for each sample type.

**Table 9. Target Parameter Suites and Individual Compounds.**

<table>
<thead>
<tr>
<th>Sample Types to be Analyzed</th>
<th>Compound Group</th>
<th>Individual Compounds</th>
<th>Acronym</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surface water, WWTP effluent, fish tissue, osprey egg</td>
<td>Perfluoroalkyl acids (PFAAs)</td>
<td>Perfluorobutanoate&lt;br&gt;Perfluoroheptanoate&lt;br&gt;Perfluorodecanoate&lt;br&gt;Perfluoroundecanoate&lt;br&gt;Perfluorododecanoate&lt;br&gt;Perfluorobutanesulfonate&lt;br&gt;Perfluorohexanesulfonate&lt;br&gt;Perfluorooctanesulfonate&lt;br&gt;Perfluoroctane sulfonamide</td>
<td>PFBA&lt;br&gt;PFPeA&lt;br&gt;PFHxA&lt;br&gt;PFHpA&lt;br&gt;PFDA&lt;br&gt;PFN&lt;br&gt;PFDoA&lt;br&gt;PFBS&lt;br&gt;PFOS&lt;br&gt;PFOSA</td>
</tr>
<tr>
<td>Surface water, WWTP effluent</td>
<td>PFAA precursors</td>
<td>Perfluorooctane sulfonamido acetic acid&lt;br&gt;N-methyl perfluorooctane sulfonamido acetic acid&lt;br&gt;N-ethyl perfluorooctane sulfonamido acetic acid&lt;br&gt;6:2 fluorotelomer carboxylic acid&lt;br&gt;8:2 fluorotelomer carboxylic acid&lt;br&gt;10:2 fluorotelomer carboxylic acid&lt;br&gt;6:2 fluorotelomer unsaturated carboxylic acid&lt;br&gt;8:2 fluorotelomer unsaturated carboxylic acid&lt;br&gt;10:2 fluorotelomer unsaturated carboxylic acid</td>
<td>FOSAA&lt;br&gt;EfFOSAA&lt;br&gt;6:2 FTCA&lt;br&gt;8:2 FTCA&lt;br&gt;10:2 FTCA&lt;br&gt;6:2 FTUCA&lt;br&gt;8:2 FTUCA&lt;br&gt;10:2 FTUCA</td>
</tr>
<tr>
<td>WWTP effluent</td>
<td>Polyfluoroalkyl phosphates (PAPs)</td>
<td>Perfluorohexylphosphonate&lt;br&gt;Perfluoroctylphosphonate&lt;br&gt;Perfluorodecylphosphonate&lt;br&gt;Bis(perfluorohexyl)phosphinate&lt;br&gt;Perfluorohexyl(perfluoroctyl)phosphinate&lt;br&gt;Bis(perfluoroctyl)phosphinate&lt;br&gt;1H, 1H, 2H, 2H-perfluoroctylphosphate&lt;br&gt;1H, 1H, 2H, 2H-perfluorodecylphosphate&lt;br&gt;Bis(1H, 1H, 2H, 2H-perfluoroctyl)phosphate&lt;br&gt;Bis(1H, 1H, 2H, 2H-perfluorodecyl)phosphate</td>
<td>PFHpA&lt;br&gt;PFPA&lt;br&gt;PFDoA&lt;br&gt;6:6 PFPi&lt;br&gt;6:8 PFPi&lt;br&gt;8:8 PFPi&lt;br&gt;6:2 monoPAP&lt;br&gt;8:2 monoPAP&lt;br&gt;6:2 diPAP&lt;br&gt;8:2 diPAP</td>
</tr>
<tr>
<td>Surface water, WWTP effluent</td>
<td>Fluorotelomer sulfonates (FTS)</td>
<td>4:2 fluorotelomer sulfonate&lt;br&gt;6:2 fluorotelomer sulfonate&lt;br&gt;8:2 fluorotelomer sulfonate</td>
<td>4:2 FTS&lt;br&gt;6:2 FTS&lt;br&gt;8:2 FTS</td>
</tr>
</tbody>
</table>
7.2 Maps or diagram

Figure 1 shows the study locations for this project.

7.3 Assumptions underlying design

The study makes the assumption that quantitation limits will be low enough to characterize PFAS contamination in areas of varying contamination potential, including those reflecting atmospheric deposition as the dominant pathway.

7.4 Relation to objectives and site characteristics

Sites were selected to capture a range of potential PFAS pathways and sources in order to support the project goal of characterizing PFAS contamination in Washington State. Table 8 describes potential sources of PFASs for each study location.

7.5 Characteristics of existing data

Limitations to the 2008 PFAS study include the relatively high quantitation limits for fish tissue analyses, a limited analyte suite, and a low number of fish tissue and osprey egg samples. This study will improve on the original 2008 study in the following ways.

- Current methods allow for lower quantitation limits for PFAAs in fish tissue and osprey eggs.
- Additional PFAS parameters will be analyzed in surface water and WWTP effluent.
- Sites and sample types were added to the study design to capture additional potential sources of PFAS contamination and help characterize fish tissue concentrations in urban areas.
8.0 Sampling Procedures

8.1 Field measurement and field sampling SOPs

The field lead and field assistants will follow the protocols described within the following Ecology SOPs:

- EAP007 – Resecting Finfish Whole Body, Body Parts, or Tissue Samples (Sandvik, 2014b)
- EAP009 – Collection, Processing, and Preservation of Finfish Samples (Sandvik, 2014a)
- EAP011 – Instantaneous Measurements of Temperature in Water (Nipp, 2006)
- EAP015 – Manually Obtaining Surface Water Samples (Joy, 2006)
- EAP031 – Collection and Analysis of pH Samples (Ward, 2014a)
- EAP032 – Collection and Analysis of Conductivity Samples (Ward, 2014b)
- EAP070 – Minimizing the Spread of Invasive Species (Parsons et al., 2012)
- EAP090 – Decontaminating Field Equipment for Sampling Toxics in the Environment (Friese, 2014)

Surface water and effluent samples will be collected in laboratory-provided pre-cleaned 1 L high density polyethylene (HDPE) containers, following the SOP listed above. Samples will be collected as near-surface grabs (15-30 cm below the water surface) from as close to the thalweg as possible for rivers. Lakes will be sampled using the same near-surface grab technique from an area as far away as possible from surface water inputs and the shoreline. Samples will be retrieved with a polyethylene and stainless steel telescopic pole sampler or stainless steel Kemmerer, if deployed from bridge. The Kemmerer will be decontaminated between sampling locations with a tap water rinse and 100% methanol wash. Deionized (DI) water will not be used to clean equipment due to the possible contamination from polytetrafluoroethylene material used in the DI water purification system.

This study is designed to mimic that of the 2008 sampling. All samples will be collected as close to the coordinates of the 2008 study as possible. The 2008 study considered multi-point, depth-integrated composite samples, but it found that the majority of historical studies characterizing PFASs in surface waters were sampled in the manner described above (Taniyasu et al., 2003; Nakayama et al., 2007; Sinclair et al., 2006).

WWTP effluent samples will be collected from final dechlorinated effluent. Samples will consist of a morning and afternoon grab composite. Grabs will be taken with a lab-provided HDPE bottle and composited into a new clean HDPE bottle.

Methods for fish collections will follow the SOP listed above, using electrofishing, netting, or angling. Fish captured by these methods will be identified to species and target species will be retained if they are in acceptable condition and target size range. Adequate numbers of fish will be collected to form one 3-5 fish composite sample for each species (fillet and liver). One bottom feeder species and one predator species will be targeted at each waterbody.

Osprey eggs will be collected following U.S. Fish and Wildlife Service (USFWS) Protocol for Bird Egg Collection, Measurement, Preparation, and Shipment for Contaminant Residue.
One warm, viable osprey egg per nest will be collected by the consulting wildlife biologist. Ecology field crew staff will assist with egg collection at the lower Columbia River site, providing boat access and field assistance. One egg will be randomly selected and removed from the nest, wrapped in aluminum foil (dull side in), and placed in a zip-closure plastic bag. A label will be placed in the plastic bag with date, nest identification, location, and collector. Eggs will be stored in protective material and placed inside a cooler with blue ice for transport to Ecology Headquarters. Samples will be stored in the walk-in cooler at Ecology Headquarters (not frozen) until processing.

Fish will be collected under Ecology’s scientific collection permits from the Washington Department of Fish and Wildlife (WDFW), USFWS, and National Oceanographic Atmospheric Administration (NOAA). Ecology will obtain the necessary permits for osprey egg collection from WDFW, Oregon Department of Fish and Wildlife, and USFWS.

### 8.2 Containers, preservation methods, holding times

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Matrix</th>
<th>Minimum Quantity Required</th>
<th>Container</th>
<th>Sample Receipt and Preservation</th>
<th>Sample Storage</th>
<th>Holding Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>PFAAs</td>
<td>Fish/Osprey tissue</td>
<td>2 g ww</td>
<td>HDPE jar</td>
<td>0 - 4°C</td>
<td>-20°C</td>
<td>1 year</td>
</tr>
<tr>
<td>PFAAs</td>
<td>SW/effluent</td>
<td>1 L</td>
<td>HDPE jar</td>
<td>0 - 4°C</td>
<td>0 - 4°C</td>
<td>60 days</td>
</tr>
<tr>
<td>PAPs</td>
<td>Effluent</td>
<td>0.5 L</td>
<td>HDPE jar</td>
<td>0 - 4°C</td>
<td>0 - 4°C</td>
<td>30 days</td>
</tr>
<tr>
<td>FTS</td>
<td>SW/effluent</td>
<td>0.1 L</td>
<td>HDPE jar</td>
<td>0 - 4°C</td>
<td>0 - 4°C</td>
<td>30 days</td>
</tr>
<tr>
<td>Precursors</td>
<td>SW/effluent</td>
<td>0.5 L</td>
<td>HDPE jar</td>
<td>0 - 4°C</td>
<td>0 - 4°C</td>
<td>30 days</td>
</tr>
</tbody>
</table>

**SW:** surface water.  **ww:** wet weight.

### 8.3 Invasive species evaluation

Field staff will follow the procedures described within SOP EAP070 – Minimizing the Spread of Invasive Species (Parsons et al., 2012). The Columbia River is considered an area of extreme concern due to the documented presence of New Zealand mudsnails (NZMS). Ecology staff will schedule these waterbodies for sampling at the end of a field run and will use the following decontamination procedure: inspection, cleaning, draining, and drying.

Inspection consists of visual inspection and physical removal of invasive species and aquatic plants. This will be performed after sampling, once at the site and again at the operations center. Motors and generators will be flushed with clean water. Gill nets, the boat hull, and the boat bilge will be cleaned with hot water (60°C). Nets will be left out to dry and the bilge will be completely drained. The exposed gear will be completely dry for 2 days before the next use. In addition, field staff will make an effort to reduce contact with sediments at the areas of extreme concern, further reducing the possibility of spreading NZMS or other invasive species.
8.4 Equipment decontamination

Equipment used to collect water samples and utensils used to process fish and osprey samples will be decontaminated using the following procedure: hand washed with Liquinox soap and hot tap water, hot tap water rinse, and a final 100% methanol rinse. All other aspects of decontamination will follow Ecology’s SOP for Decontamination of Sampling Equipment for Use in Collecting Toxic Chemical Samples (Friese, 2014).

8.5 Sample ID

Sample IDs will be assigned using MEL’s work order number followed by a consecutive number. Individual fish will be assigned unique Field IDs at the time of sample collection. After processing individual fish into composite samples in the lab, a sample ID will be given using the MEL work order number.

8.6 Chain-of-custody, if required

Chain of custody will be maintained for all samples throughout the project. Samples will be stored in a cooler or freezer in Ecology’s locked HQ chain of custody room. Ecology staff will use Manchester Environmental Laboratory’s (MEL’s) chain of custody form for shipment to the laboratory.

8.7 Field log requirements

Field data will be recorded in a bound, waterproof notebook on Rite-in-the-Rain paper. Corrections will be made with single line strikethroughs, initials, and date. An electrofishing log will be filled out at each sampling location with the following information:

- Name of project
- Date(s)
- Site name
- Field personnel
- Water quality data: temperature, conductivity, pH, and visibility
- Date, time, location, ID, and description of each sample
- Weather
- Field instrument calibrations
- Main engine hours (for electro-shock boat)
- Generator hours (for electro-shock boat)
- Electrofishing shock settings
- Fish species sighted and retained per permit requirements
- Fish lengths and weights of fish retained for analysis
- Any changes or deviations from the QAPP
- Environmental conditions
- Unusual circumstances that might affect interpretation of results
8.8 Other activities

Not applicable. Necessary activities are detailed in other sections of this QAPP.

9.0 Measurement Methods

9.1 Field procedures table/field analysis table

At the time of surface water sample collection, the following will be measured in the field: temperature, pH, and conductivity. Field measurements will be taken following guidance in Ecology’s SOPs listed in Section 8.1 using a HACH HQ40d pH meter and Orion 130A conductivity meter.

9.2 Lab procedures table

Ecology will post a solicitation for bid seeking a laboratory to carry out the analyses described Table 11. The contract will be managed through MEL. The contract laboratory will be expected to meet or exceed the quantitation limits outlined below and have established methods for the target analytes using the outlined instrumentation. The contract lab will be required to report percent moisture for osprey egg analysis.

Table 11. Lab Procedures.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Sample Matrix</th>
<th>Samples [Number, Arrival Date]</th>
<th>Expected Range of Results</th>
<th>Quantitation Limit</th>
<th>Method</th>
</tr>
</thead>
<tbody>
<tr>
<td>PFAAs</td>
<td>Water/ Effluent</td>
<td>20, May 2016 20, September 2016</td>
<td>&lt; 1.0 - 1,000 ng/L</td>
<td>1.0 - 2.0 ng/L</td>
<td>LC-MS/MS; isotopic dilution</td>
</tr>
<tr>
<td>PFAAs</td>
<td>Osprey Egg</td>
<td>12, May 2016</td>
<td>&lt; 0.5 - 1,500 ng/g ww</td>
<td>0.5 - 1.0 ng/g ww</td>
<td>LC-MS/MS; isotopic dilution</td>
</tr>
<tr>
<td>PFAAs</td>
<td>Fish Tissue</td>
<td>44, November 2016</td>
<td>&lt; 0.5 - 1,000 ng/g ww</td>
<td>0.5 - 1.0 ng/g ww</td>
<td>LC-MS/MS; isotopic dilution</td>
</tr>
<tr>
<td>PAPs</td>
<td>Effluent</td>
<td>20, May 2016 20, September 2016</td>
<td>&lt; 4.0 - 1,000 ng/L</td>
<td>4.0 - 80 ng/L</td>
<td>LC-MS/MS; isotopic dilution</td>
</tr>
<tr>
<td>FTS</td>
<td>Water/ Effluent</td>
<td>20, May 2016 20, September 2016</td>
<td>&lt; 10 - 1,000 ng/L</td>
<td>10 ng/L</td>
<td>LC-MS/MS; isotopic dilution</td>
</tr>
<tr>
<td>Precursors</td>
<td>Water/ Effluent</td>
<td>20, May 2016 20, September 2016</td>
<td>&lt; 1.0 - 1,000 ng/L</td>
<td>1.0 - 8.0 ng/L</td>
<td>LC-MS/MS; isotopic dilution</td>
</tr>
</tbody>
</table>

ww: wet weight
LC-MS/MS: liquid chromatography – tandem mass spectrometry
9.3 Sample preparation method(s)

Fish samples will be processed and homogenized according to Ecology’s SOP for Resecting Finfish Whole Body, Body Parts, or Tissue Samples (Sandvik, 2014b). Composite fish samples will be composed of 3-5 individual fish fillets. Fish livers will be identified and extracted with clean scalpels and forceps after fillets have been taken from the carcass. Small squares of fillet tissue and whole livers will be placed in clean stainless steel jars and homogenized with a decontaminated stainless steel sonicator until a consistent color and texture is reached. Homogenized samples will be placed in pre-cleaned HDPE jars, frozen, and sent to the laboratory with blue ice.

After fillets and livers are removed, the sex of the fish will be determined (when possible) and recorded. Otoliths, scales, and other aging structures will be removed from fish and sent to WDFW for age determination.

Osprey eggs will be processed and homogenized following the USFWS Protocol for Bird Egg Collection, Measurement, Preparation, and Shipment for Contaminant Residue Analysis (Buck, 2008). Eggs will be assessed for cracks, cleaned gently with a soft towel and tap water at or near the temperature of the egg, and dried. Eggs will then be weighed, measured for length at their greatest dimension, and measured for volume following the water displacement technique described by Buck (2008). Eggs will be scored around the equator of the egg until the membrane is visible. The membranes will then be cut with a scalpel and egg contents transferred to a pre-cleaned stainless steel jar for homogenizing. If possible, eggshell thickness will be measured using a dial micrometer with rounded contacts. A stainless steel sonicator will be used to homogenize the egg contents until they are of consistent color and texture. Samples will then be placed in the pre-cleaned HDPE jars, frozen, and shipped to the laboratory with blue ice.

9.4 Special method requirements

The PFAS methods required for this project are newly developed and report very low concentrations. The project manager will need to work closely with the contract laboratory and MEL’s QA officer to ensure that the methods that are used meet the needs of this study.

9.5 Lab(s) accredited for method(s)

Because the PFAS analytes are non-standard and no accreditation exists, a laboratory accreditation waiver will be obtained for this project.
10.0 Quality Control Procedures

10.1 Table of field and lab quality control (QC) required

Field QC procedures for measurements of temperature, pH, and conductivity will follow the SOPs listed in Section 8.1. Table 12 provides the laboratory QC procedures required for this study.

Table 12. Laboratory Quality Control Procedures.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Matrix</th>
<th>Field</th>
<th>Laboratory</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Blanks</td>
<td>Replicates</td>
</tr>
<tr>
<td>PFAAs</td>
<td>Tissue</td>
<td>n/a</td>
<td>n/a</td>
</tr>
<tr>
<td>PFAAs</td>
<td>Water/effluent</td>
<td>1/batch</td>
<td>1/batch</td>
</tr>
<tr>
<td>PAPs</td>
<td>Effluent</td>
<td>1/batch</td>
<td>1/batch</td>
</tr>
<tr>
<td>FTS</td>
<td>Water/effluent</td>
<td>1/batch</td>
<td>1/batch</td>
</tr>
<tr>
<td>Precursors</td>
<td>Water/effluent</td>
<td>1/batch</td>
<td>1/batch</td>
</tr>
</tbody>
</table>

Batch: 20 samples or fewer. LCS: laboratory control sample

10.2 Corrective action processes

The project manager will work closely with the contract laboratory and the MEL QA Officer conducting the data review to examine data that fall outside of QC criteria. The project manager will determine whether data should be re-analyzed, rejected, or used with appropriate qualification.

11.0 Data Management Procedures

11.1 Data recording/reporting requirements

All field data and observations will be recorded on waterproof paper kept in field notebooks. Staff will transfer information contained in field notebooks to Excel spreadsheets after they return from the field. Data entries will be independently verified for accuracy by another member of the project team. Field and laboratory data for the project will be entered into Ecology’s EIM system. Laboratory data will be uploaded into EIM using the EIM XML results template.

All fish and osprey eggs collected under scientific collection permits will be reported to appropriate state and federal agencies following instructions in the permit.
11.2 Laboratory data package requirements

The contract laboratory will deliver a Tier 4 Level data package to MEL with the complete raw laboratory dataset. After reviewing the data package from the contract laboratory, MEL will provide case narratives to the project manager with the final qualified results and a description of the quality of the contract laboratory data. Case narratives should include any problems encountered with the analyses, corrective actions taken, changes to the referenced method, and an explanation of data qualifiers. Narratives will also address the condition of samples on receipt, sample preparation, methods of analysis, instrument calibration, and results of QC tests.

11.3 Electronic transfer requirements

MEL will deliver case narratives in PDF format, and electronic data deliverables in an Excel spreadsheet format, to the project manager via email.

11.4 Acceptance criteria for existing data

Data from the 2008 survey will be used if it met MQOs from the original QAPP and data quality was determined to be sufficient to meet the needs of the study.

11.5 EIM/STORET data upload procedures

All result transmittals from laboratories must be provided in an electronic data deliverable (EDD) format that meets Ecology requirements for loading to Ecology’s Information Management (EIM) database. Data will be uploaded to Ecology EIM database following internal procedures.

12.0 Audits and Reports

12.1 Number, frequency, type, and schedule of audits

MEL and contracted laboratories must participate in performance and system audits of their routine procedures. No audits are planned specifically for this project.

12.2 Responsible personnel

Not applicable. No audits are planned for this study.
12.3 Frequency and distribution of report

A draft report of the study findings will be completed in July 2017 and a final report published on the internet in October 2017. The report will include, at a minimum, the following:

- Map showing all sampling locations and any other pertinent features of the study area.
- Coordinates of each sampling site.
- Description of field and laboratory methods.
- Discussion of data quality and the significance of any problems encountered.
- Summary tables of the chemical and physical data.
- A qualitative comparison of 2016 PFAS results and the 2008 Study results.
- PFAS concentrations relative to other studies in the U.S.
- Recommendations for follow-up actions, based on study results.

Upon study completion, all project data will be entered into Ecology’s EIM system. Public access to electronic data and the final report for the study will be available through Ecology’s Internet homepage (www.ecy.wa.gov).

12.4 Responsibility for reports

The project manager/principal investigator will be the lead responsible for the final report.

13.0 Data Verification

13.1 Field data verification, requirements, and responsibilities

Field data verification will be conducted by the project manager.

13.2 Lab data verification

Data verification involves examining the data for errors, omissions, and compliance with QC acceptance criteria. MEL’s SOPs for data reduction, review, and reporting will meet the needs of the project. Data packages will be assessed by MEL’s QA Officer using the EPA Functional Guidelines for Organic Data Review (EPA, 2014).

MEL staff will provide a written report of their data review which will include a discussion of whether (1) MQOs were met, (2) proper analytical methods and protocols were followed, (3) calibrations and controls were within limits, and (4) data were consistent, correct, and complete, without errors or omissions.

The principal investigator/project manager is responsible for the final acceptance of the project data. The complete data package, along with MEL’s written report, will be assessed for
completeness and reasonableness. Based on these assessments, the data will either be accepted, accepted with qualifications, or rejected and re-analysis considered.

Accuracy of data entered into EIM will be verified by someone other than the data engineer per the Environmental Assessment Program’s EIM data entry business rules.

13.3 Validation requirements, if necessary

Independent data validation will not be required for this project.

14.0 Data Quality (Usability) Assessment

14.1 Process for determining whether project objectives have been met

After the project data have been reviewed and verified, the principal investigator/project manager will determine if the data are of sufficient quality to make determinations and decisions for which the study was conducted. The data from the laboratory’s QC procedures will provide information to determine if MQOs have been met. Laboratory and QA staff familiar with assessment of data quality may be consulted. The project final report will discuss data quality and whether the project objectives were met. If limitations in the data are identified, they will be noted.

Some analytes will be reported near the detection capability of the selected methods. MQOs may be difficult to achieve for these results. MEL’s SOP for data qualification and best professional judgment will be used in the final determination of whether to accept, reject, or accept the results with qualification. The assessment will be based on a review of laboratory QC results. This will include assessment of laboratory precision, contamination (blanks), accuracy, matrix interferences, and the success of laboratory QC samples meeting MQOs.

14.2 Data analysis and presentation methods

A summary of the data will be presented in the final report. PFAS concentrations will be qualitatively compared to results from the 2008 study. The limited number of samples analyzed in 2008 prohibits a statistical analysis of differences between the years. See Section 12.3 for more information on how the data will be presented.

14.3 Treatment of non-detects

Laboratory data will be reported down to the method detection limit, with an associated “U” or “UJ” qualifier for non-detects. Statistical tests requiring substitution for non-detects will not be included in the published report. Summed values will include only detected concentrations.
14.4 Sampling design evaluation

The number and type of samples collected will be sufficient to meet the objectives of this project.

14.5 Documentation of assessment

Documentation of assessment will occur in the final report.

15.0 References


MDH, 2008. Minnesota Department of Health Fish Meal Advice Categories Based on Levels of PFOS in Fish. MDH Fish Consumption Advisory Program.


OECD, 2013. OECD/UNEP Global PFC Group, Synthesis Paper of per- and polyfluorinated chemicals (PFCs), Environment, Health and Safety, Environment Directorate, OECD.


16.0 Appendices
## Appendix A. Names of Target Fish Species

Table A-1. Common and Scientific Names of Target Fish Species.

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Common Name</th>
<th>Scientific Name</th>
<th>Family Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>CTT</td>
<td>Cutthroat trout</td>
<td><em>Oncorhynchus clarki</em></td>
<td>Salmonidae</td>
</tr>
<tr>
<td>LMB</td>
<td>Largemouth bass</td>
<td><em>Micropterus salmoides</em></td>
<td>Centrarchidae</td>
</tr>
<tr>
<td>LSS</td>
<td>Largescale sucker</td>
<td><em>Catostomus macrocheilus</em></td>
<td>Catostomidae</td>
</tr>
<tr>
<td>MWF</td>
<td>Mountain whitefish</td>
<td><em>Prosopium williamsoni</em></td>
<td>Salmonidae</td>
</tr>
<tr>
<td>PEA</td>
<td>Peamouth</td>
<td><em>Mylocheilus oregonensis</em></td>
<td>Cyprinidae</td>
</tr>
<tr>
<td>PS</td>
<td>Pumpkinseed</td>
<td><em>Lepomis gibbosus</em></td>
<td>Centrarchidae</td>
</tr>
<tr>
<td>RBT</td>
<td>Rainbow trout</td>
<td><em>Oncorhynchus mykiss</em></td>
<td>Salmonidae</td>
</tr>
<tr>
<td>SMB</td>
<td>Smallmouth bass</td>
<td><em>Micropterus dolomieu</em></td>
<td>Centrarchidae</td>
</tr>
<tr>
<td>TCH</td>
<td>Tench</td>
<td><em>Tinca tinca</em></td>
<td>Cyprinidae</td>
</tr>
<tr>
<td>WAL</td>
<td>Walleye</td>
<td><em>Stizostedion vitreum</em></td>
<td>Percidae</td>
</tr>
<tr>
<td>YP</td>
<td>Yellow perch</td>
<td><em>Perca flavescens</em></td>
<td>Percidae</td>
</tr>
</tbody>
</table>
## Appendix B. Sampling Locations

### Table B-1. Surface Water and Fish Tissue Sampling Locations.

<table>
<thead>
<tr>
<th>Waterbody</th>
<th>County</th>
<th>WRIA</th>
<th>Surface Water Sampling Location</th>
<th>Location Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Angle Lake</td>
<td>King</td>
<td>9</td>
<td>47.428 -122.289</td>
<td>Centroid of lake at max depth (52').</td>
</tr>
<tr>
<td>FDR Lake</td>
<td>Okanogan</td>
<td>53</td>
<td>47.948 -118.905</td>
<td>F.D.R. Lake, upstream of Grand Coulee Dam, RM 601.</td>
</tr>
<tr>
<td>Lake Meridian</td>
<td>King</td>
<td>9</td>
<td>47.363 -122.154</td>
<td>Centroid of lake at max depth (90').</td>
</tr>
<tr>
<td>Lake Washington</td>
<td>King</td>
<td>8</td>
<td>47.647 -122.302</td>
<td>Lake Washington, in Seattle, at Montlake Cut, East of University of Washington Marina.</td>
</tr>
<tr>
<td>Lower Columbia River</td>
<td>Clark</td>
<td>28</td>
<td>45.695 -122.771</td>
<td>Lower Columbia River near Vancouver, RM 98.4.</td>
</tr>
<tr>
<td>McNary Dam (Columbia River)</td>
<td>Benton</td>
<td>31</td>
<td>45.940 -119.297</td>
<td>Columbia River at McNary Dam near Umatilla, OR, RM 292.</td>
</tr>
<tr>
<td>Moses Lake</td>
<td>Grant</td>
<td>41</td>
<td>47.120 -119.339</td>
<td>Centroid of lake at max depth (36') of upper basin.</td>
</tr>
<tr>
<td>Nooksack River</td>
<td>Whatcom</td>
<td>1</td>
<td>48.937 -122.442</td>
<td>Nooksack River near Lynden, RM 18.</td>
</tr>
<tr>
<td>Puyallup River</td>
<td>Pierce</td>
<td>10</td>
<td>47.198 -122.264</td>
<td>Puyallup River at Sumner, RM 10.</td>
</tr>
<tr>
<td>Quinault River</td>
<td>Jefferson</td>
<td>21</td>
<td>47.533 -123.679</td>
<td>Quinault River in Olympic National Park, RM 47.</td>
</tr>
<tr>
<td>Snohomish River</td>
<td>Snohomish</td>
<td>7</td>
<td>47.911 -122.099</td>
<td>Snohomish River at Snohomish, behind visitor's center. RM 12.5.</td>
</tr>
<tr>
<td>South Fork Palouse River</td>
<td>Whitman</td>
<td>34</td>
<td>46.760 -117.225</td>
<td>South Fork Palouse River at Armstrong Rd, 2.8 miles northwest of Pullman.</td>
</tr>
<tr>
<td>Spokane River at Nine Mile</td>
<td>Spokane</td>
<td>54</td>
<td>47.775 -117.545</td>
<td>Upstream side of Spokane River's Nine Mile Dam, RM 58.1.</td>
</tr>
<tr>
<td>Upper Columbia River</td>
<td>Stevens</td>
<td>61</td>
<td>48.922 -117.774</td>
<td>Upper Columbia River at Northport, WA, near Canadian border. RM 735.</td>
</tr>
<tr>
<td>West Medical Lake</td>
<td>Spokane</td>
<td>43</td>
<td>47.579 -117.712</td>
<td>West Medical Lake near Medical Lake.</td>
</tr>
</tbody>
</table>

WRIA: Water Resource Inventory Area  
RM: river mile
Table B-2. Wastewater Treatment Plant Effluent Sampling Locations.

<table>
<thead>
<tr>
<th>Wastewater Treatment Plant</th>
<th>Location Description</th>
<th>Latitude</th>
<th>Longitude</th>
</tr>
</thead>
<tbody>
<tr>
<td>Marine Park (Vancouver)</td>
<td>Marine Park Wastewater Treatment Facility Effluent (Vancouver, WA); discharges to Lower Columbia River.</td>
<td>45.610</td>
<td>-122.618</td>
</tr>
<tr>
<td>Pullman</td>
<td>City of Pullman Wastewater Treatment Plant; discharges to South Fork Palouse River.</td>
<td>46.739</td>
<td>-117.191</td>
</tr>
<tr>
<td>Spokane (Riverside Park)</td>
<td>City of Spokane Wastewater Treatment Plant Effluent; discharges to Spokane River.</td>
<td>47.694</td>
<td>-117.472</td>
</tr>
<tr>
<td>Sumner</td>
<td>City of Sumner Wastewater Treatment Facility; discharges to White River upstream of confluence with Puyallup River.</td>
<td>47.200</td>
<td>-122.255</td>
</tr>
<tr>
<td>West Medical Lake</td>
<td>City of Medical Lake Reclaimed Water Facility Effluent; discharges reclaimed water to West Medical Lake.</td>
<td>47.567</td>
<td>-117.703</td>
</tr>
</tbody>
</table>
Appendix C. Acronyms, Abbreviations, and Glossary

Acronyms and Abbreviations

AFFF  aqueous film forming foam
DOH  Department of Health
Ecology  Washington State Department of Ecology
e.g.  For example
EIM  Environmental Information Management database
EPA  U.S. Environmental Protection Agency
et al.  And others
FTS  fluorotelomer sulfonate
GPS  Global Positioning System
HDPE  high density polyethylene
MEL  Manchester Environmental Laboratory
MQO  Measurement quality objective
NOAA  National Oceanic and Atmospheric Administration
PAP  polyfluoroalkyl phosphate
PBTs  persistent, bioaccumulative, and toxic chemicals
PFAS  perfluoroalkyl and perfluorooalkyl substance
QA  Quality assurance
QC  Quality Control
RM  River mile
RPD  Relative percent difference
SOP  Standard operating procedures
USFS  United States Forest Service
WAC  Washington Administrative Code
WDFW  Washington Department of Fish and Wildlife
WRIA  Water Resource Inventory Area
WWTP  Wastewater treatment plant

Units of Measurement

°C  degrees centigrade
g  gram
mm  millimeter
ng/g  nanograms per gram (parts per billion)
ng/Kg  nanograms per kilogram (parts per trillion)
ng/L  nanograms per liter (parts per trillion)
ww  wet weight
Quality Assurance Glossary

Accreditation: A certification process for laboratories, designed to evaluate and document a lab’s ability to perform analytical methods and produce acceptable data. For Ecology, it is “Formal recognition by (Ecology)...that an environmental laboratory is capable of producing accurate analytical data.” [WAC 173-50-040] (Kammin, 2010)

Accuracy: The degree to which a measured value agrees with the true value of the measured property. USEPA recommends that this term not be used, and that the terms precision and bias be used to convey the information associated with the term accuracy. (USGS, 1998)

Analyte: An element, ion, compound, or chemical moiety (pH, alkalinity) which is to be determined. The definition can be expanded to include organisms, e.g., fecal coliform, Klebsiella. (Kammin, 2010)

Bias: The difference between the population mean and the true value. Bias usually describes a systematic difference reproducible over time, and is characteristic of both the measurement system, and the analyte(s) being measured. Bias is a commonly used data quality indicator (DQI). (Kammin, 2010; Ecology, 2004)

Blank: A synthetic sample, free of the analyte(s) of interest. For example, in water analysis, pure water is used for the blank. In chemical analysis, a blank is used to estimate the analytical response to all factors other than the analyte in the sample. In general, blanks are used to assess possible contamination or inadvertent introduction of analyte during various stages of the sampling and analytical process. (USGS, 1998)

Calibration: The process of establishing the relationship between the response of a measurement system and the concentration of the parameter being measured. (Ecology, 2004)

Check standard: A substance or reference material obtained from a source independent from the source of the calibration standard; used to assess bias for an analytical method. This is an obsolete term, and its use is highly discouraged. See Calibration Verification Standards, Lab Control Samples (LCS), Certified Reference Materials (CRM), and/or spiked blanks. These are all check standards, but should be referred to by their actual designator, e.g., CRM, LCS. (Kammin, 2010; Ecology, 2004)

Comparability: The degree to which different methods, data sets and/or decisions agree or can be represented as similar; a data quality indicator. (USEPA, 1997)

Completeness: The amount of valid data obtained from a project compared to the planned amount. Usually expressed as a percentage. A data quality indicator. (USEPA, 1997)

Continuing Calibration Verification Standard (CCV): A QC sample analyzed with samples to check for acceptable bias in the measurement system. The CCV is usually a midpoint calibration standard that is re-run at an established frequency during the course of an analytical run. (Kammin, 2010)
**Control chart:** A graphical representation of quality control results demonstrating the performance of an aspect of a measurement system. (Kammin, 2010; Ecology 2004)

**Control limits:** Statistical warning and action limits calculated based on control charts. Warning limits are generally set at +/- 2 standard deviations from the mean, action limits at +/- 3 standard deviations from the mean. (Kammin, 2010)

**Data Integrity:** A qualitative DQI that evaluates the extent to which a data set contains data that is misrepresented, falsified, or deliberately misleading. (Kammin, 2010)

**Data Quality Indicators (DQI):** Commonly used measures of acceptability for environmental data. The principal DQIs are precision, bias, representativeness, comparability, completeness, sensitivity, and integrity. (USEPA, 2006)

**Data Quality Objectives (DQO):** Qualitative and quantitative statements derived from systematic planning processes that clarify study objectives, define the appropriate type of data, and specify tolerable levels of potential decision errors that will be used as the basis for establishing the quality and quantity of data needed to support decisions. (USEPA, 2006)

**Data set:** A grouping of samples organized by date, time, analyte, etc. (Kammin, 2010)

**Data validation:** An analyte-specific and sample-specific process that extends the evaluation of data beyond data verification to determine the usability of a specific data set. It involves a detailed examination of the data package, using both professional judgment, and objective criteria, to determine whether the MQOs for precision, bias, and sensitivity have been met. It may also include an assessment of completeness, representativeness, comparability and integrity, as these criteria relate to the usability of the data set. Ecology considers four key criteria to determine if data validation has actually occurred. These are:

- Use of raw or instrument data for evaluation.
- Use of third-party assessors.
- Data set is complex.
- Use of EPA Functional Guidelines or equivalent for review.

Examples of data types commonly validated would be:

- Gas Chromatography (GC).
- Gas Chromatography-Mass Spectrometry (GC-MS).
- Inductively Coupled Plasma (ICP).

The end result of a formal validation process is a determination of usability that assigns qualifiers to indicate usability status for every measurement result. These qualifiers include:

- No qualifier, data is usable for intended purposes.
- J (or a J variant), data is estimated, may be usable, may be biased high or low.
- REJ, data is rejected, cannot be used for intended purposes (Kammin, 2010; Ecology, 2004).
**Data verification:** Examination of a data set for errors or omissions, and assessment of the Data Quality Indicators related to that data set for compliance with acceptance criteria (MQOs). Verification is a detailed quality review of a data set. (Ecology, 2004)

**Detection limit** (limit of detection): The concentration or amount of an analyte which can be determined to a specified level of certainty to be greater than zero. (Ecology, 2004)

**Duplicate samples:** Two samples taken from and representative of the same population, and carried through and steps of the sampling and analytical procedures in an identical manner. Duplicate samples are used to assess variability of all method activities including sampling and analysis. (USEPA, 1997)

**Field blank:** A blank used to obtain information on contamination introduced during sample collection, storage, and transport. (Ecology, 2004)

**Initial Calibration Verification Standard (ICV):** A QC sample prepared independently of calibration standards and analyzed along with the samples to check for acceptable bias in the measurement system. The ICV is analyzed prior to the analysis of any samples. (Kammin, 2010)

**Laboratory Control Sample (LCS):** A sample of known composition prepared using contaminant-free water or an inert solid that is spiked with analytes of interest at the midpoint of the calibration curve or at the level of concern. It is prepared and analyzed in the same batch of regular samples using the same sample preparation method, reagents, and analytical methods employed for regular samples. (USEPA, 1997)

**Matrix spike:** A QC sample prepared by adding a known amount of the target analyte(s) to an aliquot of a sample to check for bias due to interference or matrix effects. (Ecology, 2004)

**Measurement Quality Objectives (MQOs):** Performance or acceptance criteria for individual data quality indicators, usually including precision, bias, sensitivity, completeness, comparability, and representativeness. (USEPA, 2006)

**Measurement result:** A value obtained by performing the procedure described in a method. (Ecology, 2004)

**Method:** A formalized group of procedures and techniques for performing an activity (e.g., sampling, chemical analysis, data analysis), systematically presented in the order in which they are to be executed. (EPA, 1997)

**Method blank:** A blank prepared to represent the sample matrix, prepared and analyzed with a batch of samples. A method blank will contain all reagents used in the preparation of a sample, and the same preparation process is used for the method blank and samples. (Ecology, 2004; Kammin, 2010)

**Method Detection Limit (MDL):** This definition for detection was first formally advanced in 40CFR 136, October 26, 1984 edition. MDL is defined there as the minimum concentration of
an analyte that, in a given matrix and with a specific method, has a 99% probability of being identified, and reported to be greater than zero. (Federal Register, October 26, 1984)

**Percent Relative Standard Deviation (%RSD):** A statistic used to evaluate precision in environmental analysis. It is determined in the following manner:

\[
\%RSD = \frac{(100 \times s)}{x}
\]

where \( s \) is the sample standard deviation and \( x \) is the mean of results from more than two replicate samples (Kammin, 2010)

**Parameter:** A specified characteristic of a population or sample. Also, an analyte or grouping of analytes. Benzene and nitrate + nitrite are all “parameters.” (Kammin, 2010; Ecology, 2004)

**Population:** The hypothetical set of all possible observations of the type being investigated. (Ecology, 2004)

**Precision:** The extent of random variability among replicate measurements of the same property; a data quality indicator. (USGS, 1998)

**Quality Assurance (QA):** A set of activities designed to establish and document the reliability and usability of measurement data. (Kammin, 2010)

**Quality Assurance Project Plan (QAPP):** A document that describes the objectives of a project, and the processes and activities necessary to develop data that will support those objectives. (Kammin, 2010; Ecology, 2004)

**Quality Control (QC):** The routine application of measurement and statistical procedures to assess the accuracy of measurement data. (Ecology, 2004)

**Relative Percent Difference (RPD):** RPD is commonly used to evaluate precision. The following formula is used:

\[
[\text{Abs}(a-b)/(a + b)/2] \times 100
\]

where “Abs()” is absolute value and \( a \) and \( b \) are results for the two replicate samples. RPD can be used only with 2 values. Percent Relative Standard Deviation is (%RSD) is used if there are results for more than 2 replicate samples (Ecology, 2004).

**Replicate samples:** Two or more samples taken from the environment at the same time and place, using the same protocols. Replicates are used to estimate the random variability of the material sampled. (USGS, 1998)

**Representativeness:** The degree to which a sample reflects the population from which it is taken; a data quality indicator. (USGS, 1998)

**Sample (field):** A portion of a population (environmental entity) that is measured and assumed to represent the entire population. (USGS, 1998)

**Sample (statistical):** A finite part or subset of a statistical population. (USEPA, 1997)
**Sensitivity:** In general, denotes the rate at which the analytical response (e.g., absorbance, volume, meter reading) varies with the concentration of the parameter being determined. In a specialized sense, it has the same meaning as the detection limit. (Ecology, 2004)

**Spiked blank:** A specified amount of reagent blank fortified with a known mass of the target analyte(s); usually used to assess the recovery efficiency of the method. (USEPA, 1997)

**Spiked sample:** A sample prepared by adding a known mass of target analyte(s) to a specified amount of matrix sample for which an independent estimate of target analyte(s) concentration is available. Spiked samples can be used to determine the effect of the matrix on a method’s recovery efficiency. (USEPA, 1997)

**Split sample:** A discrete sample that is further subdivided into portions, usually duplicates. (Kammin, 2010)

**Standard Operating Procedure (SOP):** A document which describes in detail a reproducible and repeatable organized activity. (Kammin, 2010)

**Surrogate:** For environmental chemistry, a surrogate is a substance with properties similar to those of the target analyte(s). Surrogates are unlikely to be native to environmental samples. They are added to environmental samples for quality control purposes, to track extraction efficiency and/or measure analyte recovery. Deuterated organic compounds are examples of surrogates commonly used in organic compound analysis. (Kammin, 2010)

**Systematic planning:** A step-wise process which develops a clear description of the goals and objectives of a project, and produces decisions on the type, quantity, and quality of data that will be needed to meet those goals and objectives. The DQO process is a specialized type of systematic planning. (USEPA, 2006)

**References for QA Glossary**


