APPENDIX A. DATA MANAGEMENT PLAN

Table of Contents

Tables					
Acronyms					
1 Avera	iging Laboratory Duplicate or Replicate Samples	A-1			
2 Selec	tion of Preferred Results	A-1			
3 Signi	ficant Figures and Rounding	A-2			
4 Calculating Totals		A-3			
5 Calculation of PCB Congener Toxic Equivalents					
6 Calculation of Dioxin/Furan Congener TEQs					
7 Calcu	7 Calculation of Carcinogenic Polycyclic Aromatic Hydrocarbons				
8 TOC Normalization					
9 Refer	9 References				
Tables					
Table 1.	PCB congener TEF values	C-4			
Table 2.	Dioxin/furan congener TEF values	C-5			
Table 3.	cPAH PEF values	C-6			



Acronyms

AET	apparent effects threshold	
AOC	Administrative Order on Consent	
сРАН	carcinogenic polycyclic aromatic hydrocarbon	
CSL	cleanup screening level	
DDD	dichlorodiphenyldichloroethane	
DDE	dichlorodiphenyldichloroethylene	
DDT	dichlorodiphenyltrichloroethane	
EPA	US Environmental Protection Agency	
НРАН	high-molecular-weight polycyclic aromatic hydrocarbon	
HpCDD	heptachlorodibenzo-p-dioxin	
HpCDF	heptachlorodibenzofuran	
HxCDD	hexachlorodibenzo-p-dioxin	
HxCDF	hexachlorodibenzofuran	
LAET	lowest apparent effects threshold	
LDW	Lower Duwamish Waterway	
LPAH	low-molecular-weight polycyclic aromatic hydrocarbon	
OCDD	octachlorodibenzo-p-dioxin	
OCDF	octachlorodibenzofuran	
PAH	polycyclic aromatic hydrocarbon	
РСВ	polychlorinated biphenyl	
PeCDD	pentachlorodibenzo-p-dioxin	
PeCDF	pentachlorodibenzofuran	
PEF	potency equivalency factor	
QC	quality control	
RI/FS	remedial investigation/feasibility study	
RL	reporting limit	
sco	sediment cleanup objective	
SIM	selected ion monitoring	
SMS	Washington State Sediment Management Standards	
sqs	sediment quality standards	



svoc	semivolatile organic compound	
TCDD	tetrachlorodibenzo-p-dioxin	
TCDF	tetrachlorodibenzofuran	
TEF	toxic equivalency factor	
TEQ	toxic equivalent	
тос	total organic carbon	
WAC	Washington Administrative Code	
WHO	World Health Organization	



Data management rules being followed for the AOC4 pre-design investigation are the same as those applied to the RI/FS and AOC3 pre-design studies datasets, except as noted in this appendix. Rules summarized in this appendix include those for averaging duplicate or replicate samples (Section 3.1), selecting the preferred result if more than one result is reported for a chemical (Section 3.2), handling significant figures and rounding (Section 3.3), calculating totals when results are summed for individual components (Section 3.4), calculating toxic equivalents (TEQs) for polychlorinated biphenyl (PCB) congeners and dioxin/furan congeners (Sections 3.5 and 3.6, respectively), and calculating carcinogenic polycyclic aromatic hydrocarbons (cPAHs) (Section 3.7).

1 Averaging Laboratory Duplicate or Replicate Samples

Contaminant concentrations obtained from the analysis of laboratory duplicates or replicates (i.e., two or more analyses on the same sample) will be averaged for a closer representation of the "true" concentration than that provided by the results of a single analysis. Averaging rules will be dependent on whether the individual results are detected concentrations or reporting limits (RLs) for non-detected analytes. If all concentrations are detected for a given parameter, the values will be simply averaged arithmetically. If all concentrations are non-detected for a given parameter, the minimum RL will be reported. If the concentrations are a mixture of detected concentrations and RLs, any two or more detected concentrations will be averaged arithmetically, and RLs will be ignored. If there is one detected concentration and one or more RLs, the detected concentration will be reported. The latter two rules will be applied regardless of whether the RLs are higher or lower than the detected concentration.

2 Selection of Preferred Results

In some instances, the laboratory will generate more than one result for a chemical for a given sample. Multiple results can occur for several reasons, including:

- ◆ The original result does not meet the laboratory's internal quality control (QC) guidelines, and a reanalysis is performed.
- ◆ The original result does not meet other project data quality objectives, such as a sufficiently low RL, and a reanalysis is performed.
- Two different analytical methods are used for that chemical.

In each case, a single result will be selected for use. The procedures for selecting the preferred result will differ depending on whether a single or multiple analytical methods are used for that chemical.



For the same analytical method, the results will be selected using the following guidance:

- ◆ If the results are detected and not qualified, then the result from the lowest dilution will be selected, unless multiple results from the same dilution are available, in which case the result with the highest concentration will be selected.
- If the results are a combination of estimated and unqualified detected results, then the unqualified result will be selected. This situation most commonly occurs when the original result is outside of the calibration range, thus requiring a dilution. The diluted result within the calibration range will be preferentially selected.
- ◆ If the results are all estimated, then the result will be selected using best professional judgment and considering the rationale for qualification. For example, a result qualified based on laboratory replicate results outside of QC objectives for precision will be preferred to a qualified result that is outside the calibration range.
- If the results are a combination of detected and non-detected results, then the detected result will be selected. If there are more than one detected result, the applicable rules for multiple results (as discussed above) will be followed.
- If the results are all non-detected, then the lowest RL will be selected.

For different analytical methods (i.e., when a specific chemical is analyzed in the same sample using different methods), the following rules will be applied:

- ◆ For results analyzed using the semivolatile organic compound (SVOC) full-scan (EPA 8270) and selected ion monitoring (SIM) (EPA 8270-SIM) methods, the SIM results will be selected.
- ◆ For results analyzed using EPA Method 8081A and any 8270 method (i.e., hexachlorobenzene and hexachlorocyclopentadiene), the 8081A result will be selected.

The RI/FS database rules for the selection of preferred results between two methods (as described above) were revised for the compilation of the pre-design data. In the RI/FS, the preferred result was selected based on a comparison between the methods of the detection status, RL, and data qualifiers. The revised rules select the preferred result based on a preference for method.

3 Significant Figures and Rounding

The analytical laboratories report results with various numbers of significant figures depending on the instrument, parameter, and concentration relative to the RL. The reported (or assessed) precision of each observation will be explicitly stored in the



project database as a record of the number of significant figures assigned by the laboratory. The tracking of significant figures will become important when calculating averages and performing other data summaries.

When a calculation involves addition, such as totaling PCBs or polycyclic aromatic hydrocarbons (PAHs), the calculation will be only as precise as the least precise number that goes into the calculation. For example (assuming two significant figures):

210 + 19 = 229 will be reported as 230 because 19 is only reported to 2 significant digits, and the enhanced precision of the trailing 0 in the number 210 is not significant.

When a calculation involves multiplication or division, such as carbon normalization, the original figures for each value are carried through the calculation (i.e., individual values are not adjusted to a standard number of significant figures; instead, the appropriate adjustment is made to the resultant value at the end of the calculation). The result is rounded at the end of the calculation to reflect the value with the fewest significant figures used in the calculation. For example:

 $59.9 \times 1.2 = 71.88$ will be reported as 72 because there are 2 significant figures in the number 1.2.

When rounding, if the number following the last significant figure is less than 5, the digit will be left unchanged. If the number following the last significant figure is equal to or greater than 5, the digit will be increased by 1.

4 Calculating Totals

Total PCBs, total PAHs, and total fines will be calculated by summing the detected values for the individual components (e.g., Aroclor mixtures or individual congeners for total PCBs). For samples in which none of the individual components are detected, the total value will be given as the highest RL of any individual component, and assigned a U-qualifier (no detected concentrations). No sum will be calculated in cases where 50% or less of the components are analyzed. Concentrations for analyte sums will be calculated using the following components:

- ◆ Total PCBs will be calculated, in accordance with the methods of the SMS, using only detected values for all Aroclor mixtures. For individual samples in which none of the Aroclor mixtures are detected, total PCBs will be given a value equal to the highest RL of the Aroclors and assigned a U-qualifier (no detected concentrations).
- ◆ Total low-molecular-weight PAHs (LPAHs), high-molecular-weight PAHs (HPAHs), PAHs, and benzofluoranthenes will also be calculated in accordance with the methods of the SMS. Total LPAHs will be the sum of detected concentrations for naphthalene, acenaphthylene, acenaphthene, fluorene, phenanthrene, and anthracene. Total HPAHs will be the sum of detected



concentrations for fluoranthene, pyrene, benzo(a)anthracene, chrysene, total benzofluoranthenes, benzo(a)pyrene, indeno(1,2,3,-c,d)pyrene, dibenzo(a,h)anthracene, and benzo(g,h,i)perylene. Total benzofluoranthenes will be the sum of the b (i.e., benzo(b)fluoranthene), j, and k isomers.

Because the j isomer is rarely quantified, the total benzofluoranthenes sum will be typically calculated with only the b and k isomers. In cases where the laboratory provides total benzofluoranthenes instead of or in addition to the b and k isomers, the laboratory result will be reported, and no sum will be calculated. For samples in which all individual compounds within any of the three groups described above are non-detected, the highest RL for that sample will represent the sum.

• Total fines will be calculated as the sum of clay and silt fractions (i.e. $<62.5 \mu m$).

5 Calculation of PCB Congener Toxic Equivalents

PCB congener TEQs will be calculated using the World Health Organization (WHO) consensus toxic equivalency factor (TEF) values for mammals (Van den Berg et al. 1998; Van den Berg et al. 2006), as presented in Table 1. The TEQ will be calculated as the sum of each PCB congener concentration multiplied by the corresponding TEF value. When the PCB congener concentration is reported as non-detected, then the TEF will be multiplied by one-half the RL.

Table 1. PCB congener TEF values

PCB Congener No.	TEF Value for Mammals (unitless) ^a
77	0.0001
81	0.0003
105	0.00003
114	0.00003
118	0.00003
123	0.00003
126	0.1
156	0.00003
157	0.00003
167	0.00003
169	0.03
189	0.00003

^a From Van den Berg et al. (2006).

PCB – polychlorinated biphenyl

TEF - toxic equivalency factor



6 Calculation of Dioxin/Furan Congener TEQs

Dioxin/furan congener TEQs will be calculated using the WHO consensus TEF values for mammals (Van den Berg et al. 1998; Van den Berg et al. 2006) as presented in Table 2. The TEQ will be calculated as the sum of each dioxin/furan congener concentration multiplied by the corresponding TEF value. When the dioxin/furan congener concentration is reported as non-detected, then the TEF will be multiplied by one-half the RL.

Table 2. Dioxin/furan congener TEF values

Dioxin/Furan Congener	TEF Value for Mammals (unitless) ^a
1,2,3,4,6,7,8-HpCDF	0.01
1,2,3,4,6,7,8-HpCDD	0.01
1,2,3,4,7,8,9-HpCDF	0.01
1,2,3,4,7,8-HxCDF	0.1
1,2,3,4,7,8-HxCDD	0.1
1,2,3,6,7,8-HxCDF	0.1
1,2,3,6,7,8-HxCDD	0.1
1,2,3,7,8,9-HxCDF	0.1
1,2,3,7,8,9-HxCDD	0.1
1,2,3,7,8-PeCDF	0.03
1,2,3,7,8-PeCDD	1
2,3,4,6,7,8-HxCDF	0.1
2,3,4,7,8-PeCDF	0.3
2,3,7,8-TCDF	0.1
2,3,7,8-TCDD	1
OCDF	0.0003
OCDD	0.0003

^a From Van den Berg et al. (2006).

HpCDD – heptachlorodibenzo-p-dioxin
HpCDF – heptachlorodibenzofuran
HxCDD – hexachlorodibenzo-p-dioxin
HxCDF – hexachlorodibenzo-p-dioxin
HxCDF – hexachlorodibenzofuran

TCDD – tetrachlorodibenzo-p-dioxin
TCDF – tetrachlorodibenzofuran

TCDF – tetrachlorodibenzofuran

TCDF – tetrachlorodibenzofuran

TEF – toxic equivalency factor

OCDF – octachlorodibenzofuran

7 Calculation of Carcinogenic Polycyclic Aromatic Hydrocarbons

cPAH values will be calculated using potency equivalency factor (PEF) values (California EPA 2009) based on the individual PAH component's relative toxicity to benzo(a)pyrene. PEF values are presented in Table 3. The cPAH will be calculated as the sum of each individual PAH concentration multiplied by the corresponding PEF



value. When the individual PAH component concentration are reported as non-detected, then the PEF will be multiplied by one-half the RL.

Table 3. cPAH PEF values

сРАН	PEF Value (unitless) ^a
Benzo(a)pyrene	1
Benzo(a)anthracene	0.1
Benzo(b)fluoranthene	0.1
Benzo(k)fluoranthene	0.1
Chrysene	0.01
Dibenz(a,h)anthracene	0.4
Indeno(1,2,3-cd)pyrene	0.1

PEFs for cPAHs are defined by California EPA (2009) by dividing the inhalation unit risk factor for the compound by the inhalation unit risk factor for benzo[a]pyrene.

cPAH - carcinogenic polycyclic aromatic hydrocarbon

EPA - US Environmental Protection Agency

PEF - potency equivalency factor

8 TOC Normalization

For comparison to benthic cleanup goals, sediment samples with TOC content < 0.5% or > 3.5% will not be TOC normalized for comparison to the organic carbon-normalized RALs and SMS criteria (Ecology 2015). When TOC normalization is not possible and the dry weight concentration is greater than lowest apparent effects threshold (LAET) and less than or equal to 2LAET, the concentration will be considered to be greater than sediment cleanup objectives (SCOs)¹ and less than or equal to the cleanup screening level (CSL).

9 References

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¹ SCO, as defined in the 2013 SMS Rule (Washington Administrative Code [WAC] 173-204-562), is equivalent to the term sediment quality standard (SQS) used in the RI/FS (Windward 2010; AECOM 2012).



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