Second Five-Year Review Report Hudson River PCBs Superfund Site

APPENDIX 5

PCB Aroclors Data Treatment

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SECOND FIVE-YEAR REVIEW REPORT HUDSON RIVER PCBs SUPERFUND SITE

TABLE OF CONTENTS

1	INTRODU	ICTION TO PCB MEASUREMENT CONSIDERATIONS	1-1
	1.1 PCB C	hemistry	1-2
	1.2 PCB M	letrics and Analytical Methods	1-2
	1.3 Enviror	nmental Considerations	1-6
2	SEDIMEN	Т	2-1
	2.1 Historie	cal Sediment Data Sets	
	2.2 Remedial Design Investigation, Remediation and OM&M Sediment		
	Data Se	ets	
	2.2.1	M680 Homologue Data	2-4
	2.2.2	M8082 Aroclor Data	2-6
	2.2.3	Data Ranges of Interest	2-11
	2.2.4	M680 Tri+ PCB correction factor	2-12
	2.2.5	Data Selected	2-15
	2.2.6	Model Results	2-16
	2.2.7	Update to Tri+ PCB Regression during Remedial Action	
		Monitoring Phase	2-17
	2.2.8	Congener-Specific Measurements during the 2016 OM&M	
		Sampling	2-19
	2.3 Summa	ary	2-20
3	WATER		3-1
	3.1 Paired	GE mGB and Aroclor Water Column Data Set	
	3.2 Test for Regression Outliers and High Leverage Samples		

	3.3 Validity of Using Aroclors as Predictable Variables
	3.4 Regression Model Results
	3.5 Model Selection
	3.6 Summary
4	FISH
	4.1 Discussion of NYSDEC Fish Data
	4.2 Discussion of GE Fish Data
	4.3 Aroclor Measurements and Estimation of TPCB Concentration
	4.4 Development of the Homologue Equivalent Basis 4-4
	4.5 NYSDEC Data Factors
	4.6 GE Data Factors
	4.7 Fish Species Distribution
	4.8 Summary
5	REFERENCES

SECOND FIVE-YEAR REVIEW REPORT HUDSON RIVER PCBs SUPERFUND SITE

LIST OF TABLES

Table A5-1	DB-1 Chromatograph Peaks and Corresponding PCB Congeners for
	Modified Green Bay Method
Table A5-2	GE PE sample results - PE 5 - Homolog Data
Table A5-3	Summary of GE Method 680 LCS Recovery
Table A5-4	GE PE 5 Samples - Aroclor Data
Table A5-5	GE PE 2 Samples - Aroclor Data
Table A5-6	Field Duplicate RPDs
Table A5-7	Critical Tri+ PCBs Concentration Range
Table A5-8	Lower and Upper Limits for Predicted Sediment Tri+ PCB using the
	Point-by-Point Correction Regression Model
Table A5-9	Statistics of the 2009 Regression Models
Table A5-10	Regression Coefficients Update
Table A5-11	Statistics of the 2009 and 2011 Regression Models
Table A5-12	Feature Selection Measures for Several OLS Linear Regression Models
	for Water Data
Table A5-13	List of OLS and Several Robust Regression Models of Water Data
Table A5-14	Regression Results for OLS and Several Robust Regression Models for
	Water Data
Table A5-15	Cross Validation Results and the Model Prediction Error of Water Data
Table A5-16	Water Column Data Model Validation Results using Test Data Set
Table A5-17	Regression Equations for EPA's Best Model
Table A5-18	Aroclors Reported By Year as an Average Percentage of $TPCB_{Aroclor}$ –
	NYSDEC Data
Table A5-19	Aroclors Reported By Year as an Average Percentage of $TPCB_{Aroclor}$ –
	GE Data
Table A5-20	Fish Tissue Regression Equations

- Table A5-21Coefficient (α) Summary Statistics
- Table A5-22
 Average Aroclor 1221 as Percentage of TPCB_{Aroclor} by River Mile by

 Year

SECOND FIVE-YEAR REVIEW REPORT HUDSON RIVER PCBs SUPERFUND SITE

LIST OF FIGURES

Figure A5-1	PCB Structure
Figure A5-2	Regression Results for GE 2 Model (Point by Point Correction)
Figure A5-3	Measured vs Predicted Tri+ Fraction for the GE 2 Model (Point by Point
	Correction)
Figure A5-4	Regression Residuals for GE 2 Model (Point by Point Correction)
Figure A5-5	Upper and Lower Bound for Corrected vs Predicted Tri+ PCBs using
Point-by-Point	Correction Regression
Figure A5-6	OLS regression diagnostic plots for influential points
Figure A5-7a	Comparison between the predicted and measured Tri+ PCB
	concentrations for training data set among models using the three
	correlation approach
Figure A5-7b	Comparison between the predicted and measured Tri+ PCB
	concentrations for training data set among models using the one
	correlation approach
Figure A5-8	Average Percentage Aroclors Composition in NYSDEC PCB Fish Data
	between 1990 and 2011
Figure A5-9	Average Percentage Aroclors Composition in GE PCB Fish Data
	between 2004 and 2013
Figure A5-10	TPCBHE vs TPCB Aroclor - 1991 to 2000 NYSDEC Data
Figure A5-11	TPCBHE vs TPCB Aroclor for Individual Samples - 1998 NYSDEC
	Data - Enchem Environmental Laboratory
Figure A5-12	Distribution of NYSDEC 1998 TPCBHE / TPCB Aroclor Ratio for
	Individual Samples
Figure A5-13	Wilcoxon test on 1998 NYSDEC Matched Pairs - (TPCBHE -
	TPCBAroclor) for Individual Samples

Figure A5-14	Wilcoxon test on 1998 NYSDEC Matched Pairs -
	log(TPCBHE) – log(TPCB Aroclor) for Individual Samples
Figure A5-15	TPCBHE vs. TPCB Aroclor for Individual Samples
	1999-2000 NYSDEC Data – Mississippi State Chemical Laboratories
Figure A5-16	Distribution of 1999-2000 NYSDEC TPCBHE / TPCB Aroclor Ratio
	for Individual Samples
Figure A5-17	Wilcoxon test on 1999-2000 NYSDEC Matched Pairs
	(TPCBHE - TPCB Aroclor) for Individual Samples
Figure A5-18	Wilcoxon test on 1999-2000 NYSDEC Matched Pairs
	log(TPCBHE) – log(TPCB Aroclor) for Individual Samples
Figure A5-19	TPCBHE vs. TPCB Aroclor for Individual Samples
	2004 to 2008 GE Data - Northeast Analytical Laboratory
Figure A5-20	Distribution of 2004 to 2008 GE TPCBHE / TPCB Aroclor
Figure A5-21	Wilcoxon Test on 2004-2008 GE Matched Pairs
	(TPCBHE - TPCB Aroclor) for Individual Samples
Figure A5-22	Wilcoxon Test on 2004 to 2008 GE Matched Pairs
	log(TPCBHE) – log(TPCB Aroclor) for Individual Samples
Figure A5-23	TPCBHE vs. TPCB Aroclor for Individual Samples
	2009 to 2013 GE Data - Northeast Analytical Laboratory
Figure A5-24	Distribution of 2009 to 2013 GE TPCBHE / TPCB Aroclor for
	Individual Samples
Figure A5-25	Wilcoxon Test on 2009 to 2013 GE Matched Pairs
	(TPCBHE - TPCB Aroclor) for Individual Samples
Figure A5-26	Wilcoxon Test on 2009 to 2013 GE Matched Pairs
	log(TPCBHE) – log(TPCB Aroclor) for Individual Samples
Figure A5-27	Fish Species Distribution in Matched Pair Data Set for NYSDEC Data
Figure A5-28	Fish Species Distribution in Matched Pair Data Set for GE Data

1 INTRODUCTION TO PCB MEASUREMENT CONSIDERATIONS

Measurement of PCB concentrations has evolved over time as analytical technology has improved, increasing the understanding of their physical properties and their fate in the environment. A dramatic change in analytical methods has occurred between more recent data, obtained using state-of-the-art, capillary-column, PCB congener analyses, and older analyses based on packed-column quantitation of Aroclor equivalents. Valid interpretation of trends in PCB concentrations cannot be made without consideration of the changes in analytical methods which have occurred over time. That is, a comparison is valid only when there is consistency in what is being measured. Most of the data collected for the Hudson River were analyzed using the Aroclor-based analytical method. However, EPA, the New York State Department of Environmental Conservation (NYSDEC), and General Electric Company (GE) have collected enough samples analyzed by both Aroclor-based and congener-based methods to develop a translation scheme to convert the Aroclor-based data so that it is consistent with congener-based quantitation. For the Hudson River PCBs Site, translation schemes (or regression equations) were developed for sediment, water and fish samples to convert the results from the various Aroclor-based analytical methods to a standard metric. By standardizing the PCB measurements to a congener-based equivalent quantitation, it is possible to compare the various data sets over time on a consistent basis. In this manner, variations among the various data sets through time can be attributed to temporal changes in the environment and not to changes in PCB analytical methods. These regressions also provide a standard statistical basis to estimate the uncertainties associated with these conversions when assessing temporal trends.

The purpose of this appendix is to summarize the translation schemes (using a regression factor approach) to convert Aroclor-based measurements to congener-based quantitation in sediment, water and fish. Section 1 of this appendix discusses PCB chemistry and the different analytical methods to measure PCB concentrations. Sections 2 through 4 discuss translation schemes in sediment, water and fish, respectively.

1.1 PCB Chemistry

Polychlorinated biphenyls (PCBs) are a group of man-made organic chemicals consisting of carbon, hydrogen and chlorine atoms. Each PCB consists of a biphenyl molecule with a specific number of attached chlorine atoms. The number of chlorine atoms and their location on a PCB molecule determine many of its physical and chemical properties. There are 209 distinct PCB compounds known as congeners. A PCB congener is any single, uniquely structured chemical compound in the PCB category. The name of a congener specifies the total number of chlorine substituents, and the position of each chlorine. For example: 4,4'-dichlorobiphenyl is a congener comprising the biphenyl structure with two chlorine substituents - one on each of the #4 carbons of the two rings (Figure A5-1). In 1980, a numbering system was developed which assigned a sequential number to each of the 209 PCB congeners.

1.2 PCB Metrics and Analytical Methods

PCBs were manufactured and sold as complex mixtures of several congeners with a variety of trade names, including Aroclor (e.g., Aroclor 1242). An Aroclor PCB mixture might consist of over 100 different individual PCB congeners, although 10 to 20 congeners might make up over 50 percent of the mixture. One of the most common ways to measure PCBs is a method based on the original industrial Aroclor mixtures, EPA Method 8082 [M8082]. When M8082 is employed to determine whether a sample has an Aroclor PCB mixture in it, the analytical chemist looks for a distinctive gas chromatographic pattern that is indicative of one of the Aroclors. There are nine common PCB Aroclor mixtures (1016, 1221, 1232, 1242, 1248, 1254, 1260, 1262, and 1268). Each of them has a distinctive gas chromatographic pattern.

Measuring PCBs as Aroclors relies on a relatively fixed composition of congeners in the mixture. M8082 uses a pattern recognition technique to qualitatively determine whether or not a given Aroclor mixture is present, after which that portion of the spectrum is quantified using a standard which includes the particular Aroclor. This process references certain well identified PCB peaks and compares them to the standard to determine Aroclor concentrations in the sample. Provided the sample has not been subjected to conditions that

might degrade or change the composition of the PCBs, quantitation of PCB Aroclor using M8082 will give information on the total concentration of PCBs (TPCBs¹) in the sample, but not the identity or the concentrations of the individual PCB congeners present. However, if an environmental sample has been subject to degradation, weathering or dechlorination, Aroclor-based analysis may over- or under- estimate the actual Aroclorrelated PCB concentrations since the apparent Aroclor mixture in the environmental sample may not contain the same suite of congeners or the same proportions of congeners as the standard Aroclor. In such a case, even if PCB congeners originally present in an Aroclor are present, that particular PCB Aroclor may be reported as not detected due to a lack of pattern recognition and/or the mixture may be quantified as a different Aroclor. It can also be difficult to determine a total PCB concentration using the Aroclor approach when environmental degradation or weathering has occurred. This is especially true when more than one Aroclor is determined to be present. As the individual Aroclors represent overlapping mixtures of PCB congeners, there is a possibility that "double counting" of PCBs could also occur. However, since Aroclor-based analyses do not quantitate all PCBrelated peaks in the sample chromatogram, it is also possible that Aroclor-based analysis can under-report PCB concentrations. Thus, analytical Aroclor quantitations on environmental samples are not directly comparable to actual concentrations of PCB congeners.

Environmental samples such as air samples, sediment samples, water samples and biota samples are most likely to have had their congener composition changed by environmental conditions compared to non-environmental samples. This happens because the PCB congeners with fewer chlorine atoms tend to partition into air and water more readily than those with more chlorine atoms. Biota samples can also be subjected to biodegradation with some congeners being selectively degraded or eliminated and others persisting in the animal. For samples whose congener compositions have been substantively altered,

¹ Total PCBs represents the sum of all measured PCB congeners. PCBs are a group of chemicals consisting of 209 individual compounds known as congeners. The congeners can have from one to ten chlorine atoms per molecule, each with its own set of chemical properties.

analytical testing for PCB homologues will give more reliable results than testing for Aroclors. Homologues are a way of grouping PCB congeners based on the number of chlorine substituents, which vary from one to ten. The PCB congeners that have the same number of chlorine atoms belong to the same homologue group. For example, there are 24 different tri-chloro congeners in the 3-chlorine homologue group. Laboratory results for PCB homologues will report the amount of PCBs present in the sample for each homologue group.

EPA Method 680 [M680] (PCB Homologues) is a gas chromatography/mass spectrometry (GC/MS) procedure that quantifies PCB homologues. This method has the advantage of quantifying PCBs that may not be in the form of Aroclors, as well as original Aroclor mixtures that have been weathered, or were misidentified or otherwise not detected by M8082. While M608 provides an advantage over M8082 in this regard, in some circumstances, more detailed quantification of specific congeners is needed. In those cases, PCB congeners can be accurately quantified by EPA Method 1668 [M1668] which determines the concentrations of individual congeners by a sophisticated analytical method using high-resolution gas chromatography/high resolution mass spectrometry (HRGC/HRMS) combined with isotope dilution techniques. This method requires no presumptions regarding the PCB source material; EPA M1668 identifies the presence and concentration of each of the 209 PCB congeners in a sample.

For the Hudson River PCBs Superfund Site, during the Phase 2 RI/FS investigation in the 1990s, EPA developed a program-specific method which used a dual capillary column gas chromatograph with election capture detectors (GC/ECD) to analyze PCB congeners. This method was based on the NYSDEC Analytical Services Protocol Method 91-11 (NYSDEC, 1989) for PCB congeners. This technique employs the use of two independent capillary columns with unique resolution capabilities for PCB congener separation, allowing for coelution on the first GC analytical column to be potentially resolved on the second GC analytical column. The dual column GC/ECD allows separation of a larger number of PCB congeners. EPA Phase 2 PCB analyses quantified 126 individual PCB congeners out of 209 possible PCB congeners (General Electric 1997).

GE employed an analytical technique which lies somewhere between the relatively simple M8082 and the highly sophisticated M1668. The GE technique involves the extraction of PCBs from the sample matrix, GC separation of PCB congeners on a DB-1 capillary column, and quantification with an electron capture detector. Calibration of the DB-1 column is based on the method developed by the EPA under the Green Bay Mass Balance Study (EPA, 1987). GE's modification of the original Green Bay Method [mGB] involves GC standardization using a 25:18:18 mixture of Aroclors 1232, 1248, and 1262. Individual DB-1 peak response factors (RFs) are calculated based on standard peak weight percent values originally developed by EPA (EPA, 1987). These RFs are then used to calculate PCB content of environmental samples. The DB-1 column separates PCBs into 118 unique chromatographic peaks. Several of these peaks contain multiple (coeluting) congeners. DB-1 PCB peaks have been mapped to the corresponding PCB congeners; see Table A5-1 for chromatograph peaks and the corresponding PCB congeners. Peak compositions are based on Aroclor data published in Frame et al. (1996). Another modification is due to comparison of water column PCB concentrations in samples collected by GE in 1993 with those measured by EPA as part of the Phase 2 study from the Fort Edward and Thompson Island Dam monitoring stations which suggested that the Green Bay Method results are biased low (General Electric1997). The GE study in 1997, which examined the dechlorination products, suggested that the analytical biases were manifested in individual PCB congeners, predominantly in Peak 5 (which consists of PCB congeners 2,2'dichlorobiphenyl $(BZ#4)^2$ and 2,6-dichlorobiphenyl (BZ#10). The congener distribution (predominantly peak 5 components) within the Green Bay mixed Aroclor standard was apparently miscalculated, as a revision to the calibration was later published (EPA, 1994). This error introduced systematic analytical biases in the GE data because underestimation of the Peak 5 weight percent in the DB-1 calibration standard caused measured Peak 5 values in Hudson River environmental samples to be underestimated (i.e., biased low). Since the error is in the calibration standard composition, not the PCB mass, it affects data for all DB-1 peaks (i.e., low bias in PK5 requires that other peaks are biased high). GE

² BZ refers to the numbering system for PCB congeners developed by Ballschmitter and Zell (1980)

analysis showed that biases are evident in DB-1 peaks 5, 8, and 14. Therefore, GE made a revision for the DB-1 peak response factors used to calculate PCB mass in each peak of an environmental sample. GE developed calibration error correction factors for water, sediment, and biota samples separately (HydroQual 1997). Later in 2010, as directed by EPA, these correction factors were no longer used in the Phase 2 Remedial Action Monitoring Program (RAMP) for water column samples. Instead, the mGBM has been updated to include a second column (CP-SIL5-C18) analysis for the dichlorobiphenyl congeners BZ #4 and BZ #10. The second column analysis was used to achieve a more accurate quantification for PCB congeners BZ#4 and BZ#10 (that co-elute in mGBM peak 5) by achieving full resolution and individual measurement for these two congeners. Correction factors were no longer utilized for mGBM DB-1 Peaks 8 and 14 due to their relatively minor contribution to TPCB (General Electric 2011a). GE's mGBM was used by GE through 2016 to provide congener information on a subset of samples collected for water column and fish tissue monitoring programs.

Among the analytical methods described above, the Aroclor method (M8082) is readily available from many commercial labs and relatively inexpensive. EPA M680 can be a cost-effective option for characterizing contaminated samples for PCB congeners that have undergone environmental degradation; this homologue-based method provides a more representative option for the determination of TPCBs than the Aroclor method. GE's mGBM method is available on a limited basis; only one laboratory can perform the analysis. EPA M1668 is a highly specialized analytical method and can achieve lower quantitation limits, but generally at higher cost (about five times more expensive than M8082).

1.3 Environmental Considerations

Several studies conducted in the 1980s documented PCB dechlorination in the sediments of the Upper Hudson (for example, Brown et al. 1984). Dechlorination is the chemical process of removing one or more chlorine atoms from a chemical structure, in this case a PCB molecule. The dechlorination process largely serves to change the nature of the PCBs, affecting both the geochemical and toxicological properties of the mixture. In general, dechlorination converts the PCBs to a more soluble form. The dechlorination transforms PCB congeners from those commonly found in Aroclors to lighter congeners that were virtually absent from the Aroclor composition originally manufactured.

Because of the dechlorination process in the Hudson River PCBs Site, PCB contamination within the sediments, water, and biota cannot not be accurately quantified using an Aroclorbased analytical method alone. Additionally, it was recognized that individual PCB congeners have different geochemical properties and thus would redistribute themselves among sediments, water, and biota, tending to confound the original Aroclor-related distribution. From these considerations, a congener-specific analytical method is clearly preferred, since variations in the congener mixtures could be used to identify PCB sources, as well as important geochemical transformations. As stated previously, most of the data collected for the Hudson River is in the form of Aroclors, a translation scheme is required to make the Aroclor data consistent with congener-based quantitation. Since both GE and EPA have collected enough both Aroclor-based and congener-based samples, translations schemes to convert the Aroclor-based data to the congener-based quantitation could be developed. The Aroclor samples were corrected to a common congener-based metric so that data could be accurately compared across time.

The following sections discuss the translation schemes (using a regression factor approach) to convert Aroclor-based measurements to congener-based quantitation in sediment, water and fish. Note that selection of PCB metrics for measurement, modeling, and evaluation was based on risk assessment considerations. The tri-homologue and higher (i.e., Tri+) group of compounds is expected to include the PCB congeners that are most toxic to fish, wildlife and humans, and is therefore considered to reflect a metric that captures most of the toxicity associated with PCBs. Historical quantitation of PCBs was done on an Aroclor basis; an analysis of these data show that the sum of particular Aroclors is equivalent to the trichloro and higher congeners (Tri+ PCB) (EPA 2000) and that Tri+ PCBs³ is essentially

³ Tri + PCBs represents the sum of all measured PCB congeners with three or more chlorine atoms per molecule. PCBs are a group of chemicals consisting of 209 individual compounds known as congeners.

the same as the sum of all PCBs (TPCBs) in fish tissue, the main pathway for human exposure. Thus, Tri+ PCB is a good representation of the PCBs that bioaccumulate in fish. As a result, estimates of both TPCBs and Tri+ PCB obtained from both congener and Aroclor-based analyses were used as the basis to assess PCB contamination in sediment, water and fish for the Hudson River.

The congeners can have from one to ten chlorine atoms per molecule, each with its own set of chemical properties.

2 SEDIMENT

The Hudson River is one of the most extensively monitored PCB contamination sites. The system has been studied extensively and monitored over a period of more than 30 years. The various monitoring studies provided numerous sediment data sets. This section discusses translation schemes to convert Aroclor-based measurements to congener-based quantitation in the sediment for various sediment data sets.

2.1 Historical Sediment Data Sets

Two historical large-scale sediment investigations were conducted by NYSDEC; one in 1976 to 1978 (reported in Tofflemire and Quinn, 1979), and one in 1984 (reported by Brown et al., 1988). The 1976 to 1978 sampling covered the area from Fort Edward to Troy (RM 194.8 to RM 154); whereas the 1984 sampling was restricted to the Thompson Island Pool (TIP) (RM 194.6 to RM 188.5). The 1984 sediment survey (Brown et al., 1988) represents the most comprehensive database on PCB concentrations in Thompson Island Pool sediments prior to the 2002-2005 GE remedial design investigation, with over 1,200 samples collected on a triangular grid with 125-foot centers. The spatial coverage of that effort was adequate for the purpose of estimating PCB sediment mass inventory, as discussed in Brown et al., 1988 and EPA 1997. The 1976-1978 and 1984 data were needed to identify likely areas for remediation and provide estimates of PCB mass and sediment volumes for the 2002 Record of Decision (ROD) for the Upper Hudson River.

Prior to using the 1976-1978 NYSDEC data, the comparability of these sediment data was evaluated as part of the Phase 2 RI/FS investigation. A good level of agreement between the 1976-1978 and 1994 Phase 2 conditions serves to support internal consistency of the various sediment classification data sets. TPCBs were reported by O'Brien and Gere for the 1976-1978 sediment data set. These were based on Aroclor analysis using a limited number of packed column peaks, which tended to miss the mono- and di-homologues. Based on reconstruction of the 1976-1978 total PCB results from USEPA Phase 2 sediment congener data, a regression between the Tri+ concentration and the 1977-1978 total PCB concentrations produced a zero-intercept model with which to estimate Tri+ concentrations from these data (Equation 2-1). Details of this analysis are presented in EPA 1998 and

Butcher (2000). The following relationship was used to calculate Tri+ PCB for the 1976-1978 NYSDEC data (USEPA, 2000):

$$Tri + (1977) = 1.131 \times [Aroclor 1016 + 1254]$$
 Eqn. 2-1

PCB concentrations reported by NYSDEC for the 1984 Thompson Island Pool sediment survey were dependent on the Aroclor quantitation methods used and were not equivalent to results which would be obtained using capillary column GC analysis for PCB congeners. It was thus crucial to understand what is reported in these data and estimate how well the NYSDEC-reported total represented actual total PCBs that would have been calculated by summing congener concentrations. As part of the Phase 2 RI/FS investigation, a study was made of the differences between the two techniques. This is documented in Appendix E of the Low Resolution Sediment Coring Report (EPA1998), which describes the quantitation issues relating the 1994 Phase 2 and 1984 NYSDEC PCB data. The recommendation of this analysis was to use the 1984 quantitation of total PCB as representative of the sum of congeners in the trichloro through decachloro homologue groups. A linear relationship was developed to adjust (or correct) the 1984 NYSDEC data to a basis consistent with the sum of tri- and higher-chlorinated congeners (Tri+) in the 1994 EPA data. The following equation was used to calculate Tri+ PCB for the 1984 NYSDEC data (EPA 1998):

$$Tri+(1984) = 0.944 \times 1984 Aroclor Sum$$
 Eqn. 2-2

In 1991 and 1998, GE also conducted sediment surveys and the data were reported as PCB congeners based on the mGBM. The GE 1991 survey sampled the upper river from Fort Edward to Federal Dam, while the 1998 survey sampled the TIP only. The sediment survey conducted in the TIP by GE in 1998 attempted to 'repeat' portions of the 1991 O'Brien and Gere and 1994 EPA sediment surveys GE 1999). GE 1991 and 1998 data were used in the modeling effort during Phase 2 RI/FS investigation.

During Phase 2 RI/FS investigation, EPA collected low resolution cores in 1994. These data were analyzed based on congener-specific standards as described in Section 1.2.

Rather than resurvey the entire TI Pool, the 1994 low resolution coring effort focused on replicating a representative subset of the 1984 locations. Since the EPA 1994, GE 1991, and 1998 data were reported as PCB congeners, no translation scheme was needed to calculate the Tri+ PCB concentrations.

2.2 Remedial Design Investigation, Remediation and OM&M Sediment Data Sets

During the remedial design investigation phase, GE collected sediment samples under the Sediment Sampling and Analysis Program (SSAP) in 2002-2005. These data were collected for delineating the final areas for removal. Tri+ PCB concentrations were not measured directly on the SSAP samples as a cost-saving measure for GE since the M8082 (Aroclor-based) analysis is substantially less expensive than the M680 (homologue-based) analysis. Given that the SSAP comprised some 30,000 sediment samples, EPA allowed GE to analyze all samples by M8082 and analyze only a subset by the EPA-approved M680, and then use the comparison of the paired analytical results to estimate the Tri+ PCB fraction in the remaining samples without paired analytical results.

Given the extensive the use of M8082 during the SSAP, as well as the short analytical turnaround time for M8082, GE continued to use M8082 throughout the remediation itself to satisfy the residual performance standard sampling requirements. M8082 was also used to conduct the Downstream PCB Deposition Study (DDS) during 2011 through 2013. To maintain consistency from the SSAP to the present, GE continued to use M8082 to establish the baseline sediment conditions at the beginning of the OM&M period in 2016.

In all applications of M8082, the determination of the TPCB concentration in a sample was simply the summation of all detected Aroclors in a sample. This sum was shown to be well correlated with the sum of PCB congeners determined by the mGBM as part of the SSAP (General Electric 2004). Also as part of the SSAP, GE standardized the reporting of Aroclors under M8082 to four specific Aroclors, Aroclors 1221, 1242, 1254 and 1260, and maintained consistent reporting throughout the SSAP, remediation, DDS and OM&M sampling. The basis for estimating Tri+ PCB concentrations from M8082 throughout the period 2002 to 2016 was developed from a regression relating M8082 and M680 based on

matched sample pairs obtained during the SSAP and is described below. Data quality was also tracked throughout this period and is also described below.

The quality of the regression for Tri+ determination is contingent upon the underlying data – i.e., the M680 (homologue) and M8082 (Aroclor) data. To assess the accuracy of the M680 and M8082 methods, laboratory control (LCS) and performance evaluation (PE) samples were analyzed by laboratories used by GE during the remedial design investigation. There are two measurements of "known" TPCB (sum of Aroclors) concentrations in the suite of QA/QC measurements performed by GE. One is the LCS sample analysis (A1242 spiked into a clean matrix), and the other is the analysis of PE-5 (comprised of known concentrations of A1221 and A1242). Since the LCS and PE5 results are of known composition, they can be used to evaluate the Aroclor distribution and to assess the accuracy of the analytical methods. PE-5 and LCS samples were analyzed using both M680 and M8082. In addition to the LCS and PE-5 samples, GE also prepared and analyzed additional PE samples. Sediment samples from the Upper Hudson River at 4 different locations (PE-1, PE-2, PE-3, and PE-4) were used to provide a range of PCB concentrations. Sections 2.2.1 and 2.2.2 summarize the results of QA/QC data analyzed during the remedial design investigation period.

2.2.1 M680 Homologue Data

The critical component of the M680 data is how well the reported homologue distribution (or more specifically, the Tri+ fraction) accurately reflects that of the sample analyzed and how they are compared to a known concentration sample. The M680 results are discussed below.

PE Sample Results

PE samples were analyzed concurrently with the Year 1 and Year 2 SSAP. As noted above, PE-5 is a manufactured standard, so there is information on its true or known composition. Fifteen replicates of PE5 were analyzed with the Year 1 and Year 2 SSAP. The standard deviation (SD) and relative standard deviation (RSD) were calculated for TPCBs, Tri+ PCBs, fraction Tri+, and for each homologue group (see Table A5-2). The SD and RSD are both measures of precision; the more precise data will result in smaller SDs and RSDs. As shown on Table A5-2 (M680-all samples), the RSDs are typically about 0.16 (16 percent) for most parameters. However, the RSD for monochlorobiphenyl (MonoCB) is much higher (0.345); by inspection, it can be seen that there are two anomalous low values for MonoCB (about 3 mg/kg; with the results for the other 13 samples ranging from 7 to 16 mg/kg). When these two samples are excluded (M680 - Two Samples Excluded), the MonoCB RSD becomes similar to that of the other parameters; and the Tri+ RSD reduces to 0.090.

LCS Sample Results

A total of 79 LCS samples were analyzed for PCB homologues; all homologue analyses were performed by Lab 15 (NEA). Of the 79, 28 are Year 1 and 51 are Year 2 LCS analyses. There was some difference by year, with the median Year 1 sum of homologue recovery being about 83 percent, and the median Year 2 recovery about 73 percent. The LCS data were reviewed to see if the homologue distribution might account for the low recovery of total homologs, relative to TPCBs. In virtually all the Year 1 LCS samples analyzed by GE, the homologues reported as present were di, tri, and tetra PCBs; and the median Tri+ fraction was 0.84. This fraction is in very good agreement with literature values, which suggest that about 85 percent of A1242 is Tri+. (A1242 has one percent or less monoCB, so the fact that mono was not reported as present by GE/NEA is not significant.) The homologue distribution of the Year 2 LCS samples was spot-checked; the Tri+ fraction ranged from 80 to 87 percent, with a median of 84 percent, in the nine samples checked. The overall median recovery (Year 1 and Year 2 combined) for LCS homologs is 75.8 percent.

Field Duplicate Results

The field duplicate results are presented here to show the precision of the method. A total of 79 field duplicate pairs were analyzed for homologues. As only samples considered likely to have detectable concentrations of PCBs were selected for homologue analysis (and in fact, PCBs were detected in all the samples), all of the field duplicate pairs were included in the review. Of the 79 field duplicate pairs, 65 (about 82 percent) met GE's precision criterion of \leq 40 percent RPD (relative percent difference) for TPCBs (sum of

detected homologues). The RPD for TPCBs was less than 100 percent in all but four of the samples (i.e., in 95 percent); and the median RPD for TPCBs is about 17 percent. About 78 percent of the Tri+ PCB (concentration data) met the \leq 40 percent RPD criterion, with a median value of about 17 percent (the same as for TPCBs). Precision data for individual homologs groups were also calculated for mono- through tetra-chlorobiphenyls. The median precision for each of the homologs ranged from about 16 to 20 percent RPD.

In addition to TPCBs and Tri+ PCB concentrations, the fraction Tri+ was calculated for each of the samples and the precision of the Tri+ fraction data was assessed. The data show that the Tri+ fraction is quite reproducible. The highest RPD for Tri+ fraction was 41 percent (this was the only one of the 79 samples with an RPD over 40 percent); and the RPDs were less than 20 percent for 95 percent of the samples, with a median RPD for Tri+ fraction of less than 6 percent.

<u>Summary</u>

In summary, in every instance where the Tri+ mass fraction can be calculated from sample or QC data, the results show a higher degree of precision in the estimate of this fraction than in the absolute estimate of its concentration as measured by M680. This result supports the choice of the Tri+ conversion model described previously. As described previously, LCS and PE5 sample results indicate a systematically low recovery (low reported value) relative to the known concentrations for these samples. Despite the low bias, the results for these samples as well as other QA/QC samples and the field duplicates appear to be precise with mean RSDs or RPDs on the order of 0.16 to 0.20 (16 to 20 percent) for individual parameters. Better precision for the Tri+ mass fraction as well as the major homologue fractions DiCB and TriCB was evidenced by the lower RSDs and RPDs, on the order of 0.05 (5 percent).

2.2.2 M8082 Aroclor Data

For the Aroclor data, the critical components are the accuracy and precision of TPCB concentration, and the correct assignment (identification) of Aroclors. The other critical component of the M8082 data is the assignment of PCBs to individual Aroclors.

QA/QC Sample Results

The LCS samples analyzed for this program consisted of a low concentration (typically about 1.2 mg/kg) Aroclor spike into a clean matrix. Overall, GE laboratories recovered close to 100 percent of the spiked concentration based on the median recovery value (see Table A5-3). There was some variability by laboratory, with median LCS recovery values ranging from 89 percent for Lab 1 to 117 percent for Lab 16 (Lab 16 also had the fewest LCS data points - only 35); the median Aroclor LCS recovery for Lab 15 (the lab which did all the homolog analyses) was 101 percent. The Aroclor LCS data were reviewed by year. The aggregate median recovery for the Aroclor LCS recoveries is 95 percent for Year 1 (n= 471) and 100 percent for Year 2 (n = 1528). The overall median recovery of LCS Aroclors (Year 1 and Year 2 combined) is 99.2 percent (n = 1999).

Initial analyses of PE-5 were reported in the ILCS. The "grand mean" (mean of final pool) reported in Table A5-4 is actually about 12 percent higher than the known value of PE-5. However, calculations performed by EPA, using only the reported PE data from the six laboratories which were ultimately accepted into GE's analytical program, were closer to the known value (about 4 percent high). The mean value calculated from the Year 1 and Year 2 PE-5 data (see Table A5-4) is within 1 percent of the known value. (Lab 4 data are excluded; no detected values of Aroclors have been used by GE from Lab 4. This issue is addressed in greater detail in the Year 1 DSR [Section 6.1.1] General Electric 2003].)

As with the total Aroclor quantitation, only samples of known composition can be assessed quantitatively. As the LCS was comprised of a single commercial Aroclor, the LCS data provide no information regarding the accuracy of Aroclor identification. Therefore, only the data from PE-5 provide any useful means of quantitatively assessing Aroclor identification in samples with more than one Aroclor.

Table A5-4 shows the Year 1 and Year 2 PE-5 Aroclor data. The observed mean ratio of 1221 to 1242 (just over 2.5:1) is a bit lower than the known ratio (just under 3.0:1). On an Aroclor basis, the mean concentration of A1221 is about 5 percent lower than the known

value, whereas the A1242 mean is slightly more than 10 percent higher than the known value. These relatively low errors are environmentally protective; i.e., the slight high bias toward identification of Aroclor 1242 will result in a slight high bias in Tri+ calculations (assuming that the environmental samples exhibit the same phenomena as PE-5, an assumption which is not necessarily true).

In addition to comparing the overall Aroclor concentration data to known values (to assess possible systematic bias), the precision of the data were also reviewed to judge how much error is likely to be present in any individual result. For this assessment, PE-2 data were used along with PE-5. (It was expected that there would be more variability in PE-2 results due to the fact that the Aroclor composition may not match up as well with peak patterns of Aroclor standards).

For PE-5, the RSD for TPCBs is about 0.17 (as shown on Table A5-4). Somewhat surprisingly, the precision on PE-2 was slightly better, with an RSD of about 0.15 (Table A5-5). The precision of the Aroclor identification (as "fraction A1221") was also assessed. For PE-5, the RSD is 0.042, and for PE-2, the RSD is 0.059. An example of the significance of these RSDs can be presented, assuming that the data are normally distributed and that 95 percent of the values with fall within two standard deviations of the mean. For PE-2 (for example), a sample reported to have the mean Aroclor 1221 fraction of 0.65 has a 95 percent chance that the 'true' fraction is $0.65 \pm 2^*(0.059^*0.65)$ or 0.66 ± 0.077 (i.e., the true value probably is between about 0.57 and 0.73).

Field Duplicate Results

The data discussed above represent results on QA samples. While the true values were not always known to the laboratory, it was generally evident that the laboratory knew they were QA samples and may have (consciously or unconsciously) taken extra care in the analysis and reporting of those samples. In addition, even the PE samples which were prepared from Hudson River sediment (PE-1 through 4) were well homogenized and adjusted to known moisture content, and may not have posed some of the problems that the environmental samples presented. It is difficult, if not impossible, to assess the degree to which this may be the case. The only further available measure is the performance of the laboratories on blind field duplicates; while these provide no information on the accuracy of the reported results, they do provide an indication of the precision of the results.

Field duplicate data were reported in summary form in the GE Data Summary Reports (Year 1 DSR, Phase 1 DSR, and Phase 2 DSR). Overall, about 80 percent of the detected values (3024 out of 3789 data points, including TPCBs and data for each individual Aroclor [1221, 1242, 1254, and 1260], met the established criteria (relative percent difference of 40 percent or less; or absolute difference less than two times the reporting limit for concentrations less than five times the reporting limit). A 40 percent RPD is analogous to ± 20 percent of the average of the values.

A more detailed review of the PCB (Aroclor) field duplicate data was conducted. First, the data were sorted by concentration, with the greatest focus on the samples with TPCB (sum of Aroclors) concentrations greater than or equal to 5 mg/kg in the original sample. The precision of the TPCB analysis, precision of data for A1221 and A1242 individually, and the reproducibility of the fraction of the total represented by A1221, were all reviewed.

About 723 duplicate sample pairs with concentrations greater than or equal to 5 mg/kg PCBs (sum of Aroclors) were located in the database. Not all pairs could be used for all the evaluations due to anomalies in the data. (For example, some statistics cannot be calculated for values of zero; and there were eight samples [of the 723] in which Aroclor 1221 was not detected in one or both of the analyses. There were also seven additional suspect data pairs, for which the agreement was 'too good' - the exact same result was entered for each detected Aroclor and TPCBs.)

For TPCBs, the RPD was greater than 100 percent in about 3 percent of the samples; and the RPD is less than 88 percent in 95 percent of the samples. With respect to the stated GE QA/QC criterion of 40 percent RPD, the TPCB RPD for these sample pairs was less than this value in 81 percent of the pairs. The median TPCB RPD was 16 percent. Precision was similar for A1221 and A1242 individually.

In addition to the data for TPCB concentrations, the identification of Aroclors (e.g., A1221 vs. A1242) is an important factor in the regression analysis. Even if replicate analyses are in agreement on the TPCB concentration, the Tri+ fraction will not be reproducible if the Aroclor composition of the analysis is not reproducible. To evaluate this aspect, the A1221 fraction was calculated for each sample, and the precision of duplicate pairs was assessed for this parameter. Precision for this parameter was good (even with the few poor precision samples due to A1221 not being detected in one of the analyses); the RPD was less than 50 percent in 99 percent of the samples, and was 20.1% or less in 95 percent of the samples. The median RPD for the fraction A1221 data is only 3.5 percent.

Similar assessments were also made on the set of 259 field duplicate pairs in which the TPCB (sum of Aroclors) concentration ranged from 1 to less than 5 mg/kg (see Table A5-6). Precision was somewhat less for this data set as compared to the duplicate pairs with 5 mg/kg or more TPCBs. About 70 percent of the duplicate pairs in this PCB concentration range met the \leq 40 percent RPD criterion for TPCBs (as opposed to slightly over 80 percent in the higher concentration. The median RPD for fraction 1221 in this lower-concentration group was 5.3 percent, with 95 percent of the samples having a fraction 1221 RPD of less than 31 percent.

<u>Summary</u>

The Method 8082 results agreed well with the expected values for the two "known" measurements, the LCS and PE-5. The available data suggest that the quantitation of TPCBs by M8082 is likely to be close to the true value, taken as a whole, based on available metrics. In terms of Aroclor identification, the M8082 results indicate that Aroclors have, in general, been consistently identified, with an RSD for the fraction of Aroclor 1221 in QC samples on the order of 0.05. The RPD for Aroclor identification in field duplicates is similar, with a median RPD of 5 percent or less, depending on the TPCB concentration (lower RPDs were found at higher concentrations). For both QA/QC samples as well as field duplicates, the results show a higher degree of precision in the estimation of Aroclor mass fractions relative to absolute quantitation. Mean RSDs for the sum of Aroclors in

QA/QC samples (0.15 to 0.17) were greater than those estimated for the Aroclor mass fractions (0.05 to 0.06). Nonetheless, both are acceptable for the planned use of the data. The median RPD for field duplicates for TPCBs (sum of Aroclors) in field duplicates greater than or equal to 5 ppm (16 percent) is similarly larger than the median RPD for the Aroclor mass fraction RPD of 3.5 percent. These RPDs are roughly one third higher for concentrations below 5 ppm. With respect to the M680 results, both the field duplicate precision, as well as the mass fraction precision estimates for M680, are very similar to those of M8082, further supporting the contention that the M680 bias is primarily an absolute mass underestimate and not a bias resulting from PCB identification.

Based on the LCS and PE samples results discussed above, GE's M680 TPCB results showed a systematic low bias compared to those of M8082. As a result, in the March 25, 2004 comments on GE's Draft Phase 1 Dredge Area Delineation (DAD) Report, EPA required GE to apply a regression-based correction factor to account for the systematic low bias observed in the M680 homologue PCB data relative to M8082 and mGBM results. A small subset of SSAP data was analyzed using mGBM. EPA also required additional analyses of the Tri+ PCBs regression, including an assessment of the data quality and the uncertainty in the predicted Tri+ PCB concentrations. EPA performed additional statistical analyses, which further indicated that the M680 data were biased low relative to M8082 and mGBM. In response to this concern, GE prepared a two-part correction procedure, dated June 14, 2004 (General Electric 2004), which was submitted to (and approved by) the EPA.

2.2.3 Data Ranges of Interest

As stated in the 2002 ROD, the ROD remedy is based on removal of PCBs with 3 or more chlorine atoms (Tri+ PCBs). The removal of sediments in the Hudson River was based primarily on a mass per unit area (MPA) measurement (i.e., grams of PCBs per square meter). PCB inventory in sediment is represented by samples with a MPA measurement. The ROD makes clear that, for River Sections 1 and 2, the MPA of Tri+ PCBs is to be the primary basis for identifying specific areas for dredging. The removal criteria as stated in the ROD are as follows:

- MPA of 3 g/m^2 Tri+ PCBs or greater from River Section 1;
- MPA of 10 g/m^2 Tri+ PCBs or greater from River Section 2;
- Removal of selected sediments with high concentrations of PCBs and high erosional potential (NYSDEC Hot Spots 36, 37, and the southern portion of 39) from River Section 3

Since the Tri+ PCBs concentration was estimated via a regression-based correction factor from Aroclors measurements (see Section 2.2-4 below for the regression-based correction factor), there could be biases in the estimated Tri+ PCBs concentration ranges. To address this, EPA examined three concentration ranges to assess uncertainty and potential bias and their impacts on area selection for the remedial design: 0 to 5 ppm, 5 to 80 ppm and 80 ppm or higher Tri+ PCBs. The 0 to 5 ppm Tri+ PCB (homologue) sample pairs were identified because this concentration range was unlikely to result in exceedance of removal criteria and, in the Year 1 Data Summary Report, GE described the accuracy of the homologue analyses as decaying for the lower concentration range. The 5-80 ppm range was selected because this range was critical to determining if a sample location would exceed either the surface concentrations.

2.2.4 M680 Tri+ PCB correction factor

This section discusses the development of an equation to estimate Tri+ PCB concentrations from the M680 and M8082 paired analytical results. The result is summarized here. Because of the low bias in the M680 results discussed in Section 2.2.1, a bias correction was first developed for the M680 data before the Tri+ PCB equation was developed. Several approaches were explored for correcting for the low bias in the M680 PCB homologue data. After the bias in M680 data was corrected, a multiple regression approach was used to develop the relationship between Tri+ PCB and the detected PCB Aroclors. The final model to estimate Tri+ PCB concentrations from the detected Aroclors by M8082 was:

$$Tri + PCB = 0.03*A1221 + 1.16*(A1242 + A1254)$$
 Eqn. 2-3

where A1221, A1242 and A1254 are Aroclor 1221, 1242 and 1254 concentrations reported by M8082, respectively. Uncertainty associated with the selected model and the data used to develop the model was also estimated. An uncertainty estimate based on the field duplicate precision (median variability of ± 16 to 20 percent, 95th percentile at ± 90 percent) was used in the weight of evidence approach for the dredge area delineation.

As noted above, the paired SSAP data show a low bias for TPCBs by M680 relative to TPCBs by M8082 in all concentration ranges. Thus, the correction factor is applied to all concentration ranges in the paired data. The development of the correction factors was based on point-by-point comparison of M680 and M8082 TPCB concentrations. The correction factor is calculated for each data point as follows:

$$Corr_i = \frac{Total_{8082_i}}{Total_{680_i}}$$
 Eqn. 2-4

where

 $Corr_i$ = Correction factor for sample iTotal8082 i= TPCBs by M8082 for sample iTotal680 i= TPCBs by M680 for sample i

The correction factor is then applied to the Tri+ PCB M680 concentration to yield the corrected Tri+ PCB for each sample, using the following equation:

$$Tri_{corrected_i} = Corr_i \times Tri_{680_i}$$
 Eqn. 2-5

where

 $Tri_{corrected_i}$ = the corrected Tri+ PCB value for sample i $Corr_i$ = correction factor for sample i Tri_{680_i} = the original Tri+ PCBs value by M680 for sample i

This approach presumes that the M680 results are accurate in their estimate of the fraction of Tri+ PCB present in the sample but biased low in the overall estimate of TPCB. Thus the approach increases the Tri+ PCB concentration in direct proportion to the TPCB correction.

2.2.4.1 Regression Methodology

After completing the first step of bias correction for the Tri+ PCB concentration, the second step involved the development of the regression model correlating the corrected Tri+ PCB concentrations with the M8082 Aroclor results. The regression formula represents the GE 2 draft model, which includes Aroclor 1221 and the sum of Aroclors 1242 and 1254:

$$Tri_{corrected_i} = a1 \times (A1221) + a2 \times (A1242 + A1254)$$
 Eqn. 2-6

where the a1 and a2 are regression coefficients and the other terms are defined as above.

The following sections describe the details regarding calculation of the regression coefficients and metrics of accuracy.

2.2.4.2 Algorithm

A robust regression approach based on the bisquare influence function (Holland and Welsch, 1977) was used to develop the regression equations. The robust regression method was used because it automatically identifies outliers (both regression and leverage points) and down-weights them in the derivation of the regression coefficients by minimizing the weighted error sum of squares. The computational approach used for the weighted regression in this analysis is based upon iteratively reweighted least squares regression.

The damped leverage approach used by GE is similar to the weighted robust regression method. The difference is that the weights are not calculated iteratively. Ideally, the damped leverage approach assumes that the weights are known in advance but this very seldom happens in practice. Using the available data, these weights can be computed in more than one way using classical methods. GE used the damped leverage approach to calculate the weights. However, it should be noted that when outliers are present, these weights (and all

other statistics including regression coefficients) obtained using a damped leverage procedure can get distorted; therefore, it is preferable to use robust methods which automatically identify outliers and down-weight them accordingly in the regression process.

2.2.5 Data Selected

Samples with paired data in the February 2004 database were selected for this analysis. There are 1,346 samples with paired M8082 and M680 analyses.⁴ A total of 23 sample pairs where the M8082 TPCB or M680 Tri+ PCB concentrations were nondetect or inconsistent were eliminated, and the remaining 1,323 samples were included in the regression analysis. Statistical outliers were not eliminated because the procedure is able to handle them directly. The 23 samples eliminated from the analysis included:

- 18 samples with nondetect Tri+ PCB by M680,
- 2 samples with nondetect A1221 and A1242 by M8082; and
- 3 samples where Tri+ PCB by M680 was greater than TPCB by M8082.

2.2.5.1 Training and Testing Data Sets

Two-thirds of the data were selected for the training set and used to develop the regression equations. The full set of paired data was sorted on laboratory, river section and concentration range prior to selecting a representative data subset (note that GE employed five different laboratories during the SSAP). The remaining data were used to test the regression equations. This approach allows the predictive ability of the regression equations to be tested on an independent data set.

2.2.5.2 All-Laboratories and Laboratory-Specific Equations

The influence of the different individual laboratories on the regression model was evaluated. Development of the draft Tri+ PCB regression equations indicated that coefficients for individual laboratories were generally similar to the coefficients for the single all-laboratories equation. Therefore, the regression model was constructed as a

⁴ This large data set represented the available data for analysis at the time of the negotiations between EPA and GE. Subsequent data collection efforts were reviewed and found to be consistent with the regression developed based on the February 2004 database.

single model for the entire set of data ("all-laboratories") because the uncertainty in the correction of M680 Tri+ PCB results is expected to be larger than the differences among the laboratories.

2.2.6 Model Results

The final model to determine Tri+ PCB concentrations bases on M8082 detected Aroclors was:

$$Tri + PCB = 0.03*A1221 + 1.16*(A1242 + A1254)$$
 Eqn. 2-7

This formula integrates the bias correction in the M680 data and the conversion of Aroclor results to Tri+ PCBs as described above. This model became the basis for estimating Tri+ PCB concentrations for dredge area delineation and other remedial activities. However, as described in the Consent Decree, it was anticipated that GE would update this regression based on paired mGBM and M8082 results when additional Tri+ PCB and Aroclor paired results became available (EPA, 2004).

A scatter plot of the bias-corrected measured versus the predicted Tri+ PCBs concentrations can be found in Figure A5-2 and a scatter plot of the bias-corrected measured versus the predicted Tri+ PCBs fraction can be found in Figure A5-3. The slope for the corrected measured vs. predicted Tri+ PCB fraction is close to one (with a value of 1.02) as shown in Figure A5-2, but there is a scatter around the 1 to 1 line with R² of 0.8.

Plots of the Tri+ PCB concentrations show good agreement between the predicted and corrected measured values (Figure A5-2), but the 0 to 5 ppm Tri+ PCB range is more often overestimated. The overestimation in the concentration is environmentally protective because the overestimated Tri+ PCB concentration will result in a location being identified for removal as opposed to leaving it in place. The scatter for the 5 to 80 ppm and 80 ppm or higher Tri+ PCB concentrations is more balanced. These same conclusions can be drawn from review of the residual plots in Figure A5-4.

The uncertainty in the models was estimated by the 95 percent individual confidence curves on the log of the measured and predicted Tri+ PCB concentrations. The measured and predicted results are shown on Figure A5-5 with the confidence curves by concentration range. The graphs show that, generally, the uncertainty in the data increases with increasing concentration and can be approximated by a percentage of the average concentration. To estimate this percentage, the confidence curves were approximated by linear fits by concentration range. These results are summarized in Table A5-8. For the 0 to 5 ppm Tri+ PCB concentrations, curvature in the confidence curves is evident. Ordinary least squares linear regressions on these upper and lower confidence curves, provide a poor fit to the data near 0 ppm. The trend of the data is through 0 ppm, but the linear fits are above this. No precise estimate of uncertainty was made for the 0 to 5 ppm Tri+ ppm range on this basis, but a rough approximation from examination of the data would indicate an uncertainty of approximately +70 percent and -20 percent. Estimates of uncertainty can be made on the basis of the confidence curves for the 5 to 80 ppm Tri+ PCB range (-29 percent and +38 percent) and the 80 ppm or higher Tri+ PCB range (-44 percent and +41 percent). As expected from data exhibiting heteroscedascity (i.e., data with unequal variability (scatter) across a set of second, predictor variables), the range of uncertainty is greater at higher concentrations.

2.2.7 Update to Tri+ PCB Regression during Remedial Action Monitoring Phase

As stated in the Dispute Resolution, the regression equation developed to estimate Tri+ PCB using M8082 Aroclor (see Section 2.2.6 above) may be updated using paired mGBM and M8082 results if GE performs a Tri+ PCB Study (EPA, 2004). In 2005, GE conducted a laboratory methods comparison study to compare the relative accuracy and precision of M680, M8082 and mGBM for the sediment data. That study found that the mGBM has better accuracy and precision to M680 and M8082 in measuring TPCB and Tri+ PCB (General Electric 2005). In addition, the 2005 GE study found a strong correlation between Tri+ PCB concentrations measured by the mGBM and the Aroclor concentrations measured by M8082. Using paired analyses on 150 sample extracts and 30 archived sediment samples, the following regression equation was developed (General Electric 2009): The EPA reviewed this regression and requested that GE analyze 100 additional paired samples to provide more data at low concentrations to support using an mGBM-based regression to calculate Tri+ PCB concentrations from M8082 Aroclor concentrations that will be measured on residual sediments post-dredging, which are expected to have relatively low PCB concentrations. Out of the 100 paired additional samples, 71 samples were selected from those having a M8082 Total PCB concentration less than 7 ppm, and 29 samples were selected from those having a M8082 Total PCB concentration greater than 57 ppm (General Electric 2009).

The data for the original 180 paired samples and 98 of the 100 supplemental paired samples (two samples were excluded because they did not have results for Aroclor 1221) were used to develop a refined regression model (General Electric 2009). Regression analysis was performed using the statistical software package R (http://www.R-project.org). The basic regression methodology consisted of applying a weighted least squares (i.e., "damped-leveraged) algorithm to calculate the coefficients, summary statistics and goodness of fit measures (General Electric 2009). The regression equation is:

$$Tri + PCB = 0.14*A1221 + 0.91*(A1242+A1254)$$
 Eqn. 2-9

The equation (Eqn. 2-9) is nearly identical to the equation developed from the original subset of 180 samples (Eqn. 2-8). The additional data result in tighter bounds on the coefficients (i.e., lower standard errors) and approximately the same goodness of fit statistics (Table A5-9; General Electric2009).

During remedial action period, GE continued collecting paired M8082-mGBM sediment data at a rate of 4% of the overall residual sediment samples to allow continued evaluation of the regression equation relating the Aroclor PCB concentrations measured using M8082 to the Tri+ PCB concentrations. The regression coefficients were recalculated each time

the overall data set is increased by 5% and these updated regression coefficients was used in subsequent CU evaluations (General Electric, 2009).

As documented in Corrective Action Memorandum (CAM) No. 3 – Modification of Sediment Residual Monitoring Program – Discontinuing mGBM Analysis of Sediment Samples and Updates of the Regression Coefficients (General Electric 2011c), GE performed four rounds of sampling in 2009 and 1 round of sampling in 2011 (Table A5-10) which resulted in a total of 445 paired M8082-mGBM sediment samples. The regression analysis was then repeated following the methodology described in the 2009 QAPP (General Electric 2009). The coefficients derived from the update, as well as the coefficients from the original regression and subsequent rounds of updates in 2009, are provided in Table A5-10. The coefficients changed very little as a result of the update. The additional data result in tighter uncertainty bounds on the coefficients (i.e., lower standard errors) and approximately the same goodness of fit statistics (Table A5-11). Because of the coefficients have stabilized, GE recommended discontinuing the mGBM analysis of sediment samples and subsequent update of the regression coefficients.

The latest coefficients developed in 2011 is as follows:

$$Tri + PCB = 0.13*A1221 + 0.89*(A1242 + A1254)$$
 Eqn. 2-10

Eqn. 2-10 was applied to any samples collected in all CUs that had not yet been sampled for Pass 1 residual samples as of August 5, 2011.

2.2.8 Congener-Specific Measurements during the 2016 OM&M Sampling

As part of the sediment collection conducted by GE in 2016, a subset of the samples (approximately 10 percent) were provided to EPA. At the request of the federal natural resources trustees, these samples were analyzed for PCB congeners via method 1668C (M1668C). This was done, in part, due to the anticipated transition from mGBM to a congener-specific method for the long-term OM&M program. EPA is further analyzing the relationship between the M8082 results obtained by GE and the M1668C results obtained by EPA. EPA also anticipates collecting more matched pairs of these analyses as part of

future OM&M sampling. Additionally, EPA plans to meet with the federal natural resources trustees, NYSDEC and GE to further discuss consideration related to various analytical testing methods and ongoing OM&M sampling.

2.3 Summary

This section summarizes the translation schemes (regression equations) for estimating Tri+ PCBs in the sediment for the historical data that support the EPA decisions contained in the ROD, as well as the development of relationships between Tri+ PCB and Aroclors during the remediation investigation period. The sediment data sets and the translation schemes used to support the ROD, remedial investigation and remedial action phases are summarized below:

- The 1976-1978 NYSDEC data covered the area from Fort Edward to Troy (RM 194.8 to RM 154) and was used to identify likely areas for remediation and provide estimates of PCB mass and sediment volumes for the 2002 ROD for the Upper Hudson River. The data were analyzed by Aroclor-based method. The following equation was used to calculate Tri+ PCB: $Tri + (1977) = 1.131 \times [Aroclor 1016 + 1254]$
- The 1984 NYSDEC data were restricted to TIP. Similar to the 1976-1978 NYSDEC data, the 1984 data was used support the ROD. These data sets were also dependent on the Aroclor quantitation method. The following equation was used to calculate Tri+ PCB for the 1984 NYSDEC data: *Tri+ (1984) = 0.944 × 1984 Aroclor Sum*
- GE conducted sediment surveys in 1991 and 1998. The 1991 survey sampled the upper river from Fort Edward to Federal Dam, while the 1998 survey sampled the TIP only. These data sets were used in the modeling effort during Phase 2 RI/FS investigation. Both 1991 and 1998 data were reported as PCB congeners based on mGBM. Therefore, the Tri+ PCB concentrations can be calculated directly and no translation scheme was needed.
- In 1994, the EPA collected low resolution cores and the data were analyzed based on congener-specific method. The 1994 low resolution coring effort focused on replicating a representative of the 1984 locations. Since these data were reported as

PCB congeners, no translation scheme was needed to calculate the Tri+ PCB concentrations.

- During the remedial design investigation phase, GE collected sediment samples in 2002-2005. These data were collected for delineating the final areas for removal. Given the large number of samples (approximately 30,000 sediment samples), the samples were analyzed using the M8082. A subset of the samples was analyzed by the EPA-approved M680. Using the matched-pair samples, a regression equation was developed to estimate the Tri+ PCBs from the Aroclor-based measurement. As discussed in 2.2, the regression equation to estimate the Tri+ PCB concentration integrates the bias correction in the M680 data and the conversion of Aroclor results. The following regression equation was used to estimate the Tri+ PCB: *Tri+ PCB* = 0.03*A1221 + 1.16*(A1242+A1254). This regression equation developed during the remedial investigation period was used to delineate dredge areas, to examine Tri+ PCB concentrations in residual core samples, to examine Tri+ PCB concentrations in residual core samples, to examine Tri+ PCB concentrations in residual core samples, to examine Tri+ PCB concentrations in residual core samples, to examine Tri+ PCB concentrations in residual core samples, to examine Tri+ PCB concentrations in residual core samples, to examine Tri+ PCB concentrations in period, and for the OM&M study results.
- During the remedial action phase, GE conducted a laboratory methods comparison study to compare the relative accuracy and precision of M680, M8082 and the mGBM. Based on this study, the following equation was developed to estimate the Tri+ PCB concentrations in the sediment data: *Tri+ PCB = 0.13*A1221 + 0.94*(A1242+A1254)*. Throughout the remedial action phase, GE conducted multiple rounds of paired mGBM and M8082 PCB data collection and evaluation which were used to assess the regression equation relating the Aroclor PCB concentrations measured using M8082 to the Tri+ PCB concentration. In 2011, GE recommended discontinuing the mGBM analysis of sediment samples and subsequent update of the regression coefficients because the coefficients do not change significantly and has been stabilized (General Electric 2011). The 2011 regression equation (*Tri+ PCB = 0.13*A1221 + 0.89*(A1242+A1254)*) was used for sediment samples collected after August 5, 2011.
3 WATER

Concentration of PCBs has been measured in surface water in the Upper Hudson River since 1977 by different organizations. Over the period of measurement, different lab analytical techniques have been used. Valid comparing the trends in surface water PCB concentrations can only be made when the various analytical data sets be converted to a consistent congener-based quantitation.

This section describes the analytical conversions developed for water column data in the Upper Hudson River. Since 1977, PCBs have been measured in water column in the Upper Hudson River by USGS, GE, and EPA. While the water column analyses by EPA (in 1993) and GE (starting in 1991) were routinely based on conger quantitation, the long-term record from the USGS (1977-1995) used different analytical methods that required adjustments in order to make the datasets comparable for use in the data evaluation and model calibration. It was determined that an equivalent TPCB quantitation could not be determined from the USGS data and therefore Tri+ PCB, which could be calculated in all datasets, was used for data evaluation and model calibration.

As stated in the Revised Baseline Modeling Report (RBMR) for the Upper Hudson River (USEPA, 2000), the USGS water column data represented whole water analyses, with PCBs quantified using Aroclor standards. Packed column analysis was used until 1987, when data began to be analyzed with capillary columns. Split sample analysis between USGS and EPA Phase 2 data supported use of the USGS-reported TPCB concentration from the packed column analysis as a direct measure of the Tri+ PCB. A regression relating USGS TPCB to the Tri+ sum gave a good linear fit with an intercept not significantly different from zero (TAMS Consultants, Inc. and Gradient Corp.1997). Thus, the USGS packed-column TPCB results were used directly as Tri+ through 1987. Further, the re-analysis of 60 USGS capillary column sample chromatograms by QEA (Rhea and Werth, 1999) supported use of the USGS-reported Aroclor 1242 results or, when 1242 results are not available, use of Aroclor 1248, as the best representation of Tri+ PCB concentration in the USGS data after 1987.

Beginning in 1991, water samples collected from the Hudson River by GE were routinely analyzed for PCB congeners by the modified Green Bay Method (mGBM). EPA conducted a short-term series of studies on PCB levels in Hudson River water in 1993 using its dual phase PCB congener analysis. Thus, for both GE and EPA data, congener-specific information was available to directly determine Tri+ PCB and TPCB water column concentrations. In 2011, GE asserted that the mGBM was unnecessarily specialized for compliance purposes, and they began an evaluation of the efficacy of using a standard Aroclor method (M8082) for estimating Tri+ PCB for water samples collected during the remedial construction and off-season monitoring period. Notably, the M8082 method will also be used during the OM&M period after the construction. Therefore, it is important that an accurate methodology be determined to estimate Tri+ PCB from the M8082 results so that future data can be compared to existing data and recovery trends can be accurately discerned. The procedure to determine Tri+ PCB concentrations from Aroclor data was based on the same regression methodology used to estimate Tri+ PCBs in sediments (see Section 2 above). In 2011, GE analyzed the paired data available and produced preliminary regression equations, and has used additional data collected after 2012 to refine these equations (General Electric 2012a, 2012b, 2013, 2014, 2016a). The remainder of this section describes the analyses conducted by the EPA to determine the best regression equation(s) to estimate water column concentrations of Tri+ PCBs and the associated uncertainties using all the available paired data. Different regression relationships were explored for cases where only a single Aroclor was reported, when multiple Aroclors were reported and also for the PCB concentrations determined in the samples. EPA's analysis was performed using the R[®] statistical package.

3.1 Paired GE mGB and Aroclor Water Column Data Set

A total of 249 water samples collected using automatic samplers at far-field stations in 2011 to 2013 and 2015 were analyzed by both the mGB and the Aroclor methods. These paired data were used to develop the relationship between Aroclor and Tri+ PCB. Among the 249 samples, EPA excluded 10 from the analysis because, as documented in CAM 5 and CAM 8 (General Electric 2013 and 2014, respectively), they were judged not

representative of the Hudson River or did not meet analytical QA/QC requirements. Duplicate and triplicate samples were averaged prior to incorporation into the data analysis.

3.2 Test for Regression Outliers and High Leverage Samples

Multiple linear regression analysis based on ordinary least squares (OLS) is sensitive to data points with large residuals (regression⁵) and/or high leverage.⁶ This evaluation considers whether there were influential samples which may have affected the OLS results based on Cook's distance (or Cook's D). Cook's D combines the information of leverage and residual of the observation and is commonly used to estimate the influence of individual data points. A simple operational guideline of Cook's D greater than 1 has been suggested for spotting highly influential data points (Cook and Weisberg, 1982). Figure A5-6 shows the Cook's D of the OLS regression for the three groups of samples separately and all the samples together. The results show that all the samples have Cook's D less than 1, indicating that there are no influential data that will affect the regression, EPA explored both OLS regression and several commonly-used robust regression approaches (see Section 3.4) to develop the best equation(s) for estimating Tri+ PCB in the water column data.

3.3 Validity of Using Aroclors as Predictable Variables

In the paired data sets available for this evaluation, A1221 was detected by M8082 in all 239 water samples, while A1242 and A1254 were detected in 71.5% and 7.1% of the samples, respectively; no other Aroclors were detected. For samples where A1221 was the only detectable Aroclor by M8082, EPA used A1221 as the only independent variable in the regression relating A221 and Tri+ PCB. In all other cases with multiple Aroclor detections, EPA used the concentrations of A1221 and the sum of A1242 and A1254 in the

⁵ A regression outlier is an observation with large residual. In other words, it is an observation whose dependent-variable value is unusual given its value for the predictor variables.

⁶ Leverage is a measure of how far an independent variable deviates from its mean. An observation with an extreme value on a predictor variable is a point with high leverage. High leverage points can have an inordinate effect on the estimate of regression coefficients.

regression. This section evaluates whether using individual Aroclors or total Aroclor as independent variables could produce a better correlation.

For simplicity, a test was performed using OLS linear regression for all the samples. In this approach, the best model is selected based on several feature selection measures, specifically the leave-one-out cross-validation (LOOCV) statistic (also known as PRESS, i.e., prediction residual sum of squares⁷), Akaike's Information Criterion (AIC), corrected AIC (AICc), Schwarz's Bayesian Information Criteria (BIC), and adjusted R². A small value of PRESS, AIC, AICc and BIC, or a large value of adjusted R², indicates a model with a low test error. Table A5-12 shows the combination of various linear regression model formulations considered by EPA to test the efficacy of using individual Aroclors as predictor variables. The results of this test indicate that the lowest values for PRESS, AIC, AICc and BIC and the highest value for adjusted R² can be obtained for a regression model that is similar to remedial design sediments formulated as follows:

$$Tri + PCB = a \times A1221 + b \times (A1242 + A1254)$$
 Eqn. 3-1

where a, and b are regression model regression coefficients, and A1221, A1242 and A1254 are the Aroclor concentrations reported by M8082.

In the application of this model to determine the model coefficients for the Upper Hudson River, regression analyses for three groups of data were performed as follows:

- Group 1: Samples for which Aroclor 1221 was the only detected mixture.
- Group 2: Samples with multiple Aroclors detected but limited to those for which TPCB is less than or equal to 150 ng/L, or with Aroclor ratios less than or equal to 0.2, where Aroclor ratio = (Aroclor 1242 + Aroclor 1254)/Aroclor 1221.
- Group 3: Samples with multiple Aroclors detected but limited to data with TPCB greater than 150 ng/L and Aroclor ratios greater than 0.2.

⁷ PRESS is the sum of squared prediction residuals. It is the same as the sum of the squared "leave-oneout" residuals.

3.4 Regression Model Results

In addition to the OLS regression, EPA explored several robust regression approaches to determine the best values of the model coefficients a, and b. Robust regression can provide an alternative to OLS to remediate the effects of overly influential samples without excluding them. In general, robust regression models give less weight to unusual observations that would otherwise have undue influence on the regression line. Table A5-13 lists the OLS and various robust regression methods as well as the functions applied in the statistical R[®] package to determine the regression coefficients. Table A5-14 presents the regression coefficients for each model. Figure A5-7 compares the predicted and measured Tri+ PCB concentrations for the test models. As illustrated in Figure A5-7, a comparison between predicted and measured concentrations using the training data set does not necessarily readily identify the model with the best performance. To select the best model EPA used a cross-validation approach.

3.5 Model Selection

Two model evaluation methodologies were used to determine the best model coefficients applicable to the water samples in the Hudson River. The first is a cross-validation using all available data. The second involves model validation using test data to evaluate the performance of the various regression models.

Cross-validation is primarily a way of measuring the predictive performance of a statistical model. Leave-one-out cross validation (LOOCV) was used in this study to evaluate the prediction accuracy of the regression models. LOOCV means that the regression is performed on all the data except for one point and a prediction is made for that point. This process is repeated such that every data point is included in a test set exactly once, and in a training set n-1 times (where n denotes the total number of data points). The process creates a prediction error⁸ for each point, and the average error is used to evaluate the

⁸ Prediction error or prediction residual is defined as the difference between the predicted value and the actual value for a test data point. It is generally greater than the residual from the model, which is based on the training data.

model. Table A5-15 compares the average prediction errors among the regression models. In general, the lower the average prediction error, the better the model. The results indicate that average prediction error values do not show much difference among the models. The best model with the lowest average prediction errors for each group of regression is highlighted in bold in Table A5-15. The relative prediction error of the best model was calculated as mean prediction error divided by mean of the measured Tri+ PCB concentration. The results indicate that the relative prediction error is 27, 18 and 23 percent for Group 1, Group 2 and Group 3 samples, respectively.

In the second approach, selecting the best model involved determination of the predictive performance of each statistical model using a test data set.⁹ Similar to the cross validation analysis presented above, the predictive accuracy of a model can be measured by the average prediction error on a test set. In this evaluation, the test data set contains far-field grab, near-field background and off-season in-river samples. This data set contains 12 samples from Group 1 and 23 samples from Group 2, and therefore, only Groups 1 and 2 regression could be tested. Table A5-16 provides the average prediction error for each regression model and the results indicate that the LMS and LTS models performed better for Groups 1 and 2 regressions, respectively.

Overall, based on the LOOCV results, M-Huber, OLS, and LAV appear to be the best models for Group 1, Group 2 and Group 3 samples, respectively. Based on the test set results, LMS and LTS appear to be the best models for Group 1 and Group 2 samples, respectively. Since the test data sample size is much smaller than that for the LOOCV, the best models for Group 2 and Group 3 are selected as OLS and LAV based on the LOOCV results.

3.6 Summary

The ability to predict the concentrations of Tri+ PCB accurately from Aroclor data is essential for daily compliance evaluations at the far-field monitoring stations, and for

⁹ A test data set includes samples which are not used in the development of regression models.

determining future trends in water column concentrations during the OM&M period. In this evaluation, several linear regression approaches were considered for the prediction of Tri+ PCB concentrations based on Aroclor measurements in water column samples. The models employed are Ordinary Least Square Regression (OLS), Least Absolute Values Regression (LAV), M-Estimation with Huber Weight (M-Huber), M-Estimation with Bisquare Weight (M-Bisquare), Least Trimmed Squares Regression (LTS), and Least Median Squares Regression (LMS). The models were developed using the training data set containing 239 water samples, and their predictive performance was assessed using the leave-one-out cross validation (LOOCV) and an independent test data set. The mean absolute prediction error and the relative prediction error were calculated to compare the predictive performance of the models. Based on the available data, LMS, OLS, and LAV are the best prediction models for Group 1, Group 2 and Group 3 samples, respectively. Table A5-17 provides the regression equations developed by EPA that should be used with the water column data. For comparison purposes, the table also provides the regression that GE developed and refined as documented in applicable corrective action memoranda ([i.e., CAMs] General Electric 2013 and 2014, respectively).

4 FISH

Similar to the two previous section, this section presents the analyses intended to reconcile various historical and current fish sampling efforts into a single, internally consistent series of measurements. For both sediment and water, it was necessary to separately estimate TPCB and Tri+ PCB since both measures were important to understanding fate, transport and biological exposure. For fish however, this separation is not necessary since TPCB is approximately equal to Tri+ PCB. Based on congener data obtained by EPA in the 1990s, Tri+ PCB was found to represent 90 percent or more of the total PCB burden in fish samples (EPA 1999a). This observation continued to hold true until the start of dredging in 2009, discussed further below. In all cases, the conversions developed are intended to estimate the total PCB mass in the tissue samples (TPCB_{HE}), regardless of whether the mixture is solely Tri+ PCB or includes substantive monochloro or dichloro homologues.

This section presents the procedure followed to calculate the Homologue Equivalent TPCB (TPCB_{HE}) concentrations based on the sum of Aroclor concentrations (TPCBAroclor) reported in fish samples collected by NYSDEC and GE from the Hudson River. This procedure follows the calculation process first described in the Hudson River PCBs Site remedial investigation reports and in Butcher, et al., 1998, and is necessary to facilitate year-to-year comparisons of PCB levels in fish using an equivalent basis of measurement through time. As mentioned previously, TPCB analysis by Aroclors, typically EPA M8082, is a relatively subjective analysis, relying on the discretion of the analyst as to the selection of Aroclors to be reported in a sample. As a result, the mixture of Aroclors reported, as well as the sum of Aroclors, will vary from laboratory to laboratory for equivalent samples. This is evident in the historical record of Aroclor-based measurements in fish for the Hudson River.

In addition to concerns regarding Aroclor mixture variations attributable to lab differences, it is also highly unlikely that any measured congener pattern in a fish sample will directly replicate an exact Aroclor congener distribution. This is because variations in historical Aroclor use, weathering, dechlorination, and preferential biological uptake of various congeners over others all serve to extensively modify the actual congener distribution present in fish (as described in Section 1). Thus, quantitation of PCBs in fish tissue by Aroclor-based analysis can only be an approximation of the actual TPCB mass (TPCB_{HE}). This concern applies to both the NYSDEC data and the GE data described below.

4.1 Discussion of NYSDEC Fish Data

The NYSDEC fish data examined here span the period 1990 to 2011. Prior analyses by NYSDEC are not discussed here but are covered at length in Butcher, et al. 1998. For the entire period examined, fish are reported by NYSDEC on an Aroclor basis. As presented in Figure A5-8, the selection of reported Aroclors (the Aroclor composition) in NYSDEC fish samples clearly varies through time. Annual variation is apparent through 1998, while the patterns are relatively similar from 1999 through 2011. Changes in the suite of reported Aroclors largely correspond to changes in the analytical labs performing the procedures. Table A5-18 summarizes the appearance of various Aroclor mixtures in the quantitation.

Given the dominance of the GE source of PCBs in the Upper Hudson and the large reservoir of PCB contamination in the sediments, it is unlikely that the pattern of PCBs exhibited in fish was actually varying in a significant fashion through time. This assertion is directly supported by the 1999 to 2011 period, when a single lab (Mississippi State Chemical Laboratories) was used by NYSDEC, and a uniform set of Aroclors in roughly similar proportions is consistently reported over the period.

4.2 Discussion of GE Fish Data

Aroclor concentrations reported in GE fish samples also clearly vary through time, although in a much more systematic fashion than in the NYSDEC data. As shown in Table A5-19, GE employed a single lab for PCB Aroclor analysis over the entire period (2004-2015). Prior to 2009, the GE lab consistently reported Aroclors 1248 and 1254 as the dominant fractions, with minor amounts of Aroclors 1242 and 1260. Aroclor 1221 is essentially absent during this period (see Figure A5-9). Beginning in 2009, however, the Aroclor 1221 fraction began increasing, reaching 17 to 18 percent of the total mixture for 2012 and 2013. This period is highlighted in Table A5-19.

In this instance, the change of the Aroclor distribution in fish tissues presumably is not due to a change in analytical procedures or judgment, since only one laboratory was used by GE, as shown in Table A5-19. Rather, it is most likely a direct reflection of the increased exposure of fish to congeners associated with Aroclor 1221, or more specifically to the presence of congener BZ#4 (peak 5 based on GE's mGBM) released into the water column when remedial dredging operations started in the Hudson River in 2009. An increase in the proportion of lighter congeners in the water column was extensively observed and documented in the various dredging reports issued by EPA and GE.

4.3 Aroclor Measurements and Estimation of TPCB Concentration

As noted above, the mixtures of Aroclors reported through time had varied widely but, prior to the onset of dredging, it is unlikely that the actual distribution of congeners varied so extensively with time. Indeed, even for the period where NYSDEC and GE each consistently using their own dedicated laboratory (2004 to 2011), the Aroclor mixtures reported by NYSDEC's and GE's laboratories differ significantly. This can be readily observed by comparing Figures A5-8 and A5-9 for this period. While the reporting of different Aroclor mixtures by different laboratories is clearly problematic in trying to discern changes in PCB patterns through time, it also presents a much larger issue for long-term monitoring purposes. Specifically, in reporting different Aroclors, it is highly likely that different laboratories will obtain different estimates for the mass of PCBs contained in the sample; the mass of total PCBs is estimated by summing the individual reported Aroclors.

The issue with this estimate of TPCB mass in a sample arises from disconnect between the actual PCB pattern present in the fish sample and the "standard" congener pattern present in the Aroclor standard mixtures. Included in this problem is the extensive overlap in congener content among different Aroclor mixtures. That is, many congeners are observed in several different Aroclors (*e.g.*, Aroclors 1242, 1248, 1254 and 1260 all have multiple congeners in common). As a result, reporting of multiple Aroclors with overlapping congener spectra potentially leads to "double counting" of individual congener concentrations. Different laboratories have different means of avoiding "double counting,"

again similarly leading to differences in reported concentrations; this time TPCBs as the sum of Aroclors. The presence of overlapping Aroclor spectra was extensively documented in EPA's Data Evaluation and Interpretation Report (EPA 1997) as well as in Frame et al. (1996).

Alternatively, the actual congener distributions in the fish tissue may differ widely from those found in the Aroclor standards. Variations in analytical response factors, relative contributions to total PCB mass and simply lack of pattern similarity can all result in inaccurate quantitation of the actual PCB mass, yielding both underestimates and overestimates of the TPCB concentration.

Fortunately, for all of the data sets examined here, an alternate means of quantifying PCB mass is available for a subset of samples obtained by each laboratory. Specifically, TPCBs by homologue, or by congener, (equivalent to EPA M680 or GE's mGBM, hereinafter referred to as congener-based methods) exist for each laboratory data set along with matching Aroclor-based analysis. Totals PCBs by homologue-based methods are much less subject to the analyst's judgement as to the mixture of PCBs present as a basis for quantitation. These methods simply require the quantitation of all congeners in the mixture. As a result, use of these methods should provide a consistent basis of TPCB measurement across time and laboratories.

4.4 Development of the Homologue Equivalent Basis

As noted above, congener-based PCB data are available for all of the individual laboratories but exist for only a subset of the total number of samples for each laboratory. In order to estimate TPCB concentrations as the sum of homologues (TPCB_{HE}) for the vast majority of the fish data, a series of correlation analyses were conducted using the available Aroclor and homologue-based analytical pairs in both NYSDEC and GE data sets. Ideally a simple proportionality constant should relate the Aroclor and homologue sums. Given that both the sum of Aroclors and the sum of homologues should approach zero when a sample does not contain PCBs, the translations were conducted using the following formulations:

$$TPCB_{HE} = \alpha \times TPCB_{Aroclor}$$
 Eqn. 4-1

where α is determined by a regression on the TPCB_{HE} and TPCB_{Aroclor} pairs for a given laboratory.

$$\alpha_i = \frac{TPCB_{HE}}{TPCB_{Aroclor}}$$
Eqn.4-2

where the α_i are determined for the individual samples with both TPCB_{HE} and TPCB_{Aroclor} values.

The α_i were then used to determine the arithmetic mean, median and geometric mean ratio (α) for a given laboratory and period of time. The α_i were also used in a bootstrap analysis to determine confidence intervals around the geometric mean, discussed further below.

These equations were applied for the various data sets for the period from 1998 to 2011. For NYSDEC data for the period from 1990 to 1997, the regression equations as described in the Revised Baseline Modeling Report ([RBMR] (EPA 2000) were used without modification. For the 1998 to 2013 data, the regression was prepared as follows: The limited number of nondetect results (approximately 4% of sample pairs across all post-1998 data) were eliminated for this analysis and only detected results for both Aroclor and congener-specific data were used. That is, if either the sum of Aroclors or sum of homologues was nondetect, the sample pair was not included in the regression. In all instances, the geometric mean ratio appeared to best represent the relationship between TPCB_{HE} and TPCB_{Aroclor} values across the entire range of concentrations. A summary of the resulting factors to be applied to the Aroclor sums is shown in Table A5-20, along with a brief description of the origins of the individual factors.

To provide an example of the importance of these factors in adjusting the TPCB_{Aroclor} to yield TPCB_{HE}, the NYSDEC matched pairs of TPCB_{HE} and TPCB_{Aroclor} values were plotted against each other in Figure A5-10, for the period 1991 to 2000. The great degree of scatter reflects the level of variation in the relationship between the two metrics for individual

samples. However, also note that the data cluster by year, as indicated by the colored symbols. For example, 1991 samples consistently lie below the 1-to-1 line, indicating that TPCB_{Aroclor} is consistently greater than TPCB_{HE} for this period. For the 1999 and 2000 data, the samples consistently plot above the 1-to-1 line, indicating TPCB_{Aroclor} is consistently less than TPCB_{HE} for this period. For the data collected before 1998, the EPA developed relationships between TPCB_{HE} and TPCB_{Aroclor} which were included in the documents for the ROD. Since 1998, additional data has been obtained by the NYSDEC and GE that warrants similar treatment. The development of four new relationships for the period 1998 to 2015 between TPCB_{HE} and TPCB_{Aroclor} and their associated statistical support are described below.

4.5 NYSDEC Data Factors

The currently available NYSDEC data on PCB concentrations in fish tissue post-1997 span the period 1998 to 2011. These data were analyzed by two laboratories. Enchem Environmental Laboratories was used for all fish samples collected in 1998, during which NYSDEC obtained 82 samples analyzed for both TPCB_{HE} and TPCB_{Aroclor}. These data were used to develop the formula given in Table A5-20. Table A5-21 summarizes the additional statistics examined for the 1998 data. The data are plotted as TPCB_{Aroclor} *vs*. TPCB_{HE} in Figure A5-11. In the figure, the data generally lie below the 1-to-1 line, indicating that the TPCB_{Aroclor} results tend to be higher than those for TPCB_{HE}. As a result, the estimate for α is less than 1 at 0.741. This value is similar to that obtained by arithmetic mean (0.826), median (0.783) or by the regression (0.784).

In general, the similarities among geometric mean, median and average all yielded similar values for α within a single laboratory-time period for both the NYSDEC and GE data. However, given the reduced sensitivity of the geometric mean to the occurrence of outliers relative to the average, and that statistical calculations using the geometric mean are generally simpler than those for the median, the geometric mean was selected as the basis to select α and was further examined for its uncertainty via a bootstrap technique. While the arithmetic mean and the median generally agreed well with the geometric mean, the estimate of α by regression analysis was inconsistent in its agreement across all the four

single laboratory-time period model. For this reason, the regression analysis was discontinued and is not discussed further here.

The uncertainty in the geometric mean was estimated by multiple resampling (a statistical technique referred to as "bootstrapping") of the population of matched TPCB_{Aroclor} and TPCB_{HE} pairs. Table A5-21 also contains the bootstrap output for the 2.5, 5, 95 and 97.5 percentiles confidence limits for the geometric mean. Additionally, the table contains an estimate of the uncertainty of the geometric mean expressed as a percentage of the geometric mean value itself, using the 5th and 95th percentile values. The 95 percent confidence limits on the geometric mean value for 1998 of 0.741 is \pm 8 percent. This confidence interval includes the median value, indicating that these values agree within the uncertainty. The statistical agreement between geometric mean and median occurs for all four single laboratory-time period models described here.

Besides illustrating the general correlation between TPCB_{Aroclor} and TPCB_{HE}, the symbols in Figure A5-11 are color-coded to identify the fish species analyzed for each individual sample pair. For 1998, the majority of the samples are brown bullhead and largemouth bass. Both species are similarly distributed over the range of values, indicating that the conversion from TPCB_{Aroclor} to TPCB_{HE} is similar for both species, and no species-specific conversion is needed. This observation based on the 1998 data is also supported by the 1999 to 2000 data, which also show consistent behavior between TPCB_{HE} and TPCB_{Aroclor} across the various species. The similar behavior between TPCB_{HE} and TPCB_{Aroclor} across species indicates that congener patterns of PCBs in tissues are similar across species, yielding similar Aroclor percentages across species. This consistency was also noted in the congener-specific analysis of PCB patterns in fish conducted for the ecological risk assessment for the ROD (EPA 1999b).

Figures A5-12, A5-13 and A5-14 provide additional statistical background on the estimate of the ratio (α) for the NYSDEC 1998 data. Figure A5-12 shows the detailed statistics on the range of α for the individual samples. The figure also shows that α skews right and may not be normally distributed. Figure A5-13 represents several statistical tests of significance

comparing the paired TPCB_{Aroclor} to TPCB_{HE} values. The results in this figure indicate that for 1998 data, TPCB_{Aroclor} is significantly higher than TPCB_{HE} by all tests including the Wilcoxon signed rank test, which has no requirement regarding an underlying normal distribution. Based on this test, the use of an adjustment factor (i.e., α) is statistically justified. Figure A5-14 compares the logs of TPCB_{Aroclor} and TPCB_{HE}. In this test, showing that the log values have a significant difference is mathematically equivalent to showing that the ratio of TPCB_{Aroclor} to TPCB_{HE} is statistically different than unity (1). In this test, the logs are significantly different but the difference appears relatively constant with increasing log value (*e.g.*, concentration). This result actually indicates that the ratio of TPCB_{HE} to TPCB_{Aroclor} is relatively constant, and independent of concentration. This further justifies the use of a single coefficient α to estimate TPCB_{HE} from TPCB_{Aroclor}.

The 1999 to 2000 NYSDEC data were analyzed in a parallel fashion, yielding the statistics given in Table A5-21 for this period, as well as Figures A5-15 through A5-18. These figures parallel the presentation described above for the 1998 result. Figure A5-16 presents the NYSDEC fish results, with symbol and color-coding to indicate year of collection as well as species type. Of particular note, the data appear sufficiently consistent across the more than 2 order of magnitude range to suggest that neither year (*i.e.*, 1999 *vs.* 2000) nor fish species are important factors to consider in the development of α for this period. Figures A5-16 through A5-18 present the statistical support for developing a correction factor for this period. In this instance, however, the geometric mean factor is greater than 1 (i.e., 1.17), indicating that TPCB_{Aroclor} is an underestimate of TPCB_{HE}. The signed rank tests confirm a statistically significant difference between TPCB_{HE} and TPCB_{Aroclor}. Again, the log-based analysis, shown in Figure A5-18, shows no trend with increasing concentration, although there is greater variability than for the 1998 results. This last analysis again supports the use of a single coefficient α to estimate TPCB_{HE} from TPCB_{Aroclor}, similar to the 1998 data.

One remaining concern for the NYSDEC data is the lack of further matched pairs of $TPCB_{HE}$ and $TPCB_{Aroclor}$ in the post-2000 period. Thus, there are no additional data from which to develop these factors. However, NYSDEC continued to utilize the same

laboratory from 1999 through 2011, a condition that would be expected to maintain the TPCB_{HE} to TPCB_{Aroclor} relationship observed in 1999-2000. Consistent with this, the Aroclor mixtures reported by the NYSDEC laboratory during this period have remained similar through the period (see Table A5-18 and Figure A5-8 for the period 1999 to 2011). These considerations justify the continued use of the 1999 to 2000 coefficient α value of 1.17 to estimate TPCB_{HE} from TPCB_{Aroclor} for the post-2000 period.

The NYSDEC data factors represents the Upper Hudson and near-Albany Lower Hudson conditions. It is unclear if these factors will be applicable to Lower Hudson conditions characterized by relatively lower PCB concentrations in fish.

4.6 GE Data Factors

The GE data for 2004 to 2013 were analyzed in a manner parallel to the NYSDEC data described above. Note that analysis by the mGBM was performed on 5 percent of the total number of fish samples during odd years (2005, 2007, etc.) by GE. GE did not perform mGBM in 2015, but performed it in 2016, as requested by the EPA (EPA 2016b). The shift to 2016 was to allow sample-pair data one year after dredging was finished. Since GE has only employed a single laboratory for fish analysis to date, it was anticipated that a single coefficient α would be needed to estimate TPCB_{HE} from TPCB_{Aroclor} for the entire period. However, initial review of the GE data showed that the mixture of Aroclors reported by the GE laboratory was not constant, but rather changed substantially over the period of record, particularly after the start of the Phase 2 dredging program in 2011. Specifically, the reported laboratory results changed from trace amounts of Aroclor 1221 to an average of 17 percent of the total Aroclor mixture (see Figure A5-9 and Table A5-19). The coincidence of this change with the more extensive dredging effort and substantial increase in monochloro- and dichloro-homologue water column loads is almost certainly not by chance. The likely causative relationship between water column loads and fish body burdens of Aroclor 1221 is further illustrated in Table A5-22, which presents the average Aroclor 1221 content in fish tissue as a function of both year and river mile.

In particular, later years tend to have higher fractions. However, the table has been highlighted to show the coincidence between the major areas of dredging during the 2009 to 2015 periods and fish sampling. Evident from the table is the higher fractions of reported Aroclor 1221 in the areas of dredging and in areas downstream. This evidence suggests that the change in Aroclor mixture noted in the GE data is likely a reflection of an increase in the monochloro- and dichloro-homologue fractions in fish tissue, resulting in higher reported Aroclor 1221 fractions. However, the higher Aroclor 1221 fraction reported in 2009 and later would also indicate that α is likely to change in response. Based on this, the GE data were split into two periods for analysis: 2004 to 2008, and 2009 to 2013. Estimates of α for these periods are given in Table A5-21 along with the associated statistics and uncertainty estimates.

The GE data for the period 2004 to 2008 are shown in Figures A5-19 to A5-22. These figures parallel the same analyses performed for the NYSDEC data, and yield similar conclusions concerning the statistical significance of α . Given the apparent lack of species-specific differences in the relationship between TPCB_{HE} from TPCB_{Aroclor} for the NYSDEC data, EPA did not repeat the analysis for the GE data. Rather, for the GE data, the data in Figure A5-19 (and Figure A5-23 discussed below) are color-coded by year of collection. The data examined in this fashion do not suggest substantial variation in α from year to year within the 2004 to 2008 interval.

One additional observation is worth noting, however. Specifically, as can be observed in Figure A5-19 the correlation between TPCB_{HE} and TPCB_{Aroclor} is much stronger for the GE data relative to the NYSDEC data. This is reflected in the narrower range of variability in the GE data, as well as in smaller estimates of variance, *i.e.*, the standard deviation and the geometric mean confidence levels (see Table A5-21).

The analysis for the period 2009 to 2013 is shown in Figures A5-23 to A5-26, again paralleling the analyses conducted for the NYSDEC data. Like the 2004 to 2008 period, the GE results show reduced variance as compared to the NYSDEC data. Like the previous results, the GE data for 2009 to 2013 support the use of α for estimating TPCB_{HE} from

TPCB_{Aroclor}. Of note, α for the 2009 to 2013 period based on the geometric mean 0.784, is similar to the value obtained for the 2004 to 2008 period (0.831). However, as can be inferred from the lack of overlap between the 5th and 95th percent confidence intervals, the difference in α between the two periods is considered to be statistically different. This analysis confirms EPA's decision to treat these periods differently, although the difference in the ratios is less than 6 percent.

Similar to NYSDEC data factors, the GE data factors represents the Upper Hudson and near-Albany Lower Hudson conditions. It is unclear if these factors will be applicable to Lower Hudson conditions characterized by relatively lower PCB concentrations in fish.

4.7 Fish Species Distribution

The NYSDEC analyses described above considered the possibility of variations in α related to fish species but did not find any strong evidence for the need to consider fish species in the analysis. For completeness, the available distribution of fish species as a percentage of the total number of paired samples for both the NYSDEC and GE data are summarized in Figures A5-27 and A5-28. Notably, the NYSDEC paired data for 1998 to 2000 represent fewer species than the GE data (2004 to 2015).

4.8 Summary

This section reviews the basis for estimating TPCB_{HE} from TPCB_{Aroclor} for the historical data that supported the EPA decisions contained in the ROD, as well as the development of new relationships between TPCB_{HE} and TPCB_{Aroclor} for fish data collected by NYSDEC and GE during the post-ROD period. The following conclusions were drawn from this analysis.

 PCB analysis by Aroclor is subject to significant variations in the types of Aroclors reported as well as in the sum of Aroclors as a basis to estimate total PCB mass in a sample. This is readily apparent in the NYSDEC data, which were generated by multiple laboratories, but is an issue for all measurements based on Aroclors (TPCB_{Aroclor}).

- When compared to estimates of total PCB mass obtained by homologue or congener-based methods (TPCB_{HE}), the sum of Aroclors (TPCB_{Aroclor}) can either exceed or fall below the TPCB_{HE} value.
- Despite this limitation, internal to a single laboratory, the TPCB_{Aroclor} can represent a sufficiently precise measurement so as to permit the estimation of TPCB_{HE} for a given period of time.
- Based on these observations, coefficients (α) to estimate TPCB_{HE} from TPCB_{Aroclor} were developed for four lab-period pairs.
 - For NYSDEC data, these were 1998 with the Enchem laboratory, and 1999-2000, with the Mississippi State Chemical Laboratories. The results for the latter period (1999 to 2000) were applied to all subsequent NYSDEC data since no additional TPCB_{HE} / TPCB_{Aroclor} sample pairs were available and NYSDEC did not change laboratories during the subsequent period.
 - For GE data, the laboratory-period pairs were 2004 to 2008 with NEA, and 2009 to 2013 with NEA. In this instance, two periods were needed because the Aroclor pattern in fish changed between periods, likely due to the increased presence of lighter congeners in the water column released via dredging associated with the remediation.
- While there is variability in the ratio of TPCB_{HE} to TPCB_{Aroclor} in all the matched pairs evaluated, the data sets are sufficiently large as to provide well-constrained estimates of the ratio (expressed as α). The maximum uncertainty (95 percent confidence interval) obtained by a bootstrap analysis of variation in the geometric mean estimate expressed as a percentage of α was ± 8 percent.
- In all instances, the α values were shown to be statistically significant and therefore, their application as a basis to estimate TPCB_{HE} from TPCB_{Aroclor} is statistically supported.
- Finally, the absolute magnitude of the α values is sufficient that they represent a substantive, as well as a statistically significant adjustment to the TPCB_{Aroclor} in order to obtain a more accurate estimate of TPCB_{HE}. Failure to account for variations in the TPCB_{HE} to TPCB_{Aroclor} is likely to introduce significant

uncertainties and potential trend artifacts unrelated to actual changes in fish body burdens in any time-based trend analysis.

- While these factors represent Upper Hudson and near-Albany Lower Hudson conditions, there may be an issue with extrapolation of these coefficients to Lower Hudson conditions characterized by relatively lower PCB concentrations in fish.
- The range of the geometric mean of α across data sets was 0.74 to 1.17. These values represented the two NYSDEC lab-period pairs. The range of α values for the GE laboratory-period pairs was much tighter (i.e., 0.79 to 0.854).

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Second Five-Year Review Report Hudson River PCBs Superfund Site

APPENDIX 5

PCB Aroclors Data Treatment

Tables and Figures

Prepared by: Louis Berger US, Inc.

May 2017

Table A5-1 DB-1 Chromatograph Peaks and Corresponding PCB Congeners for Modified Green Bay Method

DB-1	Congener	Chlorination	DB-1	Congener	Chlorination									
Peak #	BZ #	Structure	Peak #	BZ #	Structure									
1	0	biphenyl	31	52	2 2'5 5'	53	90	2 2'3 4'5	74	105	2 3 3 4 4	109	201	2 2'3 3'4 5 5'6'
2	1	2	31	73	2 3'5'6	53	101	2 2 4 5 5	74	132	2 2'3 3'4 6'	110	196	2 2'3 3'4 4'5 6'
3	2	3	32	49	2 2 4 5	54	99	2 2 4 4 5	75	153	2 2 4 4 5 5	110	203	2 2'3 4 4'5 5'6
1 4	3	4	33	47	2 2 4 4	55	112	2 3 3 5 6	76	168	2 3'4 4'5'6	111	189	2 3 34 45 5
5	4	2 2	34	48	2 2'4 5	55	119	2 3 4 4 6	77	141	2 2'3 4 5 5'	112	195	2 2'3 3'4 4'5 6
5	10	26	34	75	2 4 4 6	55	150	2 2'3 4'6 6'	78	179	2 23 35 6 6	113	208	2 2'3 3'4 5 5'6 6'
6	7	24	35	62	2346	56	83	2 2'3 3'5	79	130	2 2'3 3'4 5'	114	207	2 23 34 4566
6	9	25	35	65	2356	56	109	2 3 3 4 6	80	137	2 2'3 4 4'5	115	194	2 2'3 3'4 4'5 5'
7	6	2 3	36	35	3 3 4	57	86	2 2'3 4 5	81	176	2 2'3 3'4 6 6'	116	205	2 3 3 4 4 5 5 6
8	5	23	37	44	2 2 3 5	57	97	2 2'3 4'5	82	138	2 2'3 4 4'5'	117	206	2 2'3 3'4 4'5 5'6
8	8	2 4	37	104	2 2 4 6 6	57	152	2 2 3 5 6 6	82	163	2 3 3 4 5 6	118	209	2 2'3 3'4 4'5 5'6 6'
9	14	35	38	37	344	58	87	2 2'3 4 5'	83	158	2 3 3 4 4 6	NQ	20	2 3 3
10	19	2 2'6	38	42	2 2'3 4'	58	111	2 3 3'5 5'	84	129	2 2'3 3'4 5	NQ	38	345
11	30	246	38	59	2 3 3 6	58	115	23446	85	178	2 2 3 3 5 5 6	NQ	41	2 2'3 4
12	11	33	39	64	2346	59	85	2 2'3 4 4'	86	166	234456	NQ	43	2 2'3 5
13	12	34	39	71	2 3 4 6	59	116	23456	87	175	2 2'3 3'4 5'6	NQ	69	2 3 4 6
13	13	34	40	68	2 3 4 5	60	136	2 2'3 3'6 6	88	182	2 2'3 4 4'5 6	NQ	72	2 3'5 5'
	15	4 4	41	96	2 2 3 6 6	61	11	3 3 4 4	88	187	2 2'3 4'5 5'6	NQ	78	3 3 4 5
	18	225	42	40	2 2 3 3	61	110	23346	89	128	2 2 3 3 4 4	NQ	79	3 3 4 5
15		224	43	57	2335	62	154	2 2 4 4 5 6	90	183	2 2 3 4 4 5 6	NQ	80	3 3 5 5
16	24	236	43	103	2 2 4 5 6	63	82	2 2 3 3 4	91	167	2 34 45 5	NQ	81	3445
16	21	236	44	6/	2 3 4 5	64	151	2 2 3 5 5 6	92	185	2 23 4 55 6	NQ	88	22346
11	16	223	44	100	2 2 4 4 6	65	124	2 3 4 5 5	93	174	2 23 34 56	NQ	102	2 2 4 5 6
11/	32	246	45	58	2 3 3 5	65	135	2 2 3 3 5 6	93	181	2 2 3 4 4 5 6	NQ	113	2 3 3 5 6
18	23	235	45	63	2 3 4 5	66	144	2 2 3 4 5 6	94	177	2 23 34 56	NQ	117	23456
19	34	2 35	46	74	2445	67	107	2 3 3 4 5	95	156	2 3 3 4 4 5	NQ	120	2 3 4 5 5
19	54	2 26 6	46	94	2 2 3 5 5	67	108	23345	95	171	2 2'3 3'4 4'6	NQ	121	2 3 4 5 6
20	29	245	47	61	2345	67	147	2 2 3 4 5 6	96	202	2 2'3 3'5 5'6 6	NQ	125	2 3 4 5 6
21	26	2 3 5	47	70	2 3 4 5	68	123	2 3 4 4'5'	97	157	2 3 3 4 4 5	NQ	126	3 3 4 4 5
22	25	234	47	76	2 3 4 5	69	106	23345	98	173	2 2'3 3'4 5 6	NQ	127	3 3 4 5 5
23	31	245	40	00	2344	69	118	2 3 4 4 5	99	200	2 2 3 3 4 5 6 6	NQ	142	223456
	28	244	40	93	22356	69	149	223450	39	204	22344566	NQ	145	223465
	50	2240	40	80	22356	/0	139	223440	100	1/2	2233455	NQ	148	223456
25	21	234	49	55	2334	70	140	2 2 3 4 46	100	192	2 3 3 4 5 5 6	NQ	159	2 3 3 4 5 5
25	33	234	49	91	2 2 3 4 6		114	23445	101	197	2 2 3 3 4 4 6 6	NQ	160	2 3 3 4 5 6
25	23	2250	49	98	22346	1 4	134	223356	102	180	2 2 3 4 4 5 5	NG	162	2 3 3455
20	22 E1	2245	50		2334	1 2	143	223450	103	193	2 3 3 4 5 5 6	NG	164	233456
20	51	2226	50		2 3 4 4	12	122	2 3 3 4 5	104	191	2 3 3 4 4 5 6	NG	165	2 3 3 5 5 6
21	40	2230	51	64	22336	12	131	223346	105	199	22334566	NG	169	3 3 4 4 5 5
20	30	3 3 5	51	92	22355	1/2	133	2 2 3 3 5 5	106	1/0	2233445	NG	184	2234466
29	46	2230	51	155	224406	13	146	223455	107	190	2334456	NQ	186	2234566
30	39	345	52	69	22340	73	161	2 3 3 4 5 6	108	198	22334556	NQ	188	2234566

Note: NQ = Not quantified in DB-1 method

Source: GE, 1997

Development of Corrections for Analytical Biases in the 1991-1997 GE Hudson River PCB Database. 1997. Prepared by HydroQual, Inc. for General Electric.

Table A5-2 GE PE sample results PE 5 - Homolog Data

Method 680 Results											
FIELD_SAMPLE_ID	PE ID	LAB	Total PCB: Tri-	+ PCBs	% Tri+	MonoCB	DiCB	TriCB	TetraCB	PentaCB	HexaCB
RS1-8988-PE030827-E01	PE5	Lab 15	29.15	8.35	28.6%	12	8.8	4.3	3.5	0.55	
RS1-8988-PE030827-E02	PE5	Lab 15	25.74	7.74	30.1%	11	7.0	3.8	3.4	0.54	
RS1-8988-PE030827-E03	PE5	Lab 15	26.58	7.68	28.9%	11	7.9	4.2	3.3	0.18	
RS1-8988-PE030827-E04	PE5	Lab 15	32.90	10.20	31.0%	13	9.7	4.7	4.5	1.00	
RS1-9089-PE021030-C01	PE5	Lab 15	20.79	9.69	46.6%	2.9	8.2	4.6	4.1	0.99	
RS1-9291-PE021024-C02	PE5	Lab 15	30.52	11.52	37.7%	7	12	6.1	5.1	0.32	
RS1-9291-PE030625-C01	PE5	Lab 15	29.23	8.53	29.2%	12	8.7	4.2	3.6	0.73	
RS2-8483-PE030605-A02	PE5	Lab 15	20.15	8.85	43.9%	3.1	8.2	4.2	3.8	0.82	0.034
RS2-8887-PE030918-B01	PE5	Lab 15	37.23	11.23	30.2%	16	10	5.7	4.8	0.73	
RS2-8887-PE030918-C01	PE5	Lab 15	29.59	8.69	29.4%	12	8.9	4.3	3.7	0.69	
RS3-7170-PE030717-B01	PE5	Lab 15	28.62	7.72	27.0%	14	6.9	3.8	3.3	0.62	
RS3-7271-PE031009-A04	PE5	Lab 15	26.48	7.48	28.2%	11	8.0	4.0	3.2	0.28	
RS3-7271-PE031023-A01	PE5	Lab 15	24.14	6.94	28.7%	9.8	7.4	3.6	3.0	0.34	
RS3-7978-PE030806-C01	PE5	Lab 15	32.57	10.07	30.9%	13	9.5	4.6	4.5	0.97	
RS3-7978-PE030806-C02	PE5	Lab 15	28.34	7.74	27.3%	12	8.6	4.2	3.2	0.34	
		Mean	28.14	8.83	31.9%	10.65	8.65	4.42	3.80	0.61	
		Median	28.62	8.53	29.4%	12.00	8.60	4.20	3.60	0.62	
S	tandard	Deviation	4.471	1.410	6.0%	3.681	1.301	0.678	0.650	0.272	
	Relative	e Std Dev	0.159	0.160	0.189	0.345	0.150	0.153	0.171	0.449	

Method 680 Results-two samples (low MonoCB values) excluded

FIELD_SAMPLE_ID	PÉ ID	LAB	Total PCBs T	ri+ PCBs	% Tri+	MonoCB	DiCB	TriCB	TetraCB	PentaCB	HexaCB
RS1-8988-PE030827-E01	PE5	Lab 15	29.15	8.35	28.6%	12	8.8	4.3	3.5	0.55	
RS1-8988-PE030827-E02	PE5	Lab 15	25.74	7.74	30.1%	11	7.0	3.8	3.4	0.54	
RS1-8988-PE030827-E03	PE5	Lab 15	26.58	7.68	28.9%	11	7.9	4.2	3.3	0.18	
RS1-8988-PE030827-E04	PE5	Lab 15	32.90	10.20	31.0%	13	9.7	4.7	4.5	1.00	
RS1-9291-PE021024-C02	PE5	Lab 15	30.52	11.52	37.7%	7	12	6.1	5.1	0.32	
RS1-9291-PE030625-C01	PE5	Lab 15	29.23	8.53	29.2%	12	8.7	4.2	3.6	0.73	
RS2-8887-PE030918-B01	PE5	Lab 15	37.23	11.23	30.2%	16	10	5.7	4.8	0.73	
RS2-8887-PE030918-C01	PE5	Lab 15	29.59	8.69	29.4%	12	8.9	4.3	3.7	0.69	
RS3-7170-PE030717-B01	PE5	Lab 15	28.62	7.72	27.0%	14	6.9	3.8	3.3	0.62	
RS3-7271-PE031009-A04	PE5	Lab 15	26.48	7.48	28.2%	11	8.0	4.0	3.2	0.28	
RS3-7271-PE031023-A01	PE5	Lab 15	24.14	6.94	28.7%	9.8	7.4	3.6	3.0	0.34	
RS3-7978-PE030806-C01	PE5	Lab 15	32.57	10.07	30.9%	13	9.5	4.6	4.5	0.97	
RS3-7978-PE030806-C02	PE5	Lab 15	28.34	7.74	27.3%	12	8.6	4.2	3.2	0.34	
		Mean	29.31	8.76	29.8%	11.83	8.72	4.42	3.78	0.56	
		Median	29.15	8.35	29.2%	12.00	8.70	4.20	3.50	0.55	
S	Deviation	3.466	1.500	2.7%	2.130	1.391	0.728	0.697	0.261		
	Relative	e Std Dev	0.118	0.171	0.090	0.180	0.159	0.165	0.184	0.466	

Table A5-3 Summary of GE Method 680 LCS Recovery

LAB_SAMPLE_ID	Туре	Method	Analysis Date	Unit	LCS Conc	LCS rec	GE Rec %	TAMS Rec %	Count
Year 1 LCS Samples									
RS1-9594-LABQCCOC02110052-AF10920L	LCS	GEHR680	11/9/2002	mg/Kg	1.36	0.93	68.4	68.4%	
RS1-9493-LABQCCOC02110055-AF10974L	LCS	GEHR680	11/14/2002	mg/Kg	1.24	1.10	88.7	88.7%	
RS1-9392-LABQCCOC02110056-AF10991L	LCS	GEHR680	11/15/2002	mg/Kg	0.82	0.53	64.6	64.6%	
RS1-9392-LABQCCOC02110054-AF10953L	LCS	GEHR680	11/17/2002	mg/Kg	1.24	1.10	88.7	88.7%	
RS1-9392-LABQCCOC02110088-AF11187L	LCS	GEHR680	11/19/2002	mg/Kg	0.82	0.61	74.0	74.4%	
RS1-9493-LABQCCOC02110089-AF11211L	LCS	GEHR680	11/21/2002	mg/Kg	1.24	1.00	80.6	80.6%	
RS1-9493-LABQCCOC02110091-AF11259L	LCS	GEHR680	11/21/2002	mg/Kg	1.24	0.87	70.2	70.2%	
RS1-9392-LABQCCOC02110090-AF11235L	LCS	GEHR680	11/22/2002	mg/Kg	1.24	1.20	96.8	96.8%	
RS1-9493-LABQCCOC02110106-AF11402L	LCS	GEHR680	11/29/2002	mg/Kg	1.33	1.20	90.2	90.2%	
RS1-9392-LABQCCOC02110109-AF11473L	LCS	GEHR680	12/1/2002	mg/Kg	1.24	1.20	96.8	96.8%	
RS1-9493-LABQCCOC02110128-AF11662L	LCS	GEHR680	12/1/2002	mg/Kg	1.21	0.90	74.4	74.4%	
RS2-8584-LABQCCOC02110129-AF11684L	LCS	GEHR680	12/2/2002	mg/Kg	1.2	0.91	75.8	75.8%	
RS2-8685-LABQCCOC02110107-AF11425L	LCS	GEHR680	12/2/2002	mg/Kg	1.3	1.10	84.6	84.6%	
RS1-9291-LABQCCOC02110104-AF11355L	LCS	GEHR680	12/4/2002	mg/Kg	1.24	0.99	79.8	79.8%	
RS1-9392-LABQCCOC02110110-AF11496L	LCS	GEHR680	12/4/2002	mg/Kg	1.24	0.78	62.9	62.9%	
RS2-8584-LABQCCOC02120012-AF12215L	LCS	GEHR680	12/4/2002	mg/Kg	1.35	1.00	74.1	74.1%	
RS1-9392-LABQCCOC02110111-AF11507L	LCS	GEHR680	12/6/2002	mg/Kg	1.24	1.10	88.7	88.7%	
RS1-9190-LABQCCOC02120014-AF12248L	LCS	GEHR680	12/7/2002	mg/Kg	1.24	1.60	129.0	129.0%	
RS2-8584-LABQCCOC02120022-AF12301L	LCS	GEHR680	12/8/2002	mg/Kg	1.33	1.20	90.2	90.2%	
RS2-8584-LABQCCOC02120023-AF12325L	LCS	GEHR680	12/10/2002	mg/Kg	1.23	1.20	97.6	97.6%	
RS2-8483-LABQCCOC02120024-AF12348L	LCS	GEHR680	12/11/2002	mg/Kg	1.3	1.20	92.3	92.3%	
RS1-9190-LABQCCOC02120025-AF12371L	LCS	GEHR680	12/13/2002	mg/Kg	1.24	1.10	88.7	88.7%	
RS1-9392-LABQCCOC02120026-AF12394L	LCS	GEHR680	12/13/2002	mg/Kg	1.32	1.00	75.8	75.8%	
RS1-9089-LABQCCOC02120028-AF12441L	LCS	GEHR680	12/14/2002	mg/Kg	1.24	1.00	80.6	80.6%	
RS2-8584-LABQCCOC02120027-AF12417L	LCS	GEHR680	12/14/2002	mg/Kg	1.29	1.20	93.0	93.0%	
RS1-9190-LABQCCOC02120029-AF12464L	LCS	GEHR680	12/16/2002	mg/Kg	1.24	1.20	96.8	96.8%	
RS1-9291-LABQCCOC02120030-AF12487L	LCS	GEHR680	12/16/2002	mg/Kg	1.24	0.92	74.2	74.2%	
RS2-8986-LABQCCOC03010033-AG00169L	LCS	GEHR680	1/15/2003	mg/Kg	1.34	1.00	74.6	74.6%	
Year 1 LCS - Median					1.24	1.05	82.6	82.6%	28
Year 2 LCS Analyses - First Half (4/5 through 9	/18/03)								
RS1-9392-LABQCCOC03040003-AG01635L	LCS	GEHR680	4/5/2003	mg/Kg	1.32	0.85	64.4	64.4%	
RS2-8483-LABQCCOC03040004-AG01653L	LCS	GEHR680	4/7/2003	mg/Kg	1.33	1.20	90.0	90.2%	
RS2-8584-LABQCCOC03060235-AG06101L	LCS	GEHR680	6/26/2003	mg/Kg	1.36	0.97	71.3	71.3%	
RS2-8584-LABQCCOC03060236-AG06115L	LCS	GEHR680	7/1/2003	mg/Kg	1.24	0.95	76.6	76.6%	
RS2-8483-LABQCCOC03070063-AG07346L	LCS	GEHR680	7/11/2003	mg/Kg	1.24	0.88	71.0	71.0%	
RS1-9291-LABQCCOC03070064-AG07369L	LCS	GEHR680	7/13/2003	mg/Kg	1.33	1.10	82.7	82.7%	
RS1-9089-LABQCCOC03070065-AG07392L	LCS	GEHR680	7/14/2003	mg/Kg	1.24	0.90	72.6	72.6%	
RS1-8988-LABQCCOC03070129-AG07913L	LCS	GEHR680	7/21/2003	mg/Kg	1.32	1.00	80.0	75.8%	

Table A5-3 Summary of GE Method 680 LCS Recovery

LAB_SAMPLE_ID	Туре	Method	Analysis Date	Unit	LCS Conc	LCS rec C	GE Rec %	TAMS Rec %	Count
RS1-8988-LABQCCOC03070130-AG07937L	LCS	GEHR680	7/21/2003	mg/Kg	1.24	0.96	77.4	77.4%	
RS1-8988-LABQCCOC03070203-AG08680L	LCS	GEHR680	7/26/2003	mg/Kg	1.24	0.72	58.1	58.1%	
RS1-9089-LABQCCOC03070195-AG08554L	LCS	GEHR680	7/26/2003	mg/Kg	1.24	1.10	88.7	88.7%	
RS1-9190-LABQCCOC03070128-AG07890L	LCS	GEHR680	7/26/2003	mg/Kg	1.24	1.60	129.0	129.0%	
RS1-9190-LABQCCOC03070202-AG08656L	LCS	GEHR680	7/28/2003	mg/Kg	1.24	0.45	36.3	36.3%	
RS1-9291-LABQCCOC03070240-AG09069L	LCS	GEHR680	7/29/2003	mg/Kg	1.24	0.34	27.4	27.4%	
RS1-9291-LABQCCOC03070241-AG09092L	LCS	GEHR680	7/30/2003	mg/Kg	1.24	0.39	31.5	31.5%	
RS3-6766-LABQCCOC03070242-AG09116L	LCS	GEHR680	7/30/2003	mg/Kg	1.24	0.98	79.0	79.0%	
RS3-7877-LABQCCOC03070328-AG10030L	LCS	GEHR680	8/7/2003	mg/Kg	1.24	1.20	96.8	96.8%	
RS3-7978-LABQCCOC03070329-AG10053L	LCS	GEHR680	8/10/2003	mg/Kg	1.3	0.96	73.8	73.8%	
RS3-7069-LABQCCOC03080065-AG10744L	LCS	GEHR680	8/11/2003	mg/Kg	1.24	0.90	72.6	72.6%	
RS3-8281-LABQCCOC03080066-AG10768L	LCS	GEHR680	8/12/2003	mg/Kg	1.24	0.98	79.0	79.0%	
RS3-7473-LABQCCOC03080134-AG11428L	LCS	GEHR680	8/24/2003	mg/Kg	1.24	0.98	79.0	79.0%	
RS3-7776-LABQCCOC03080135-AG11452L	LCS	GEHR680	8/24/2003	mg/Kg	1.36	0.83	61.0	61.0%	
RS3-7069-LABQCCOC03080201-AG12173L	LCS	GEHR680	8/26/2003	mg/Kg	1.29	0.94	72.9	72.9%	
RS3-7069-LABQCCOC03080204-AG12207L	LCS	GEHR680	8/27/2003	mg/Kg	1.24	0.74	59.7	59.7%	
RS3-7170-LABQCCOC03080136-AG11475L	LCS	GEHR680	8/27/2003	mg/Kg	1.24	1.00	80.6	80.6%	
RS3-5958-LABQCCOC03080205-AG12230L	LCS	GEHR680	8/28/2003	mg/Kg	1.24	0.90	72.6	72.6%	
RS3-6463-LABQCCOC03080241-AG12687L	LCS	GEHR680	9/2/2003	mg/Kg	1.24	0.98	79.0	79.0%	
RS3-6968-LABQCCOC03080242-AG12711L	LCS	GEHR680	9/5/2003	mg/Kg	1.24	1.10	88.7	88.7%	
RS3-5958-LABQCCOC03080243-AG12734L	LCS	GEHR680	9/6/2003	mg/Kg	1.24	0.95	76.6	76.6%	
RS3-6766-LABQCCOC03090032-AG13156L	LCS	GEHR680	9/9/2003	mg/Kg	1.24	0.97	78.2	78.2%	
RS3-7069-LABQCCOC03090031-AG13133L	LCS	GEHR680	9/9/2003	mg/Kg	1.35	1.10	81.5	81.5%	
RS3-6766-LABQCCOC03090033-AG13180L	LCS	GEHR680	9/13/2003	mg/Kg	1.24	0.98	79.0	79.0%	
RS3-7069-LABQCCOC03090082-AG13789L	LCS	GEHR680	9/15/2003	mg/Kg	1.24	0.99	79.8	79.8%	
RS3-6463-LABQCCOC03090083-AG13813L	LCS	GEHR680	9/18/2003	mg/Kg	1.24	0.97	78.2	78.2%	
Year 2 - First half - Median								77.0%	34
Year 2 LCS Analyses - Second Half (9/1/9 throu	gh 11/12/0	3)							
RS3-6766-LABQCCOC03090084-AG13837L	LCS	GEHR680	9/19/2003	mg/Kg	1.24	0.80	64.5	64.5%	
RS2-8685-LABQCCOC03090138-AG14571L	LCS	GEHR680	9/22/2003	mg/Kg	1.36	0.88	64.7	64.7%	
RS2-8685-LABQCCOC03090139-AG14596L	LCS	GEHR680	9/26/2003	mg/Kg	0.99	0.72	72.7	72.7%	
RS2-8382-LABQCCOC03090140-AG14620L	LCS	GEHR680	9/30/2003	mg/Kg	1.24	0.93	75.0	75.0%	
RS2-8685-LABQCCOC03090185-AG15145L	LCS	GEHR680	10/5/2003	mg/Kg	1.35	1.00	74.1	74.1%	
RS1-9291-LABQCCOC03090186-AG15168L	LCS	GEHR680	10/7/2003	mg/Kg	1.35	0.84	62.2	62.2%	
RS3-6766-LABQCCOC03090187-AG15190L	LCS	GEHR680	10/9/2003	mg/Kg	1.24	0.86	69.4	69.4%	
RS3-7170-LABQCCOC03100026-AG15968L	LCS	GEHR680	10/11/2003	mg/Kg	1.24	0.86	69.4	69.4%	
RS3-7372-LABQCCOC03100027-AG15992L	LCS	GEHR680	10/11/2003	mg/Kg	1.32	0.96	72.7	72.7%	
RS3-6059-LABQCCOC03100028-AG16015L	LCS	GEHR680	10/14/2003	mg/Kg	1.24	0.89	71.8	71.8%	
RS1-9291-LABQCCOC03100072-AG16535L	LCS	GEHR680	10/21/2003	mg/Kg	1.32	0.84	63.6	63.6%	

Table A5-3Summary of GE Method 680 LCS Recovery

LAB_SAMPLE_ID	Туре	Method	Analysis Date	Unit	LCS Conc	LCS rec	GE Rec %	TAMS Rec %	Count
RS2-8483-LABQCCOC03100144-AG17460L	LCS	GEHR680	10/24/2003	mg/Kg	1.3	0.79	60.8	60.8%	
RS2-8887-LABQCCOC03100143-AG17436L	LCS	GEHR680	10/24/2003	mg/Kg	1.24	0.89	71.8	71.8%	
RS2-8786-LABQCCOC03100145-AG17484L	LCS	GEHR680	10/26/2003	mg/Kg	1.24	0.80	64.5	64.5%	
RS2-8887-LABQCCOC03100176-AG17870L	LCS	GEHR680	10/29/2003	mg/Kg	1.3	0.63	48.5	48.5%	
RS2-8786-LABQCCOC03100177-AG17894L	LCS	GEHR680	11/8/2003	mg/Kg	1.24	0.59	47.6	47.6%	
RS3-6463-LABQCCOC03110017-AG18506L	LCS	GEHR680	11/12/2003	mg/Kg	1.24	0.67	54.0	54.0%	
Year 2 - second half only - Median								64.7%	17
Year 2 - complete - Median					1.24	0.93	72.7	72.7%	79
								TAMS	S/Earthtech

Table A5-4 GE PE 5 Samples Aroclor Data

			Aroclor	Concentrati	ions		Fraction	Calculated	Literature
LAB_SAMPLE_ID	PE ID	LAB	1221	1242	1254	Total PCB	1221	Tri+ PCBs	Tri+
RS1-8988-PE030827-N46939-21	PE5	Lab 1	27.7	11.2		38.9	0.712	0.355	0.316
RS1-9089-PE021029-N25559-22	PE5	Lab 1	27.2	11.6		38.8	0.701	0.368	0.324
RS1-9291-PE021022-N25115-17	PE5	Lab 1	39.9	12.3		52.2	0.764	0.296	0.277
RS1-9291-PE030625-N42266-9	PE5	Lab 1	23.9	9.78		33.7	0.710	0.358	0.318
RS1-9594-PE021015-N24646-4	PE5	Lab 1	32.8	11.4		44.2	0.742	0.321	0.293
RS2-8483-PE030605-N40750-1	PE5	Lab 1	27.9	13.1		41	0.680	0.391	0.340
RS2-8887-PE030918-N48618-16	PE5	Lab 1	17.2	6.64		23.8	0.721	0.345	0.309
RS3-7170-PE030717-N43957-13	PE5	Lab 1	26.1	11		37.1	0.704	0.365	0.322
RS3-7271-PE031009-N50386-4	PE5	Lab 1	20.6	8.16		28.8	0.716	0.351	0.313
RS3-7271-PE031023-N51499-1	PE5		26.2	10.8		37	0.708	0.360	0.319
RS3-7978-PE030806-N45396-1	PE5		26.5	10.6		37.1	0.714	0.353	0.314
RS1-0900-FE030027-4111200 RS1 0100 RE030605 4057651	PES	Lab 14	24	12		40	0.721	0.345	0.309
RS1-9190-FE030005-4057051 RS1-0201_PE021101_3032816	PE0 DE5	Lab 14	24	96		30.6	0.007	0.407	0.335
RS1-9291-FE021101-3932010 RS1-9291-PE030625-4072109	PE5	Lab 14	21	9.0 10		30.0	0.000	0.303	0.333
PS1_0302_PE021018_3022707	DE5	Lab 14	20	14		50	0.737	0.346	0.237
RS1-9493-PE021011-3917819	PE5	Lab 14	43	17		50 60	0.720	0.350	0.313
RS2-8483-PE021025-3928521	PE5	Lab 14		13		42	0.690	0.380	0.332
RS2-8887-PE030918-4125239	PE5	Lab 14	31	12		43	0.721	0.345	0.309
RS3-6766-PE030806-4096032	PE5	Lab 14	26	10		36	0.722	0.344	0.308
RS3-7170-PE030717-4084503	PE5	Lab 14	33	13		46	0.717	0.349	0.312
RS3-7271-PE031009-4142257	PE5	Lab 14	27	11		38	0.711	0.357	0.317
RS3-7271-PE031023-4150066	PE5	Lab 14	27	11		38	0.711	0.357	0.317
RS1-8988-PE030827-AG12595	PE5	Lab 15	33	11		44	0.750	0.313	0.288
RS1-9089-PE030606-AG04382	PE5	Lab 15	23	11		34	0.676	0.396	0.343
RS1-9291-PE021101-AF10570	PE5	Lab 15	29	10		39	0.744	0.320	0.292
RS1-9291-PE030625-AG06003	PE5	Lab 15	33	11		44	0.750	0.313	0.288
RS1-9392-PE021018-AF09487	PE5	Lab 15	35	12		47	0.745	0.319	0.291
RS1-9493-PE021004-AF08274	PE5	Lab 15	28	10		38	0.737	0.327	0.297
RS1-9493-PE021011-AF08745	PE5	Lab 15	31	11		42	0.738	0.326	0.296
RS1-9493-PE030317-AG01181	PE5	Lab 15	29	10		39	0.744	0.320	0.292
RS2-8483-PE021025-AF09981	PE5	Lab 15	27	10		37	0.730	0.335	0.303
RS2-8887-PE030918-AG14534	PE5	Lab 15	30	10		40	0.750	0.313	0.288
RS2-8986-PE030417-AG02189	PE5	Lab 15	25	11		36	0.694	0.375	0.329
RS3-7170-PE030717-AG08613	PE5	Lab 15	31	10		41	0.756	0.306	0.283
RS3-7271-PE031009-AG16592	PE5	Lab 15	30	11		41	0.732	0.333	0.301
R53-7271-PE031023-AG17951	PE0	Lab 15	30	10		41	0.732	0.333	0.301
RS3-7877-PEU31118-AG19392	PE0 DE5	Lab 15	20	10		30	0.722	0.344	0.308
RS3-7976-FE030600-AG10000	PES	Lab 15	22	12		40	0.733	0.331	0.300
RS1-9291-FE021024-300099	PE5	Lab 10	20	0.2		30.2	0.700	0.309	0.325
RS1_8988_PE030827_C3H280124022	PE5	Lab 10	21	J.Z 11		38	0.033	0.357	0.320
RS1-9089-PE021028-C2.1290172016	PE5	Lab 6	22	13	17	36.7	0.599	0.483	0.407
RS1-9291-PE030625-C3E260341011	PE5	Lab 6	32	13	1.7	46.3	0.691	0.379	0.336
RS1-9392-PE021021-C2J220181016	PE5	Lab 6	28	12		40	0 700	0.369	0.325
RS1-9493-PE021007-C2J080284011	PE5	Lab 6	28	14		42	0.667	0.407	0.350
RS1-9594-PE021014-C2J150280012	PE5	Lab 6	38	15		53	0.717	0.350	0.312
RS2-8483-PE030605-C3F060207001	PE5	Lab 6	28	14	0.95	43.0	0.652	0.423	0.364
RS2-8887-PE030918-C3I190390017	PE5	Lab 6	28	9.5		37.5	0.747	0.316	0.290
RS3-7271-PE031009-C3J110201011	PE5	Lab 6	17	8.9		25.9	0.656	0.418	0.358
RS3-7271-PE031023-C3J240278001	PE5	Lab 6	21	9		30	0.700	0.369	0.325
RS3-7776-PE030717-C3G180160009	PE5	Lab 6	33	14		47	0.702	0.367	0.323
RS3-7877-PE030806-C3H070274004	PE5	Lab 6	31	13		44	0.705	0.364	0.322
		Mean	28.43	11.32	1.32	39.83	0.713	0.35	0.32
		Median	28.00	11.00	1.30	39.00	0.716	0.35	0.31
	Standar	d Deviation	5.110	1.808	0.375	6.520	0.030	0.034	0.023
	Relati	ve Std Dev	0.182	0.164	0.289	0.167	0.042	0.097	0.074
Lab 4 Data (excluded)				4 -			0.050	45.05	1
RS1-9089-PE021030-1A2J0P830018			29	15		44	0.659	15.65	
R51-9392-PEU21016-1A2J0P496018		Lab 4	15	0.5		21.5	0.698	1.025	
RS1-3034-FEU21009-TA2JUF000004		LaD 4	19	9.0 10		20.3 26	0.071	9.000	
102-0700-1 LUZ 1020-1A200F019001			24	14		50	0.007	12.0	

"Calculated" Tri + based on (.03*Ar1221+(1.16*(Ar1242+1254))/(Ar1221+1242+1254) Literature Tri+ based on (0.1 *Ar1221)+(.85*Ar1242)+(.99*Ar1254)

Table A5-5 GE PE 2 Samples Aroclor Data

			Aroclor	Concentra	ations		Fraction	Calculated	Literature
LAB_SAMPLE_ID	PE ID	LAB	1221	1242	1254	Total PCB	1221	Tri+ PCBs	Tri +
RS1-9089-PE021030-N25643-21	PE2	Lab 1	17.6	8.21		25.81	0.682	0.389	0.339
RS1-9392-PE021016-N24752-12	PE2	Lab 1	11.1	5.8		16.90	0.657	0.418	0.357
RS1-9493-PE021009-N24212-10	PE2	Lab 1	14	7.41	0.618	22.03	0.636	0.442	0.377
RS2-8584-PE030529-N40160-1	PE2	Lab 1	11.2	5.58		16.78	0.667	0.406	0.349
RS3-6968-PE030730-N44844-11	PE2	Lab 1	11.1	5.62	0.577	17.30	0.642	0.435	0.373
RS3-7978-PE031002-N49731-1	PE2	Lab 1	9.82	5.5		15.32	0.641	0.436	0.369
RS1-9089-PE021029-3929926	PE2	Lab 14	9.8	5.3		15.10	0.649	0.427	0.363
RS1-9392-PE021015-3919355	PE2	Lab 14	9.4	5.3		14.70	0.639	0.437	0.370
RS1-9392-PE021022-3924871	PE2	Lab 14	11	5.8		16.80	0.655	0.420	0.359
RS1-9493-PE021008-3914767	PE2	Lab 14	11	6.5		17.50	0.629	0.450	0.379
RS2-8584-PE030529-4053262	PE2	Lab 14	11	6.5		17.50	0.629	0.450	0.379
RS3-7069-PE030730-4091703	PE2	Lab 14	11	6.5		17.50	0.629	0.450	0.379
RS3-7978-PE031002-4136357	PE2	Lab 14	9	5.6		14.60	0.616	0.463	0.388
RS1-9190-PE021028-AF10153	PE2	Lab 15	13	5.2		18.20	0.714	0.353	0.314
RS1-9392-PE021021-AF09641	PE2	Lab 15	15	7.5		22.50	0.667	0.407	0.350
RS1-9392-PE030317-AG01052	PE2	Lab 15	14	5.8		19.80	0.707	0.361	0.320
RS1-9392-PE030529-AG03661	PE2	Lab 15	12	5.4		17.40	0.690	0.381	0.333
RS1-9493-PE021007-AF08331	PE2	Lab 15	11	5.2		16.20	0.679	0.393	0.341
RS1-9594-PE021014-AF08824	PE2	Lab 15	12	5.6		17.60	0.682	0.390	0.339
RS2-8483-PE030317-AG01263	PE2	Lab 15	13	5.3		18.30	0.710	0.357	0.317
RS2-8986-PE021211-AF12606	PE2	Lab 15	15	6.1		21.10	0.711	0.357	0.317
RS2-8986-PE021218-AF12814	PE2	Lab 15	15	6.3		21.30	0.704	0.364	0.322
RS3-6968-PE030730-AG09977	PE2	Lab 15	12	6		18.00	0.667	0.407	0.350
RS3-7877-PE031211-AG19678	PE2	Lab 15	11	5		16.00	0.688	0.383	0.334
RS3-7978-PE031002-AG15884	PE2	Lab 15	13	5.8		18.80	0.691	0.379	0.331
RS1-9190-PE021025-386567	PE2	Lab 16	11	5.5	0.69	17.19	0.640	0.437	0.376
RS1-9392-PE021101-388327	PE2	Lab 16	8.5	4	0.48	12.98	0.655	0.420	0.364
RS1-9493-PE021011-383281	PE2	Lab 16	11	5.1	0.58	16.68	0.659	0.415	0.360
RS1-9392-PE021017-C2J180225018	PE2	Lab 6	9.6	6.2	0.7	16.50	0.582	0.503	0.420
RS1-9392-PE021031-C2K010155013	PE2	Lab 6	15	7.7		22.70	0.661	0.413	0.354
RS1-9392-PE030529-C3E300201007	PE2	Lab 6	12	6.6	0.74	19.34	0.620	0.459	0.390
RS1-9493-PE021010-C2J110279021	PE2	Lab 6	12	6.3		18.30	0.656	0.419	0.358
RS2-8584-PE021024-C2J250174001	PE2	Lab 6	9.6	6.5	0.71	16.81	0.571	0.515	0.428
RS3-6766-PE030730-C3G310148023	PE2	Lab 6	12	8.3	0.73	21.03	0.571	0.515	0.427
RS3-7877-PE031002-C3J030129006	PE2	Lab 6	9	5.5	0.68	15.18	0.593	0.490	0.412
		Mean	11.79	6.01	0.65	17.99	0.65	0.42	0.36
		Median	11.10	5.80	0.69	17.50	0.66	0.42	0.36
:	Standard	Deviation	2.067	0.929	0.084	2.687	0.038	0.043	0.030
	Relativ	e Std Dev	0.186	0.160	0.122	0.154	0.059	0.104	0.085
Lab 4 Data (excluded)									
RS1-9190-PE021028-TA2J0P768018	PE2	Lab 4	14	8.7		22.70	0.617	0.463	
RS1-9392-PE021021-TA2J0P599010	PE2	Lab 4	9	4.6		13.60	0.662	0.412	
RS1-9594-PE021007-TA2J0P236006	PE2	Lab 4	8.7	4.9		13.60	0.640	0.437	
RS1-9594-PE021014-TA2J0P418015	PE2	Lab 4	8	4.8		12.80	0.625	0.454	

"Calculated" Tri + based on (.03*Ar1221+(1.16*(Ar1242+1254))/(Ar1221+1242+1254) Literature Tri+ based on (0.1 *Ar1221)+(.85*Ar1242)+(.99*Ar1254)

Table A5-6 Field Duplicate RPDs

				Homologs					
		> 5 mg/kg			1 - 4.9 mg/kg	All Data			
	Total PCB	Ar1221	% Ar1221	Total PCB	Ar1221	% Ar1221	Total PCB	Tri+ PCB	% Tri +
Number of pairs	723	720	720	259	259	259	79	79	79
95th percentile RPD	88	97.7	20.1	115	129	30.5	90	94	25
90th percentile RPD	60	69.8	13.4	78	84.4	21	60	65	15
80th percentile RPD	39	42.9	8.0	52.7	57.6	13.5	37	43	11
Median (50th) RPD	16.3	17.5	2.8	22.2	25.4	5.3	16.8	16.8	5.7
Percent RPD <40	81	77.6	99	70.7	64.3	96.1	82.3	78.5	99

Table A5-7

		6-in Segment	22-in Segment
River Section 1 - MPA Threshold	3 g/ m ²		
Finer Grained Areas			
Bulk Density	g/cc	0.85	0.85
Tri+ PCB needed to exceed			
MPA criteria	ppm	23	6.4
Coarser Grained Areas			
Bulk Density	g/cc	1.8	1.8
Tri+ PCB needed to exceed			
MPA criteria	ppm	11	3
River Sections 2 & 3 - MPA Three	shold 10 g/ m ²		
Finer Grained Areas			
Bulk Density	g/cc	0.85	0.85
Tri+ PCB needed to exceed			
MPA criteria	ppm	77	21

Critical Tri+ PCBs Concentration Range

Notes:

- 1. The MPA is the product of the Tri+ PCB concentration, length and bulk density. The concentration in finer-grained areas must be higher to have the same MPA as a coarse grained area. The table above shows the range of critical concentrations.
- 2. In River Section 1, the MPA threshold is 3 g/m². For coarse-grained areas of River Section 1 and thicker segments, the concentration would only need to be 3 ppm or higher to exceed the threshold.
- 3. In River Sections 2 and 3, the MPA threshold is 10 g/m². For finer-grained areas and shorter segments, the concentration would need to be 77 ppm or higher to exceed the threshold. 80 ppm or higher data tend to be widely scattered, but are likely to result in exceedance of the removal criteria unless there is a pronounced bias in the predicted values.

Table A5-8 Lower and Upper Limits for Predicted Sediment Tri+ PCB using the Point-by-Point Correction Regression Model

Corrected Tri+ PCB Range	Lower Limit	Upper Limit	
(mg/kg)	(%)	(%)	
0-5	NA	NA	
5-80	-29	38	
80+	-44	41	

Model:

1) Tri+ Concentration = a[A21]+b[A42+A54]

Table A5-9Statistics of the 2009 Regression Models

Statistic	Model using original 180	Model using full data set
A1221 Coefficient	0.126	0.140
Std. Error of A1221 Coefficient	0.0114	0.0105
A1242+A1254 Coefficient	0.944	0.913
Std. Error of A1242+A1254 Coefficient	0.0293	0.0253
Multiple R ²	0.970	0.961

Table A5-10
Regression Coefficients Update

Round	Regression Date	Number of Samples		Tri+ PCB = a[A1221] + b[A1242 + A1254]		
		Additional	Total	а	b	
Original	In QAPP 2009		278	0.14	0.91	
2009 Round 1	8/14/2009	21	299	0.14	0.91	
2009 Round 2	9/3/2009	63	362	0.13	0.90	
2009 Round 3	9/17/2009	21	383	0.13	0.90	
2009 Round 4	10/26/2009	20	403	0.13	0.89	
2011	7/29/2011	42	445	0.13	0.89	
	Original			2011		
---	----------	---------	---------	---------	---------	--------
Statistic	Original	Round 1	Round 2	Round 3	Round 4	2011
A1221 Coefficient	0.140	0.140	0.134	0.128	0.127	0.125
Std. Error of A1221 Coefficient	0.0105	0.0099	0.0083	0.0079	0.0076	0.0071
A1242+A1254 Coefficient	0.913	0.911	0.893	0.895	0.892	0.886
Std. Error of A1242+A1254 Coefficient	0.0253	0.0235	0.0209	0.0203	0.0195	0.0183
Multiple R2	0.961	0.962	0.959	0.959	0.959	0.960

Table A5-11Statistics of the 2009 and 2011 Regression Models

Table A5-12

Feature Selection Measures for Several OLS Linear Regression Models for Water Data

Models	Linear Regression Formula	LOOCV - PRESS	AIC	AICc	BIC	Adjusted R ²
	$Tri+PCB = a \times A1221 + b \times$					
GE Model	(A1242+A1254)	100	791	791	801	0.96
	$Tri+PCB = a \times A1221 + b \times A1242 + c \times$					
Test Model 1	A1254	101	1101	1102	1115	0.95
Test Model 2	$Tri+PCB = a \times (A1221+A1242+A1254)$	188	1252	1252	1258	0.91

Note:

LOOCV: leave-one-out cross-validation

PRESS: prediction residual sum of squares

AIC: Akaike's Information Criterion

AICc: corrected AIC

BIC: Schwarz's Bayesian Information Criteria

Table A5-13 List of OLS and Several Robust Regression Models of Water Data

Method Acronym	Method Name	R function
OLS	Ordinary Least Square Regression	lm()
LAV	Least Absolute Values Regression	rq() in package quantreg
M-Huber	M-Estimation with Huber Weight	rlm() in package MASS
M-Bisquare	M-Estimation with Bisquare Weight	rlm(,psi = psi.bisquare) in package MASS
LTS	Least Trimmed Squares Regression	ltsreg() in package lqs
LMS	Least Median Squares Regression	lmsreg() in package lqs

Table A5-14 **Regression Results for OLS and Several Robust Regression Models for Water Data**

	Group 1 (n=68) ¹	¹ Group 2 (n=136) ²		Group 3	(n=35) ³	All Samples (n=239) ⁴		
Model	а	а	b	а	b	а	b	
OLS	0.236	0.126	0.757	0.191	0.696	0.143	0.749	
LAV	0.247	0.129	0.709	0.188	0.721	0.146	0.694	
M-Huber	0.240	0.128	0.722	0.185	0.721	0.144	0.721	
M-Bisquare	0.245	0.128	0.708	0.185	0.720	0.145	0.699	
LTS	0.258	0.168	0.384	0.185	0.749	0.223	-0.037	
LMS	0.274	0.134	0.612	0.188	0.721	0.227	-0.021	

Note: all the regression analysis was based on the formula of $Tri+PCB = a \times A1221 + b \times (A1242+A1254)$

Group 1 corresponds to the first correlation in GE's model.
 Group 2 corresponds to the second correlation in GE's model.

(3) Group 3 corresponds to the third correlation in GE's model.

(4) Models were developed using all available samples.

Table A5-15
Cross Validation Results and the Model Prediction Error of Water Data

Model		Group 1 (n=68) ¹	Group 2 (n=136) ²	Group 3 (n=35) ³	All Samples (n=239) ⁴
OLS	CV (ng/L)	4.912	7.160	16.343	9.988
LAV	CV (ng/L)	4.924	7.363	15.912	9.992
M-Huber	CV (ng/L)	4.897	7.167	16.297	10.011
M-Bisquare	CV (ng/L)	4.915	7.195	16.306	10.048
LTS	CV (ng/L)	5.060	9.064	15.952	17.583
LMS	CV (ng/L)	5.738	7.668	16.440	16.625
Mean of Measured	18	41	68	38	
Mean of Prediction Error f	4.897	7.160	15.912	9.988	
Relative Prediction Err	or for the Best Model	27%	18%	23%	26%

Note: CV-mean of the estimated prediction errors.

(1) Group 1 corresponds to the first correlation in GE's model.

(2) Group 2 corresponds to the second correlation in GE's model.

(3) Group 3 corresponds to the third correlation in GE's model.

(4) Models were developed using all available samples.

Table A5-16

Water Column Data Model Validation Results using Test Data Set

	Model	Group 1 (n=12)	Group 2 (n=23)	All Samples (n=35)
GE Model	Average prediction error	2.447	5.085	NA
OLS	Average prediction error	3.174	5.337	5.761
LAV	Average prediction error	2.892	5.154	5.545
M-Huber	Average prediction error	3.054	5.192	5.638
M-Bisquare	Average prediction error	2.944	5.144	5.555
LTS	Average prediction error	2.630	4.661	6.069
LMS	Average prediction error	2.396	5.413	6.608
Mean of	Measured Tri+ PCB (ng/L)	10	34	26
Mean of Predicti	on Error for the Best Model (ng/L)	2.396	4.661	5.545
Relative Pred	iction Error for the Best Model	24%	14%	22%

Table A5-17

Regression Equations for EPA's Best Model.

Model	First Regression	Second Regression	Third Regression
EPA's Best Model ¹	0.274×A1221 (LMS Model)	0.126×A1221+0.757×(A1242+A1254) (OLS Model)	0.188×A1221+0.721×(A1242+A1254) (LAV Model)
GE weighted least squares (i.e., "damped- leveraged) algorithm	0.27×A1221	0.12×A1221+0.73×(A1242+A1254)	0.16×A1221+0.85×(A1242+A1254)

Note: (1) Best Model was selected among OLS and several robust regression models discussed in this work (LAV, M-Huber, M-Bisquare, LTS, LMS)

			Aro	clors R	eporteo	1		
Year	1016	1221	1242	1248	1254	1260	1254/ 1260	Laboratory as reported in NYSDEC Database
1000	53%						47%	Hale Creek
1990	28%	0.2%			72%			Hazleton Laboratories
1001	52%				10%		38%	Dept. of Health
1991	47%				17%		37%	Hale Creek
1002	54%				44%	0.3%	2%	Hale Creek
1992			4%	12%	74%	11%		Hazleton Laboratories
1002	58%					42%		Hale Creek
1995			3%	32%	53%	12%		Hazleton Laboratories
1994			2%	27%	59%	12%		Hazleton Laboratories
1995				29%	53%	17%		Hazleton Laboratories
1006	86%						14%	Hale Creek
1990				31%	54%	16%		Hazleton Laboratories
1007	0.004%			31%	46%	23%		EnChem Environmental Laboratories
1997				5%	70%	25%		Hazleton Laboratories
1998				41%	38%	21%		EnChem Environmental Laboratories
1999			21%	30%	28%	21%		Mississippi St. Chem. Laboratories
2000			19%	27%	28%	25%		Mississippi St. Chem. Laboratories
2001			18%	23%	30%	29%		Mississippi St. Chem. Laboratories
2002			20%	24%	33%	23%		Mississippi St. Chem. Laboratories
2003			22%	26%	32%	20%		Mississippi St. Chem. Laboratories
2004			21%	24%	35%	21%		Mississippi St. Chem. Laboratories
2005			10%	14%	41%	35%		Mississippi St. Chem. Laboratories
2006			10%	13%	42%	35%		Mississippi St. Chem. Laboratories
2007			13%	18%	40%	29%		Mississippi St. Chem. Laboratories
2008			5%	16%	44%	34%		Mississippi St. Chem. Laboratories
2009			12%	21%	32%	35%		Mississippi St. Chem. Laboratories
2010			10%	20%	37%	32%		Mississippi St. Chem. Laboratories
2011			12%	18%	38%	31%		Mississippi St. Chem. Laboratories

 Table A5-18

 Aroclors Reported By Year as an Average Percentage of TPCB_{Aroclor} – NYSDEC Data

Note, for example, the different mixtures reported by different laboratories in the same year of collection. Additionally, Aroclor 1016 and an unresolved mixture of Aroclors 1254/1260 are reported only in 1996 and prior (primarily by the Hale Creek lab), while Aroclor 1242 appears very infrequently among all labs through 1998. Aroclor 1221, which can be inferred to indicate the presence of monochloro and dichloro homologues in the samples, was only reported in 1990 by Hazelton Laboratories.¹

¹ The lack of reported Aroclor 1221 in these data are further evidence of the lack of monochloro and dichloro homologues in fish tissue, as reported in the various documents supporting the EPA ROD.

 Table A5-19

 Aroclors Reported By Year as an Average Percentage of TPCB_{Aroclor} – GE Data

		Aro	clors Repo	rted		
Year	1221	1242	1248	1254	1260	Laboratory
2004	1.47%	0.66%	56.65%	38.17%	3.05%	
2005	1.11%	0.38%	56.31%	39.54%	2.65%	
2006	0.38%	0.32%	56.45%	38.00%	4.85%	
2007	0.49%	0.32%	57.88%	36.38%	4.93%	
2008	0.74%	0.23%	59.50%	38.08%	1.45%	
2009	4.26%	1.73%	53.50%	38.18%	2.33%	Northeast
2010	3.27%	1.88%	46.04%	44.03%	4.78%	Analytical
2011	12.38%	4.20%	45.21%	34.69%	3.52%	lnc.
2012	19.78%	0.15%	42.52%	30.99%	6.55%	
2013	19.56%	0.56%	43.88%	27.29%	8.72%	
2014	16.21%	0.09%	46.51%	30.39%	6.80%	
2015	12.17%	0.14%	44.43%	32.23%	11.03%	
2016	3.87%	0.00%	49.71%	34.20%	12.23%	

Highlighted area represents the remedial dredging period and a change in Aroclors as reported by GE

Data Source	Period of Available Data	Applicable Laboratory Codes	Equation to Obtain the Homologue Equivalent Total PCB Concentration (TPCB _{HE})	Equation Source	Period of Application
	1990	HES, U	1.3070 * (Aroclor 1016 + Aroclor 1254)	USEPA. 2000. Further Site Characterization and Analysis, Revised Baseline Modeling Report (RBMR), Hudson River PCBs Reassessment RI/FS	1990
	1990-1993, 1996	HC, DOH	Deb pyEquation to Obtain the Homologue Equivalent Total PCB Concentration (TPCB _{HE})Equation SourcePe AppJ1.3070 * (Aroclor 1016 + Aroclor 1254)USEPA. 2000. Further Site Characterization and Analysis, Revised Baseline Modeling Report (RBMR), Hudson River PCBs Reassessment RI/FS Volume 2D. Prepared for USEPA Region 2 and USACE, Kansas City District by TAMS Consultants, Inc., Limno-Tech, Inc., Menzie-Cura & Associates, Inc., and Tetra-Tech, Inc. January 2000.1990.ES,0.8754 * (Aroclor 1248 + Aroclor 1254 + Aroclor 1260)Geometric mean of TPCB _{HE} /TPCB _{Aroclor} 1998 NYSDEC. See Figure 4.19I1.1743 * TPCB _{Aroclor} Geometric mean of TPCB _{HE} /TPCB _{Aroclor} 2004-2008 GE data. See Figure 12.200.788 * TPCB _{Aroclor} Geometric mean of TPCB _{HE} /TPCB _{Aroclor} 2009-2013 GE data. See Figure 16.20	Volume 2D. Prepared for USEPA Region 2 and USACE, Kansas City District by TAMS Consultants, Inc., Limno-Tech, Inc., Menzie-Cura	
Data Source A	1992-1997	ENC, HES, HC	0.8754 * (Aroclor 1248 + Aroclor 1254 + Aroclor 1260)	& Associates, Inc., and Tetra-Tech, Inc. January 2000.	1992-1997
	1998	ENC	$0.7407 * TPCB_{Aroclor}$	Geometric mean of TPCB _{HE} /TPCB _{Aroclor} 1998 NYSDEC. See Figure 4.	1998
	1999-2000	MSC	$1.1743 * TPCB_{Aroclor}$	Geometric mean of TPCB _{HE} /TPCB _{Aroclor} 1999-2000 NYSDEC. See Figure 8.	1999-2011
	2004-2008	NEA	$0.8542 * TPCB_{Aroclor}$	Geometric mean of TPCB _{HE} /TPCB _{Aroclor} 2004-2008 GE data. See Figure 12.	2004-2008
GE	2009-2013 NEA 0.788 * TPCB _{Aroclor}		$0.788 * \mathrm{TPCB}_{\mathrm{Aroclor}}$	Geometric mean of $\text{TPCB}_{\text{HE}}/\text{TPCB}_{\text{Aroclor}}$ 2009-2013 GE data. See Figure 16.	2009-2013

Table A5-20Fish Tissue Regression Equations

Notes:

TPCB_{Aroclor} refers to the sum of detected Aroclor concentrations in the sample. See text for discussion.

Lab Code Key

DOH Dept. of Health ENC Enchem Environmental Laboratories HES Hazelton Laboratories HC Hale Creek MSC Mississippi St. Chem. Laboratories NEA Northeast Analyical Laboratories U Unknown

Coefficient (a) Summary Statistics															
D .		Statistics for Determination of α									Confidence Interval for the Geometric Mean (Bootstrap Analysis)				ic Mean
Source	Period	Number of Samples	Arithmetic Mean	Std. Dev.	Median	Geometric Mean	Regression Estimate	Minimum Observed	Maximum Observed	2.5%	5% ¹	95% ¹	97.5%	95% CI expressed as a Percentage of the Mean	
NYSDEC	1998	82	0.826	0.535	0.783 *	0.741	0.784 *	0.129	4.840	0.672	0.684	0.802	0.813	-7.7%	8.2%
NYSDEC	1999-2000	173	1.284	0.637	1.181 *	1.174	0.871	0.285	5.996	1.104	1.114	1.236	1.248	-5.1%	5.3%
GE	2004-2008	259	0.893	0.334	0.831 *	0.854	0.703	0.151	4.210	0.826	0.830	0.880	0.885	-2.8%	3.0%
GE	2009-2013	140	0.823 **	0.244	0.784 *	0.788	0.851	0.101	2.226	0.746	0.752	0.822	0.828	-4.5%	4.3%

Table A5-21 Coefficient (α) Summary Statistics

Note:

1 Value used in estimated 95% CI as percentage of the geometric mean.

* Value falls between 5th and 95th percent confidence interval for the geometric mean.

** Value falls between 2.5th and 97.5th percent confidence interval for the geometric mean.

River Mile	2004	2005	2006	2007	2008	2009	2010	2011	2012	2013
206	1.1	0.8	0	0	0	0	0	2.2	4.3	6.1
194	1.9	1.8	0	0	0	12.2	0	0.6	1.9	2.6
193	6.7	3.0	0	0	0	4.9	6.0	18.2	8.9	8.1
192	7.1	0	0	0	0	6.7	2.1	7.8	35.5	2.4
191	0	0	4.6	2.0	0	2.1	2.1	11.7	26.4	7.2
190	4.9	0	0	0	0	10.9	3.0	6.5	25.3	15.2
188	0	0	0	0	NA	0	2.8	16.0	25.7	9.0
187	0	0	0	0	NA	0	0	13.1	25.0	13.4
186	0	9.0	0	1.5	2.2	3.1	0	16.3	21.8	21.1
184	1.5	1.5	0	0	3.0	0.9	9.9	18.3	26.3	36.9
182	0	0	0	0	0	0	3.8	14.7	15.6	23.1
178	0	0	2.0	2.4	0	7.9	3.7	17.0	22.1	22.4
178	0	1.5	0	0	0	0	0	9.0	10.3	13.0
173	0	0	2.3	0	3.9	4.8	4.1	21.0	23.2	34.8
169	0	4.1	0	3.9	0	8.2	7.7	18.6	27.5	34.0
148	0	0	0	0	0	5.0	5.9	16.7	24.4	28.9
115	NA	NA	NA	NA	NA	NA	0.3	5.5	4.2	13.8

 $Table \ A5-22 \\ Average \ Aroclor \ 1221 \ as \ Percentage \ of \ TPCB_{Aroclor} \ by \ River \ Mile \ by \ Year$

Notes:

Highlighted cells indicate fish sample results obtained from river subsections in which dredging had occurred that year.

In 2013, dredging also occurred at CU-99 at RM 159 and CU-100 at RM 154.3. These were small areas and are not indicated on the table.

NA Not Available





Figure A5-2 Regression Results for GE 2 Model (Point by Point Correction)



Figure A5-3 Measured vs Predicted Tri+ Fraction for the GE 2 Model (Point by Point Correction)



Figure A5-4 Regression Residuals for GE 2 Model (Point by Point Correction)



Figure A5-5 Upper and Lower Bound for Corrected vs Predicted Tri+ PCBs using Point-by-Point Correction Regression

















Quantiles			Summary Statistics		
100.0%	maximum	4.84	Mean	0.83	
99.5%		4.84	Std Dev	0.54	
97.5%		2.25	Std Err Mean	0.06	
90.0%		1.11	Upper 95% Mean	0.94	
75.0%	quartile	0.92	Lower 95% Mean	0.71	
50.0%	median	0.78	Ν	82.00	
25.0%	quartile	0.59			
10.0%		0.46			
2.5%		0.31			
0.5%		0.13			
0.0%	minimum	0.13			



Distribution of NYSDEC 1998 TPCB_{HE} / TPCB $_{Aroclor}$ Ratio for Individual Samples



TPCB HE	10.4749	t-Ratio	-4.24056
IPCB Aroclors	13.1468	DF	81
Mean Difference	-2.6719	Prob > t	<.0001*
Std Error	0.63009	Prob > t	1.0000
Upper 95%	-1.4182	Prob < t	<.0001*
Lower 95%	-3.9256		
Ν	82		
Correlation	0.76951		

Wilcoxon Signed Rank				
	TPCB _{HE} – TPCB _{Aroclors}			
Test Statistic S	-1193.5			
Prob> S	<.0001*			
Prob>S	1.0000			
Prob <s< td=""><td><.0001*</td></s<>	<.0001*			



Wilcoxon test on 1998 NYSDEC Matched Pairs (TPCB_{HE} - TPCB $_{Aroclor}$) for Individual Samples













TP TP	PCB _{HE}	9.27859	t-Ratio	2.844704
	Aroclors	8.18697	DF	172
Me	an Difference	1.09162	Prob > t	0.0050 *
Sto	l Error	0.38374	Prob > t	0.0025 *
Up	per 95%	1.84907	Prob < t	0.9975
Lo	wer 95%	0.33418		
Ν		173		
Co	rrelation	0.9268		
	Wilcoxon Signed	Rank		
		TPCB _{HE} – TPCB _{Aroc}	lors	
	Test Statistic S	36	600.50	
	Prob> S	<.0001*		
	Prob>S	<.0001*		
	Prob <s< td=""><td>1.0000</td><td></td><td></td></s<>	1.0000		





Log (TPCB _{HE}) Log (TPCB ,)	0.79532 0.72556	t-Ratio DF	5.075368 172
Interence	0.06976	Prob > t	<.0001*
Std Error	0.01375	Prob > t	<.0001*
Upper 95%	0.09689	Prob < t	1.0000
Lower 95%	0.04263		
Ν	173		
Correlation	0.89763		

Wilcoxon Signed Rank					
	Log (TPCB _{HE})– Log (TPCB _{Aroclors})				
Test Statistic S		3681.50			
Prob> S	<.0001*				
Prob>S	<.0001*				
Prob <s< td=""><td>1.0000</td><td></td></s<>	1.0000				







Quantiles			Summary Statistics	
100.0%	maximum	4.21	Mean	0.89
99.5%		3.79	Std Dev	0.33
97.5%		1.70	Std Err Mean	0.02
90.0%		1.16	Upper 95% Mean	0.93
75.0%	quartile	0.97	Lower 95% Mean	0.85
50.0%	median	0.83	Ν	259.00
25.0%	quartile	0.73		
10.0%		0.65		
2.5%		0.54		
0.5%		0.24		
0.0%	minimum	0.15		





2.12049	t-Ratio	-5.49086
2.70432	DF	258
-0.5838	Prob > t	<.0001*
0.10633	Prob > t	1.0000
-0.3744	Prob < t	<.0001*
-0.7932		
259		
0.94907		
	2.12049 2.70432 -0.5838 0.10633 -0.3744 -0.7932 259 0.94907	2.12049 t-Ratio 2.70432 DF -0.5838 Prob > t 0.10633 Prob > t -0.3744 Prob < t -0.7932 259 0.94907 ↓

Wilcoxon Signed Rank				
	TPCB _{HE} – TPCB _{Aroclors}			
Test Statistic S		-12659		
Prob> S	<.0001*			
Prob>S	1.0000			
Prob <s< td=""><td><.0001*</td><td></td></s<>	<.0001*			





Log (TPCB _{HE})	-0.0239	t-Ratio	-8.92025
Log (TPCB Araclars)	0.04453	DF	258
	-0.0684	Prob > t	<.0001*
Std Error	0.00767	Prob > t	1.0000
Upper 95%	-0.0533	Prob < t	<.0001*
Lower 95%	-0.0835		
Ν	259		
Correlation	0.98472		

Wilcoxon Signed Rank				
	Log (TPCB _{HE})– Log (TPCB _{Aroclors})			
Test Statistic S	-	-11438		
Prob> S	<.0001*			
Prob>S	1.0000			
Prob <s< td=""><td><.0001*</td><td></td></s<>	<.0001*			



May 2017





Quantiles			Summary Statistics		
100.0%	maximum	2.23	Mean	0.82	
99.5%		2.23	Std Dev	0.24	
97.5%		1.40	Std Err Mean	0.02	
90.0%		1.11	Upper 95% Mean	0.86	
75.0%	quartile	0.94	Lower 95% Mean	0.78	
50.0%	median	0.78	Ν	140.00	
25.0%	quartile	0.69			
10.0%		0.61			
2.5%		0.47			
0.5%		0.10			
0.0%	minimum	0.10			



Distribution of 2009 to 2013 GE $\mbox{TPCB}_{\rm HE}$ / $\mbox{TPCB}_{\rm Aroclor}$ for Individual Samples



TPCB _{HE} TPCB _{Aroclors}	2.27202 2.87119	t-Ratio DF	-5.92639 139
Mean Difference	-0.5992	Prob > t	<.0001*
Std Error	0.1011	Prob > t	1.0000
Upper 95%	-0.3993	Prob < t	<.0001*
Lower 95%	-0.7991		
Ν	140		
Correlation	0.97011		

Wilcoxon Signed Rank					
	TPCB _{HE} – TPCB _{Aroclors}				
Test Statistic S		-4036.0			
Prob> S	<.0001*				
Prob>S	1.0000				
Prob <s< td=""><td><.0001*</td><td></td></s<>	<.0001*				





	Log (TPCB _{HE})	-0.018 0.08546	t-Ratio DF	-8.98789 139
LOg (TF	LOg (IPCD Aroclors)	-0.1035	Prob > t	<.0001*
	Std Error	0.01151	Prob > t	1.0000
	Upper 95%	-0.0807	Prob < t	<.0001*
	Lower 95%	-0.1262		
	Ν	140		
	Correlation	0.97326		








