US ERA ARCHIVE DOCUMENT

CATALOG DOCUMENTATION EMAP-ESTUARIES PROGRAM LEVEL DATABASE LOUISIANIAN PROVINCE 1991-1994 SEDIMENT CHEMISTRY DATA

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APPENDIX A.

APPENDIX B.

- 1. DATA SET IDENTIFICATION
 - 1.1 Title

EMAP-Estuaries Province Level Database Louisianian Province Sediment Chemistry Data

1.2 Catalog Author

Virginia Engle, U.S. Environmental Protection Agency - NHEERL/GED Linda Harwell, U.S. Environmental Protection Agency - NHEERL/GED Tom Heitmuller, U.S. Geological Survey - BRD/GBPO

1.3 Catalog Revision Date

June 16, 1999

1.4 Data Set Name

SED_CHEM

1.5 Task Group

ESTUARIES

1.6 Data set identification code

00045, 00085, 00125, 00165

1.7 Version number for a data set

003, 003, 004, 002

1.8 Requested acknowledgment

If you plan to publish these data in any way, EPA requires a standard statement for work is has supported:

"Although the data described in this article have been funded wholly or in part by the U.S. Environmental Protection Agency through its EMAP Estuaries Program, it has not been subjected to Agency review, and therefore does not necessarily reflect the views of the Agency and no official endorsement should be inferred."

2. INVESTIGATOR INFORMATION

2.1 Principal Investigator

John M. Macauley U.S. Environmental Protection Agency NHEERL - GED

2.2 Sample Collection Investigator

John M. Macauley U.S. Environmental Protection Agency NHEERL - GED

2.3 Sample Processing Investigator

Tom Heitmuller U.S. Geological Survey BRD - GBPO

2.4 Data Analysis Investigator

Virginia D. Engle U.S. Environmental Protection Agency NHEERL - GED

2.5 Additional Investigators

N/A

3. DATA SET ABSTRACT

3.1 Abstract of the Data Set

The Sediment Chemistry data file presents a suite of chemical concentrations derived from analyses of a surface sediment sample collected at a station in the Louisianian Province. Individual and summed analyte concentrations are presented.

A code for each compound is given under ANALYTE. These include inorganics, organics, organic concentration sums, total organic carbon (TOC) and acid volatile sulfides (AVS). Concentrations are recorded in dry weight. Units are reported under a separate attribute, CHMUNITS, as ug/g, ng/g, % or µmoles/g. Quality Assurance/Quality Control issues are coded. Depending on the QA code, only a detection limit may be reported.

3.2 Keywords for the Data Set

Contaminants, DDT, metals, inorganic analytes, organic analytes, alkanes, isoprenoids, PAH, PCB, pesticides, QA Code, sediment, sediment chemistry, TOC and AVS.

4. OBJECTIVES AND INTRODUCTION

4.1 Program Objective

The Environmental Monitoring and Assessment Program (EMAP) was designed to periodically estimate the status and trends of the Nation's ecological resources on a regional basis. EMAP provides a strategy to identify and bound the extent, magnitude and location of environmental degradation and improvement on a regional scale based on randomly located station sites. Only the randomly located Base Sampling Sites were included in this data set.

4.2 Data Set Objective

The objective of the Sediment Chemistry data file is to present the concentrations of a suite of analytes and compounds measured from a surface sediment sample collected at a station in the Louisianian Province. The sample was an homogenate composed of the surficial 2 cm from several grabs.

4.3 Data Set Background Information

The presence of contaminants in estuaries has been identified in both the scientific and popular press as a major problem contributing to degraded ecological resources and restricted harvest of fish and shellfish resources due to human health concerns. Reducing contaminant inputs and concentrations, therefore, is often a major focus of regulatory programs for estuaries. Contaminants include both inorganic (primarily metals) and organic forms originating from many sources, including atmospheric deposition, freshwater inputs, land runoff and point sources. These sources are poorly characterized, except in the most well-studied estuaries. Most contaminants that are potentially toxic to indigenous biological resources tend to bind to particles, which

ultimately are deposited at the bottom of estuaries. This binding changes the form of contaminants and removes them from the water column; consequently, contaminants accumulate in estuarine sediments.

Sediment contaminant concentrations were measured to aid in the interpretation of the spatial patterns observed in the condition of biological resources in the estuaries of the Louisianian Province.

EMAP monitoring efforts have focused on sediment contaminants rather than measurement of water column contaminants because concentrations of contaminants in sediments are less variable and the sediment integrates contaminant inputs to estuaries over time (i.e., months and years).

Metals in the sediment are derived from anthropogenic sources or from the natural geochemical processes of weathering and erosion of the earth's crust. The difficulty arises in identifying which portion of the total metal content of the sediment was due to natural processes and which was due to human activities. Several methods can be used to determine whether measured metal concentrations in estuarine sediments represent anthropogenically-enriched or natural conditions.

4.4 Summary of Data Set Parameters

The organic and inorganic compound concentrations measured generally included: 15 major and trace elements, 25 Alkanes, 2 Isoprenoids, 44 individual Poly-Aromatic Hydrocarbon (PAH) compounds, the pesticide, DDT, and its metabolites, 19 pesticides other than DDT, 20 individual Poly-Chlorinated Biphenyl (PCB) congeners, mono-, diand tri-butyltin (MBT, DBT, TBT), Total Organic Carbon (TOC) and acid volatile sulfides (AVS). This suite of analytes is similar to that measured in the National Oceanic and Atmospheric Administration's (NOAA) National Status and Trends (NS&T) program with the addition of aliphatic hydrocarbons and several PAHs. Values in this data file include individual inorganic and organic compound concentrations and concentrations summed for several major groups: total PAHs, Low and High Molecular Weight PAHs, PCBs, DDTs, BHCs, Chlordanes, Alkanes, and Isoprenoids.

Concentrations of all sediment chemistry analytes are reported on a dry weight basis.

4.5 Year-Specific Information about Data

During the first three years of the program (1991-93), all sediment chemistry was analyzed by the Geochemical and Environmental Research Group (GERG) at Texas A& M University. The quantitative list of analytes listed in 3.4 reflects the list of analytes provided by this laboratory in exceedance of the requirements of EMAP. In addition to this list, in 1993, two organophosphates (Chlorpyrifos and Dicofol) were reported. In 1994 the sediment chemistry analyses were conducted by the University System of Georgia, Skidaway Institute of Oceanography, Savannah, GA. In 1991-92, both TOC and AVS were analyzed by the Department of Biology at the University of Mississippi. In 1993, TOC was analyzed by the

Department of Biology at the University of Mississippi and AVS was analyzed by the Chemistry Department at the Gulf Coast Research Lab.

In 1994, AVS was analyzed at GCRL; TOC analyses were subcontracted through GCRL to the Department of Geology at the University of South Carolina.

In 1994, sediment samples were no longer analyzed for alkanes.

5. METHODS

5.1 Data Acquisition

5.1.1 Sampling Objective

Collect sediment samples suitable for the analysis of organic and inorganic compounds, TOC and AVS. One sediment sample was expected to be collected at each station.

5.1.2 Sample Collection Methods Summary

The grab sampler was lowered through the water column such that travel through the last 5 meters was no faster than 1 m/sec. The grab penetrated the sediment by gravity releasing a trigger allowing the jaws to close. When the grab was pulled from the sediment using the winch, the jaws closed, encapsulating the sediment sample. The chance of sampling the exact same location twice was minimized. After three grabs were taken, the boat was moved five meters downstream by letting out the appropriate length of anchor line.

Stainless steel utensils were used to remove the top two cm of sediment from a grab. The sediment was removed to a stainless steel bowl and placed in a cooler of ice to remain cold, but The grab sampler was rinsed and re-deployed. procedure was repeated until approximately 3,000 cc of sediment was collected. The sediment was mixed by hand until thoroughly homogenized, and aliquots were placed immediately into precleaned glass jars (for organics) or plastic containers (for inorganics and AVS). In 1992-1994, the AVS subsample was taken independently from the composited, homogenated sediment. For AVS, a small plug was taken at a depth of 2-3 (cm) from each of the 3-4 grabs; the plugs were combined into a 60-ml wide-mouth container until the container was nearly devoid of any This procedure was adopted to avoid the loss of volatile sulfides by aeration or release during the homogenization process with the bulk composite sediment. sample container was held on ice between the collection of individual plugs. The completed samples were immediately stored on ice following collection. The remainder of the sediment was split between grain size and sediment toxicity samples.

5.1.3 Beginning Sampling Date

09 July 1991

08 July 1992

06 July 1993

06 July 1994

5.1.4 Ending Sampling Date

10 September 1991

- 11 September 1992
- 19 August 1993
- 15 September 1994

5.1.5 Sampling Platform

Each team was supplied with a 25-foot SeaArk work boat equipped with a 7.5 L gas engine fitted with a Bravo outdrive, an "A" frame boom assembly and hydraulic winch. On-board electronics consist of: a Loran C unit, GPS (beginning in 1993), radar unit, 2 VHF radios, cellular phone, compass, a depth finder and a pump-driven seawater washdown system. Each vessel was provided with an extensive tool kit and all required safety equipment. One completely outfitted spare boat was stored at the Field Operations Center (EPA Lab) as backup.

5.1.6 Sampling Equipment

A 1/25~m2, stainless steel, Young-modified Van Veen Grab sampler was used to collect sediments. This grab sampled an area of 440 cm2 and a maximum depth of penetration in the sediment of 10 cm.

5.1.7 Manufacturer of Sampling Equipment

5.1.8 Key Variables

5.1.9 Sampling Method Calibration

Field technicians were trained to follow Standard Operating Procedures to insure the collection of representative, uncontaminated and high quality samples. QA/QC measures were taken in the field to avoid or reduce contamination and insure the collection of representative samples. These included: use of stainless steel instruments, thorough cleaning of the sampler between grabs and use of pre-cleaned containers for sediment storage.

5.1.10 Sample Collection Quality Control

A successful grab had relatively level, intact sediment over the entire area of the grab and a sediment depth of 7-10 centimeters. Unacceptable grabs included those: containing no sediments, which were partially filled or had shelly substrates or grossly slumped surfaces. Grabs completely filled to the top, where the sediment was oozing out of the hinged top, were also unacceptable.

5.1.11 Sample Collection Method Reference

Macauley, J. M. 1991. Environmental Monitoring and Assessment Program-Near Coastal Louisianian Province: 1991 Monitoring Demonstration. Field Operations Manual. EPA/600/X-91/XXX. U.S. Environmental Protection Agency, Office of Research and Development, Environmental Research Laboratory, Gulf Breeze, FL 32561.

Macauley, J. M. 1992. Environmental Monitoring and AssessmentProgram: Louisianian Province: 1992 Sampling: Field Operations Manual. EPA/ERL-GB No. SR-119. U.S. Environmental Protection Agency, Office of Research and Development, Environmental Research Laboratory, Gulf Breeze, FL 32561.

Macauley, J. M. 1993. Environmental Monitoring and Assessment Program: Louisianian Province: 1993 Sampling: Field Operations Manual. EPA/ERL-GB No. SR-XXX. U.S. Environmental Protection Agency, Office of Research and Development, Environmental Research Laboratory, Gulf Breeze, FL 32561.

Macauley, J. M. 1994. Environmental Monitoring and Assessment Program: Louisianian Province: 1993 Sampling: Field Operations Manual. EPA/ERL-GB No. SR-XXX. U.S. Environmental Protection Agency, Office of Research and Development, Environmental Research Laboratory, Gulf Breeze, FL 32561.

5.1.12 Sample Collection Method Deviations

None

5.2 Data Preparation and Sample Processing

5. 2. 1 Date Preparation Objective

Process sediment samples for characterization of contaminants.

5.2.2 Data Processing Methods Summary

Upon receipt at the laboratory, the samples were frozen pending analysis, except for those collected for TOC analysis which were refrigerated at 4 C. The frozen sediment samples were thawed and thoroughly homogenized prior to analysis. Separate aliquots of the homogenized sediment were removed. The aliquots were processed for several types of chemical analyses. These included: inorganic analyses (major and trace elements); butyltins (MBT, TBT and DBT)); and organic analyses (PAHs, PCBs, aliphatic hydrocarbons, and pesticides).

5. 2. 2. 1 Inorganic Analytes

Analysis of the sediment for major and trace elements involved a total digestion (i.e., complete dissolution) of the sediment matrix using HNO3/HC1O3 in pressurized Teflon bombs heated at 130E (conventional oven) followed with additions of HF. Instrument analysis for metals Al, Cu, Fe, Mn, and Zn was by flame atomic absorption (FAA); for metals Ag, As, Cd, Cr, Ni,

Pb, Se and Sn, graphite furnace atomic absorption (GFAA) for 1991-1993 and by Inductively Coupled Plasma (ICP) method for 1994; and for mercury (Hg), cold vapor atomic absorption.

Sediment concentrations of butyltin compounds were determined using high resolution gas chromatography and flame photometric detection.

5. 2. 2. 2 Organi c Analytes

The analyses of organic contaminants involved extraction and cleanup followed by instrumental analysis. This included the following basic steps: Soxhlet extraction with methylene chloride, extract drying using sodium sulfate, extract concentration using Kuderna-Danish apparatus, removal of elemental sulfur with activated copper, removal of organic interferents with silica gel/alumina column purification. Following extraction and cleanup, PAH compounds were analyzed using gas chromatography/mass spectrometry (GC/MS). The pesticides and PCB congeners were analyzed using gas chromatography/electron capture detection (GC/ECD).

5. 2. 2. 3 Other Analytes

The concentration of total organic carbon (TOC) in each sediment sample was determined by TOC analyzer using combustion/non- dispersive infrared gas analysis. Acid volatile sulfides (AVS) were measured using a sulfide ion-specific electrode following reaction of the sediment with hydrochloric acid and subsequent trapping of the evolved hydrogen sulfide in solution.

6. DATA MANIPULATIONS

6.1 Name of New or Modified Values

PCB_TOT, PAH_TOT, PAH_LMW, PAH_HMW, DDT_TOT and TOT_ANAL

- 6.2 Data Manipulation Description
 - 6. 2. 1 PCB_TOT, PAH_TOT, PAH_LMW, PAH_HMW, DDT_TOT

Summary values were calculated for groups of organic analytes. The values under a summed analyte are the sum of the concentrations of a specific set of compounds.

6. 2. 2 TOT_ANAL

The number of analytes under TOT_ANAL include only those with a concentration.

6.3 Data Manipulation Examples

The following groups must be summed in order to have consistency across Provinces:

6.3.1 Total Poly-aromatic Hydrocarbons

PAH_TOT = sum of concentrations of biphenyl, fluorene, 1-methyl naphthal ene, 2-methyl naphthal ene, naphthal ene, 2, 6-di methyl naphthal ene, acenaphthene, phenanthrene, anthracene, acenaphthlyene, 2, 3, 5-tri methyl naphthal ene, 1-methyl phenanthrene, fluoranthene, pyrene, benz(a) anthracene, chrysene, benzo(b, k) fluoranthene or individual compounds, ideno(1, 2, 3-c, d) pyrene, benzo(g, h, i) perylene, perylene, benzo(a) pyrene, benzo(e) pyrene and dibenz(a, h) anthracene.

6.3.2 Low Molecular Weight PAHs

PAH_LMW = sum of concentrations of biphenyl, fluorene, 1-methyl naphthal ene, 2-methyl naphthal ene, naphthal ene, 2, 6-di methyl naphthal ene, anthracene, acenaphthl yene, 2, 3, 5-tri methyl naphthal ene.

6.3.3 High Molecular Weight PAHs

PAH_HMW = sum of concentrations of fluoranthene, pyrene, benz(a) anthracene, chrysene, benzo(b, k) fluoranthene or individual compounds, ideno(1, 2, 3-c, d) pyrene, benzo(g, h, i) perylene, perylene, benzo(a) pyrene, benzo(e) pyrene and dibenz(a, h) anthracene, acenaphthene, phenanthrene and 1-methyl phenanthrene.

6.3.4 Total DDT

DDT_TOT = sum of concentrations of OPDDE, OPDDD, OPDDT, PPDDE, PPDDD, PPDDT.

6.3.5 Total Polychlorinated Biphenyls

PCB_TOT = sum of concentrations of the following PCB congeners: 8, 18, 28, 52, 44, 66, 101, 118, 153, 105, 138, 187, 128, 180, 170, 195, 206 and 209.

- 6.4 Data Manipulation Computer Code Set
- 6.5 Data Manipulation Computer Code Language
- 6.6 Data Manipulation Computer Code

7. DESCRIPTION OF PARAMETERS

7.1 Description of Parameters

See Appendix A for a list of analytes.

7.1.1 Parameter Name

Data	Max Field		Vari abl e
Type	Len	Format	Field Label
Char	8	8.	The Station Identifier
Num	8	YMMDD6.	The Date the Sample was Collected
Char	8	8.	Analyte Code
Num	13	13. 6	Conc. of Analyte (dry wt.)
Char	12	12.	Concentration Units
Char	15	15.	Quality Assurance Code for Data
Num	8	3.	Analytes (#) Included in Summed Conc.
Num	13	13. 6	Method Detection Limit for Analyte
Char	15	15.	General Category for Group of Analytes
Char	25		Full Chemical Name of Analyte
	Type Char Num Char Num Char Char Num Num Num Num Char	Type Len Char 8 Num 8 Char 8 Num 13 Char 12 Char 15 Num 8 Num 13 Char 15	Data Field Type Len Format Char 8 8. Num 8 YMMDD6. Char 8 8. Num 13 13.6 Char 12 12. Char 15 15. Num 8 3. Num 13 13.6 Char 15 15.

7.1.6 Precision to which values are reported

The sediment chemistry concentrations presented are in a format of 6 decimal places. This format is necessary because some concentrations are in ug/g and some concentrations are in ng/g. However, the concentrations are only valid FOR THREE SIGNIFICANT FIGURES (not necessarily three decimal places), e.g., 345.67 ug/g is 346 ug/g but 0.00235 ng/g remains as 0.00235 ng/g.

- 7.1.7 Minimum Value in Data Set by Analyte
- 7.1.8 Maximum Value in Data Set by Analyte

7.2 Data Record Example

7. 2. 1 Column Names for Example Records

OBS STA_NAME ANALYTE CONC DETLIMIT QA_CODE TOT_ANAL CHMUNITS _LABEL_ ANAL_CAT VST_DATE

7.2.2 Example Data Records

OBS	STA_NAME	ANALYTE	CONC	DETLI MI T	QA_CODE
1	LA91LR01	ACENTHE	0. 090000	4. 500000	СН- В
2	LA91LR01	ACENTHY	0.040000	3. 700000	СН- В
3	LA91LR01	AG	0. 020000	•	
4	LA91LR01	AL	1700. 000000	3000. 000000	СН- В
5	LA91LR01	ALDRI N		0. 160000	CH- A
OBS	TOT_ANA	AL CHMUNITS	.	LABEL_	ANAL_CAT
1		. ng/g	A	CENAPHTHENE	PAH
2		. ng/g	A	CENAPHTHYLENE	PAH
3		. ug/g	S	I LVER	METAL
4		. ug/g	A	LUMI NUM	METAL
5		ng/g	A	LDRI N	PESTI CI DE

- OBS VST_DATE
 - 1 910721
 - 2 910721
 - 3 910721
 - 4 910721
 - 5 910721
- 7.3 Related Data Sets
 - 7.3.1 Related Data Set Name
 - 7.3.2 Related Data Set Identification Code
- 8. GEOGRAPHIC AND SPATIAL INFORMATION
 - 8.1 Minimum Longitude
 - -97 Degrees 27 Minutes 13.20 Decimal Seconds
 - 8.2 Maxi mum Longi tude
 - -82 Degrees 39 Minutes 28.20 Decimal Seconds
 - 8.3 Maximum Latitude
 - 30 Degrees 48 Minutes 30.00 Decimal Seconds
 - 8.4 Minimum Latitude
 - 26 Degrees 02 Minutes 55.80 Decimal Seconds
 - 8.5 Name of the area or region

Louisianian Province - Coastal distribution of sampling is along the Gulf of Mexico from the Rio Grande, TX to Anclote Key, FL. States represented: Texas, Louisiana, Alabama, Mississippi, Florida

8.6 Direct Spatial Reference Method

Poi nt

8.7 Horizontal Coordinate System Used

Universal Transverse Mercator

8.8 Resolution of Horizontal Coordinates

0.5

8.9 Units for Horizontal Coordinates

Meters

8.10 Vertical Coordinate System

N/A

8.11 Resolution of Vertical Coordinates

N/A

8.12 Units for Vertical Coordinates

N/A

9. QUALITY CONTROL AND QUALITY ASSURANCE

Because of the complexity and importance of sediment contaminant data, EMAP has expended a tremendous effort in the Quality Assurance of these data as is reflected in the detail provided in this section.

9.1 Measurement Quality Objectives

Measurement Quality Objectives (MQOs) for the Louisianian Province sediment chemistry analyses were defined in the Louisianian Province Quality Assurance Project Plan. This plan required each laboratory to analyze the following quality control (QC) samples along with every batch or "set" of field chemistry samples: laboratory reagent blank, calibration check standards, laboratory fortified sample matrix, laboratory duplicate, and Laboratory Control Material (LCM). Results for these QC samples had to fall within certain pre-established control limits.

9.1.1 Sample Processing Method Calibration

For the sediment analyses, a Standard or Certified Reference Material (SRM or CRM) typically was used as the Laboratory Control Material (LCM). SRMs and CRMs have known or "certified" concentrations of the analytes being measured and therefore, are useful for assessing both accuracy and precision. The QA Plan required the laboratory's percent recovery (relative to the certified concentration in the reference material) to fall within the range of 80 to 120 % for each inorganic analyte and 65 to 135 % for each organic analyte. If the laboratory consistently failed to meet these accuracy goals for the CRM or SRM, the values reported for the failed analytes were considered to be suspect and were flagged.

9.1.2 Sample Processing Quality Control

Each laboratory was required to analyze the following quality control (QC) samples along with every batch or "set" of field chemistry samples: laboratory reagent blank, calibration check standards, laboratory fortified sample matrix, laboratory duplicate and Laboratory Control Material (LCM). Results for these QC samples had to fall within certain pre-established control limits for the analysis of a batch of samples to be considered acceptable.

9.2 Quality Assurance/Control Methods

If results for these QC samples did not fall within certain pre-established control limits, the analysis of a batch of samples was not considered acceptable. These and other quality control

issues are coded in four data qualifier codes (QA_CODE) or "flags" used in the Louisianian Province sediment chemistry data file:

CH- A CODE

The "CH-A" code indicates that an analyte was not detected. When the "CH-A" code is used, the concentration field is left blank and the method detection limit for the analyte in that particular sample is reported under DETLIMIT.

CH-B CODE

It is sometimes possible for a laboratory to detect an analyte and report its concentration at a level which is below the calculated method detection limit for the sample. In these situations, the analyst is confident that the analyte was present in the sample, but there is a high degree of uncertainty in the reported concentration. The "CH-B" code is used to flag reported values which are below the calculated method detection limit for the sample. Such values are considered estimates only and should be used with discretion.

CH-C CODE

The CH-C code indicates that the laboratory experienced minor deficiencies meeting the QC requirements, but the overall data quality is judged to be reliable for EMAP assessments.

CH-I CODE

Some analytes are difficult to quantify because they co-elute with other closely related analytes. This phenomenon is called "matrix interference". When this occurs, the suspect analyte(s) are given a "CH-I" code and concentration is left blank.

CH- M CODE

Samples were lost or destroyed.

CH- X CODE

In favor of expediency, a laboratory may elect to cease reporting some of the analytes. EMAP protocol only requires that the laboratory analyze a given list of chemicals; when they go beyond this list and report additional chemicals, we include them in our data. The "CH-X" code indicates that an analyte has been excluded from a given set of data

CH-Z CODE

Some of the analytes listed represent the sum of concentrations of similar analytes (e.g., PCB_TOT is the sum of the concentrations of all PCB congeners). In the event that the concentrations for all of the individual analytes included in the sum are non-detects (have CH-A code) the sum is missing. This is not technically a non-detect, but a sum of non-detects, hence the CH-Z code.

Only "unflagged", CH-B and CH-C coded values are considered valid and useful for most assessment purposes.

Note: In 1994, the method detection limits (field value DETLIMIT) for PAH analytes changed during the analysis due to equipment upgrade. There are two detection limits for all PAH analytes for this year.

9.3 Actual Measurement Quality

Results of QC sample analyses are stored in the Louisianian Province database and are available upon request.

9.3.1 1991-1993 Analyses for Chemical Contaminants in Sediment

Inorganic Contaminants

All analytical data related to inorganic contaminants (trace elements and metals) in 1991-93 Louisianian Province (LP) sediment samples were acceptable for EMAP assessments without further qualification. Results of the inorganic SRM analyses (primary QC check for relative accuracy) are summarized in Tables 9-1(a-c).

Organi c Contami nants

Data quality for the analyses of organic contaminants (organochlorine pesticides, organophosphorus pesticides, PCBs, PAHs, and butyltin compounds) in 1991-93 LP sediment samples routinely met the quality criteria and the analytical results for most analytes were acceptable for EMAP assessments; results of the SRM/LCM analyses (primary QC checks for relative accuracy) are summarized in Tables 9-2(a-c). There were, however, isolated cases in which the laboratory failed to meet the quality criteria for certain organic analyses resulting in those data being rejected or otherwise flagged with an appropriate qualifier code, as follows:

```
1991-
     4, 4' - DDD -
                          CH-C code (minor deficiencies)
                          rejected
     Dieldrin -
     PCB 52 -
                          rej ected
     PCB 170-
                          rejected
1992-
     PCB 52 -
                          CH-C code
                          CH-C code
     PCB 170 -
     Dieldrin -
                          rejected
1993-
     PCB 206 -
                          CH-C code
     PCB 209 -
                          CH-C code
     Al kanes (C13, C14
     C15, C16, C29,
     and C26) -
                          CH-C code
     PCB 110/77 -
                          rejected
     PCB 170 -
                          rej ected
```

Dieldrin - rejected Biphenyl (PAH) - rejected

The data from all other 1991-93 organic analyses of sediment samples (i.e., organochlorine pesticides, organophosphorus pesticides, PCBs, PAHs, and butyltins) were acceptable without further qualification.

9.3.2 1994 Analyses for Chemical Contaminants in Sediments

In 1994, the cooperative agreement for analytical chemistry supporting Louisianian Province Monitoring was awarded to laboratory different from that responsible for the previous analyses of 1991-93 EMAP-LP samples. Laboratory selections for both the initial and subsequent cooperative agreements were based on competitive proposals and were awarded for 3-year terms.

Inorganic Analyses - 1994

The laboratory routinely met or exceeded the required QC criteria related to the analyses of inorganic contaminants in sediment samples and all 1994 results for inorganic contaminants were acceptable for EMAP assessments without further qualification. See Table 9-1d for a summary of the laboratory's 1994 SRM results for inorganic analytes in sediments.

Organic Analyses - 1994

The laboratory generally met the QC requirements for most of the analyses of organic contaminants in sediments; see Table 9-2d for a summary of the laboratory's 1994 SRM results for organic analytes in sediments. However, some analytes, particularly within the organochlorine classes (pesticides and PCBs) proved to be problematic; as a result, the analytical data for those analytes that consistently failed to meet the QC criteria were dropped from the database (see below). All data for OC pesticides was flagged with a "CH-C" qualifier code because the entire set of OC pesticide data was characterized by minor data quality deficiencies, however, these data were still judged to be acceptable for most EMAP assessments. The following summary details the data quality for organic analyses with 1994 sediments:

Organochlorine (OC) pesticides:

Endrin - rejected 2, 4-DDT - rejected

All other OC pesticides were flagged with the "CH-C" code (minor deficiencies, but generally acceptable for EMAP assessments).

PCBs:

PCB 77- rejected PCB 8 - CH-C code PCB 52 - CH-C code

All other PCB data were acceptable for EMAP assessments without further qualification.

PAHs:

All PAH data were acceptable for EMAP assessments without further qualification.

Organophosphorus (OP) Compounds

All OP data were acceptable for EMAP assessments without further qualification.

Butyltin Compounds:

All butyltin data were acceptable without further qualification.

9.3.3 Total Organic Carbon (TOC) Analyses

All QC results for the analysis of TOC in sediment sample from the 1991-94 Louisianian Province Monitoring consistently met the required quality criteria for both accuracy and precision. The Certified Reference Material BCSS-1 (standard marine sediment) supplied by the National Research Council of Canada was analyzed along with each batch of 10-20 field samples; also, along with each sample batch, a duplicated sample and a blank were included. For all four years, the percent recovery of TOC from the CRM remained >90% and the relative percent difference for duplicate analyses remained <10%. All 1991-94 sediment TOC data were acceptable for EMAP assessments without further qualification.

9. 3. 4 Acid Volatile Sulfides (AVS)

The QC data for the analysis of AVS in sediment for the 1991-94 Louisianian Province Monitoring consistently met the required quality criteria for accuracy and precision. The percent recovery of sulfide for calibration check standards, laboratory fortified blanks, and laboratory fortified matrix samples routinely fell within 85-115% of the known concentration; the relative percent difference between duplicate samples routinely met the control limit of 20%. All AVC analyses for 1991-94 were judged acceptable for EMAP assessments.

It should be noted that the 1991 AVS analysis was conducted differently from the 1992-94 analyses. For 1991, only AVS was measured, no simultaneously extracted metals (SEMs) were analyzed; in 1992-94 both AVS and SEMs were analyzed. Also, in 1991, the field collected samples for AVS analysis were taken from a composited sediment sample (consisting of the top 2-3 cm of surficial sediment combined from 5-6 separate sediment grabs). The

composited sediment was well mixed to ensure homogeneity before subsamples for the various sediment analyses were collected. After the fact, it was suggested that AVS sediment samples should be collected in as undisturbed manner as possible to avoid the release of sulfides. In subsequence monitoring (1992-94), at each sampling site, small 2-3 cm cores of sediment were carefully collected from individual grabs and combined into a single sample container with little disruption to the sediment. Comparison of cumulative distribution frequencies for AVS concentration across the Louisianian Province on year-to-year basis showed no significant difference between the 1991 AVS values and the other years; therefore, the inadvertent sediment mixing that occurred in 1991 had little apparent effect on the release of volatile sulfides from the samples.

- 9.4 Sources of Error
- 9.5 Known Problems with the Data
- 9.6 Confidence Level/Accuracy Judgement

Appendix B (Tables)

- 9.7 Allowable Minimum Values
- 9.8 Allowable Maximum Values
- 9.9 QA Reference Data
- 10. DATA ACCESS
 - 10.1 Data Access Procedures

A Data Request Package can be requested from a contact under Section 10.3. Data can be downloaded from the WWW site.

10.2 Data Access Restrictions

Data can only be accessed from the WWW site.

10.3 Data Access Contact Persons

Dr. J. Kevin Summers
Technical Director, EMAP-Estuaries
U. S. Environmental Protection Agency
National Health and Environmental Effects Lab
Gulf Ecology Division
1 Sabine Island Dr.
Gulf Breeze, FL 32561
(904) 934-9244
(904) 934-9201 (FAX)
summers. kevin@epa. gov (E-MAIL)

John M. Macauley
Province Manager, EMAP-E Louisianian Province
U.S. Environmental Protection Agency
National Health and Environmental Effects Lab
Gulf Ecology Division
1 Sabine Island Dr.
Gulf Breeze, FL 32561
(904) 934-9353
(904) 934-9201 (FAX)
macauley.john@epa.gov (E-MAIL)

10.4 Data Set Format

Data can be transmitted in a variety of formats derived from SAS data files when a Data Request Form is submitted.

10.5 Information Concerning Anonymous FTP

Not accessible

- 10.6 Information Concerning Gopher
- 10.7 Information Concerning World Wide Web

Data can be downloaded from the WWW

10.8 EMAP CD-ROM Containing the Data set

Data not available on CD-ROM

11. REFERENCES

11.1 EMAP References

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11.2 Background References

Engle, V.D., J.K. Summers, G.R. Gaston. 1994. A Benthic Index of Environmental Condition of Gulf of Mexico Estuaries. Estuaries: 17:372-384.

Summers, J. Kevin, John F. Paul, Andrew Robertson. 1995. Monitoring the Ecological Condition Of Estuaries In The United States. U.S. Environmental Protection Agency, Office of Research and Development, Environmental Research Laboratory, Gulf Breeze, FL 32651.

12. GLOSSARY AND TABLE OF ACRONYMS

- 12.1 Acronym used in the Detailed Documentation
- 12.2 Definition of Acronym

13. PERSONNEL INFORMATION

Louisianian Province Manager John M. Macauley U. S. EPA NHEERL-GED 1 Sabine Island Dr. Gulf Breeze, FL 32561 (904) 934-9353 (Tel.) (904) 934-9201 (FAX) macauley.john@epa.gov

EMAP-Estuaries Quality Assurance Coordinator P. Thomas Heitmuller U.S.G.S. - BRD Gulf Breeze Project Office 1 Sabine Island Dr. Gulf Breeze, FL 32561 (904) 934-9373 (Tel.) (904) 934-2495 (FAX) heitmuller.tom@epa.gov

EMAP-Estuaries Data Analyst Virginia D. Engle U. S. EPA NHEERL-GED Gulf Breeze Project Office 1 Sabine Island Dr. Gulf Breeze, FL 32561 (904) 934-9354 (Tel.) (904) 934-9201 (FAX) engl e. vi rgi ni a@epa. gov

Appendix A - List of Analytes

----- General Category for Group of Analytes=ALKANES-----

ANALYTE	_LABEL_
ALKANE_T	TOTAL ALKANES
N_DECANE	N-DECANE ALIPHATIC HYDROCARBONS
N_DOCOSN	N-DOCOSANE ALIPHATIC HYDROCARBONS
N_DODECA	N-DODECANE ALIPHATIC HYDROCARBONS
N_DOTRCT	N-DOTRIACONTANE ALIPHATIC HYDROCARBONS
N_EI COSN	N-EICOSANE ALIPHATIC HYDROCARBONS
N_HENESN	N-HENEI COSANE ALI PHATI C HYDROCARBONS
N_HENTCT	N-HENTRI ACONTANE ALI PHATI C HYDROCARBONS
N_HEPADC	N-HEPTADECANE ALIPHATIC HYDROCARBONS
N_HEPTSN	N-HEPTACOSANE ALIPHATIC HYDROCARBONS
N_HEXADC	N-HEXADECANE ALIPHATIC HYDROCARBONS
N_HEXASN	N-HEXACOSANE ALIPHATIC HYDROCARBONS
N_NONADC	N-NONADECANE ALIPHATIC HYDROCARBONS
N_NONASN	N-NONACOSANE ALIPHATIC HYDROCARBONS
N_OCTADC	N-OCTADECANE ALIPHATIC HYDROCARBONS
N_OCTASN	N-OCTACOSANE ALIPHATIC HYDROCARBONS
N_PENTDC	N-PENTEDECANE ALIPHATIC HYDROCARBONS
N_PENTSN	N-PENTACOSANE ALIPHATIC HYDROCARBONS
N_TETRCT	N-TETRATRIACONTANE ALIPHATIC HYDROCARBON
N_TETRDC	N-TETRADECANE ALIPHATIC HYDROCARBONS
N_TETRSN	N-TETRACOSANE ALIPHATIC HYDROCARBONS
N_TRI ACT	N-TRIACONTANE ALIPHATIC HYDROCARBONS
N_TRI CSN	N-TRICOSANE ALIPHATIC HYDROCARBONS
N_TRI DC	N-TRIDECANE ALIPHATIC HYDROCARBONS
N_TRITCT	
N_UNDECA	N_UNDECANE ALIPHATIC HYDROCARBONS
General	Category for Group of Analytes=AVS

ANALYTE _LABEL_

AVS_CON ACID-VOLATILE SULFIDES

```
General Category for Group of Analytes=ISOPRENOID ------
                         ANALYTE
                                    LABEL
                                    TOTAL ISOPRENOIDS
                         I SOPRN_T
                         PHYTANE
                                    PHYTANE
                         PRI STANE
                                    PRISTANE
------ General Category for Group of Analytes=METAL
                             ANALYTE
                                        _LABEL_
                             AG
                                        SILVER
                             AL
                                        ALUMI NUM
                             AS
                                        ARSENI C
                             CD
                                        CADMI UM
                             CR
                                        CHROMI UM
                             CU
                                        COPPER
                             FΕ
                                        I RON
                             HG
                                        MERCURY
                             MN
                                        MANGANESE
                             NI
                                        NI CKEL
                             PB
                                        LEAD
                             SB
                                        ANTI MONY
                                        SELENI UM
                             SE
                             SN
                                        TIN
                             ZN
                                        ZINC
               General Category for Group of Analytes=ORGANOPHOS-----
                           ANALYTE
                                        _LABEL_
                           CARBOFEN
                                        CARBOFENOTHI ON
                           CHLORPYR
                                        CHLORPYRIFOS
                           DI AZI NON
                                        DI AZI NON
                           DI SULFOT
                                        DI SULFOTON
                           ETHI ON
                                        ETHI ON
                           TERBUFOS
                                        TERBUFOS
                   General Category for Group of Analytes=PAH------
                    ANALYTE
                                   _LABEL_
                    ACENTHE
                                   ACENAPHTHENE
                    ACENTHY
                                   ACENAPHTHYLENE
                    ANTHRA
                                   ANTHRACENE
                    BENANTH
                                   BENZO(A) ANTHRACENE
                    BENAPY
                                   BENZO(A) PYRENE
                    BENEPY
                                   BENZO(E) PYRENE
                    BENZOBFL
                                   BENZO(B) FLUORANTHENE
                    BENZOKFL
                                   BENZO(K) FLUORANTHENE
                    BENZOP
                                   BENZO(G, H, I) PERYLENE
                    BI PHENYL
                                   BI PHENYL
                    C1CHRYS
                                   C1-CHRYSENES
```

```
General Category for Group of Analytes=PAH (cont.)-----
 C1DI BENZ
                 C1-DI BENZOTHI OPHENES
 C1FLRAN
                 C1-FLUORANTHENE PYRENE
 C1FLUOR
                 C1-FLUORENES
 C1NAPH
                 C1-NAPHTHALENES
 C1PHENAN
                 C1-PHENANTHRENES
 C2CHRYS
                 C2-CHRYSENES
 C2DI BENZ
                 C2-DI BENZOTHI OPHENES
 C2FLUOR
                 C2-FLUORENES
 C2NAPH
                 C2-NAPHTHALENES
 C2PHENAN
                 C2-PHENANTHRENES
 C3CHRYS
                 C3-CHRYSENES
 C3DI BENZ
                 C3-DI BENZOTHI OPHENES
 C3FLU0R
                 C3-FLUORENES
 C3NAPH
                 C3-NAPHTHALENES
 C3PHENAN
                 C3-PHENANTHRENES
 C4CHRYS
                 C4- CHRYSENES
 C4NAPH
                 C4-NAPHTHALENES
 C4PHENAN
                 C4-PHENANTHRENES
 CHRYSENE
                 CHRYSENE
 DI BENZ
                 DI BENZO (A, H) ANTHRACENE
                 DI BENZOTHI OPHENE
 DI BENZO
 DI METH
                 2, 6- DI METHYLNAPHTHALENE
 FLUORANT
                 FLUORANTHENE
 FLUORENE
                 FLUORENE
                 (I) 1, 2, 3-C, D-PYRENE
 I NDENO
 MENAP1
                 1 - METHYLNAPHTHALENE
                 2-METHYLNAPHTHALENE
 MENAP2
 MEPHEN 1
                 1 - METHYLPHENANTHRENE
                 NAPHTHALENE
 NAPH
 PAHTOT_L
                 TOTAL PAHS - LA PROVINCE
                 HIGH MOLECULAR WT - TOT. PAH
 PAH_HMW
 PAH_LMW
                 LOW MOLECULAR WT - TOT. PAH
 PAH_TOT
                 TOTAL PAHS
 PERYLENE
                 PERYLENE
 PHENANTH
                 PHENANTHRENE
 PYRENE
                 PYRENE
 TRI METH
                 2, 3, 5-TRI METHYLNAPHTHALENE
General Category for Group of Analytes=PCB -----
   ANALYTE
                 LABEL
   PCB101
                 PCB 101
                 PCB 105
   PCB105
   PCB110
                 PCB 110/77
   PCB118
                 PCB 118/108/149
   PCB126
                 PCB 126
   PCB128
                 PCB 128
   PCB138
                 PCB 138
   PCB153
                 PCB 153
   PCB170
                 PCB 170
                 PCB 18
   PCB18
```

PCB 180

PCB 195

PCB 187/182/159

PCB180

PCB187

PCB195

```
----- General Category for Group of Analytes=PCB (cont.)-----
                                   PCB 200
                     PCB200
                     PCB206
                                   PCB 206
                                  PCB 209
                     PCB209
                     PCB28
                                   PCB 28
                     PCB29
                                   PCB 29
                     PCB44
                                   PCB 44
                     PCB52
                                  PCB 52
                     PCB66
                                   PCB 66
                     PCB8
                                   PCB 8
                     PCB87
                                   PCB 87
                     PCBTOT_L
                                   TOTAL PCBS - LA PROVINCE
                     PCB_TOT
                                  TOTAL PCBS
              General Category for Group of Analytes=PESTICIDE -----
                       ANALYTE
                                             _LABEL_
                                             ALDRIN
                       ALDRI N
                       ALPHABHC
                                             ALPHA-BHC
                                             ALPHA- CHLORDANE
                       ALPHACHL
                       BETABHC
                                             BETA-BHC
                                             TOTAL BHC
                       BHC_TOT
                       CHL_TOT
                                             TOTAL CHLORDANE
                       CI SNONA
                                             CIS-NONACHLOR
                                             OP-DDD + PP-DDD
                       DDD
                       DDE
                                             OP-DDE + PP-DDE
                       DDT
                                             OP-DDT + PP-DDT
                       DDT\_TOT
                                             TOTAL DDT
                       DELTABHC
                                             DELTA-BHC
                       DI COFOL
                                             DI COFOL
                       DI ELDRI N
                                             DI ELDRI N
                       ENDOSUL1
                                             ALPHA-ENDOSULFAN
                       ENDOSUL2
                                             BETA - ENDOSULFAN
                       ENDRIN
                                             ENDRIN
                       GAMMACHL
                                             GAMMA- CHLORDANE
                       HEPTACHL
                                             HEPTACHLOR
                       HEPTAEPO
                                             HEPTACHLOR- EPOXI DE
                       HEXACHL
                                             HEXACHLOROBENZENE
                       LI NDANE
                                             LINDANE (GAMMA-BHC)
                       MI REX
                                             MI REX
                       OPDDD
                                             O, P' DDD
                       OPDDE
                                             O, P' DDE
                       OPDDT
                                             O, P' DDT
                       OXYCHL
                                             OXYCHLORDANE
                       OXYFL
                                             OXYFLUORFEN
                       PPDDD
                                             P, P' DDD
                       PPDDE
                                             P, P' DDE
                       PPDDT
                                             P, P' DDT
                       TNONCHL
                                             TRANS-NONACHLOR
```

TOXAPHENE

TOXAPHEN

General	Category for	Group of Analytes=TBT
	ANALYTE	_LABEL_
	DBT MBT TBT	DI - BUTYL TIN MONO BUTYL TIN TRI - BUTYL TIN
General	Category for	Group of Analytes=TOC
ANA	ALYTE _LAB	EL_
TO	C	TOTAL ORGANIC CARBON

Appendix B

ANALYTE

TABLE 9-1a. Relative accuracy for the 1991 EMAP-LP analyses of inorganic contaminants in sediments based on the laboratory's analytical results of a certified reference materials (CRMs), the National Research Council of Canada's (NRC) BCSS-1, MESS-1, and BEST-1 (for mercury, only); the CRMs were analyzed on a per batch basis (batch = <=20 samples). Relative accuracy was computed by comparing the laboratory's result for an analyte against the NRC's certified value to calculate percent recovery. The accuracy goal for EMAP-LP inorganic analyses was that the Laboratory's averaged value to be within +-20% agreement to the CRM values. (note: this pertained only to those analytes with NRC-reported concentrations >= 10X the laboratory's reported MDL).

RANGE

(% recov)

RECOVERY

(%)

	(10)	(,, 1000.)
BCSS-1 (n=7)		
Arseni c	88	81 - 99
Cadmi um	94	72 - 108
Chromi um	90	85 - 97
Copper	85	75 - 95
Lead	100	86 - 119
Manganese	96	89 - 106
Ni ckel	93	89 - 95
Sel eni um	93	46 - 140
Ti n	106	86 - 135
Zi nc	90	84 - 103
MESS-1 (n=7)		
Arseni c	90	78 - 106
Cadmi um	100	95 - 107
Chromi um	84	76 - 90
Copper	89	76 - 98
Lead	89	84 - 96
Manganese	96	92 - 103
Ni ckel	85	77 - 94
Sel eni um	76	33 - 100
Ti n	73	55 - 88
Zi nc	95	89 - 106
BEST-1 (n=7)		
Mercury	92	78 - 100

TABLE 9-1b. Relative accuracy for the 1992 EMAP-LP analyses of inorganic contaminants in sediments based on the laboratory's analytical results of a certified reference materials (CRMs), the National Research Council of Canada's (NRC) BCSS-1 and BEST-1 (for mercury, only); the CRMs were analyzed on a per batch basis (batch = <=20 samples). Relative accuracy was computed by comparing the laboratory's result for an analyte against the NRC's certified value to calculate percent recovery. The accuracy goal for EMAP-LP inorganic analyses was that the Laboratory's averaged value (n=18) to be within +-20% agreement to the CRM values. (note: this pertained only to those analytes with NRC-reported concentrations >= 10X the laboratory's reported MDL).

ANALYTE	RECOVERY (%)	RANGE (% recov)
Arseni c	104	90 - 153
Cadmi um	116	104 - 140
Chromi um	87	78 - 96
Copper	90	85 - 96
Lead	97	86 - 109
Manganese	104	99 - 108
Mercury	101	90 - 111
Ni ckel	100	89 - 115
Sel eni um	102	0 - 130
Ti n	110	85 - 147
Zi nc	97	92 - 103

TABLE 9-1c. Relative accuracy for the 1993 EMAP-LP analyses of inorganic contaminants in sediments based on the laboratory's analytical results of a certified reference material (CRM), the National Research Council of Canada's (NRC) MESS-1; the CRMs were analyzed on a per batch basis (batch = <=20 samples). Relative accuracy was computed by comparing the laboratory's result for an analyte against the NRC's certified value to calculate percent recovery. The accuracy goal for EMAP-LP inorganic analyses was that the Laboratory's averaged value (n=15) to be within +-20% agreement to the CRM values. (note: this pertained only to those analytes with NRC-reported concentrations >= 10X the laboratory's reported MDL).

ANALYTE	RECOVERY (%)	RANGE (% recov)
Arseni c	97	83 - 117
Cadmi um	98	88 - 108
Chromi um	93	85 - 99
Copper	92	85 - 109
Lead	89	83 - 95
Manganese	99	93 - 102
Mercury	99	89 - 108
Ni ckel	96	86 - 107
Sel eni um	100	89 - 107
Silver	101	83 - 117
Ti n	86	81 - 97
Zinc	95	86 - 102

TABLE 9-1d. Relative accuracy for the 1994 EMAP-LP analyses of inorganic contaminants in sediments based on the laboratory's analytical results of a certified reference material (CRM), the National Research Council of Canada's (NRC) BCSS-1; the CRM was analyzed on a per batch basis (batch = <=20 samples). Relative accuracy was computed by comparing the laboratory's result for an analyte against the NRC's certified value to calculate percent recovery. The accuracy goal for EMAP-LP inorganic analyses was that the Laboratory's averaged value (n=11) to be within +-20% agreement to the CRM values. (note: this pertained only to those analytes with NRC-reported concentrations >= 10X the laboratory's reported MDL).

ANALYTE	RECOVERY	RANGE
	(%)	(% recov)
Arseni c	95	78 - 119
Cadmi um	111	96 - 136
Chromi um	92	70 - 103
Copper	103	97 - 112
Lead	96	83 - 121
Manganese	98	93 - 109
Mercury	100	100
Ni ckel	101	89 - 114
Sel eni um	97	77 - 119
Silver	98	58 - 136
Ti n	101	77 - 134
Zinc	94	84 - 106

TABLE 9-2a. Summary of 1991 laboratory results for the analyses of the organics-in-sediment laboratory control material (LCM). The LCM used was the National Institute of Standards and Technology (NIST) issued Standard Reference Material 1941 (SRM 1941) - Organics in Marine Sediment; the LCM was analyzed with each batch run of samples. Results are reported as the laboratory's average percent recovery (n=19) based on the NIST-reported "true" concentrations for the analytes of interest. The EMAP-E performance-based accuracy goal for organic analyses was that the laboratory percent recovery consistently fall within +- 30% the NIST concentration (note: this pertained only to those analytes with NIST-reported concentrations >= 10X the laboratory's reported MDL).

ANALYTE	RECOVERY	RANGE	
	(%)	(% recov)	
PAHs:			
Acenaphthene	101	(93 - 109)	
Acenaphthyl ene	100	(90 - 115)	
Anthracene	99	(92 - 100)	
Benzo (a) anthracene	100	(94 - 105)	
Benzo (a) pyrene	99	(92 - 100)	
Benzo (e) pyrene	101	(83 - 124)	
Benzo (b) fluoranthene	100	(98 - 100)	
Benzo (k) fluoranthene	107	(81 - 143)	
Benzo (g, h, , i) peryl ene	99	(85 - 114)	
Bi phenyl	101	(100 - 107)	
Chrysene	108	(78 - 128)	
2, 6- Di methyl napthal ene	101	(94 - 109)	

TABLE 9-2a. (cont.)		
ANALYTE	RECOVERY	RANGE
	(%)	(% recov)
Fluoranthene	100	(98 - 108)
Fluorene	102	(89 - 110)
(i) 1, 2, 3-c, d-pyrene	99	(77 - 116)
1- methyl naphthal ene	100	(89 - 114)
2- methyl naphthal ene	100	(85 - 109)
1-methyl phenanthrene	98	(77 - 110)
Naphthal ene	101	(86 - 115)
Peryl ene	102	(96 - 112)
Phenanthrene	99	(79 - 106)
Pyrene	99	(92 - 103)
PCBs:		
PCB 105	110	(77 - 170)
PCB 153	126	(69 - 182)
PCB 180	117	(58 - 177)
PCB 206	79	(26 - 147)
PCB 28	81	(62 - 98)
Organochl ori ne pesti ci des:		
al pha- Chl ordane	74	(37 - 99)
Di el dri n	298	(0 - 476)
Heptachl or-epoxi de	308	(0 - 688)
p, p-DDD	61	(33 - 115)
p, p- DDE	101	(63 - 202)
p, p- DDT	64	(0 - 155)
trans-Nonachl or	57	(0 - 89)

TABLE 9-2b. Summary of 1992 laboratory results for the analyses of the organics-in-sediment laboratory control material (LCM). The LCM used was the National Institute of Standards and Technology (NIST) issued Standard Reference Material 1941 (SRM 1941) - Organics in Marine Sediment; the LCM was analyzed with each batch run of samples. Results are reported as the laboratory's average percent recovery (n=14) based on the NIST-reported "true" concentrations for the analytes of interest. The EMAP-E performance-based accuracy goal for organic analyses was that the laboratory percent recovery consistently fall within +- 30% the NIST concentration (note: this pertained only to those analytes with NIST-reported concentrations >= 10X the laboratory's reported MDL).

ANALYTE	RECOVERY	RANGE	
PAHs:	(%)	(% recov)	
Acenaphthene	69	(57 - 90)	
Acenaphthyl ene	77	(66 - 93)	
Anthracene	100	(99 - 100)	
Benzo (a) anthracene	101	(98 - 106)	
Benzo (a) pyrene	98	(89 - 100)	
Benzo (e) pyrene	106	(90 - 122)	

TABLE 9-2b. (cont.) ANALYTE	RECOVERY (%)	RANGE (% recov)
- 42		
Benzo (b) fluoranthene	100	(96 - 100)
Benzo (k) fluoranthene	157	(131 - 180)
Benzo (g, h, , i) peryl ene	98	(73 - 112)
Bi phenyl	86	(76 - 100)
Chrysene	140	(111 - 162)
2, 6- Di methyl napthal ene Fl uoranthene	88 99	(77 - 100) (92 - 100)
Fl uoranthene Fl uorene	99 74	(92 - 100) (61 - 87)
(i) 1, 2, 3- c, d- pyrene	99	(69 - 116)
1- methyl naphthal ene	81	(74 - 90)
2- methyl naphthal ene	84	(76 - 90)
1- methyl phenanthrene	89	(69 - 100)
Naphthal ene	71	(64 - 78)
Peryl ene	79	(69 - 100)
Phenanthrene	93	(79 - 100)
Pyrene	99	(90 - 100)
PCBs:		
PCB 101	121	(103 - 146)
PCB 105	118	(56 - 224)
PCB 118	115	(80 - 159)
PCB 138	96	(68 - 119)
PCB 153	125	(101- 152)
PCB 18	60	(44 - 82)
PCB 180	118	(73 - 152)
PCB 187	111	(73 - 152) (84 - 126)
PCB 195	190	(0 - 310)
PCB 206	92	(41 - 156)
PCB 209	101	(54 - 172)
PCB 28	84	(59 - 101)
PCB 52	226	(157 - 298)
PCB 66	67	(43 - 77)
Organochl ori ne pesti ci des:		
•	440	(00 171)
al pha- Chl ordane	112	(62 - 174)
Di el dri n	451	(286 - 707)
Heptachl or- epoxi de	250	(0 - 494)
p, p- DDD	65	(40 - 100)
p, p- DDE	100	(86 - 124)
p, p- DDT trans- Nonachl or	18	(0 - 100)
trans-nonachi or	87	(55 - 125)

TABLE 9-2c. Summary of 1993 laboratory results for the analyses of the organics-in-sediment laboratory control material (LCM). The LCM used was the National Institute of Standards and Technology (NIST) issued Standard Reference Material 1941a (SRM 1941a) - Organics in Marine Sediment; the LCM was analyzed with each batch run of samples. Results are reported as the laboratory's average percent recovery (n=10) based on the NIST-reported "true" concentrations for the analytes of interest. The EMAP-E performance-based accuracy goal for organic analyses was that the laboratory percent recovery consistently fall within +- 30% the NIST concentration (note: this pertained only to those analytes with NIST-reported concentrations >= 10X the laboratory's reported MDL).

ANALYTE	RECOVERY	RANGE
PAHs:	(%)	(% recov)
Acenaphthene Acenaphthyl ene Anthracene Benzo (a) anthracene Benzo (a) pyrene Benzo (e) pyrene Benzo (b) fluoranthene Benzo (k) fluoranthene Benzo (g, h, , i) peryl ene Bi phenyl Chrysene Di benzo (a, h) anthracene Di benzothi ophene Fluoranthene Fluorene (i) 1, 2, 3-c, d-pyrene 1-methyl phenanthrene Naphthal ene Peryl ene Phenanthrene Pri stane Pyrene	99 115 76 87 82 82 90 104 82 43 111 107 72 78 71 88 85 80 60 75 142 74	(96 - 100) (100 - 134) (55 - 100) (73 - 100) (71 - 92) (71 - 98) (68 - 101) (62 - 159) (57 - 100) (36 - 57) (87 - 142) (100 - 117) (58 - 89) (64 - 100) (57 - 95) (69 - 100) (71 - 112) (66 - 100) (46 - 88) (61 - 100) (100 - 205) (61 - 83)
PCBs: PCB 8 PCB 18 PCB 28 PCB 44 PCB 52 PCB 66 PCB 101 PCB 105 PCB 110 PCB 118 PCB 128 PCB 128 PCB 138 PCB 153 PCB 170 PCB 180 PCB 187	133 223 87 92 119 95 88 71 168 71 73 71 64 595 106 91	(96 - 241) (145 - 302) (75 - 100) (73 - 100) (95 - 162) (69 - 100) (67 - 108) (41 - 100) (100 - 247) (52 - 92) (0 - 193) (57 - 100) (44 - 87) (268 - 1560) (97 - 134) (65 - 100)

TABLE 9-2c. (continued)

p, p-DDE

p, p-DDT

trans-Nonachlor

ANALYTE	RECOVERY (%)	RANGE (% recov)
PCB 206	41	(14 - 88)
PCB 209	53	(36 - 87)
Organochl ori ne pesti ci des:		
al pha- Chl ordane	77	(42 - 100)
Di el dri n	140	(100 - 217)
Hexachl orobenzene	97	(86 - 100)
o, p- DDE	70	(0 - 149)
p, p- DDD	90	(69 - 106)
• • <u> </u>		1

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89

TABLE 9-2d. Relative accuracy for the 1994 EMAP-LP analyses of organics in sediments based on the laboratory's analytical results of a standard reference material (SRM), the National Institute of Technology and Standards (NIST) SRM 1941a; the SRM was analyzed on a per sample batch basis. Relative accuracy was computed by comparing the laboratory's value for an analyte against either end of the range of values (i.e., 95% confidence limits) reported by NIST for the analyte. The accuracy goal for EMAP-LP organic analyses was that the laboratory's averaged value (n=10) to be within +- 35% agreement to the SRM values.

(42 - 97)

(0 - 55)

(61 - 100)

ANALYTE	RECOVERY	RANGE
PAHs:	(%)	(% recov)
Acenaphthene Acenaphthylene Anthracene Benzo (a) anthracene Benzo (a) pyrene	110 118 107 108 94	(102 - 118) (102 - 135) (98 - 115) (101 - 117) (81 - 100)
Benzo (e) pyrene Benzo (b) fluoranthene Benzo (k) fluoranthene Benzo (g, h, , i) perylene	102 95 123 100	(87 - 109) (70 - 111) (97 - 163) (88 - 113)
Bi phenyl Chrysene Di benzo (a, h) anthracene Fl uoranthene	85 104 109 102	(69 - 102) (98 - 109) (85 - 134) (95 - 105)
Fluorene (i) 1,2,3-c,d-pyrene 1-methylphenanthrene Naphthalene	96 115 92 101	(85 - 116) (93 - 140) (78 - 113) (85 - 109)
Peryl ene Phenanthrene Pyrene	87 105 101	(68 - 107) (99 - 115) (93 - 107)

TABLE 9-2d. (continued)

ANALYTE	RECOVERY (%)	RANGE (% recov)
PCBs:		
PCB 8	158	(106 - 284)
PCB 18	248	(209 - 317)
PCB 28	99	(97 - 100)
PCB 44	113	(100 - 144)
PCB 52	111	(100 - 140)
PCB 66	148	(123 - 203)
PCB 101	102	(94 - 119)
PCB 105	167	(137 - 205)
PCB 118	99	(89 - 118)
PCB 138	110	(98 - 137)
PCB 153	74	(58 - 100)
PCB 170	134	(112 - 169)
PCB 180	131	(100 - 158)
PCB 187	99	(94 - 100)
PCB 206	141	(94 - 355)
PCB 209	84	(65 - 100)
Organochl ori ne pesti ci des:		

al pha- Chl ordane	66	(0 - 100)
Di el dri n	104	(63 - 133)
Hexachl orobenzene	74	(46 - 100)
o, p- DDE	163	(0 - 232)
p, p- DDD	91	(79 - 102)
p, p- DDE	89	(69 - 112)
p, p- DDT	96	(73 - 131)
trans-Nonachlor	115	(100 - 151)
0xychl ordane	15	(0 - 72)