

NATIONAL DIOXIN STUDY TIER 4 — COMBUSTION SOURCES

Final Test Report — Site 4 Black Liquor Boiler BLB — A

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FOREWORD

This report is the result of a cooperative effort between the Office of Research and Development's Hazardous Waste Engineering Research Laboratory (HWERL) and the Office of Air Quality Planning and Standard's Monitoring and Data Analysis Division (MDAD). The overall management of Tier 4 of the National Dioxin Study was the responsibility of MDAD. In addition, MDAD provided technical guidance for the source test covered by this report. HWERL was directly responsible for the management and technical direction of the source test.

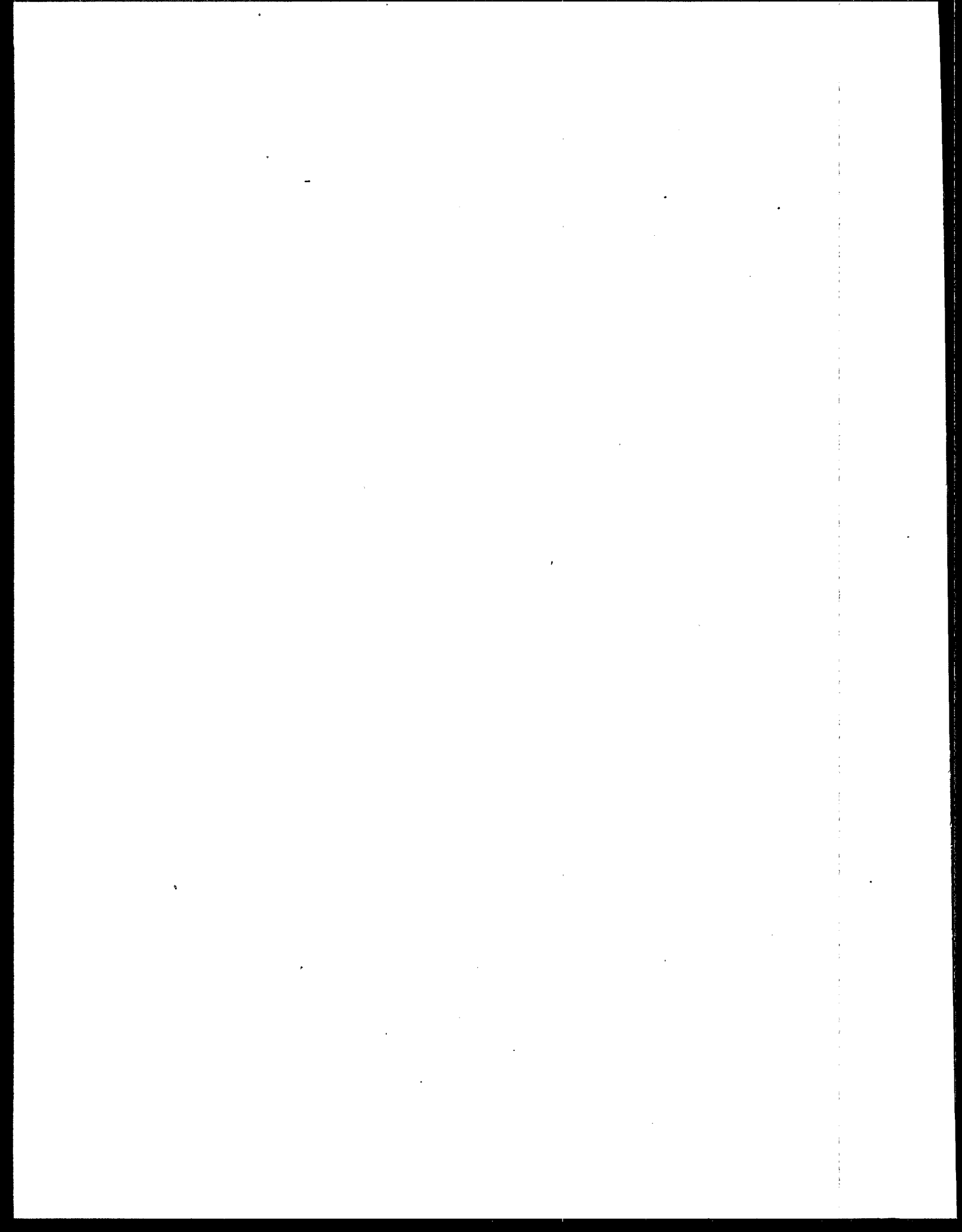


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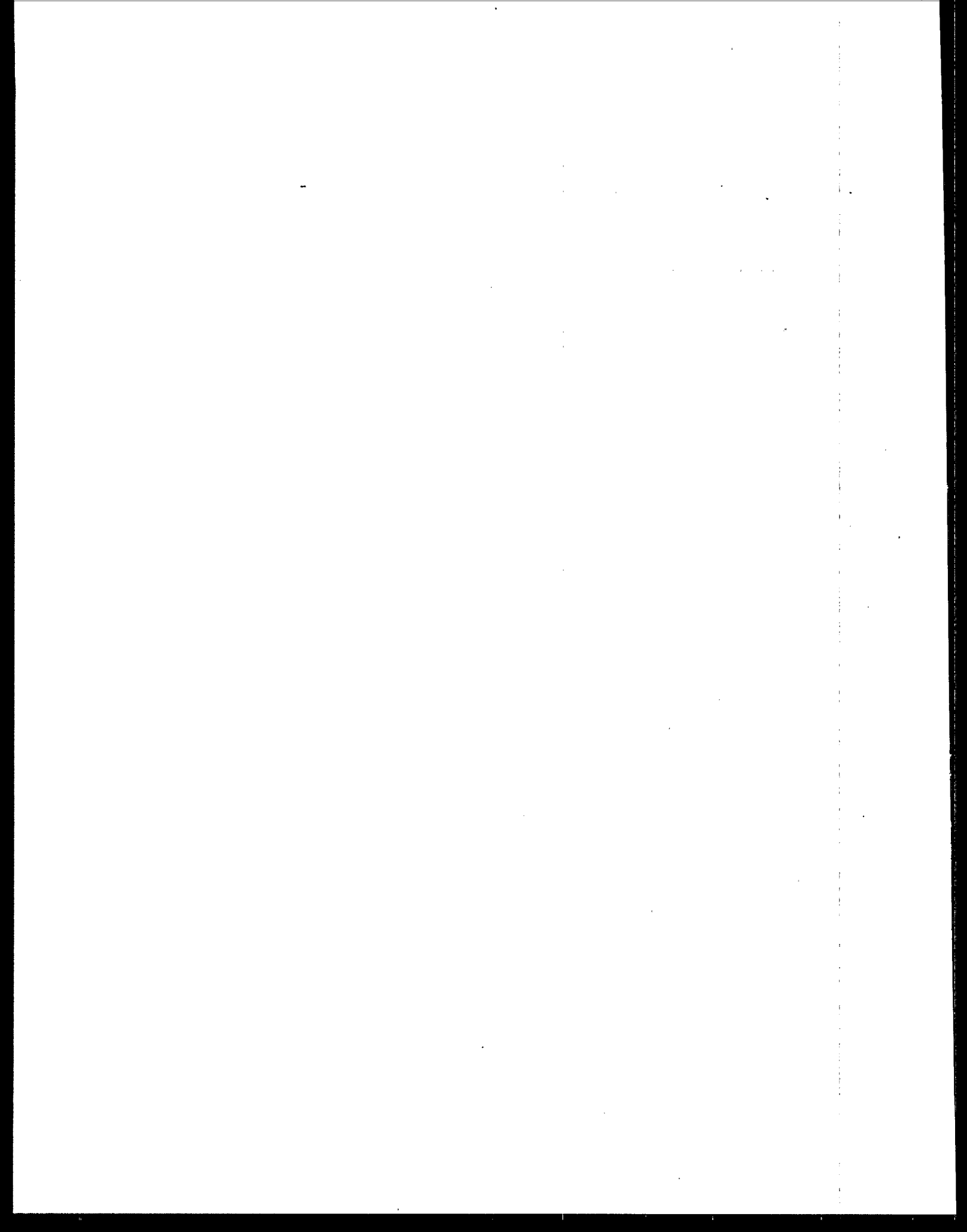
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1.0 INTRODUCTION

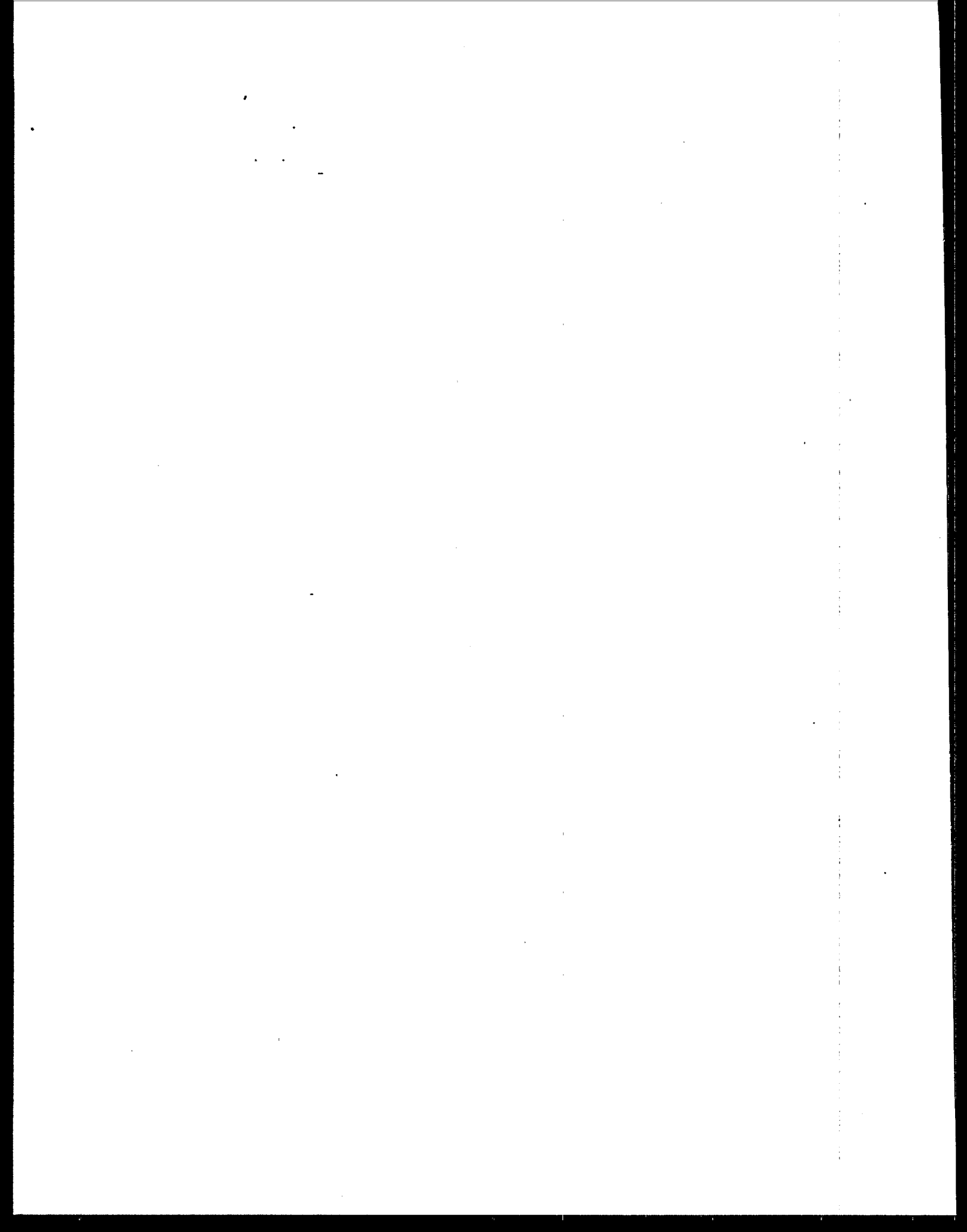
This report summarizes the results of a dioxin/furan^a emissions test of a black liquor recovery boiler equipped with a wet-bottom electrostatic precipitator for particulate matter emissions control. Black liquor recovery boilers are used at Kraft pulp mills to produce process steam and to reclaim inorganic chemicals from spent wood pulping liquors. The test is the fourth in a series of several dioxin/furan emissions tests being conducted under Tier 4 of the National Dioxin Study. The primary objective of Tier 4 is to determine if various combustion sources are sources of dioxin and/or furan emissions. If any of the combustion sources are found to emit dioxin or furan, the secondary objective of Tier 4 is to quantify these emissions.

Black liquor recovery boilers are one of 16 combustion source categories being considered in the Tier 4 program. The tested black liquor boiler, hereafter referred to as boiler BLB-A, was selected for this test after an initial information screening and a one-day pretest survey visit.

Boiler BLB-A is considered representative of new black liquor recovery boilers built in the last 5 to 10 years. The amount of chlorides present in the black liquor circuit at this site is typical of that found at other Kraft pulp mills.

This test report is organized as follows. A summary of test results and conclusions is provided in Section 2.0, followed by a detailed process description in Section 3.0. The source sampling and analysis plan is outlined in Section 4.0, and the dioxin test data are presented in Section 5.0. Sections 6.0 through 9.0 present various testing details. These include descriptions of the sampling locations and procedures (Section 6.0), descriptions of the analytical procedures (Section 7.0), and a summary of the quality assurance/quality control results (Section 8.0). The appendices contain data generated during the field sampling and analytical activities.

^aThe term "dioxin/furan" and the acronyms PCDD and PCDF as used in this report refer to the polychlorinated dibenzo-p-dioxin and dibenzofuran homologues with four or more chlorine atoms.



2.0 SUMMARY

2.1 SOURCE SAMPLING AND ANALYSIS OVERVIEW

The host plant (Site 04) is a Kraft pulp mill that produces pulp and paper products. Black liquor recovery boiler BLB-A combusts concentrated spent liquor from the pulping process and recovers the inorganic chemicals used to produce pulp from wood chips. Particulate matter emissions from black liquor boiler BLB-A are controlled by a wet bottom electrostatic precipitator. A simplified process flow diagram of the system is shown in Figure 2-1.

The gaseous, liquid, and solid sampling performed during the test program is summarized in Table 2-1. Sampling for dioxin and furan was performed simultaneously at the electrostatic precipitator outlet exhaust stack and the electrostatic precipitator inlet location (i.e., black liquor boiler outlet) in each of a series of three test runs conducted on December 11 through 14, 1984. The dioxin/furan sampling was based on the October 1984 draft of the Modified Method 5 (MM5) procedure developed by the American Society of Mechanical Engineers (ASME) for measuring emissions of chlorinated organic compounds from municipal waste incinerators. Modifications to the ASME protocol are discussed in Section 6. MM5 train components and train rinses were analyzed for dioxins and furans by EMSL-TRP and ECL-BSL, two of three EPA laboratories collectively known as Troika. The dioxin/furan analysis quantified the 2378-TCDD* isomer and the tetra- through octa- dioxin/furan homologues present in the samples.

Dioxin/furan and dioxin/furan precursor analyses were performed on samples of the concentrated black liquor fed to the boiler. The black liquor dioxin/furan analyses were performed by EPA ECL-BSL and EMSL-RTP laboratories, and the dioxin precursor analyses were performed by Radian. The specific dioxin precursors analyzed for were chlorophenols, chlorobenzenes, polychlorinated biphenyls, and total chlorine. Samples of black liquor circuit intermediates (caustic, white liquor, and weak black liquor) and ClO_2

*The terms TCDD and TCDF as used in this report refer to tetrachloro dibenzo-p-dioxin and tetrachlorodibenzo furan.

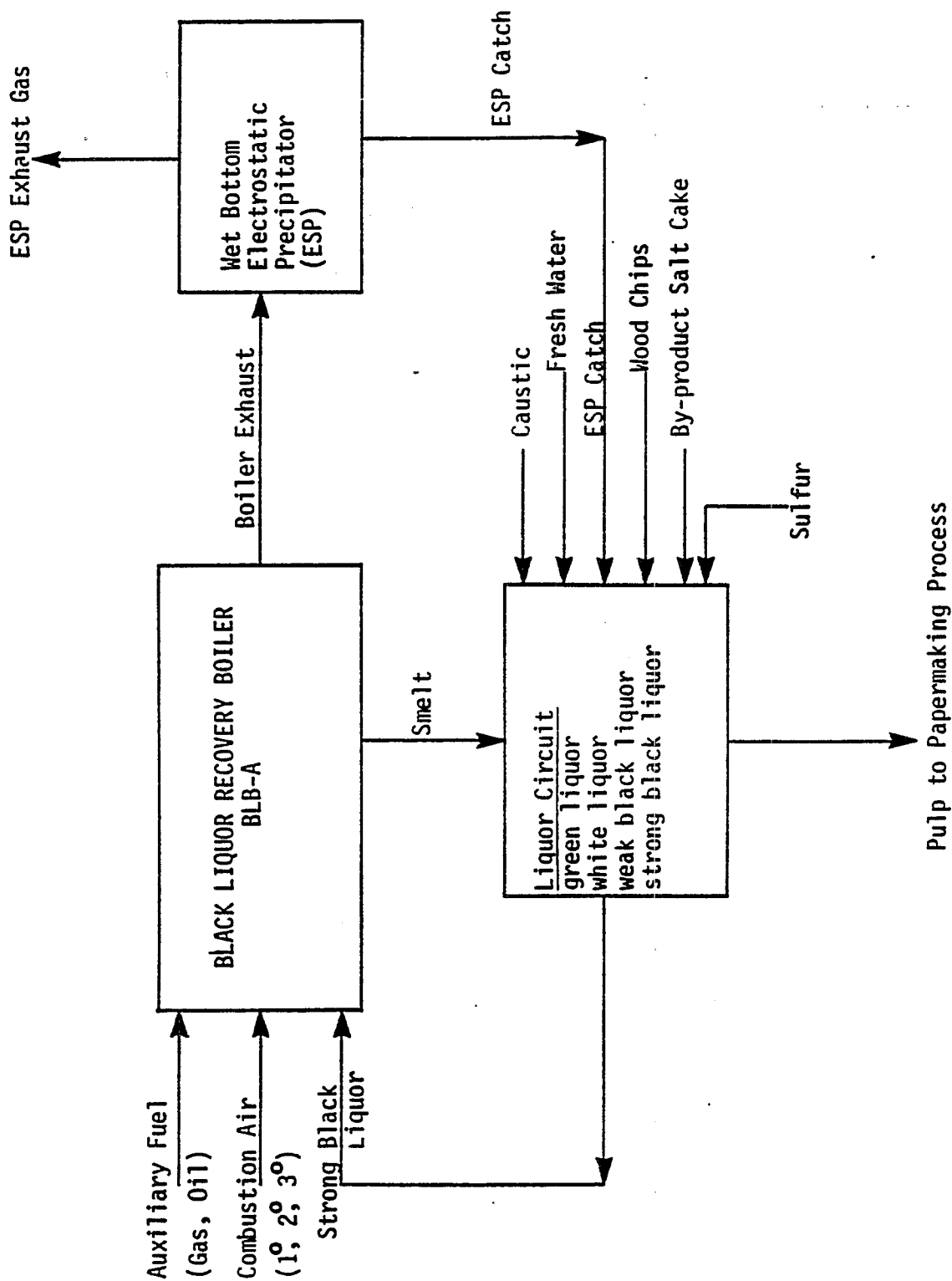


Figure 2-1. Simplified Process Flow Diagram of Black Liquor Recovery Boiler System BLB-A

TABLE 2-1. SOURCE SAMPLING AND ANALYSIS OVERVIEW

Item	Item Description
1. Number of test runs	- Three identical test runs (Runs 1, 2, 3).
2. Gaseous Sampling	<ul style="list-style-type: none"> - MM5 sampling at black liquor boiler outlet and ESP outlet exhaust stack (Runs 1, 2, 3). Dioxin/furan analysis. - EPA Reference Methods 2 and 4 at black liquor boiler outlet and ESP outlet exhaust stack (Runs 1, 2, 3). Gas velocity and moisture. - Integrated bag sampling at black liquor boiler outlet and ESP outlet exhaust stack (Runs 1, 2, 3). CO₂, O₂, N₂ analysis for molecular weight determination. - HCl sampling at ESP Outlet exhaust stack (Runs 1, 2, 3). HCl analysis. - Continuous monitoring of CO, CO₂, O₂, SO₂, and NO_x, total hydrocarbons at black liquor boiler outlet (Runs 1, 2, 3).
3. Liquid Sampling	<ul style="list-style-type: none"> - Strong black liquor sampling^a (Runs 1, 2, 3). Dioxin/furan analysis, dioxin/furan precursor analysis, and total chlorine analysis. - Caustic sampling (Runs 1, 2, 3). Total chlorine analysis. - White liquor sampling (Runs 1, 2, 3). Total chlorine analysis. - Weak black liquor sampling (Runs 1, 2, 3). Total chlorine analysis. - ClO₂ generation system by-product salt cake sampling (Runs 1, 2, 3). Total chlorine analysis.
4. Solids Sampling	- Soil sampling ^a (one composite sample from 10 locations). Potential dioxin/furan analysis.

^aSample(s) to be analyzed pending evaluation of the dioxin/furan emissions data from the MM5 sampling train.

generation system by-product salt cake that is fed to the black liquor circuit were taken and analyzed by Radian for total chlorine. The total chlorine analyses of these streams were used to quantify the major chlorine inputs to the black liquor circuit. A single composite soil sample was also taken, but analysis of this sample was deferred pending evaluation of the dioxin/furan emissions data.

Continuous emissions monitoring (CEM) was performed at the electrostatic precipitator inlet location for CO, CO₂, NO_x, SO₂, total hydrocarbons (THC), and O₂. Total reduced sulfur (TRS) monitoring data taken regularly by the plant at the ESP outlet exhaust stack were also obtained. Plant personnel calibrated the TRS monitoring instrument daily during the test program. The continuous monitoring data were used in conjunction with process data to document the stability of combustion conditions during the test.

2.2 SUMMARY OF RESULTS

The data obtained at Site BLB-A during the Tier 4 test program are summarized in Figure 2-2. Only the hepta- and octa-CDD species were consistently detected in the stack gas at the outlet from the ESP. For the furan species, the tetra-, hexa-, hepta- and octa-CDF homologues were consistently detected in the ESP outlet gas stream. As shown in Table 2-2, average as-measured stack gas concentrations of the total PCDD and total PCDF at the ESP outlet were 0.67 ng/dscm and 0.51 ng/dscm, respectively. The average hourly emission rates were 150 ug/hr for total PCDD and 114 ug/hr for total PCDF. Octa-CDD was the most prevalent of the tetra- through octa-chlorinated dioxin homologues, while the furans were fairly evenly distributed among the tetra- through octa-chlorinated furan homologues. The ESP appeared to have positive control for reducing dioxin/furan emissions although analytical uncertainties inherent in GC/MS analysis limited the ability to quantify the control efficiency accurately.

Analysis of the flue gas samples taken at the ESP inlet did not detect any 2378 TCDD or 2378 TCDF. Detectable quantities of PCDD and PCDF homologues were found in the inlet flue gas to the ESP. Average as-measured stack gas concentrations of total PCDD and total PCDF at the ESP inlet were 1.59 ng/dscm

Dioxin/Furan Emissions Data

Species	Concentration (ng/dscm @ 3% O ₂)	Emission Rate (ug/hr)	Emissions Factor (pg/kg feed, dry)
INLET:			
2378 TCDD	ND	--	--
Total PCDD	1.79	339	40-6
Total PCDF	1.47	283	50-5
OUTLET:			
2378 TCDD	ND	--	--
Total PCDD	0.75	150	3
Total PCDF	0.57	114	2

NR = not reported; ND = not detected.

Black Liquor Boiler Operating Data (A)

Feedrate	395 gpm
Operating hours	8,760 hrs/yr

ESP Operating Data (B)

Inlet temperature	178°C
Inlet gas flowrate:	
East duct	1,800 dscmm
West duct	1,800 dscmm
Total	3,600 dscmm

Black Liquor Feed Precursor Data (C)

Chlorobenzenes	ND
PCB's	ND
Chlorophenols	trace
Total chlorides	896 ppm

Continuous Monitoring Data (C)

O ₂	6.0 %vol, dry
CO ₂	15.6 %vol @ 3% O ₂ , dry
CO	136.4 ppmv @ 3% O ₂ , dry
THC	3.5 ppmv @ 3% O ₂ , wet
SO ₂	94.4 ppmv @ 3% O ₂ , dry
NO _x	83.2 ppmv @ 3% O ₂ , dry

Flue Gas Parameter Data

INLET:	Flowrate 3,600 dscmm
	Temperature 178°C
	Moisture 28 vol%
OUTLET:	Flowrate 3,700 dscmm
	Temperature 173°C
	Moisture 25 vol%

Liquor Circuit Inputs (E) (F)

Stream	Flowrate	Total chlorides Concentration (ppm, wet basis)	Average Mass Flowrate of Chloride (lb/yr)
Caustic	25-30 gpm	1300	22.2
By-product salt cake	24 TPD	700	1.4

Chloride Emissions Data (C)

Train Component	Concentration (mg/dscm @ 3% O ₂)	Emission Rate (kg/hr)	Emission Factor (mg/kg feed, dry)
Front half	1.5	0.3	0.9
Back half	108.9	21.5	614.8
Train total	110.4	21.8	615.7

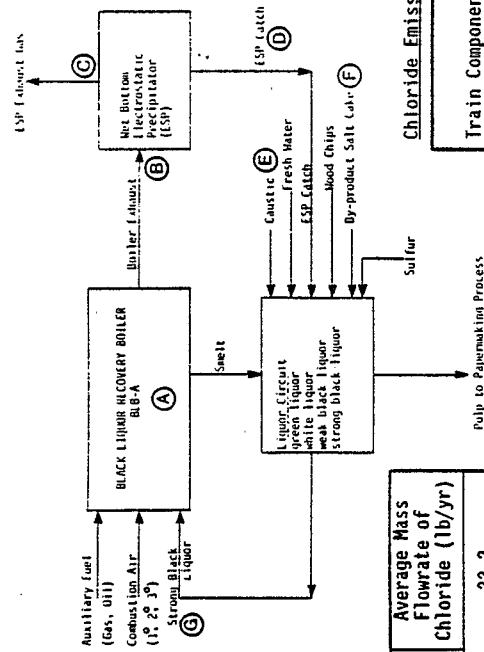


Figure 2-2. Data summary for Site BLB-A.

TABLE 2-2. SUMMARY OF MEAN DIOXIN AND FURAN EMISSIONS DATA FOR SITE BLB-A

Parameter	2378 TCDD ^a	Total PCDD	Total PCDF
INLET:			
<u>Emissions Concentration</u> (ng/dscm)			
As-measured	ND	1.59	1.31
Corrected to 3% O ₂	--	1.79	1.47
<u>Emissions Rate</u> (ug/hr)	--	339	283
OUTLET:			
<u>Emissions Concentration</u> (ng/dscm)			
As-measured	ND	0.67	0.51
Corrected to 3% O ₂	--	0.75	0.57
<u>Emissions Rate</u> (ug/hr)	--	150	114

^aValues in parenthesis are detection limits expressed in the corresponding units.

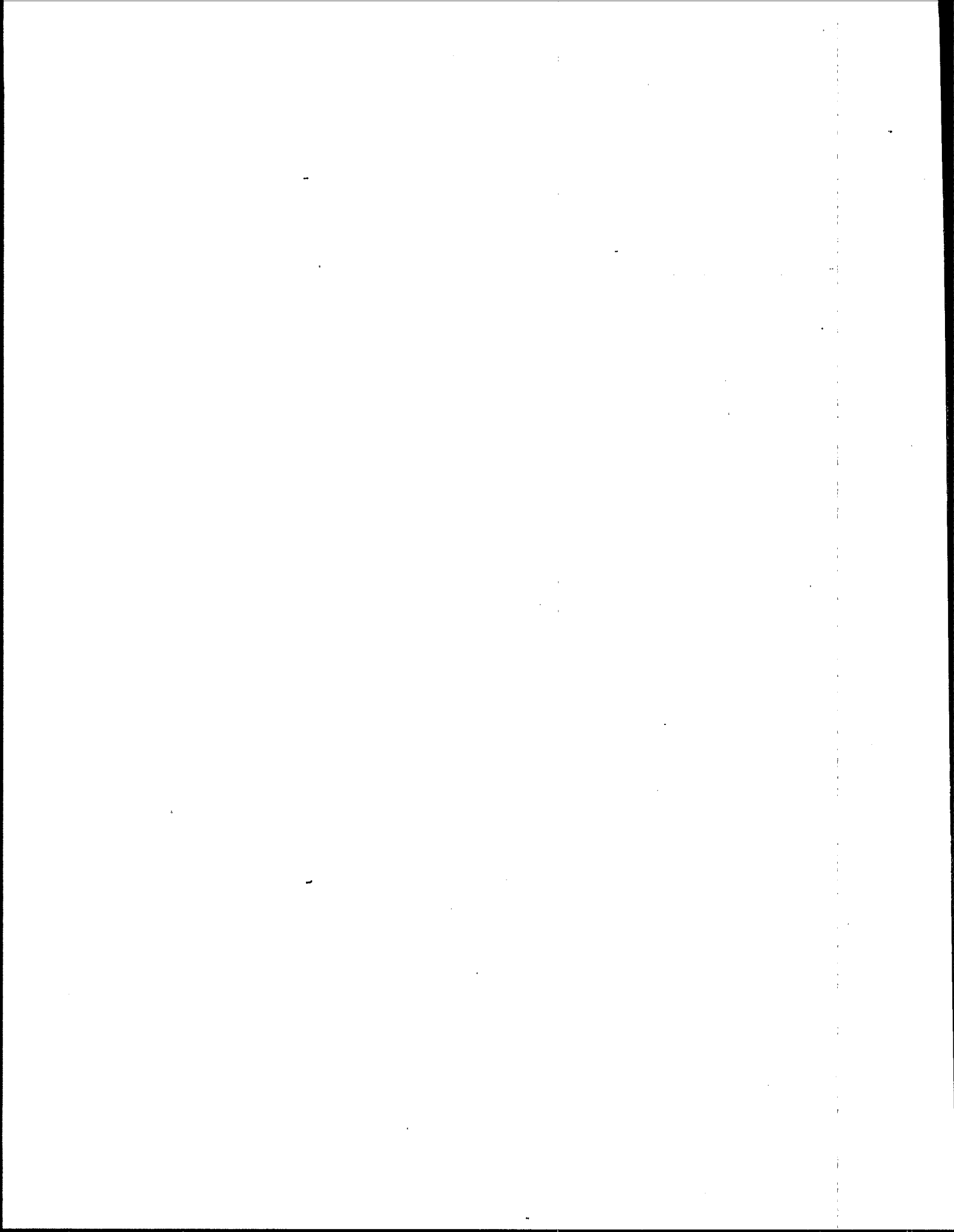
ND - Not detected.

and 1.31 ng/dscm respectively. The hourly emission rates were 339 ug/hr for total PCDD and 283 ug/hr for total PCDF. The octa-chlorinated dioxin homologue was the predominant dioxin species while the furan species were distributed fairly evenly among the tetra- through hepta-CDF homologues.

Chloride emissions at the ESP outlet were measured at 98.1 mg/dscm which corresponds to 110.4 ng/dscm @ 3% O₂. The average chlorides emission factor was calculated to be 615.7 mg chloride emitted per kilogram of black liquor fired on a dry basis.

The black liquor feed rate to Boiler BLB-A was 395 gpm during the test period. Precursor analysis of the black liquor did not detect chlorobenzenes or polychlorinated biphenyls. A trace amount of chlorophenols were detected, and the black liquor contained 896 ppm of total chlorides.

The ESP treated an average of 3,600 dscmm at a temperature of 178°C. At the ESP outlet, the measured flowrate was 3,700 dscmm at a temperature of 173°C. Average flue gas concentrations measured at the ESP outlet by the Radian continuous emissions monitoring system were: O₂, 6.0 vol%; CO₂, 15.6 vol% @ 3% O₂ (dry); CO, 136.4 ppmv @ 3% O₂; THC, 3.5 ppmv @ 3% O₂, wet; SO₂, 94.4 ppmv @ 3% O₂, dry; and NO_x, 83.2 ppmv @ 3% O₂, dry.



3.0 PROCESS DESCRIPTION

This section describes the host site and the black liquor recovery boiler/electrostatic precipitator system tested. Data summarizing the operation of the boiler and the precipitator during the test periods are presented in Section 5.0.

3.1 FACILITY DESCRIPTION

The host plant (Site 04) is a typical Kraft pulp and paper mill with a rated capacity of 855 Mg/day (940 TPD) of air-dried pulp. The plant maintains three black liquor recovery boilers with rated capacities of 365 Mg/day (400 TPD), 590 Mg/day (650 TPD), and 815 Mg/day (900 TPD) of air-dried, unbleached pulp. Two of the three black liquor boilers were operating during the dioxin/furan tests, while the third was being modified for future use. These boilers are typically base-loaded. Two combination boilers firing bark, wood refuse, and non-condensable gases from the black liquor evaporators are swing-loaded to cover variations in process steam demand.

Pine logs and pine wood chips are transported to Site BLB-A by truck. To the best knowledge of plant personnel, none of the wood processed in the pulping circuit has been stored in salt water or treated with pentachlorophenol (PCP). Approximately 80 digester batches per day are processed in the 10 digesters at the pulping plant. Each batch consists of 55 Mg (60 tons) of wood chips, 42 cubic meters (1500 ft³) of white liquor, and a small amount of weak black liquor.

The contents of the finished digester batches are sent to a "blow tank" for intermittent storage and then to a countercurrent "brown stock washer" system. Pulp is separated from the rest of the finished digester batch in the brown stock washers. Dirty water from the brown stock washers (i.e., weak black liquor) is sent to an evaporator system for solids concentration prior to being fired in the black liquor boiler. Typical process flow rates of various streams in the pulp mill circuit are shown in Table 3-1.

TABLE 3-1. TYPICAL PROCESS FLOW RATES OF VARIOUS
PULP MILL MATERIALS AT SITE BLB-A

Material	Process Flow Rate	Other Information
-By-product salt cake	24 tons/day	Predominantly sodium sulfate
-White liquor	620 gpm	Sodium sulfide/ sodium hydroxide
-Weak black liquor	1800-2000 gpm	18 wt % solids density 8.7 lb/gal
-Strong black liquor	395 gpm ^a	65 wt % solids density 11.8 lb/gal
-Caustic	15-30 gpm ^b	20% Sodium Hydroxide

a. No. 3 recovery boiler (BLB-A), 260 gpm; No. 1 recovery boiler, 135 gpm.

b. Caustic flow rate during tests was 25-30 gpm.

Cleaned pulp from the brown stock washer is bleached, pressed, and rolled into a variety of paper products. Bleaching is performed using chlorine dioxide (ClO_2), which is produced on-site using the the R3 process. Approximately 22 Mg/day (24 tons/day) of by-product salt cake (predominantly sodium sulfate, Na_2SO_4) from this process is fed to the black liquor circuit as a make-up material for sodium and sulfur losses. According to plant personnel, the atomic chlorine content of the by-product salt cake ranges from approximately 0.2 to 1.5 weight percent. Most of this chlorine is present in the form of chloride ion.

3.2 BLACK LIQUOR RECOVERY BOILER DESCRIPTION

Black liquor recovery boiler BLB-A is a low odor Babcock and Wilcox recovery boiler with a rated capacity of 815 Mg/day (900-TPD) unbleached pulp. A schematic diagram of the pulping process is shown in Figure 3-1. The boiler is typically baseloaded and operates at a steady black liquor firing rate. Particulate matter emissions from the boiler are controlled by a wet bottom electrostatic precipitator.

The solids content of the concentrated black liquor fired in Boiler BLB-A is approximately 65 percent by weight. Concentration of the weak black liquor produced by the brown stock washers is accomplished using a three-stage multiple effect evaporator system and a "concentrator," which is essentially a large single stage evaporator. Noncondensable gases collected by the evaporator system are fired in one of the two combination power boilers at the plant. Strong black liquor from the concentrator flows through the bottom of the electrostatic precipitator that controls particulate matter emissions from boiler BLB-A. The precipitator catch is mixed with the strong black liquor before the liquor is fired in the boiler.

According to plant personnel, the chloride content of the concentrated black liquor fed to boiler BLB-A is typically 0.5 weight percent chlorine (wt % Cl), with a range of approximately 0.1 to 0.7 wt % Cl. Chlorine enters the black liquor circuit as inorganic chloride ions primarily through the ClO_2 generation system byproduct salt cake that is added to the weak black liquor

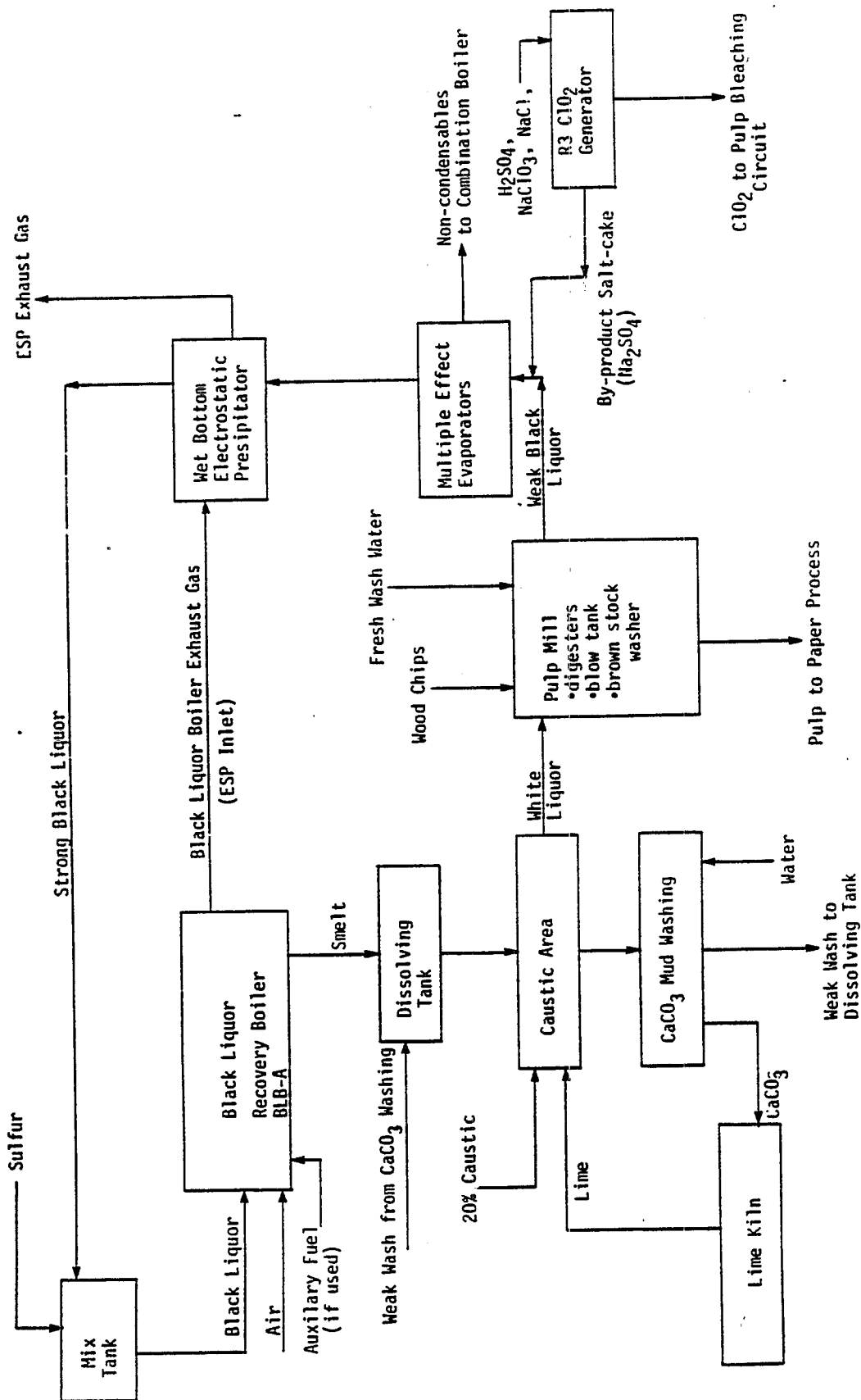


Figure 3-1. Process Flow Diagram for Site BLB-A

prior to the evaporators. The by-product salt cake contributes makeup sodium and sulfur to the black liquor circuit. The chlorine content of the by-product salt cake ranges from about 0.2 to 1.5 wt % Cl. Other potential sources of chlorine in the black liquor circuit are sodium hydroxide used in the conversion of green liquor to white liquor, wood chips fed to the pulping process, and fresh water used for CaCO_3 mud washing and pulp washing. Data on the relative amounts of chlorine added to the black liquor circuit through each of these sources have been developed from total chlorine analyses performed on process samples taken during the dioxin/furan tests.

Concentrated black liquor is sprayed into the combustion zone of boiler BLB-A at a temperature of approximately 105°C (250°F) using four oscillating "guns". Primary, secondary, and tertiary combustion air is supplied to the boiler by forced draft fans. The primary:secondary:tertiary air ratio is approximately 1.4:1.8:1.0. The combustion air supply is computer controlled using flue gas oxygen monitoring and black liquor feed rate monitoring. Oxygen is continuously monitored at the economizer section of the boiler and at the electrostatic precipitator outlet stack, while carbon monoxide, opacity, and total reduced sulfur (TRS) are continuously monitored only at the electrostatic precipitator outlet stack. The target flue gas oxygen at the economizer section is 3.0 to 3.5 percent O_2 . Opacity at the outlet stack is typically 15 percent. The Federal New Source Performance Standards (NSPS) TRS emissions limit for black liquor recovery boiler stacks is 5 ppm corrected to 8 percent oxygen. Plant personnel reported that boiler BLB-A is in compliance with the NSPS.

3.3 ELECTROSTATIC PRECIPITATOR DESCRIPTION

Exhaust gases from black liquor recovery boiler BLB-A pass through a two-chamber Wheelabrator-Frye wet bottom electrostatic precipitator (ESP) for particulate removal. Each chamber of the precipitator handles approximately half of the total boiler exhaust gas flowrate. The ESP was designed for a total gas flowrate of 110,000 acmm (387,000 acfm) at an operating temperature of 175°C (350°F). Total plate collection area is $17,150 \text{ m}^2$ ($184,570 \text{ ft}^2$), corresponding to a design specific collection area of approximately 0.16

m^2/acmm ($0.5 \text{ ft}^2/\text{acfm}$). An energy management system trims the power supply to the ESP to maintain a stack opacity of approximately 15 percent. The design particulate matter removal efficiency of the ESP is 99.6 percent when operated at full electrical power. Under typical power supply conditions, the particulate matter removal efficiency of the ESP is estimated by plant personnel to be 98.6 to 99.0 percent. The most recent particulate matter emissions data generated by the host plant showed a particulate grain loading of 0.005 gr/dscf in the precipitator exhaust gas.

4.0 TEST DESCRIPTION

This section describes the field sampling, process monitoring, and analytical activities that were performed for test Site 04. The purpose of this section is to provide sufficient descriptive information about the test so that the test data presented in Section 5.0 can be easily understood. Specific testing details (specific sampling locations and procedures) are presented in Section 6.0.

This section is divided into three parts. Section 4.1 summarizes field sampling activities, Section 4.2 summarizes process monitoring activities, and Section 4.3 summarizes analytical activities performed during the test program.

4.1 FIELD SAMPLING

Table 4-1 shows the source sampling and analysis matrix for test Site BLB-A. Three dioxin/furan emissions tests (Runs 01, 02, 03) were performed. Samples were taken simultaneously at the electrostatic precipitator (ESP) inlet location and the ESP outlet exhaust stack. These locations are shown as Points B and C on Figure 4-1. Dioxin/furan sampling followed the Modified Method 5 (MM5) sampling protocol developed by the American Society of Mechanical Engineers (ASME) for measuring emissions of chlorinated organic compounds. Testing was performed at the ESP outlet exhaust stack for a period corresponding to 240 minutes of on-line sampling. Testing was performed during the same time period at the ESP inlet location, but due to a number of filter changes and sample port changes the on-line sampling period for the inlet train was typically about 140 minutes.

Concentrations of HCl in the flue gas were determined for each test day at the ESP outlet exhaust stack using another modification of EPA Method 5. The sampling train was identical to that of Method 5 except that water in the impingers was replaced with 0.1 M NaOH or 0.1 M KOH. Sodium hydroxide (NaOH) was used in the impinger during Run 01 according to typical HCl sampling procedures. The impinger solution was changed to potassium hydroxide for

TABLE 4-1. SOURCE SAMPLING AND ANALYSIS MATRIX FOR SITE 04

Sample Location	Sample Type or Parameter	Sampling Method	Analytical Method	Number of Samples or Frequency
1. Electrostatic precipitator inlet/black liquor boiler outlet (Point C, Figure 4-1)	Dioxin, furan	Modified EPA Method 5 (MM5)	Gas chromatograph/mass spectrometer	Three test runs; one per test day
	Volumetric flow	EPA Method 2	S-type pitot	Once per MM5 test run
	Molecular weight	EPA Method 3	Gas chromatography/thermal conductivity detector	Two integrated bag samples per MM5 test run
	Moisture	EPA Method 4	Gravimetric balance	Once per MM5 test run
	HCl	Modified EPA Method 5 (MM5/HCl)	Ion chromatography	Once per MM5 test run
2. Electrostatic precipitator inlet/black liquor boiler outlet (Point B, Figure 6-1)	Dioxin, furan	Modified EPA Method 5	Gas chromatograph/mass spectrometer	Three test runs; one per test day
	Volumetric flow	EPA Method 2	S-type pitot	One per MM5 test run
	Molecular weight	EPA Method 3	Gas chromatography/thermal conductivity detector	Two integrated bag samples per MM5 test run
	Moisture	EPA Method 4	Gravimetric balance	Once per MM5 test run
	CO/CO ₂	In-stack filter probe and heat-traced Teflon sample line	Nondispersive infrared analyzer	Continuously during MM5 test runs
	O ₂	Same as CO/CO ₂	Paramagnetic analyzer	Continuously during MM5 test runs
	NO _x	Same as CO/CO ₂	Chemiluminescent analyzer	Continuously during MM5 test runs
	SO ₂	Same as CO/CO ₂	Pulsed fluorescence analyzer	Continuously during MM5 test runs
	Total hydrocarbons (THC)	Same as CO/CO ₂	Flame ionization analyzer	Continuously during MM5 test runs
	Strong black liquor mix tank for dioxin/furan, dioxin/furan precursors, and total chloride analyses	Grab samples	Gas chromatography/mass spectrometer, ion chromatography	Three identical samples of hourly sample composites for each MM5 test run
4. Caustic line from storage area to liquor circuit	20% caustic for total chloride analysis	Tap valve samples	Ion chromatography	One composite of two samples taken during each test run
5. White liquor storage tank	White liquor for total chloride analysis	Tap valve samples	Ion chromatography	One composite of two samples taken during each test run

TABLE 4-1. SOURCE SAMPLING AND ANALYSIS MATRIX FOR SITE 04 (Con't.)

Sample Location	Sample Type or Parameter	Sampling Method	Analytical Method	Number of Samples or Frequency
6. Weak black liquor line leading to concentrators	Weak black liquor for total chloride analysis	Tap valve samples	Ion chromatography	One composite of two samples taken during each test run
7. Byproduct salt cake rotary filter in ClO ₂ plant	Byproduct salt cake for total chloride analysis	Grab samples	Ion chromatography	One composite of two samples taken during each test run
8. Plant property	Soils for dioxin, furan	Grab samples	Gas chromatograph/mass spectrometer	One composite of ten samples taken at various locations

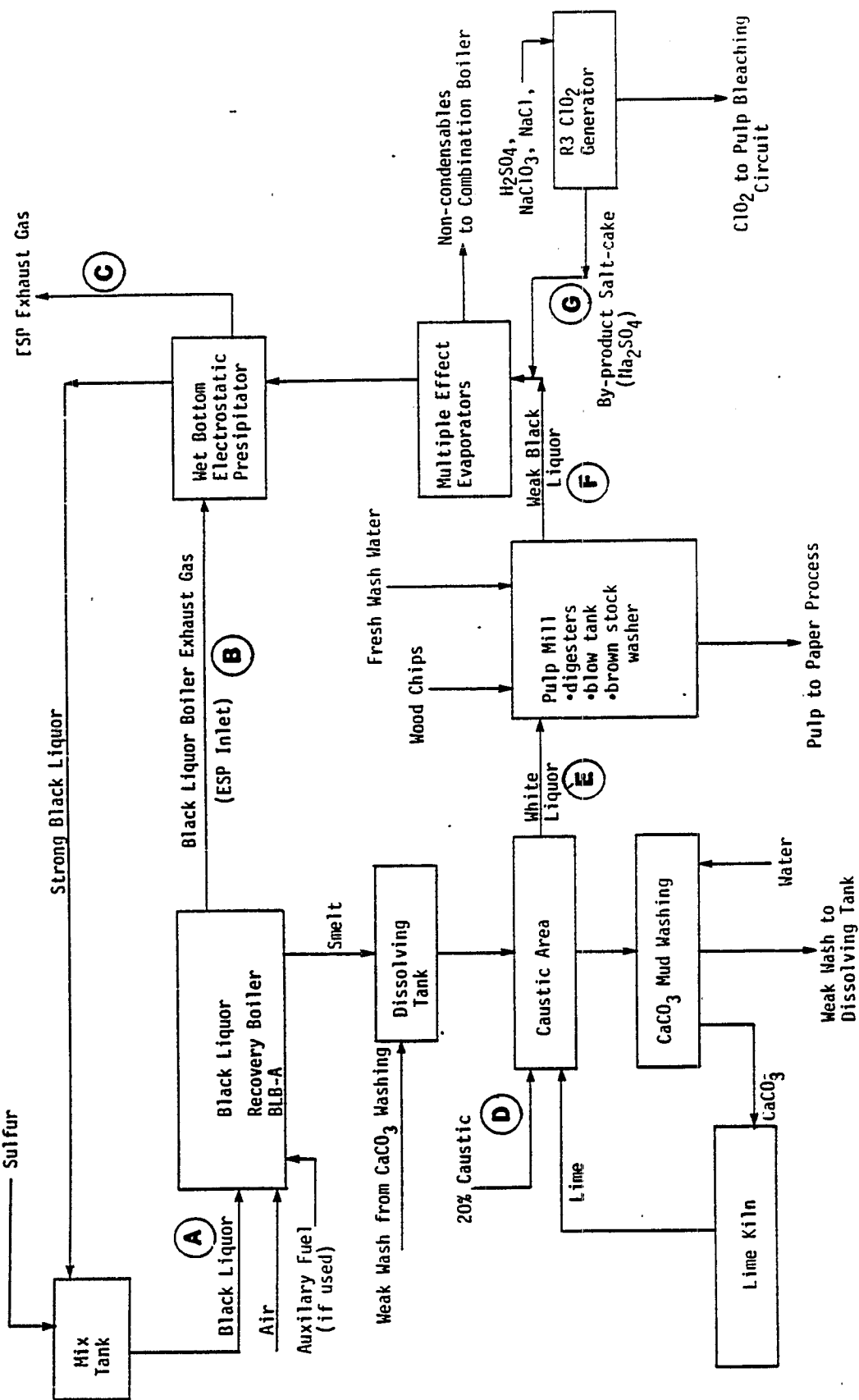


Figure 4-1. Sample Point Diagram for Black Liquor Recovery Boiler System BLB-A

subsequent runs at the suggestion of the National Council for Air and Stream Improvement (NCASI). The substitution of KOH in the impinger was performed in an effort to eliminate sodium chloride (NaCl) interferences in the HCl measurement.

Continuous emissions monitoring (CEM) of O_2 , CO, CO_2 , SO_2 , NO_x , and total hydrocarbons (THC) was performed during the three MM5 test runs. These data were obtained to assess variations in combustion during the sampling periods. Instantaneous concentration values for each species monitored were determined every five minutes by the CEM system.

Five types of process samples were taken during the MM5 test periods: strong black liquor, caustic (sodium hydroxide), white liquor, weak black liquor, and ClO_2 generation system by-product salt cake. The strong black liquor samples were taken to characterize dioxin/furan and dioxin/furan precursor contents of the material fed to the boiler. Three identical composites of hourly strong black liquor samples were prepared: one for dioxin/furan analysis by Troika, one for dioxin/furan precursor analysis by Radian/RTP, and one for total chloride analysis by Radian/Austin. The caustic, white liquor, weak black liquor, and ClO_2 generation system by-product salt cake samples were taken to indicate the major contributors of chloride to the mill liquor circuit. These samples were taken twice during each test day, and a single sample composite for each test run was sent to Radian/Austin for total chloride analysis.

Soil samples were collected from ten locations at the plant site. The ten samples were combined into a single composite, which was held for potential dioxin/furan analysis pending evaluation of the MM5 dioxin/furan emissions data.

4.2 PROCESS DATA COLLECTION

Process data were collected to characterize the operation of the black liquor boiler and electrostatic precipitator (ESP) during the MM5 test periods. A computer system in the black liquor boiler control room was used to print out a graphic display of boiler load, boiler exhaust gas oxygen content, black liquor solids content, and stack CO concentration during the

test runs. In addition, hourly average steam flow, boiler O₂, combustion air flow and temperature, black liquor characteristics, ESP gas flow and temperature, and ESP exhaust gas composition data were printed out by the computer in hourly log reports. Electrostatic precipitator voltage and current data were recorded manually at intermittent times throughout the test period. The process data are used in Section 5.1 with the CEM data to evaluate and compare combustion conditions during the three MM5 test periods.

4.3 LABORATORY ANALYSES

Laboratory analyses performed on samples from test Site BLB-A included dioxin/furan analyses, dioxin/furan precursor analyses and chloride, sulfate, and sodium analyses. Samples analyzed for dioxin/furan are discussed in Section 4.3.1 and samples analyzed for dioxin precursors are discussed in Section 4.3.2. Samples analyzed for chloride (Cl⁻) are discussed in Section 4.3.3.

4.3.1 Dioxin/Furan Analyses

All dioxin/furan analyses for Site 04 samples were performed by the EPA ECL-BSL and EMSL-RTP laboratories, two of the three laboratories collectively known as the Troika. Field samples requiring dioxin/furan analysis were prioritized by Tier 4 based on their relative importance to the Tier 4 program objectives. The priority levels, in order of decreasing importance, were designated Priority 1 through Priority 3.

Priority 1 samples were sent to Troika with instructions to perform immediate extraction and analysis. These included the MM5 train components for the electrostatic precipitator inlet and outlet sampling locations, and an MM5 train field blank.

Priority 2 samples sent to Troika were to be analyzed for dioxin/furan pending the results of the Priority 1 analyses. Priority 2 samples included only the strong black liquor samples.

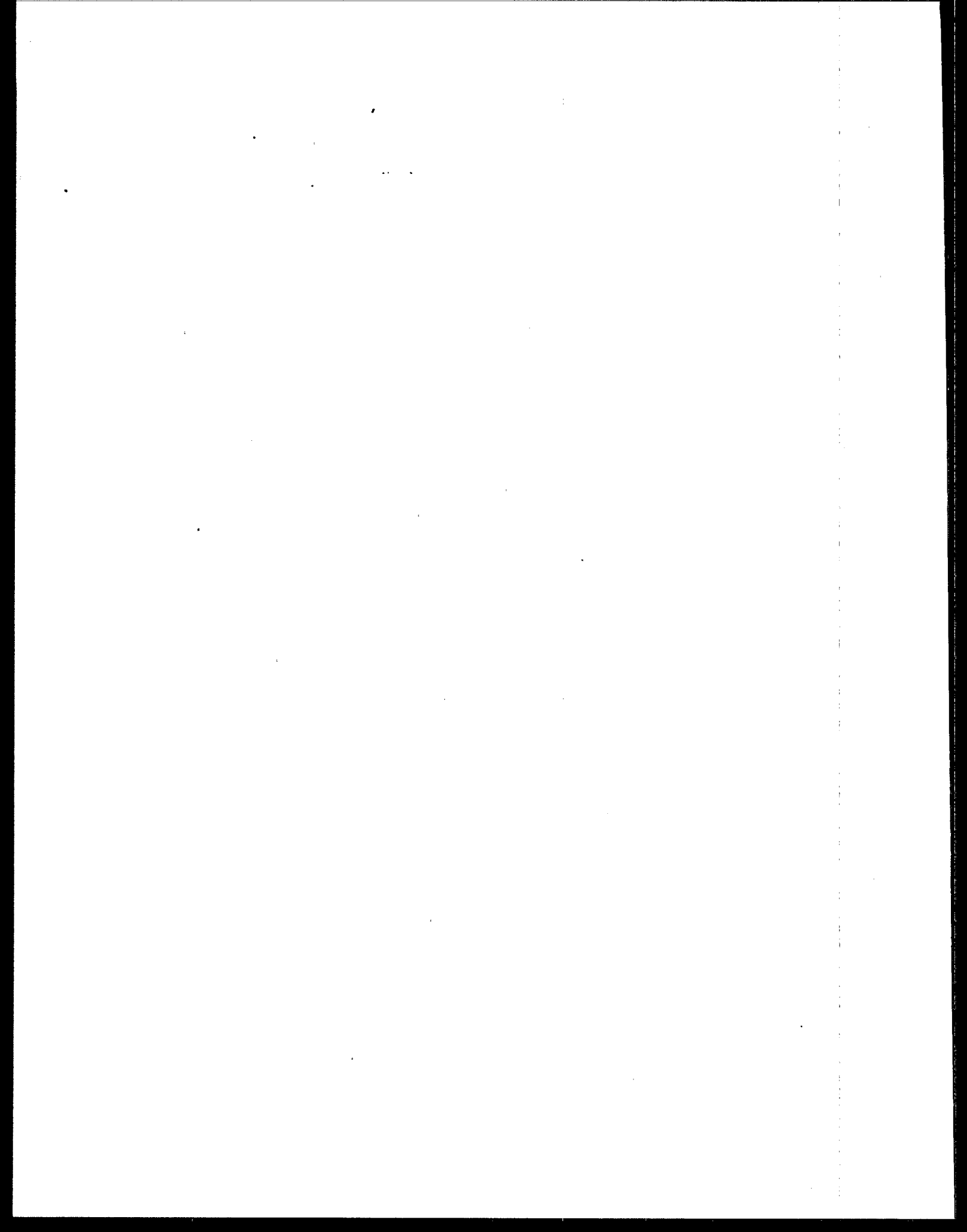
Priority 3 samples included only the composite soil sample. The soil sample is being held by Radian's Research Triangle Park (RTP), N.C. laboratory pending evaluation of the Priority 1 and 2 analyses.

4.3.2 Dioxin/Furan Precursor Analysis

Dioxin/furan precursor analyses of strong black liquor samples were performed by Radian/RTP. The specific dioxin/furan analyzed for included chlorophenols, chlorobenzenes, PCB's and total chlorine.

4.3.3 Total Chloride Analysis

Chloride analysis was performed on the combined probe rinse/filter sample and on the back-half rinse/impinger solution sample for each HCL train (i.e., front-half and back-half analysis). Chloride analysis was also performed on the strong black liquor, caustic, white liquor, weak black liquor, and by-product salt cake.



5.0 TEST RESULTS

The results of the Tier 4 dioxin/furan emissions test of black liquor boiler BLB-A are presented in this section. The individual test runs are designated as Runs 1-3.

Process data obtained during the test runs are presented in Section 5.1, and flue gas parameter data are contained in Section 5.2. Continuous monitoring results for O_2 , CO, CO_2 , NO_x , SO_2 , and THC are presented in Section 5.3. The dioxin/furan emissions data are contained in Section 5.4. Strong black liquor and dioxin/furan precursor analyses are presented in Section 5.5. Results of HCl train sampling at the electrostatic precipitator (ESP) outlet and chlorine analysis data of various process samples are presented in Sections 5.6 and 5.7.

5.1 PROCESS DATA

Process data were obtained to document black liquor boiler and electrostatic precipitator operation during the test runs. The boiler operating data are summarized in Section 5.1.1, and the electrostatic precipitator operating data are summarized in Section 5.1.2. In general, the data show that process operations were very stable during the three test runs.

5.1.1 Black Liquor Boiler Operating Data

Plant-maintained data summarizing the mean operating conditions of black liquor boiler BLB-A during the three MM5 test runs are shown in Table 5-1. Continuous data on boiler load, flue gas oxygen content in the economizer section of the boiler, strong black liquor solids content, and flue gas carbon monoxide content at the ESP outlet exhaust stack are shown for each run in Figures 5-1, 5-2, and 5-3. These data plots were generated for the dioxin/furan test program by a computer system in the black liquor boiler control room. The data show that the incinerator was operated similarly during the individual test runs. There were no process upsets during any of the runs that resulted in sampling interruptions. The one hour data gap in Figure 5-3 is due to a brief malfunction in the data acquisition system. The process was stable during this time period.

TABLE 5-1. MEAN VALUES^a OF VARIOUS BOILER OPERATING PARAMETERS MONITORED BY HOST PLANT AT SITE BLB-A

Run	Boiler Load ^b (10 ³ lb/hr)	Boiler Oxygen ^c (%)	Black Liq. Flow (gpm)	Black Liq. % Solids (wt%)	Stack Oxygen ^d (%)	Stack CO ^e (ppmv)	Stack TRS ^f (ppmv) @ 8% O ₂)	Stack Opacity (%)
Run 1	389	4.1	256	65.6	6.4	3.7	4.0	15.0
Run 2	389	3.6	254	65.6	6.0	17	4.5	15.0
Run 3	389	3.1	264	63.7	5.5	65	3.5	15.0
Mean Value	389	3.6	258	65.0	6.0	29	4.0	15.0
Runs 1-3								

a. Values shown in units used by host plant.

To convert from:

1b/hr

kg/hr

multiply by:

0.454

0.00379

6.893

0.556

0.454

cumeter/min

psi

°R

°K

1360°R

1452 Btu/lb)

1452 Btu/lb)

b. Boiler load in 10³lb/hr steam @ 914.7 psi, 1360°R (1452 Btu/lb)

c. Boiler oxygen measured at economizer section (wet basis)

d. Stack oxygen measured at ESP outlet stack (dry basis)

e. Stack CO measured at ESP outlet stack (wet basis, O₂ as measured)

f. Stack TRS measured at ESP outlet stack (dry basis, corrected to 8% O₂)

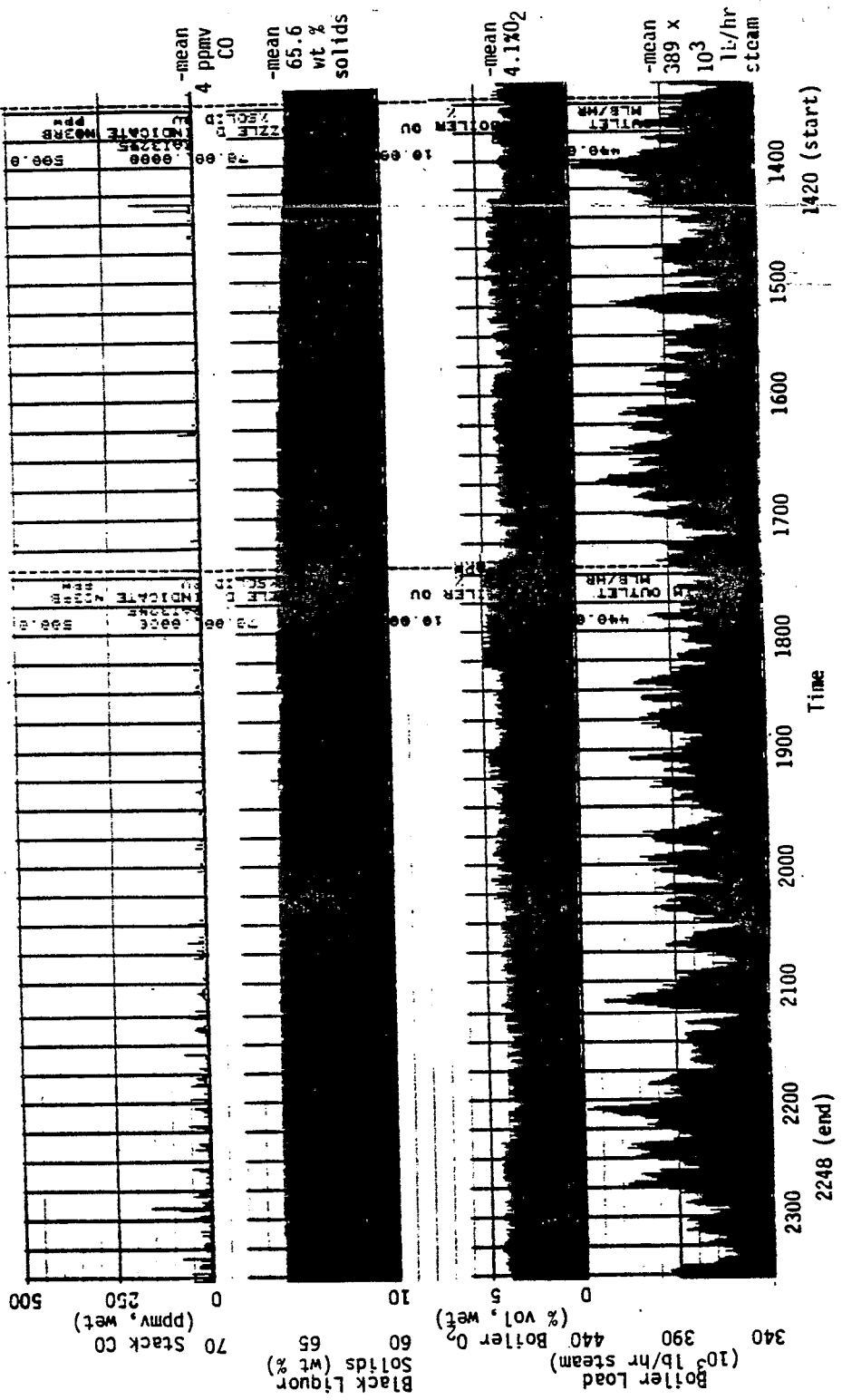


Figure 5-1. Continuously Monitored Boiler Operating Parameters During Run 1

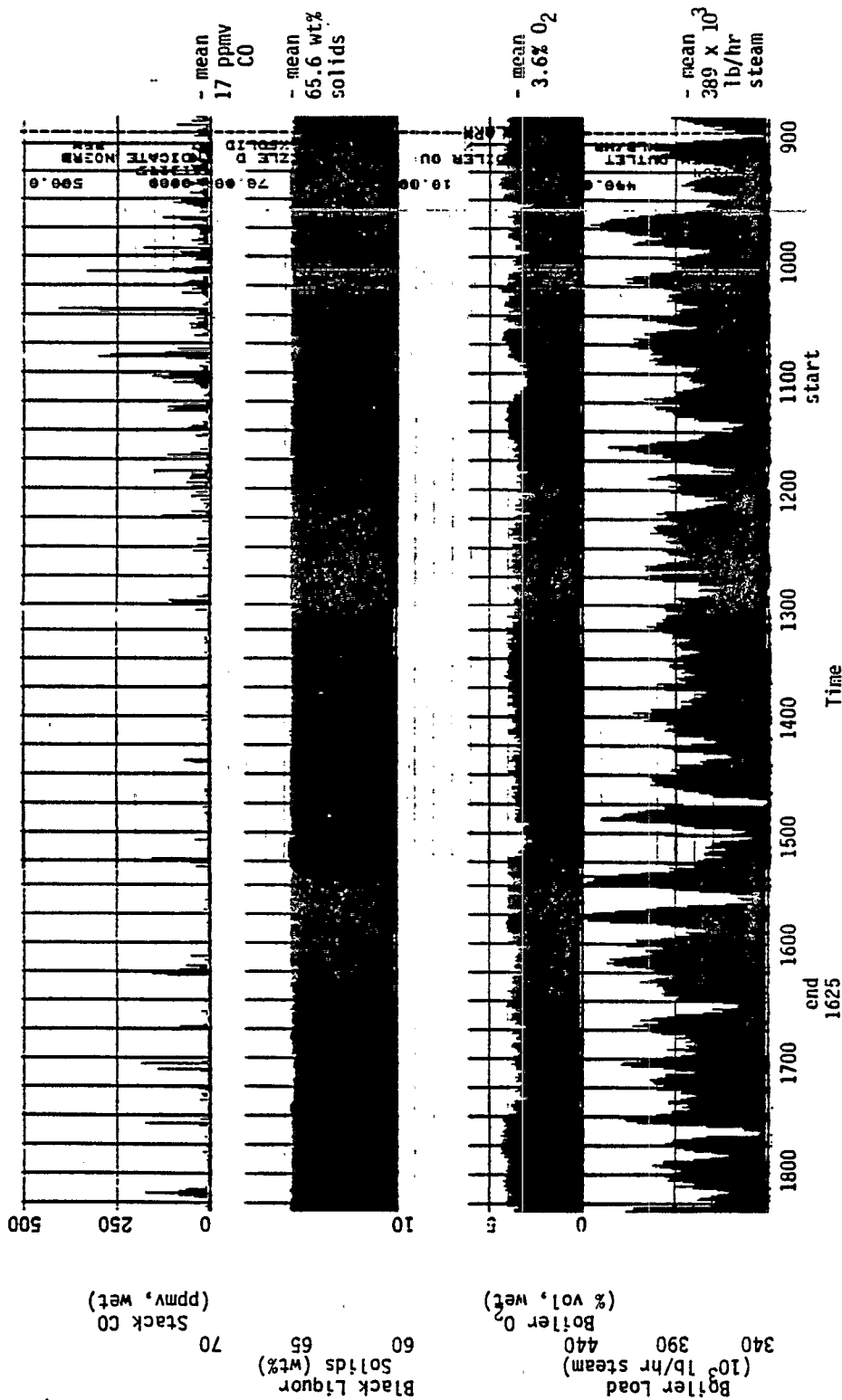


Figure 5-2 Continuously Monitored Boiler Operating Parameters During Run 2

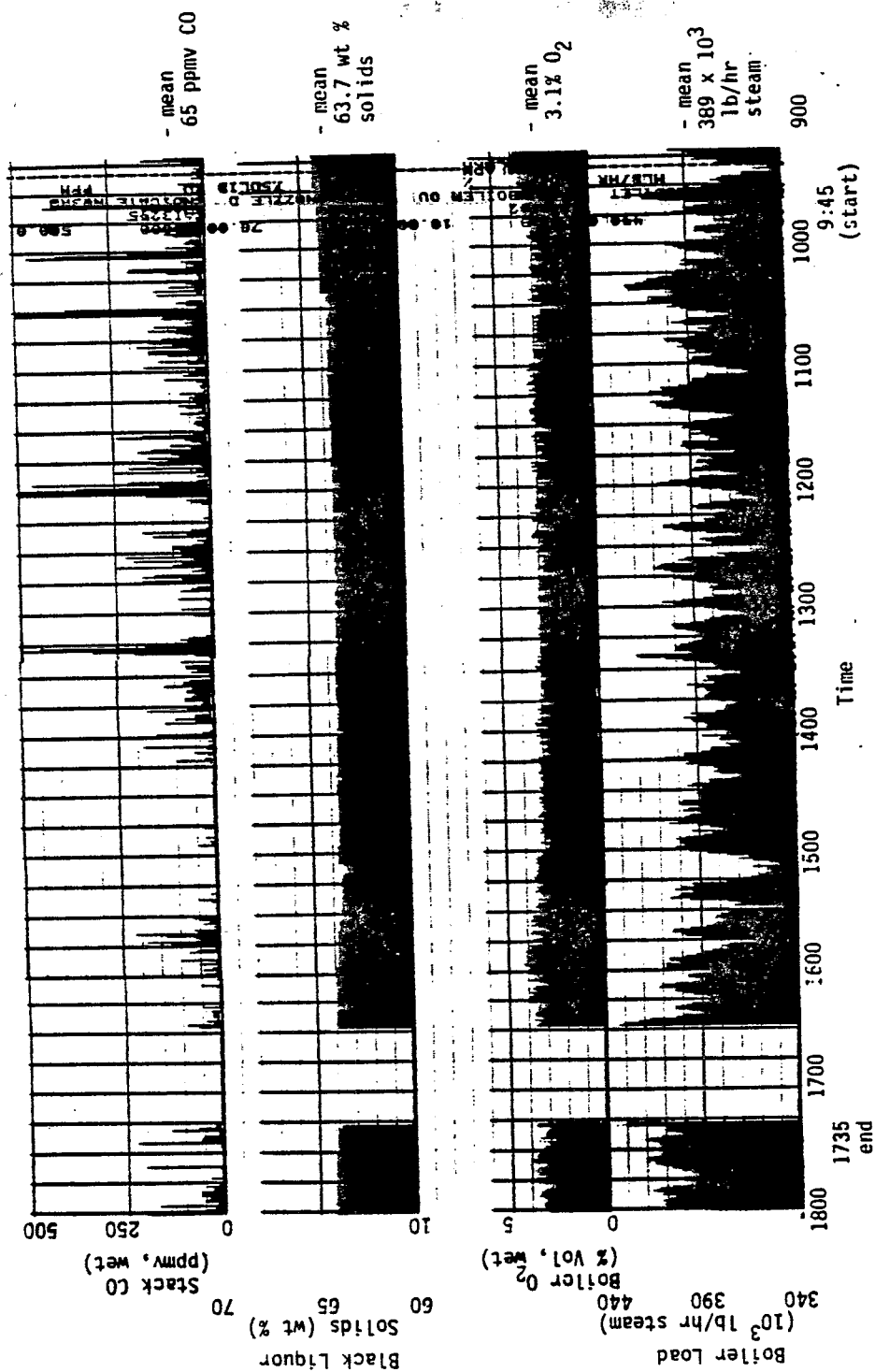


Figure 5-3. Continuously Monitored Boiler Operating Parameters During Run 3

The mean boiler load was identical for the three test runs: 176,000 kg/hr (389,000 lb/hr) steam corrected to 6305 kPa and 755⁰K (915 psi and 1360⁰R). The enthalpy of steam at these temperatures and pressure conditions is approximately 2210 kJ/kg (950 Btu/lb) relative to liquid water at the saturation temperature of the steam (552⁰K, or 994⁰F). Thus, the steam output of the boiler on an energy basis averaged 110 MW (370 MMBtu/hr) for each of the three test runs. The maximum within-run deviation of the instantaneous boiler load values shown in Figures 5-1 through 5-3 is approximately ± 13 percent of the mean value for each run.

The mean flue gas oxygen values in the economizer section of boiler BLB-A were 4.1, 3.6, and 3.1 percent O₂ (wet basis) for Runs 1-3 respectively. This corresponds to mean excess air values of approximately 30, 25, and 21 percent respectively. The maximum deviation of the instantaneous flue gas oxygen values shown in Figures 5-1 through 5-3 is approximately ± 1 percent O₂, which corresponds to a within-run excess air variability of approximately ± 10 percent excess air. Carbon monoxide data generated by the host plant are consistent with the oxygen monitoring data and show typical inverse CO/O₂ behavior. Run 1, which showed the highest boiler flue gas oxygen value (4.1% O₂), also showed the lowest flue gas carbon monoxide value (4.6 ppmv @ 3% O₂). Run 3, which showed the lowest boiler flue gas oxygen value (3.1% O₂, wet), showed the highest flue gas carbon monoxide value (76 ppmv @ 3% O₂). The plant continuous monitoring data are compared with the Radian continuous monitoring data in Section 5.2.

The solids content of the strong black liquor combusted in boiler BLB-A showed little variability during the test runs, although the mean value for Run 3 (63.7 wt % solids) was slightly lower than the mean values for Runs 1 and 2 (65.6 wt % solids). The black liquor flow rate also showed little variability. The flow rate was approximately 3.5 percent higher for Run 3 than for Runs 1 and 2. The low black liquor solids content and the high black liquor flow rate for Run 3 are probably related. In order to produce a constant amount of steam, the feed forward boiler control system responded to the lower solids content of the black liquor (i.e., lower heating value) by increasing the black liquor flow to the burners.

The TRS monitoring data showed an average of 4.0 ppmv TRS @ 8% O₂, which is in compliance with the Federal New Source Performance Standard (NSPS) for black liquor boilers. TRS values did not seem to track other combustion-related parameters such as O₂ or CO. Stack opacity averaged 15.0 percent for each of the three test runs. The opacity values depend on electrostatic precipitator operation as well as combustion conditions in the boiler.

5.1.2 Electrostatic Precipitator Operating Data

Electrostatic precipitator (ESP) operating data maintained by the host plant are summarized in Table 5-2 for test runs 01, 02, and 03. The data show that the ESP was operated similarly during the three runs.

The ESP was set in the opacity control mode during each run, which is typical of normal operation at Site 04. Under the opacity control mode, the power supplied to the ESP is adjusted to maintain a stack outlet opacity of 15 percent. The total power consumption rate of the precipitation during the test runs was approximately 50 kw, split equally between the east and west halves of the unit. The mean outlet gas temperature ranged from 174⁰ to 175⁰ C (345⁰ to 347⁰ F), and the gas flow rates to the two halves of the unit were nearly identical. Opacity monitors stationed at the outlet from each half of the ESP varied considerably from east to west, but the east half and west half values were nearly constant between runs. Plant personnel were not certain whether the higher opacity values measured in the east half outlet were due to a difference in particulate matter removal efficiencies between the east and west halves, or whether the east half opacity monitor was not working correctly. In any event, the total stack opacity of the combined streams at the outlet exhaust stack was nearly constant at 15.0 percent. The low stack opacity values were consistent with visual observation.

The gas flow rate measured by Radian at the inlet to the ESP ranged from 7930 acmm to 8160 acmm (280,000 acfm to 288,000 acfm). Using a plate area of 17,150 m² (184,570 ft²), this corresponds to a specific collection area of approximately 0.20 m²/acmm (0.65 ft²/acfm) which is higher than the design value of 0.16 m²/acmm (0.5 ft²/acfm).

TABLE 5-2. SUMMARY OF PLANT-MAINTAINED ELECTROSTATIC
PRECIPITATOR OPERATING DATA FOR SITE BLB-A

Parameter	Run 01	Run 02	Run 03
Control Mode	Opacity Control	Opacity Control	Opacity Control
Gas Flow Rate			
1. East Precip Flow Indicator (10 ³ acfm)	107	106	106
2. West Precip Flow Indicator (10 ³ acfm)	110	108	109
Outlet Temperature			
3. East Precip Outlet Temperature (°F)	346	345	346
4. West Precip Outlet Temperature (°F)	347	345	346
Opacity			
5. East Precip Opacity (%)	25.7	24.8	24.7
6. West Precip Opacity (%)	13.0	13.9	13.6
7. Stack Opacity (%)	15.0	15.0	15.0
Power Consumption ^a			
8. East Precip (kw)	23.6	25.3	27.9
9. West Precip (kw)	26.6	28.5	24.6

^a Power consumption numbers are based on limited data (7 voltage-current pairs during the 3 tests)

5.2 FLUE GAS PARAMETER DATA

Table 5-3 summarizes flue gas temperature, moisture, volumetric flowrate, and oxygen concentration data obtained at Site BLB-A. These parameters were fairly consistent among test runs. The average flue gas temperature and moisture content measured at the ESP inlet and outlet were 178°C, 27.8%, and 173°C, 24.8%, respectively. The average gas flowrates for the ESP inlet and outlet under actual stack temperature and moisture conditions were 8100 acmm (285,900 acfm) and 7,600 acmm (268,300 acfm), respectively. The average dry standard flowrate was 3,600 dscmm (127,100 dscfm) for the inlet and 3,700 dscmm (130,600 dscfm) for the outlet. Standard EPA conditions are 20°C (68°F) and 1 atm.

Flue gas oxygen concentration data were obtained from the plant O₂ analyzer, the Radian CEM system, and integrated bag samples (EPA Method 3). The average ESP inlet O₂ concentration of the flue gas measured by EPA Method 3, the Radian CEM, and the plant O₂ analyzer was 5.2, 6.0, and 4.8 vol%, respectively. The ESP outlet O₂ concentration was 5.0 vol%, as measured by EPA Method 3.

5.3 CONTINUOUS EMISSIONS MONITORING DATA

Mean values and standard deviations of the continuously monitored combustion gases at the ESP inlet location (O₂, CO, CO₂, SO₂, NO_x and THC) are shown for each MM5 test run in Table 5-4. The data show that most of the runs have similar mean concentration values for the individual gases. The overall mean values for the three test runs are as follows: oxygen, 6.0 percent by volume (dry); carbon monoxide, 136 ppmv (dry @ 3% O₂); carbon dioxide, 15.6 percent by volume (dry @ 3% O₂); sulfur dioxide, 94 ppmv (dry @ 3% O₂); nitrogen oxides, 83 ppmv (dry @ 3% O₂); and total hydrocarbons, 4 ppmv (wet @ 3% O₂, as propane).

The mean oxygen concentration measured by the Radian CEM system at the ESP inlet was fairly consistent between runs, but was slightly higher for Run 03 (6.5%) than for Runs 01 and 02 (5.9 and 5.7%, respectively). This was not consistent with the trend of the measured oxygen content data obtained by the plant at the economizer section of the boiler and at the stack. Table 5-5

TABLE 5-3. FLUE GAS PARAMETERS AT SITE BLB-A^a

Flue Gas Parameter	Run 01		Run 02		Run 03		Average	
	Inlet	Outlet	Inlet	Outlet	Inlet	Outlet	Inlet	Outlet
Temperature (°C)	177.0	175.0	178.0	173.0	178.0	172.0	178.0	173.0
Moisture (Vol %)	25.0	24.9	27.0	23.9	31.5	25.7	27.8	24.8
<u>Volumetric Flowrate</u>								
Actual (acmm)	8200	7800	8100	7600	8000	7500	8100	7600
Dry standard (dscmm)	3800	3800	3700	3700	3400	3600	3600	3700
<u>Oxygen Content (Vol%)</u>								
EPA Method 3 ^b	-	-	-	5.6	5.2	4.3	5.2	5.0
Radian CEM	5.9	-	5.7	-	6.5	-	6.0	-
Plant CEM ^c	5.5	6.4	4.8	6.0	4.1	5.5	4.8	6.0

^aMetric units are reported for all of the flue gas measurement data.

To convert to alternate units: °F = 1.8(°C) + 32; cfm = cmm x 35.5.

^bVolume % concentration on a dry basis.

^cThe inlet values were measured at the economizer; adjusted to a dry basis using 25% H₂O. The outlet values were measured at the stack; reported on a wet basis.

TABLE 5-4. MEAN VALUES AND STANDARD DEVIATIONS OF CONTINUOUSLY MONITORED COMBUSTION GASES DURING DIOXIN/FURAN TESTS AT SITE BLB-A

Parameter ^a	Run 01	Run 02	Run 03	Overall Mean
O ₂ (% vol)	5.9 (0.3)	5.7 (0.4)	6.5 (0.5)	6.0
CO (ppmv @ 3% O ₂)	84.4 (65.2)	159.4 (79.4)	165.5 (149.2)	136.4
CO ₂ (% Vol @ 3% O ₂)	16.3 (0.8)	14.7 (1.3)	15.9 (1.4)	15.6
SO ₂ (ppmv @ 3% O ₂)	112.1 (102.1)	54.1 (43.0)	117.0 (85.2)	94.4
NO _x (ppmv @ 3% O ₂)	75.1 (7.4)	89.1 (4.5)	85.4 (4.3)	83.2
THC (ppmv @ 3% O ₂)	2.9 (3.6)	2.8 (2.2)	4.8 (6.5)	11.9

^aAll concentrations expressed on a dry volume basis except for total hydrocarbon concentrations, which are expressed on a wet volume basis.

^bTotal hydrocarbon data are expressed in units of ppmv (wet) as propane.

summarizes the Radian CEM data at the ESP inlet location, the Radian Method 3 data at the ESP inlet and stack outlet locations, the plant CEM data at the economizer section of the boiler and the plant CEM data at the outlet stack location. While the data are not conclusive, it appears that a minor leakage may have developed in the Radian CEM sampling system during the tests.

Instantaneous concentration values obtained at 5-minute intervals for each of the continuously monitored combustion gases are tabulated in Appendix A-2 and are shown graphically as functions of time in Figures 5-4 through 5-9. These graphs show that in general the measured O_2 and NO_x values were fairly constant within runs. The CO and CO_2 data show considerable random scatter, much of which is attributable to instrument noise. However, the CO data do verify the trend toward higher CO levels for Runs 02 and 03 than for Run 01, as was also indicated by the plant CO monitor at the stack. The THC data do not seem to track either O_2 or CO data. THC peaks in the 20 ppm range were observed for each test run, but these peaks could not be correlated with variations in process conditions in the boiler.

The SO_2 data show a distinctive cyclical behavior which is believed to be associated with the make-up sulfur addition rate in the black liquor mix tank. Peaks as high as 400 ppmv SO_2 @ 3% O_2 were observed during Runs 01 and 03, with lower peaks observed during Run 02.

5.4 MM5 DIOXIN/ FURAN EMISSIONS DATA

5.4.1 Electrostatic Precipitator Inlet

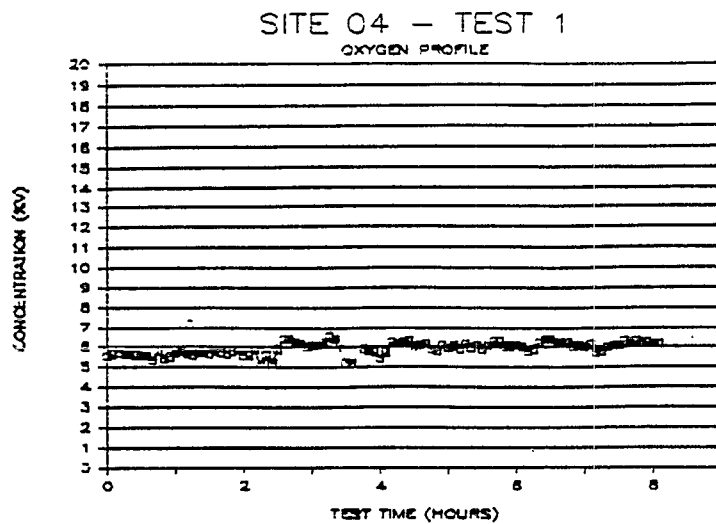
Emission concentrations and emissions rate data measured at the electrostatic precipitator (ESP) inlet are shown in Table 5-6 and 5-7 for the 2378 TCDD, total PCDD, and total PCDF species. The data include dioxin and furan collection in the entire MM5 train, including filter, XAD sorbent trap, impingers, and sample train clean-up rinses.

Average as-measured ESP inlet concentrations of total PCDD, and PCDF species were 1.58 ng/dscm total PCDD and 1.31 ng/dscm total PCDF. When corrected to 3% O_2 using the Radian CEM oxygen concentration data, these values correspond to 1.79 ng/dscm @ 3% O_2 and 1.47 ng/dscm @ 3% O_2 , respectively. Average emission rates were 340 ug/hr total PCDD, and 280 ug/hr

TABLE 5-5. COMPARISON OF MEASURED OXYGEN CONCENTRATION
VALUES (VARIOUS LOCATIONS AND METHODS)

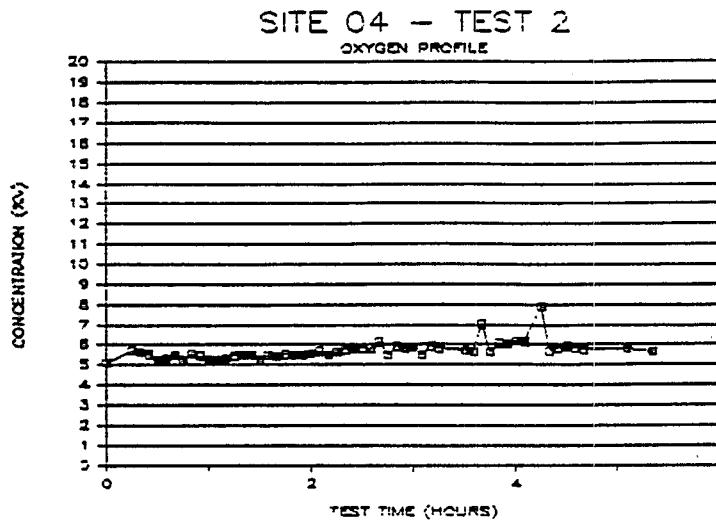
Run Number	Economizer		Precipitator Inlet		Outlet Stack	
	Plant CEM at Economizer (wet)	(dry) ^a	Radian CEM	Radian Method 3	Radian Method 3	Plant CEM at stack
01	4.1	5.5	5.9	-	-	6.4
02	3.6	4.8	5.7	-	5.6	6.0
03	3.1	4.1	6.5	5.2	4.3	5.5

^aAssuming 25% moisture.



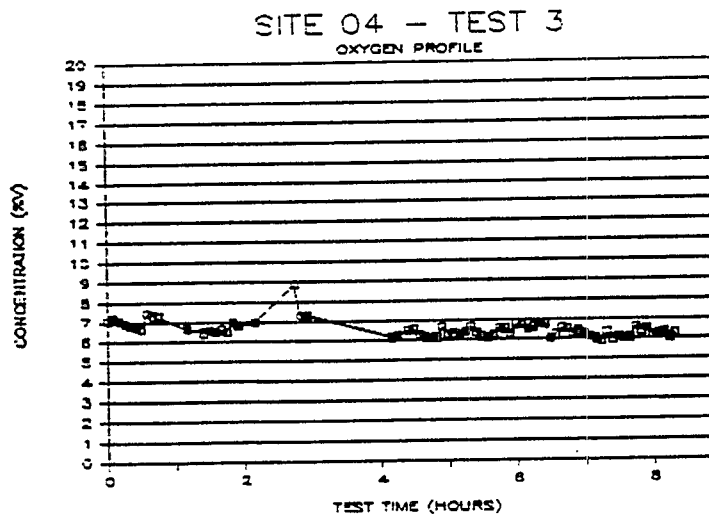
mean = 5.9% O_2

std dev = 0.3% O_2



mean = 5.7% O_2

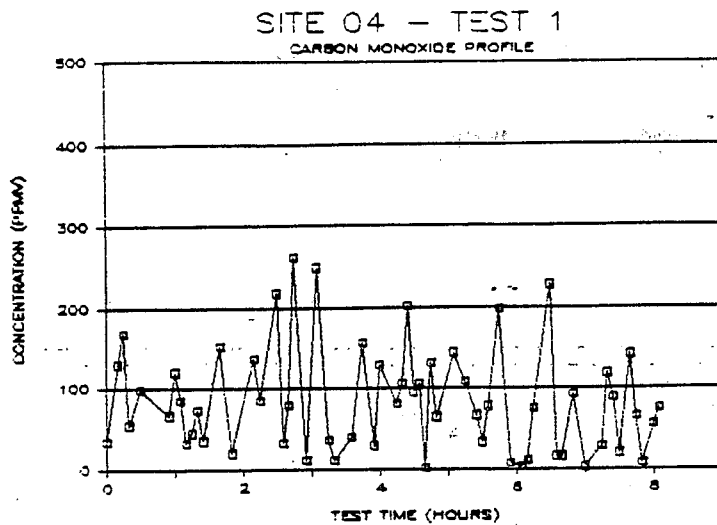
std dev. = 0.4% O_2



mean = 6.5% O_2

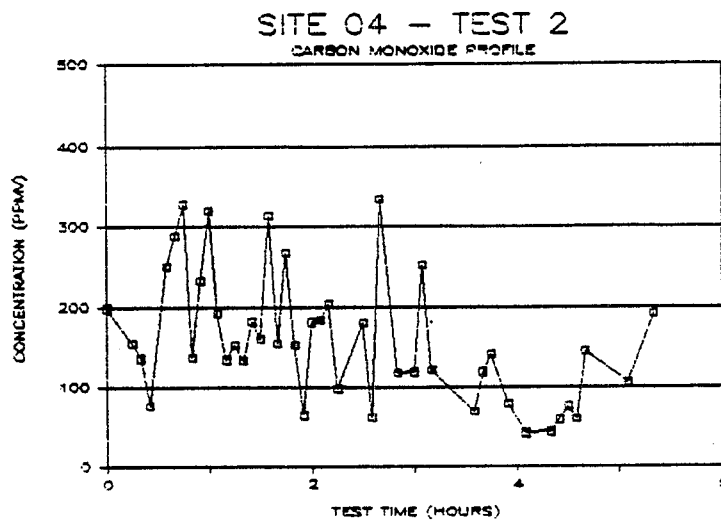
std dev. = 0.5% O_2

Figure 5-4 Oxygen Concentration History at the Electrostatic Precipitator Inlet Location (dry vol %)



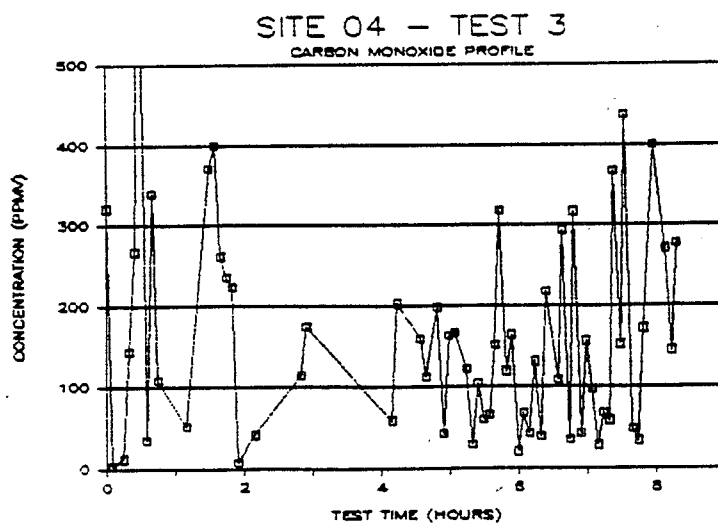
mean = 84.4 ppmv CO
@ 3% O₂

std dev = 65.2 ppmv CO
@ 3% O₂



mean = 159.4 ppmv CO
@ 3% O₂

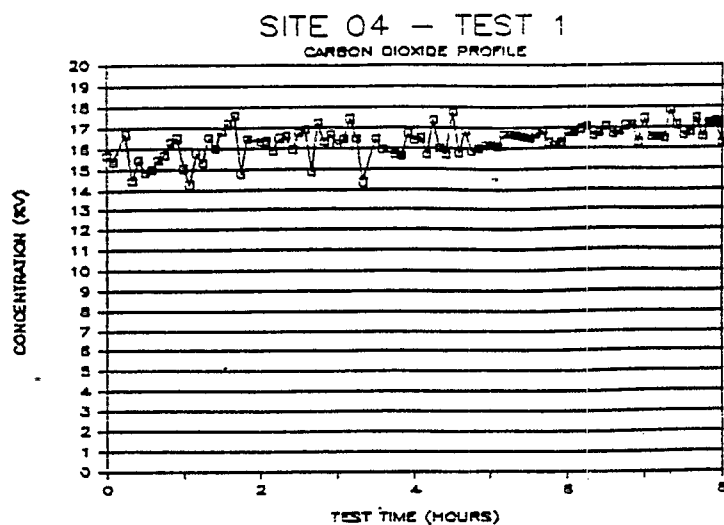
std dev = 79.4 ppmv CO
@ 3% O₂



mean = 165.5 ppmv CO
@ 3% O₂

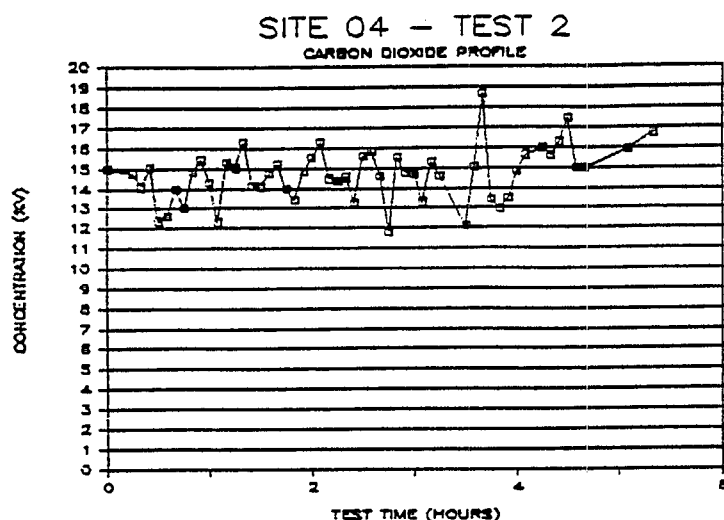
std dev = 149.2 ppmv CO
@ 3% O₂

Figure 5-5. Carbon Monoxide Concentration History at the Electrostatic Precipitator Inlet Location
(dry ppmv @ 3% O₂)



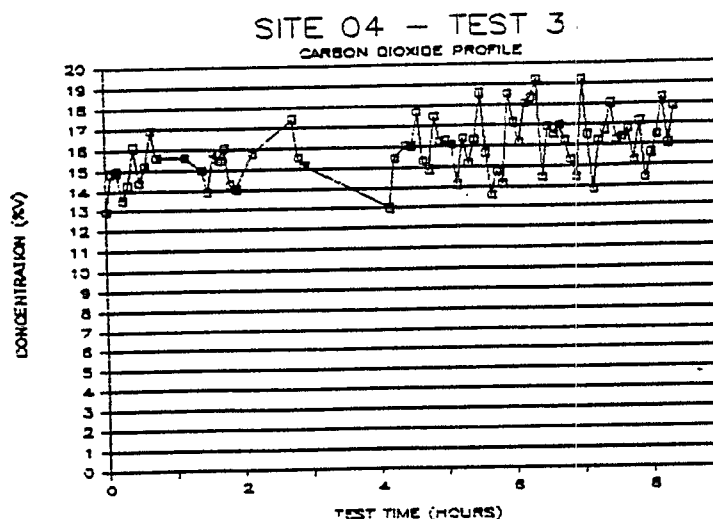
mean = 16.3% CO_2
@ 3% O_2

std dev = 0.8% CO_2
@ 3% O_2



mean = 14.7% CO_2
@ 3% O_2

std dev = 1.3% CO_2
@ 3% O_2



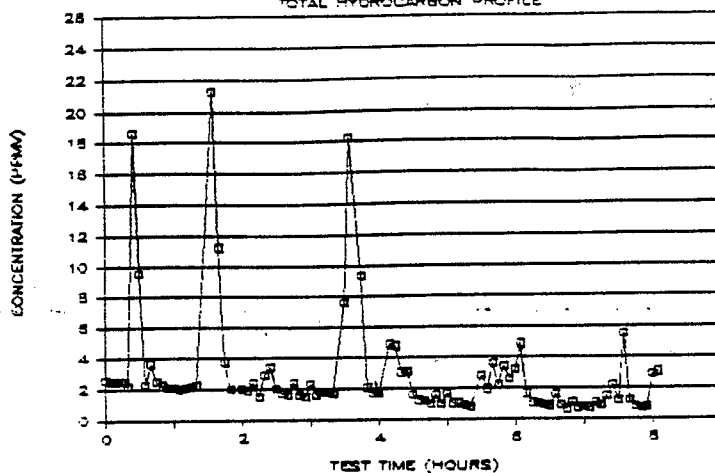
mean = 15.9% CO_2
@ 3% O_2

std dev = 1.4% CO_2
@ 3% O_2

Figure 5-6 Carbon Dioxide Concentration History at the Electrostatic Precipitator Inlet Location (dry vol % @ 3% O_2)

SITE 04 - TEST 1

TOTAL HYDROCARBON PROFILE

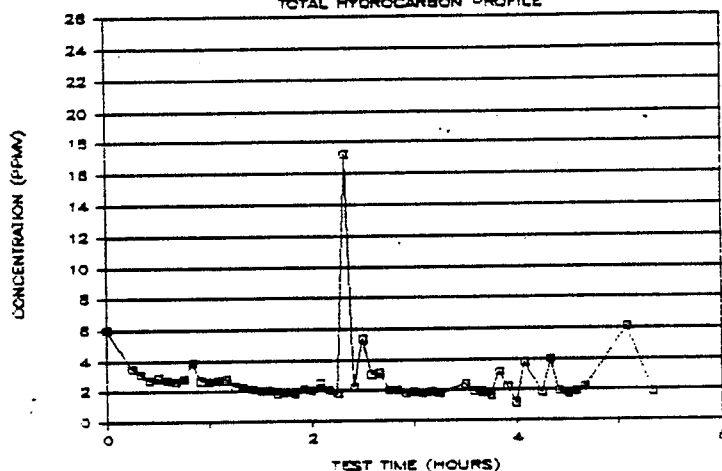


mean = 2.9 ppmv THC
@ 3% O₂

std dev = 3.6 ppmv THC
@ 3% O₂

SITE 04 - TEST 2

TOTAL HYDROCARBON PROFILE

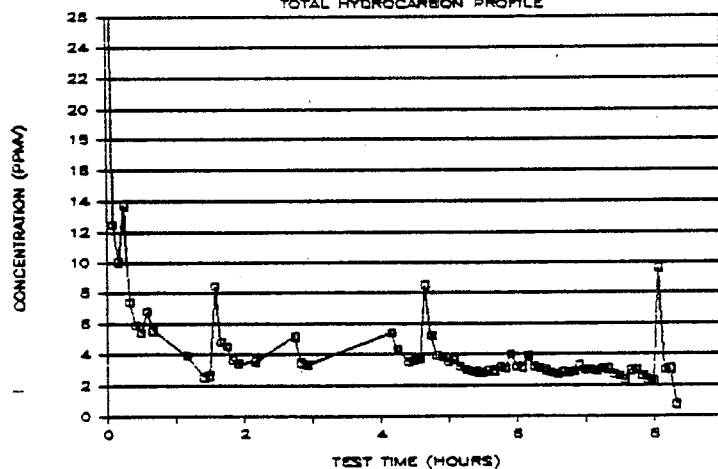


mean = 2.8 ppmv THC
@ 3% O₂

std dev = 2.2 ppmv THC
@ 3% O₂

SITE 04 - TEST 3

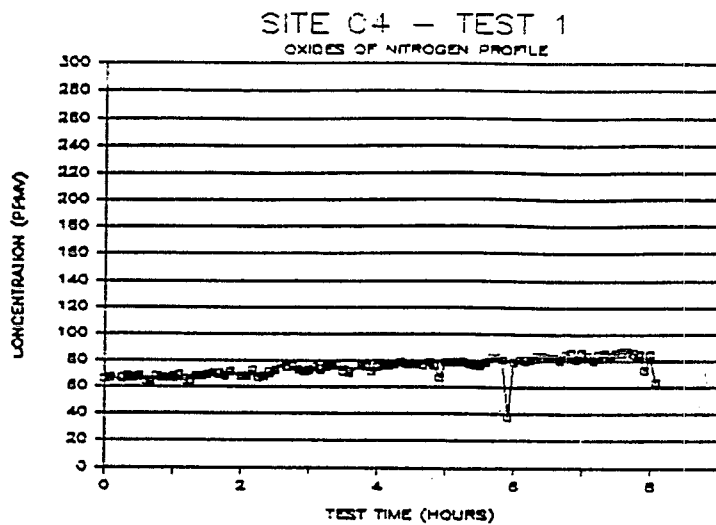
TOTAL HYDROCARBON PROFILE



mean = 4.8 ppmv THC
@ 3% O₂

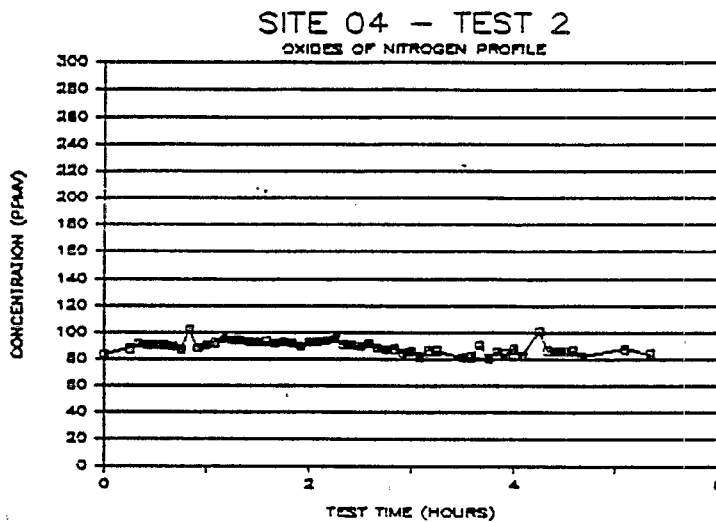
std dev = 6.5 ppmv THC
@ 3% O₂

Figure 5-7 Total Hydrocarbon Concentration History at the Electrostatic Precipitator Inlet Location (wet ppmv as propane @ 3% O₂)



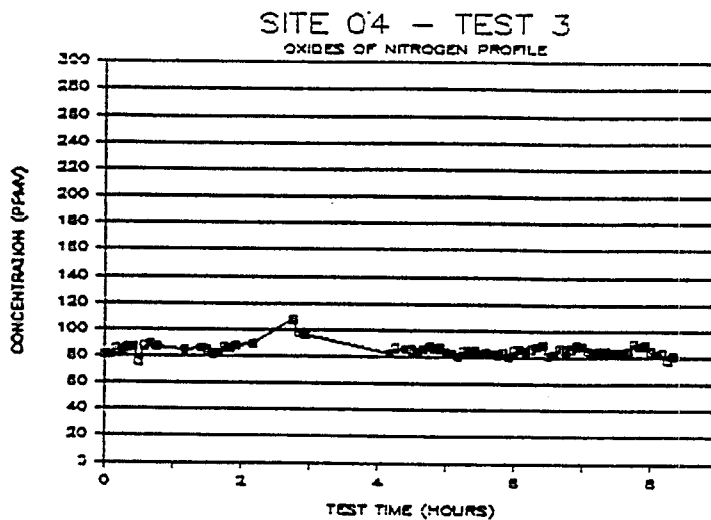
mean = 75.1 ppmv NO_x
@ 3% O_2

std dev = 7.4 ppmv NO_x
@ 3% O_2



mean = 89.1 ppmv NO_x
@ 3% O_2

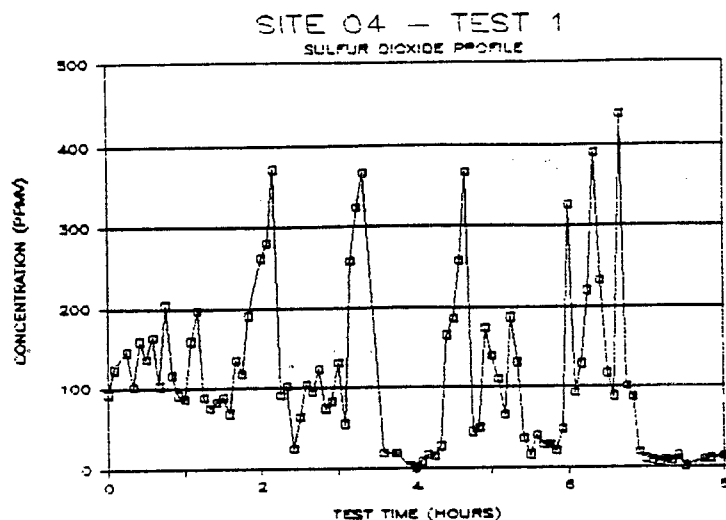
std dev = 4.5 ppmv NO_x
@ 3% O_2



mean = 85.4 ppmv NO_x
@ 3% O_2

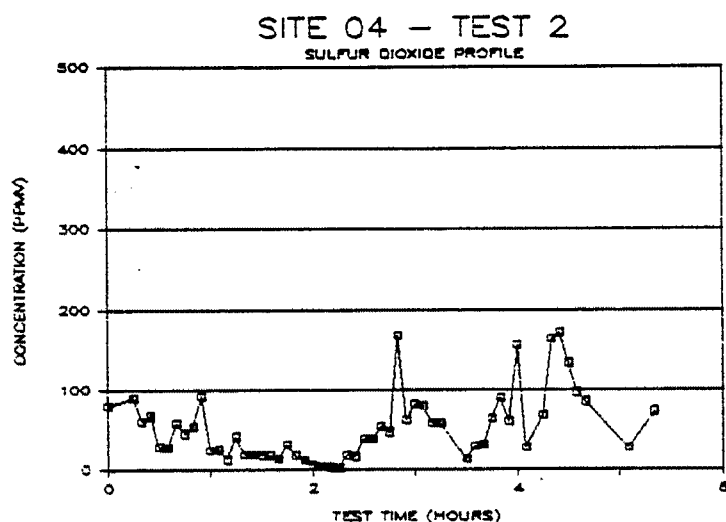
std dev = 4.3 ppmv NO_x
@ 3% O_2

Figure 5-8 Nitrogen Oxides Concentration History at the Electrostatic Precipitator Inlet Location (dry ppmv @ 3% O_2)



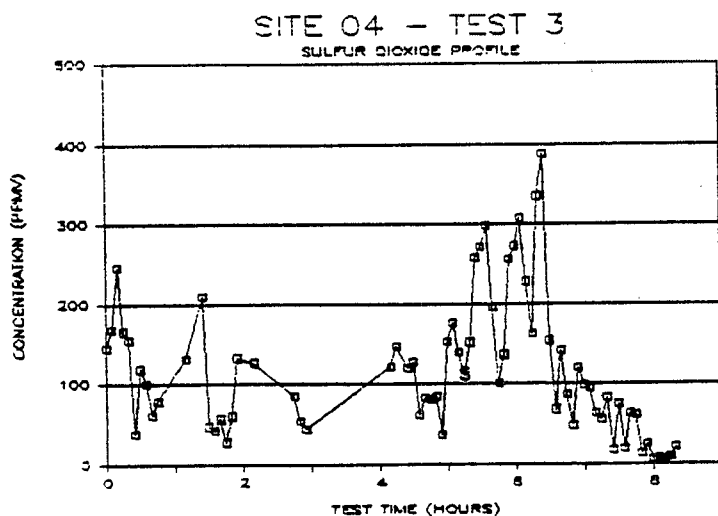
mean = 112.1 ppmv SO_2
@ 3% O_2

std dev = 102.1 ppmv SO_2
@ 3% O_2



mean = 54.1 ppmv SO_2
@ 3% O_2

std dev = 43.0 ppmv SO_2
@ 3% O_2



mean = 117.0 ppmv SO_2
@ 3% O_2

std dev = 85.2 ppmv SO_2
@ 3% O_2

Figure 5-9 Sulfur Dioxide Concentration History at the
Electrostatic Precipitator Inlet Location
(dry ppmv @ 3% O_2)

TABLE 5-6. OVERVIEW OF DIOXIN/FURAN EMISSIONS CONCENTRATION DATA
FOR SITE BLB-A (ELECTROSTATIC PRECIPITATOR INLET)

Run Number	2378 TCDD	Total PCDD	Total PCDF
<u>Emissions Concentration</u> (as measured), ng/dscm			
Run 01	ND	1.21	1.31
Run 02	ND	0.93	1.03
Run 03	ND	2.61	1.58
Average	--	1.59	1.31
<u>Emissions Concentration</u> (corrected to 3% O ₂), ng/dscm @ 3% O ₂			
Run 01	ND	1.37	1.47
Run 02	ND	1.05	1.16
Run 03	ND	2.94	1.77
Average	--	1.79	1.47

ND = not detected. See Tables 5-8 and 5-9 for detection limits.

TABLE 5-7. SUMMARY OF DIOXIN AND FURAN EMISSIONS RATE DATA
FOR SITE BLB-A (ELECTROSTATIC PRECIPITATOR INLET)

Run Number	Dioxin/Furan Emission Rate, ug/hr		
	2378 TCDD	Total PCDD	Total PCDF
Run 01	ND	280	302
Run 02	ND	206	228
Run 03	ND	530	320
Average	--	339	283

ND = not detected. See Table 5-8 for detection limits.

total PCDF. ~~Emissions of 2378 TCDD~~ ^{was not detectable} ~~were consistently below the detection limit.~~ The total PCDD and PCDF emissions were fairly consistent between runs. The maximum deviation of any individual run from the overall average was approximately 56 percent for total PCDD emissions and 19 percent for total PCDF emissions.

Isomer and homologue-specific emission concentration data are summarized in Table 5-8 and 5-9 for the three test runs. Run-specific data tables showing homologue emission concentrations in both ng/dscm and parts-per-trillion units and homologue emission rates in ug/hr units are included in Appendix D. Detectable quantities were found for roughly two-thirds of the isomers and homologues analyzed for at the ESP inlet of Site BLB-A. Figure 5-10 is a histogram that shows the relative distributions of the 2378 TCDD/TCDF isomers and the tetra- through octa-PCDD/PCDF homologues in the inlet ESP emissions (mole basis). The distribution of dioxin species was extremely non-uniform among the various homologues. Only the hepta- and octa-CDD homologues were consistently detectable in the ESP inlet emissions. The octa-CDD homologue accounted for roughly 80 percent of the total dioxins detected. The hepta-CDD homologues accounted for the remaining 20 percent of the total dioxins in the ESP inlet emissions. The furan species were more uniformly distributed than the dioxin species, with the hepta-CDF homologue being the largest single contributor to the total PCDF emissions. The contributions of the tetra- through octa-chlorinated furan homologues to the total PCDF were: tetra, 19-26%; penta, 0-24%; hexa, 33-50%; hepta, 12-24%; and octa, 0-8%.

Emission factors for the ESP inlet at site BLB-A are shown in Table 5-10. Average emission factors for total PCDD and total PCDF were 0.006 ug total PCDD emitted per Kg feed and 0.005 ug total PCDF emitted per Kg feed (dry solids feed basis). Emission factors for the various dioxin and furan homologues varied considerably between runs.

TABLE 5-8. SUMMARY OF DIOXIN/FURAN EMISSIONS DATA FOR SITE BLB-A INLET
(AS-MEASURED CONCENTRATIONS)

Dioxin/Furan Isomer	Isomer Concentration in Flue Gas (ng/dscm)			
	Run 01	Run 02	Run 03	Avg.
DIOXINS				
2378 TCDD	ND(1.94E-02)	ND(1.98E-02)	ND(7.47E-02)	.00E+00
Other TCDD	ND(1.94E-02)	1.98E-02	ND(7.47E-02)	6.61E-03
Penta-CDD	ND(7.52E-02)	ND(2.38E-02)	ND(9.96E-02)	.00E+00
Hexa-CDD	1.21E-01	7.94E-02	ND(3.90E-01)	6.69E-02
Hepta-CDD	2.91E-01	1.98E-01	5.81E-01	3.57E-01
Octa-CDD	8.01E-01	6.35E-01	2.03E+00	1.16E+00
Total PCDD	1.21E+00	9.33E-01	2.61E+00	1.59E+00
FURANS				
2378 TCDF	ND(1.46E-02)	ND(7.94E-02)	ND(1.66E-01)	.00E+00
Other TCDF	2.43E-01	1.98E-01	4.15E-01	2.85E-01
Penta-CDF	3.16E-01	2.38E-01	ND(6.14E-01)	1.85E-01
Hexa-CDF	4.37E-01	3.97E-01	7.88E-01	5.41E-01
Hepta-CDF	2.43E-01	1.19E-01	3.73E-01	2.45E-01
Octa-CDF	7.28E-02	7.94E-02	ND(2.61E-01)	5.07E-02
Total PCDF	1.31E+00	1.03E+00	1.58E+00	1.31E+00

NOTE: Isomer concentrations shown are at as-measured oxygen conditions.

ND = not detected (detection limit in parentheses).

ng = 1.0E-09g

8760 operating hours per year

TABLE 5-9. SUMMARY OF DIOXIN/FURAN EMISSIONS DATA FOR SITE BLB-A INLET
(CONCENTRATIONS CORRECTED TO 3 PERCENT OXYGEN)

Dioxin/Furan Isomer	Isomer Concentration in Flue Gas (ng/dscm @ 3% oxygen)			
	Run 01	Run 02	Run 03	Avg.
DIOXINS				
2378 TCDD	ND(2.18E-02)	ND(2.23E-02)	ND(8.40E-02)	.00E+00
Other TCDD	ND(2.18E-02)	2.23E-02	ND(8.40E-02)	7.44E-03
Penta-CDD	ND(8.46E-02)	ND(2.68E-02)	ND(1.12E-01)	.00E+00
Hexa-CDD	1.37E-01	8.93E-02	ND(4.39E-01)	7.53E-02
Hepta-CDD	3.28E-01	2.23E-01	6.54E-01	4.01E-01
Octa-CDD	9.01E-01	7.14E-01	2.29E+00	1.30E+00
Total PCDD	1.37E+00	1.05E+00	2.94E+00	1.79E+00
FURANS				
2378 TCDF	ND(1.64E-02)	ND(8.93E-02)	ND(1.87E-01)	.00E+00
Other TCDF	2.73E-01	2.23E-01	4.67E-01	3.21E-01
Penta-CDF	3.55E-01	2.68E-01	ND(6.91E-01)	2.08E-01
Hexa-CDF	4.92E-01	4.46E-01	8.87E-01	6.08E-01
Hepta-CDF	2.73E-01	1.34E-01	4.20E-01	2.76E-01
Octa-CDF	8.19E-02	8.93E-02	ND(2.94E-01)	5.71E-02
Total PCDF	1.47E+00	1.16E+00	1.77E+00	1.47E+00

NOTE: Isomer concentrations shown are corrected to 3% oxygen.

ND = not detected (detection limit in parentheses).

ng = 1.0E-09g

8760 operating hours per year

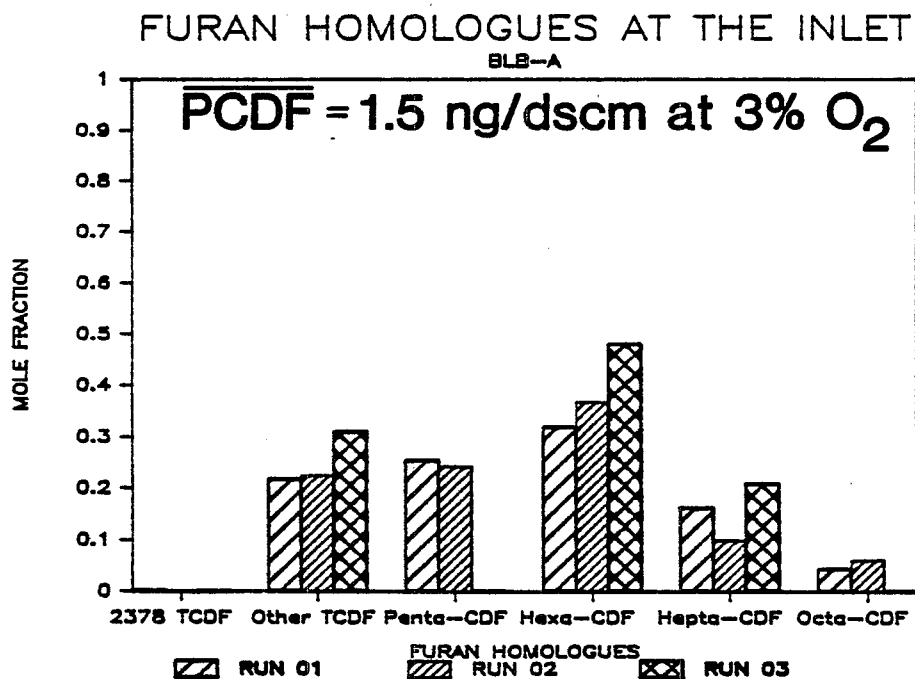
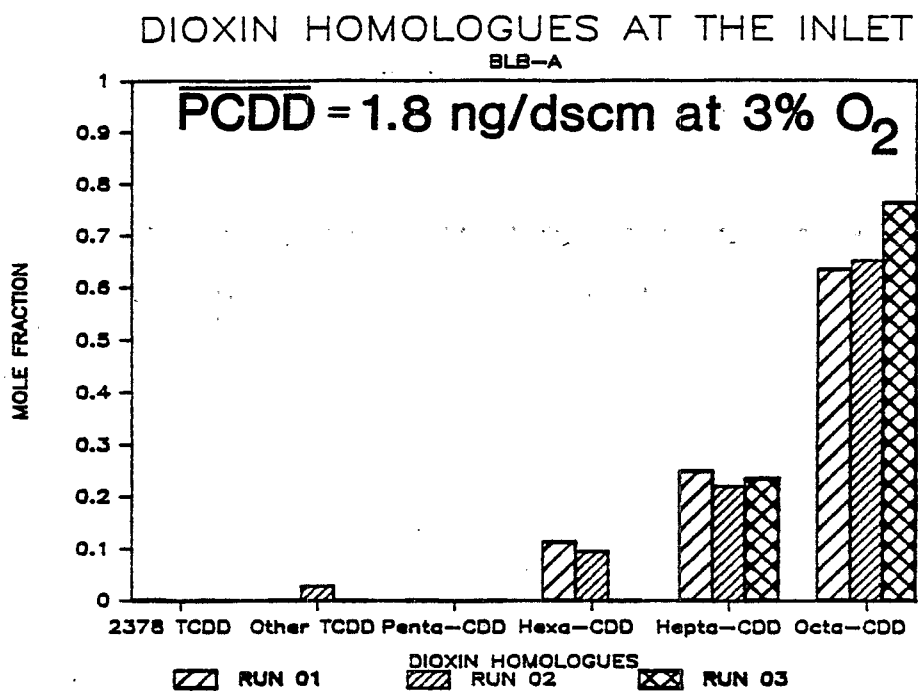


Figure 5-10. Dioxin and furan homologue distributions of the electrostatic precipitator inlet emissions for Site BLB-A.

TABLE 5-10. DIOXIN/FURAN EMISSION FACTORS FOR SITE BLB-A INLET

Dioxin/Furan Isomer	Dioxin/Furan Emission Factors (ug/kg)			
	Run 01	Run 02	Run 03	Avg.
DIOXINS				
2378 TCDD	ND(8.28E-05)	ND(8.18E-05)	ND(2.80E-04)	.00E+00
Other TCDD	ND(8.28E-05)	8.18E-05	ND(2.80E-04)	2.73E-05
Penta-CDD	ND(3.21E-04)	ND(9.81E-05)	ND(3.73E-04)	.00E+00
Hexa-CDD	5.18E-04	3.27E-04	ND(1.46E-03)	2.82E-04
Hepta-CDD	1.24E-03	8.18E-04	2.18E-03	1.41E-03
Octa-CDD	3.42E-03	2.62E-03	7.62E-03	4.55E-03
Total PCDD	5.18E-03	3.84E-03	9.80E-03	6.27E-03
FURANS				
2378 TCDF	ND(6.21E-05)	ND(3.27E-04)	ND(6.22E-04)	.00E+00
Other TCDF	1.04E-03	8.18E-04	1.55E-03	1.14E-03
Penta-CDF	1.35E-03	9.81E-04	ND(2.30E-03)	7.76E-04
Hexa-CDF	1.86E-03	1.64E-03	2.95E-03	2.15E-03
Hepta-CDF	1.04E-03	4.91E-04	1.40E-03	9.75E-04
Octa-CDF	3.11E-04	3.27E-04	ND(9.80E-04)	2.13E-04
Total PCDF	5.59E-03	4.25E-03	5.91E-03	5.25E-03

ND = not detected (detection limit in parentheses).

ug = 1.0E-06g

8760 operating hours per year

NOTE: Emission factors were calculated using the black liquor dry solids feed rate.

5.4.2 Electrostatic Precipitator Outlet

Emission concentrations and emissions rate data measured at the electrostatic precipitator (ESP) outlet are shown in Tables 5-11 and 5-12 for the 2378 TCDD, total PCDD, and total PCDF species. The data include dioxin and furan collected in the entire MM5 train, including filter, primary XAD sorbent trap, impingers, and sample train clean-up rinses.

Average as-measured outlet emission concentrations of total PCDD and PCDF were 0.67 ng/dscm total PCDD and 0.51 ng/dscm total PCDF. When corrected to 3% O₂ using the EPA Method 3 oxygen concentration data, these values correspond to 0.75 ng/dscm @ 3% O₂ and 0.57 ng/dscm @ 3% O₂, respectively. Average emission rates were 150 ug/hr total PCDD, and 110 ug/hr total PCDF. Emissions of 2378 TCDD were non-detectable for all three runs. The total PCDD and PCDF outlet emissions showed considerable variability between runs. The maximum deviation of any individual run from the overall average was approximately 80 percent for total PCDD and PCDF emissions.

Isomer and homologue specific outlet emission concentration data are summarized in Tables 5-13 and 5-14 for the three test runs. Run-specific data tables showing homologue emission concentrations in ng/dscm and parts-per-trillion units and homologue emission rates in ug/hr units are included in Appendix D. Detectable quantities were found for roughly half of the isomers and homologues analyzed for at the ESP outlet of Site BLB-A. Figure 5-11 is a histogram that shows the relative distributions of the 2378 TCDD/TCDF isomers and the tetra- through octa- PCDD/PCDF homologues in the ESP outlet emissions (mole basis). The distribution of dioxin species was non-uniform among the various homologues. Only the hepta and octa-CDD homologues were consistently detectable in the ESP outlet emissions. The octa-CDD homologue accounted for roughly 70 percent of the total dioxins found while the hexa-CDD isomer accounted for most of the remaining PCDD. The total PCDF emissions were more evenly distributed than the dioxin species with the contributions of the tetra- through octa-chlorinated furan homologues to the total PCDF being as follows: tetra, 10-33%; penta, 0-11%; hexa, 0-15%; hepta, 22-50%; and octa, 9-50%.

Emission factors for the ESP outlet at site BLB-A are shown in Table 5-15. Average outlet emission factors for total PCDD and total PCDF

TABLE 5-11. OVERVIEW OF DIOXIN AND FURAN EMISSIONS CONCENTRATION DATA
FOR SITE BLB-A (ELECTROSTATIC PRECIPITATOR OUTLET)

Run Number	2378 TCDD	Total PCDD	Total PCDF
<u>Emissions Concentration</u> (as measured), ng/dscm			
Run 01	ND	1.18	0.89
Run 02	ND	0.51	0.52
Run 03	ND	0.31	0.10
Average	--	0.67	0.51
<u>Emissions Rate Concentration</u> (corrected to 3% O ₂), ng/dscm @ 3% O ₂			
Run 01	ND	1.33	1.00
Run 02	ND	0.59	0.61
Run 03	ND	0.33	0.11
Average	--	0.75	0.57

TABLE 5-12. SUMMARY OF DIOXIN AND FURAN EMISSIONS RATE DATA FOR
SITE BLB-A (ELECTROSTATIC PRECIPITATOR OUTLET)

Run Number	Dioxin/Furan Emission Rate, ug/hr		
	2378 TCDD	Total PCDD	Total PCDF
Run 01	ND	268	202
Run 02	ND	113	117
Run 03	ND	68	22
Average	--	150	114

TABLE 5-13. SUMMARY OF DIOXIN/FURAN EMISSIONS DATA FOR SITE BLB-A OUTLET
(AS-MEASURED CONCENTRATIONS)

Dioxin/Furan Isomer	Isomer Concentration in Flue Gas (ng/dscm)			
	Run 01	Run 02	Run 03	Avg.
DIOXINS				
2378 TCDD	ND(6.00E-02)	ND(1.69E-02)	ND(1.88E-02)	.00E+00
Other TCDD	ND(6.00E-02)	5.06E-02	ND(1.88E-02)	1.69E-02
Penta-CDD	ND(2.51E-02)	ND(2.02E-02)	ND(3.77E-02)	.00E+00
Hexa-CDD	1.55E-01	ND(5.40E-02)	ND(4.11E-02)	5.16E-02
Hepta-CDD	3.09E-01	8.43E-02	8.56E-02	1.60E-01
Octa-CDD	7.16E-01	3.71E-01	2.23E-01	4.36E-01
Total PCDD	1.18E+00	5.06E-01	3.08E-01	6.65E-01
FURANS				
2378 TCDF	ND(7.74E-03)	ND(3.37E-02)	ND(1.71E-02)	.00E+00
Other TCDF	3.29E-01	5.06E-02	3.42E-02	1.38E-01
Penta-CDF	1.55E-01	ND(3.88E-02)	ND(2.23E-02)	5.16E-02
Hexa-CDF	1.35E-01	5.06E-02	ND(8.73E-02)	6.20E-02
Hepta-CDF	1.93E-01	1.69E-01	5.14E-02	1.38E-01
Octa-CDF	7.74E-02	2.53E-01	1.71E-02	1.16E-01
Total PCDF	8.90E-01	5.23E-01	1.03E-01	5.05E-01

NOTE: Isomer concentrations shown are at as-measured oxygen conditions.

ND = not detected (detection limit in parentheses).

ng = 1.0E-09g

8760 operating hours per year

TABLE 5-14. SUMMARY OF DIOXIN/FURAN EMISSIONS DATA FOR SITE BLB-A OUTLET
(CONCENTRATIONS CORRECTED TO 3 PERCENT OXYGEN)

Dioxin/Furan Isomer	Isomer Concentration in Flue Gas (ng/dscm @ 3% oxygen)			
	Run 01	Run 02	Run 03	Avg.
DIOXINS				
2378 TCDD	ND(6.75E-02)	ND(1.97E-02)	ND(2.03E-02)	.00E+00
Other TCDD	ND(6.75E-02)	5.91E-02	ND(2.03E-02)	1.97E-02
Penta-CDD	ND(2.83E-02)	ND(2.37E-02)	ND(4.05E-02)	.00E+00
Hexa-CDD	1.74E-01	ND(6.31E-02)	ND(4.42E-02)	5.80E-02
Hepta-CDD	3.48E-01	9.86E-02	9.21E-02	1.80E-01
Octa-CDD	8.05E-01	4.34E-01	2.40E-01	4.93E-01
Total PCDD	1.33E+00	5.91E-01	3.32E-01	7.50E-01
FURANS				
2378 TCDF	ND(8.70E-03)	ND(3.94E-02)	ND(1.84E-02)	.00E+00
Other TCDF	3.70E-01	5.91E-02	3.68E-02	1.55E-01
Penta-CDF	1.74E-01	ND(4.53E-02)	ND(2.40E-02)	5.80E-02
Hexa-CDF	1.52E-01	5.91E-02	ND(9.40E-02)	7.05E-02
Hepta-CDF	2.18E-01	1.97E-01	5.53E-02	1.57E-01
Octa-CDF	8.70E-02	2.96E-01	1.84E-02	1.34E-01
Total PCDF	1.00E+00	6.11E-01	1.11E-01	5.74E-01

NOTE: Isomer concentrations shown are corrected to 3% oxygen.

ND = not detected (detection limit in parentheses).

ng = 1.0E-09g

8760 operating hours per year

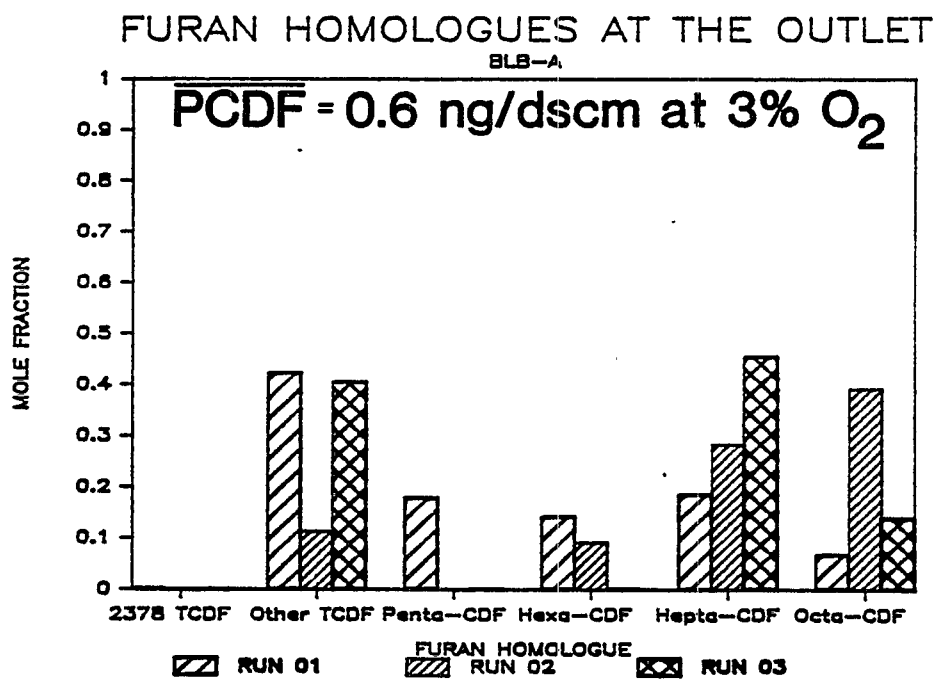
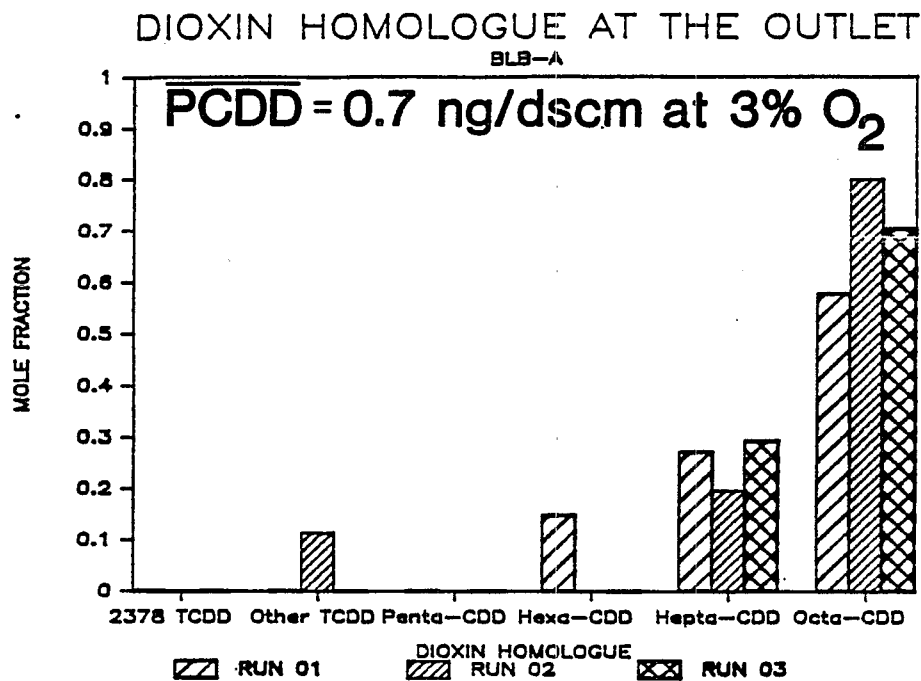


Figure 5-11. Dioxin and furan homologue distributions of the electrostatic precipitator outlet emissions for Site BLB-A.

TABLE 5-15. DIOXIN/FURAN EMISSION FACTORS FOR SITE BLB-A OUTLET

Dioxin/Furan Isomer	Dioxin/Furan Emission Factors (ug/kg)			
	Run 01	Run 02	Run 03	Avg.
DIOXINS				
2378 TCDD	ND(2.52E-04)	ND(7.06E-05)	ND(7.54E-05)	.00E+00
Other TCDD	ND(2.52E-04)	2.12E-04	ND(7.54E-05)	7.06E-05
Penta-CDD	ND(1.06E-04)	ND(8.47E-05)	ND(1.51E-04)	.00E+00
Hexa-CDD	6.51E-04	ND(2.26E-04)	ND(1.64E-04)	2.17E-04
Hepta-CDD	1.30E-03	3.53E-04	3.43E-04	6.66E-04
Octa-CDD	3.01E-03	1.55E-03	8.91E-04	1.82E-03
Total PCDD	4.96E-03	2.12E-03	1.23E-03	2.77E-03
FURANS				
2378 TCDF	ND(3.26E-05)	ND(1.41E-04)	ND(6.85E-05)	.00E+00
Other TCDF	1.38E-03	2.12E-04	1.37E-04	5.77E-04
Penta-CDF	6.51E-04	ND(1.62E-04)	ND(8.91E-05)	2.17E-04
Hexa-CDF	5.70E-04	2.12E-04	ND(3.49E-04)	2.60E-04
Hepta-CDF	8.14E-04	7.06E-04	2.06E-04	5.75E-04
Octa-CDF	3.26E-04	1.06E-03	6.85E-05	4.84E-04
Total PCDF	3.74E-03	2.19E-03	4.11E-04	2.11E-03

ND = not detected (detection limit in parentheses).

ug = 1.0E-06g

8760 operating hours per year

NOTE: Emission factors were calculated using the black liquor dry solids feed rate.

were 0.003 ug total PCDD emitted per Kg feed and 0.002 ug total PCDF emitted per Kg feed (dry solids feed basis). The outlet emission factors for the individual dioxin and furan homologues varied considerably between test runs.

5.4.3 Reduction of Dioxin/Furan Concentrations Due to the ESP.

The dioxin/furans which condense on particulate in the stack gas are removed from the stack gas along with the particulate matter by the pollution control device. The dioxin/furan removal efficiency of the control device is calculated from the difference of the inlet and outlet concentration of each dioxin/furan homologue divided by the inlet concentration of each homologue. Each concentration value may have an analytical uncertainty of $\pm 50\%$. An analysis of the uncertainty of the control device efficiency (contained in Appendix F) indicates that with a measured efficiency of greater than 67%, the true removal efficiency is most likely positive. With measured efficiencies between 67% and -200%, a definite conclusion cannot be drawn concerning the true removal efficiency, and below -200%, the true removal efficiency is most likely negative.

The measured ESP removal efficiencies for each dioxin/furan homologue at Site BLB-A are summarized in Table 5-16. The PCDD/PCDF removal efficiencies across the ESP were widely scattered for the respective homologues and between runs. Run 03 indicates positive removal efficiency across the ESP while no conclusions can be made from the efficiencies calculated for Runs 01 and 02. The overall data is inconclusive concerning the true removal efficiency for the ESP.

5.4.4 Economizer Ash Results

During the pre-survey site visit a single economizer ash sample was collected. This sample was analyzed for dioxin/furan to determine if it was necessary to sample the economizer ash during the actual site testing. Hexa-CDF and octa-CDD were the only homologues detected in concentrations of 0.01 ng/g (ppb) for each respective homologue. The detection limits for the homologues which were not detected were between 0.01 ng/g (ppb) and 0.02 ng/g (ppb). Due to the minimal concentrations of the dioxin/furan homologues found in this sample, the economizer ash was not sampled during the site testing.

TABLE 5-16. MEASURED ESP REMOVAL EFFICIENCIES AT SITE BLB-A^a

Homologue	ESP Removal Efficiency, (%)		
	Run 1	Run 2	Run 3
Dioxins			
2378 TCDD	--	--	--
Other TCDD	--	- 165	--
Penta-CDD	--	--	--
Hexa-CDD	-27.0	+ 100	--
Hepta-CDD	- 6.1	+55.8	+85.9
Octa-CDD	+10.7	+39.2	+89.5
Total PCDD	+ 2.9	+43.7	+88.7
Furans			
2378 TCDF	--	--	--
Other TCDF	-35.5	+73.5	+92.1
Penta-CDF	+51.0	+ 100	--
Hexa-CDF	+69.1	+86.7	+ 100
Hepta-CDF	+20.1	-47.0	+98.0
Octa-CDF	- 6.2	- 232	--
Total PCDF	+32.0	+47.3	+93.7

NOTE: dash (-) indicates that the species was not detected at the ESP inlet location.

^aESP removal efficiency values were calculated using concentration data corrected to 3% O₂ (Table 5-9 and 5-14).

5.5 BLACK LIQUOR PRECURSOR DATA

As discussed in Section 4.3.2, the strong black liquor was sampled at Site BLB-A. These samples were analyzed for chlorinated benzenes, chlorinated biphenyls, and chlorinated phenols.

Table 5-17 summarizes the results of the precursor analyses. Trace levels of pentachlorophenols were detected for Run 02, but overall the specific precursors analyzed for (i.e., chlorobenzenes, chlorophenols, and chlorinated biphenyls) were not detected. This suggests that either (i) the specific precursors analyzed for were not present in the samples, or (ii) the precursors were not easily detected using the GC/MS procedure.

Table 5-18 presents the results of the strong black liquor and weak black liquor total chloride analyses. The chloride concentration was consistent among the three test runs. The greatest deviation of any run from the overall average was 36 percent for the strong black liquor and 27 percent for the weak black liquor. As seen from Table 5-18, the strong black liquor was roughly 2.5 times as concentrated as the weak black liquor with respect to total chloride content. This increase in chloride content follows from the process since the dirty water from the brown stock washers (i.e., weak black liquor) is sent to an evaporation system for solids concentration prior to being fired in the boiler.

5.6 AUXILIARY PROCESS SAMPLE ANALYSES

In addition to the chloride analysis performed on the black liquor samples, chloride analysis was also performed on white liquor, by-product saltcake, and caustic samples collected at Site BLB-A. The results of the strong and weak black liquor chloride analyses are presented above in Section 5.5. Table 5-19 summarizes the data obtained from the total chloride analyses of the other process samples mentioned above. The majority of the chlorine entering the black liquor circuit comes from the caustic and the by-product salt cake with each contributing 122 and 7.7 ug chlorine/Mg of solids burned, respectively.

During the pre-survey plant visit a grab sample of the economizer ash was taken for dioxin/furan analysis. All of the homologues were below the detection limit for the method except for the octa-CDD and hexa-CDF

TABLE 5-17. SUMMARY OF DIOXIN PRECURSOR DATA FOR SITE BLB-A FEED SAMPLES

Precursor Categories	Precursor Concentration, ug/g (ppm)			
	Black Liquor Feed			Average
	Run 01	Run 02	Run 03	
Total Chlorinated Benzenes	ND	ND	ND	--
Total Chlorinated Biphenyls	ND	ND	ND	--
Total Chlorinated Phenols	ND	trace	ND	--

ND = not detected.

TABLE 5-18. TOTAL CHLORIDE ANALYSES OF THE BLACK LIQUOR SAMPLES FOR SITE BLB-A

Liquor	Run No.	Total Chloride Concentration (ug/g)
Strong Black Liquor	01	578
	02	951
	03	1159
	Average	896
Weak Black Liquor	01	320
	02	506
	03	369
	Average	398

TABLE 5-19. TOTAL CHLORIDE ANALYSES OF AUXILIARY
LIQUOR CIRCUIT SAMPLES AT SITE BLB-A
(ug/g as Cl⁻)

Test Run	Caustic	By-Product Salt Cake	White Liquor
01	2900	1300	1600
02	600	120	1300
03	320	NS	1700
Average	1300	700	1500

NS = Not sampled. By-product salt cake was not being produced during Run 03.

homologues. Ten parts per trillion of each of these homologues were detected in the economizer ash sample. The detection limits for the ash analysis ranged from 5 to 20 parts per trillion. The doixin/furan contamination in the economizer ash was obviously minimal.

5.7 HCl TRAIN CHLORIDE EMISSIONS DATA

Table 5-20 summarizes HCl train chloride emissions data measured at the electrostatic precipitator outlet sampling location. The data are reported as "front-half," "back-half," and "train-total" chloride emissions. The front-half emissions represent chlorides captured in the probe rinse/filter fraction of the HCl train, which may include metal chlorides contained in the particulate matter. The back-half emissions represent chlorides captured in the HCl sample train impingers, which would include HCl and any metal chlorides that pass through the sample train filter. The train-total emissions represent the sum of the front-half and back-half emissions.

As shown in Table 5-20, the average as-measured train-total chloride emissions concentration was approximately 98 mg/dscm (1.5 grains/dscf). Corrected to 3% O₂ using the Radian CEM data, this corresponds to approximately 110 mg/dscm @ 3% O₂ (1.7 gr/dscf @ 3% O₂). The average train total chloride mass emission rate from the ESP outlet exhaust stack was about 22 Kg/hr (0.97 lb/hr). Chloride emissions were predominantly captured in the back half of the HCl sample train.

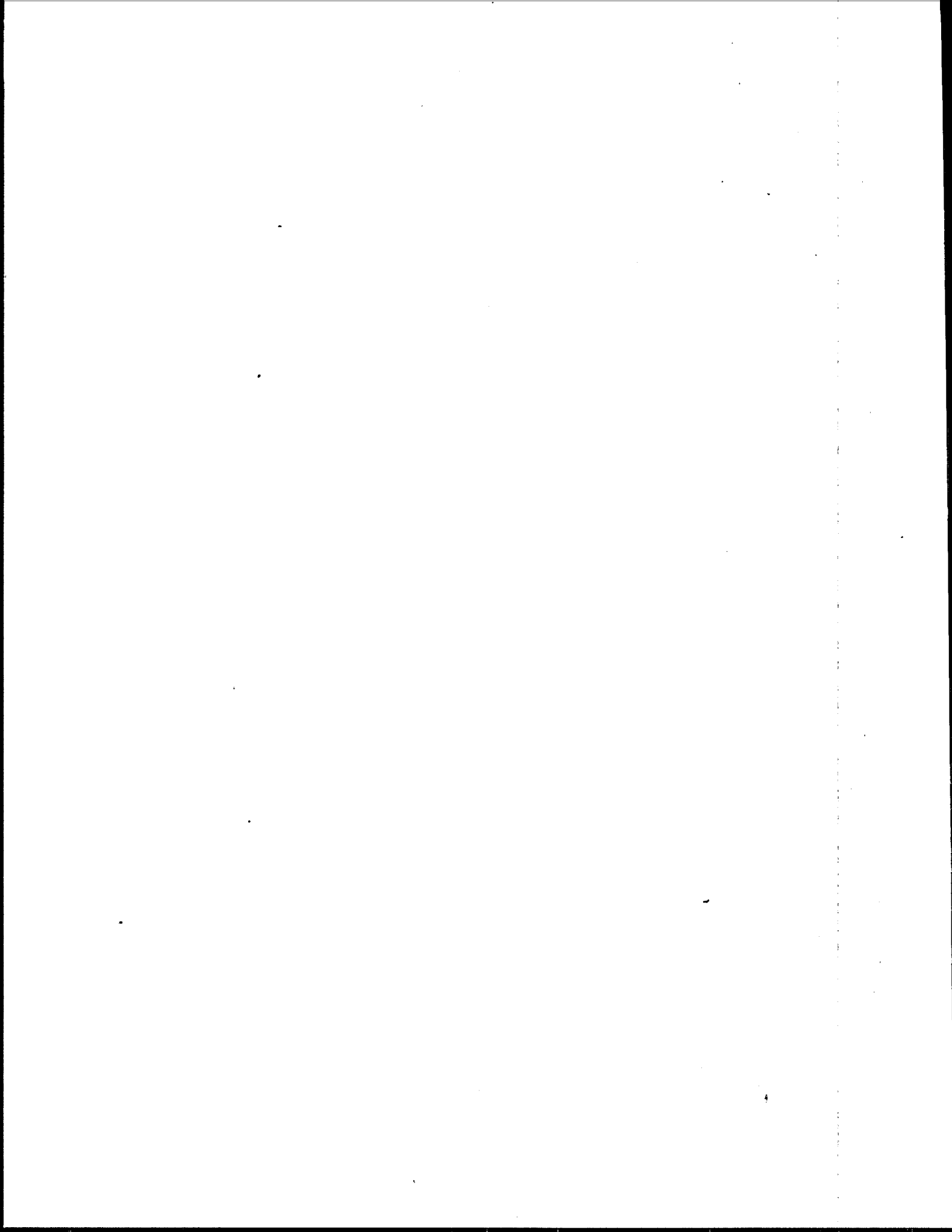
TABLE 5-20. HCl TRAIN CHLORIDE EMISSIONS DATA FOR SITE BLB-A
AT THE ELECTROSTATIC PRECIPITATOR OUTLET

Parameter	Run 01a	Run 01b	Run 02	Run 03	Average
Total Chloride Concentration (mg/dscm, as measured)					
Front Half	1.2	1.2	1.5	1.6	1.4
Back Half	108	91.2	95.3	91.6	96.7
Train Total	109	92.4	96.9	93.2	98.1
Total Chloride Concentration (mg/dscm, corrected to 3% Oxygen) ^a					
Front Half	1.4	1.3	1.7	1.8	1.5
Back Half	122	103	107	103	109
Train Total	124	104	109	105	110
Total Chloride Mass Emission Rate (Kg/hr)					
Front Half	0.3	0.3	0.3	0.4	0.3
Back Half	24.7	20.4	20.6	20.2	21.5
Train Total	25	20.7	20.9	20.5	21.8

^aConcentration corrected to 3% O₂ using the equation:

$$[Cl^-] @ 3\% O_2 = [Cl^-], \text{ as measured } \times (20.9 - 3)/(20.9 - \%O_2)$$

where: %O₂ = oxygen concentration in stack gas as measured by EPA Method 3
(See Table 5-5).



6.0 SAMPLING LOCATIONS AND PROCEDURES

Samples were collected from eight different locations at the test site. Two of the locations were for gaseous sampling, four were for liquid sampling, and two were for solid sampling. The source sampling and analysis matrix in Table 4-1 lists the sample locations, measured parameters, sampling methods, and analytical methods that were used.

Details on the sampling locations and methods are discussed in Section 6.1 through 6.3. Analytical procedures for continuous monitoring of CO, CO₂, O₂, NO_x, SO₂, and THC are included in Section 6.1. All other analytical procedures are discussed in Section 7.

6.1 GASEOUS SAMPLING

Four types of gaseous samples were taken during this test program: Modified Method 5 (MM5), HCl, EPA Method 3, and continuous monitoring (CEM). The sampling locations and methods are further discussed in this section.

6.1.1. Gaseous Sampling Locations

6.1.1.1 Electrostatic Precipitator Outlet Exhaust Stack.

The electrostatic precipitator (ESP) outlet exhaust stack sampling location is shown as point C in Figure 4-1. This location was used for dioxin sampling and HCl sampling using MM5 procedures described in Section 6.1.2.1 and 6.1.2.2. Gas velocity, molecular weight, and moisture were determined using EPA Methods 1 through 4. In addition, total reduced sulfur (TRS), CO, O₂, and opacity monitoring were performed by the plant at this location during the test periods.

Dimensions of the ESP outlet exhaust stack sampling location are shown in Figure 6-1. The diameter of the circular stack is 10.5 feet. Sampling ports were located approximately 4 duct diameters downstream of the connection between the ESP outlet ducting and the stack, and approximately 7.2 duct diameters upstream of the top of the stack. Based on EPA Method 1, a total of 24 traverse points were required for velocity determination at this location.

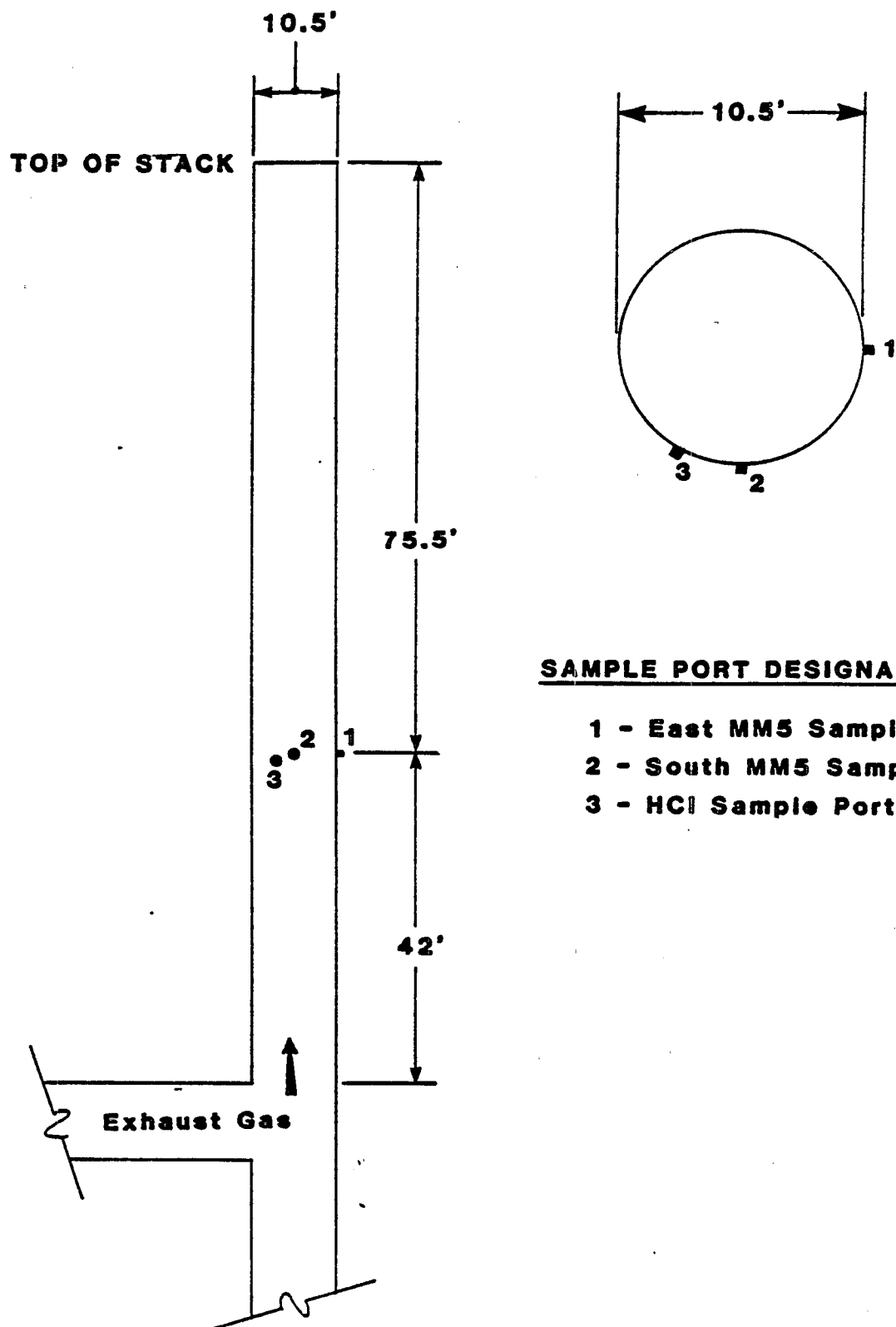


Figure 6-1 Dimensions of Electrostatic Precipitator Outlet Exhaust Stack

6.1.1.2 Black Liquor Boiler Outlet.

The black liquor boiler outlet sampling location (electrostatic precipitator inlet) is shown as point B in Figure 4-1. It consists of two identical rectangular ductwork sections that direct approximately equal volumetric flowrates of boiler exhaust gases to the East and West chambers of the ESP. Each of the two rectangular ductwork sections has four sampling ports. A schematic diagram of the ductwork is shown in Figure 6-2. This location was used for dioxin/furan sampling using MM5 procedures and for continuous monitoring of CO, CO₂, O₂, total hydrocarbons (THC), NO_x, and SO₂. Gas velocity, molecular weight, and moisture were determined using EPA Methods 1 through 4. The dimensions of the black liquor boiler outlet sampling locations relative to the nearest flow disturbances are shown in Figure 6-3. The sampling ports were located less than 1 duct diameter downstream of a 45° bend and less than 1 duct diameter upstream of the inlet to the ESP. Seven of the ports were used for MM5 sampling, and the eighth port was dedicated to the continuous monitoring probe. Based on EPA Method 1, a total of 49 traverse points were used for velocity determination at this location.

6.1.2 Gaseous Sampling Procedures

Gaseous sampling procedures used during the testing are listed in Table 6-1. These procedures are discussed in detail in the Tier 4 Quality Assurance Project Plan (QAPP). A brief description of each method and any necessary deviations from the procedures outlined in the QAPP are provided in the following section.

6.1.2.1 Modified Method 5 (MM5).

Gas sampling for dioxins and furans was conducted according to the October 1984 draft of the ASME chlorinated organic compound sampling protocol. Minor deviations from the ASME protocol are discussed later in this section. This sampling method is a modified version of EPA Method 5 that includes a solid sorbent module for trapping vapor phase organics. The only differences in the sampling protocol which were not discussed in the Tier 4 QAPP are:

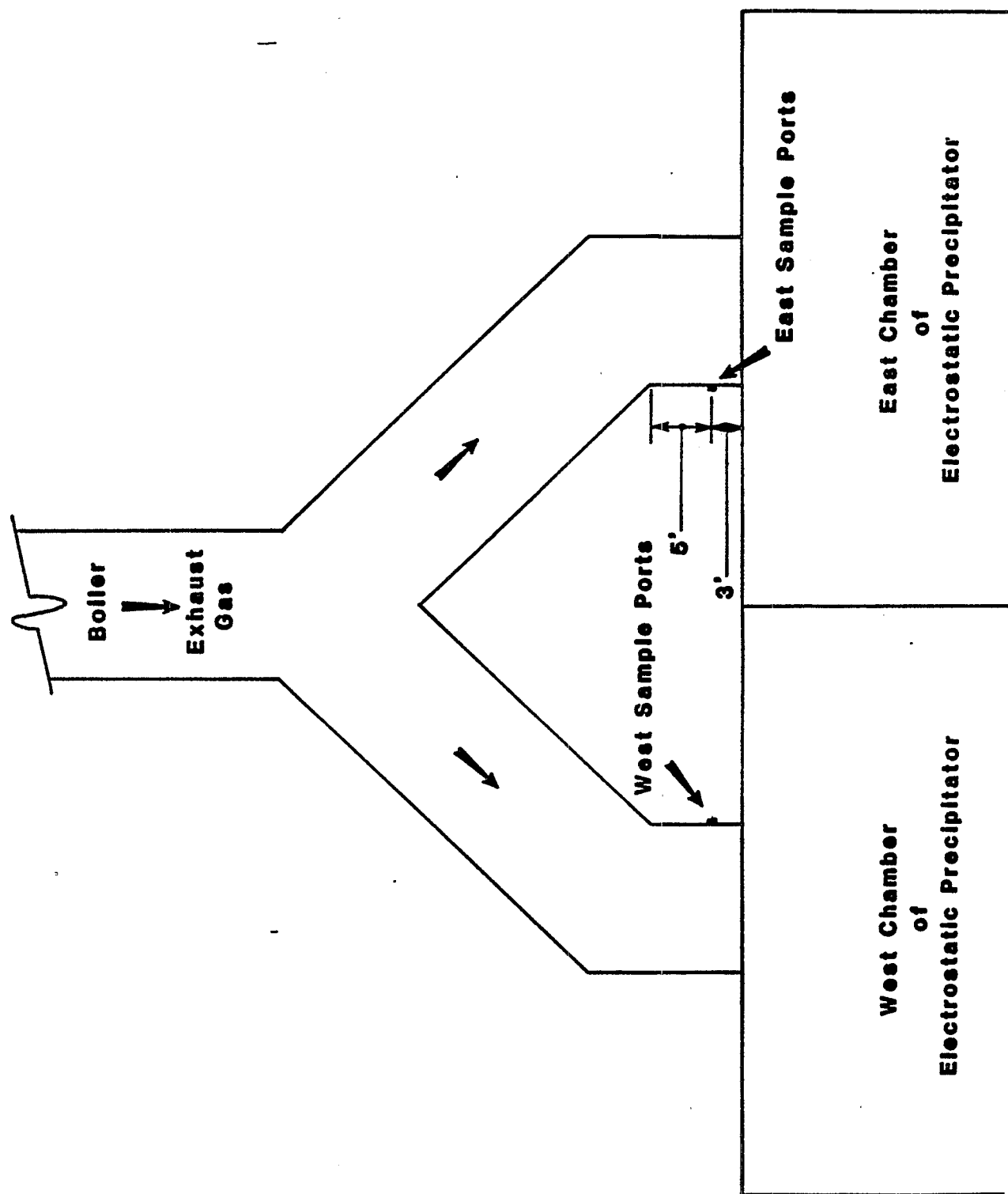


Figure 6-2 Schematic Diagram of Black Liquor Boiler Outlet Ductwork

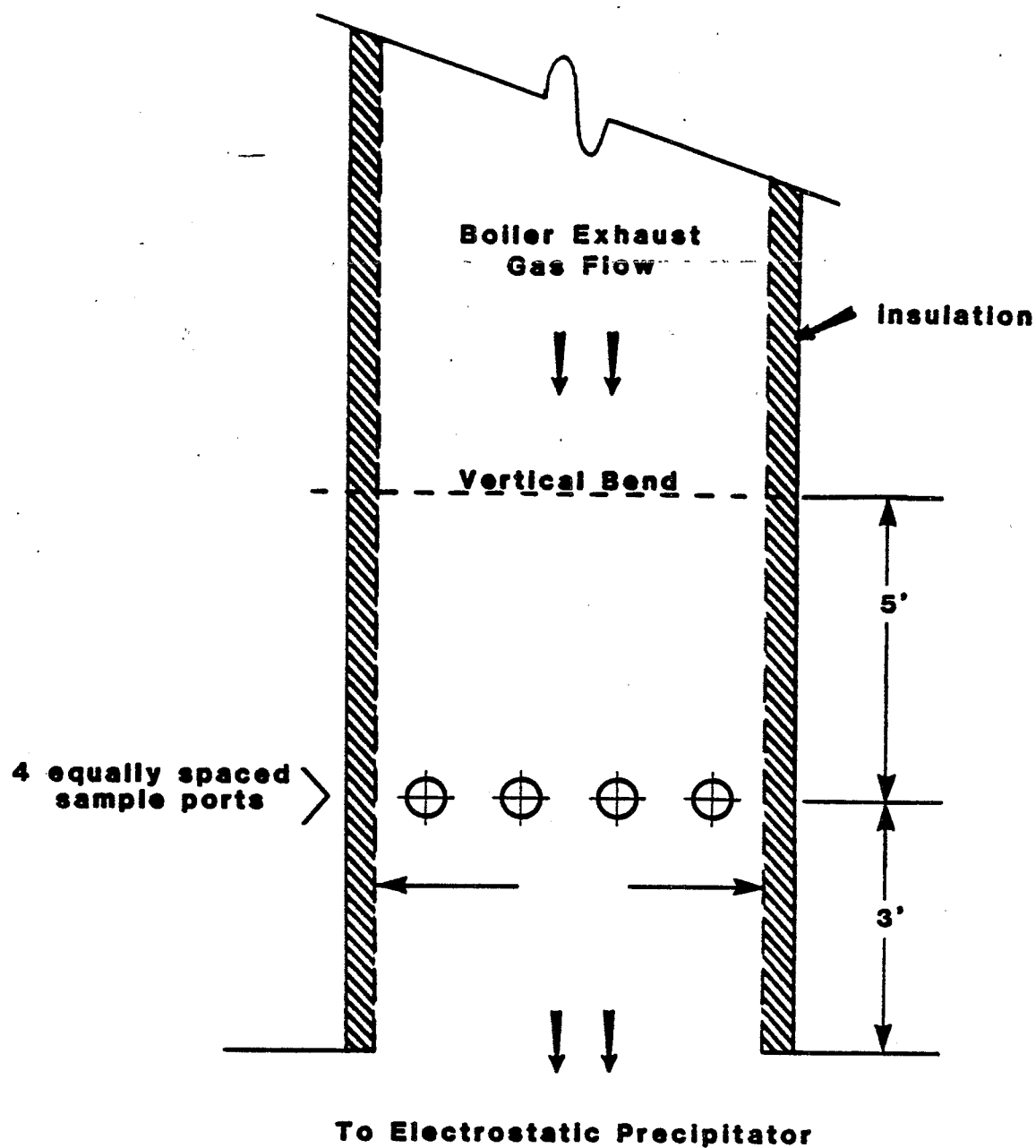


Figure 6-3 Dimensions of Black Liquor Boiler Outlet Sampling Location

TABLE 6-1. SUMMARY OF GAS SAMPLING METHODS FOR SITE 04

Sample Location	Sample Type or Parameter	Sample Collection Method
ESP outlet exhaust stack (Point C on Figure 4-1)	Dioxin	Modified EPA Method 5
	Volumetric Flow	EPA Method 2
	Molecular Weight	EPA Method 3
	Moisture	EPA Method 4
	HCl	HCl train
Black liquor boiler outlet (Point B on Figure 4-1)	Dioxin	Modified EPA Method 5
	Volumetric flow	EPA Method 2
	Molecular weight	EPA Method 3
	Moisture	EPA Method 4
	CO, CO ₂ , O ₂ , SO ₂ , NO _x and THC monitoring	Continuous monitors

- (1) Benzene was substituted for hexane or toluene as both the cleanup and extractant solvent for both the MM5 filters and the XAD-2 resin. This was caused by a discrepancy between the draft ASME sampling protocol and the draft ASME analytical protocol. (November 16, 1985)
- (2) Methylene chloride was substituted for hexane as the final field rinse solvent for the MM5 train. Methylene chloride was also substituted for hexane in the glassware cleaning procedure. This was caused by a high field blank train. (February 27, 1985)

The MM5 sampling train was used to collect samples at the ESP outlet exhaust stack and at the black liquor boiler outlet sampling location. Following sample recovery, the various parts of the sample (filter, solvent rinses, sorbent trap, etc.) were sent to the EPA's Troika laboratories to quantify 2378-TCDD, tetra- through octa-dioxin homologues, and tetra- through octa-furan homologues. A total of three MM5 test runs were conducted simultaneously at each of the two sampling locations, with one test run being conducted at each location per test day. The MM5 samples were collected isokinetically over a 240-minute on line sampling period at the ESP outlet with a sample flow rate of approximately 0.75 scfm. Sampling was performed during the same time period at the ESP inlet location, but the actual on line sampling period was approximately 140 minutes. Multiple filter changes were required at this location because of the high particulate loading and the "sticky" nature of the particulate. The sample flow rate at the ESP inlet was approximately 0.6 scfm.

A schematic diagram of the MM5 sampling train is shown in Figure 6-4. Flue gas is pulled from the stack through a nozzle and a heated glass probe. Particulate matter is removed from the gas stream by means of a glass fiber filter housed in a teflon-sealed glass filter holder maintained at $248 \pm 25^{\circ}\text{F}$. The gas passes through a sorbent trap similar to that illustrated in Figure 6-5 for removal of organic constituents. The trap consists of separate sections for (1) cooling the gas stream, and (2) adsorbing the organic compounds on Amberlite XAD-2^R resin (XAD). A chilled impinger train following the sorbent trap is used to remove water from the flue gas, and a dry gas meter is used to measure the sample gas flow.

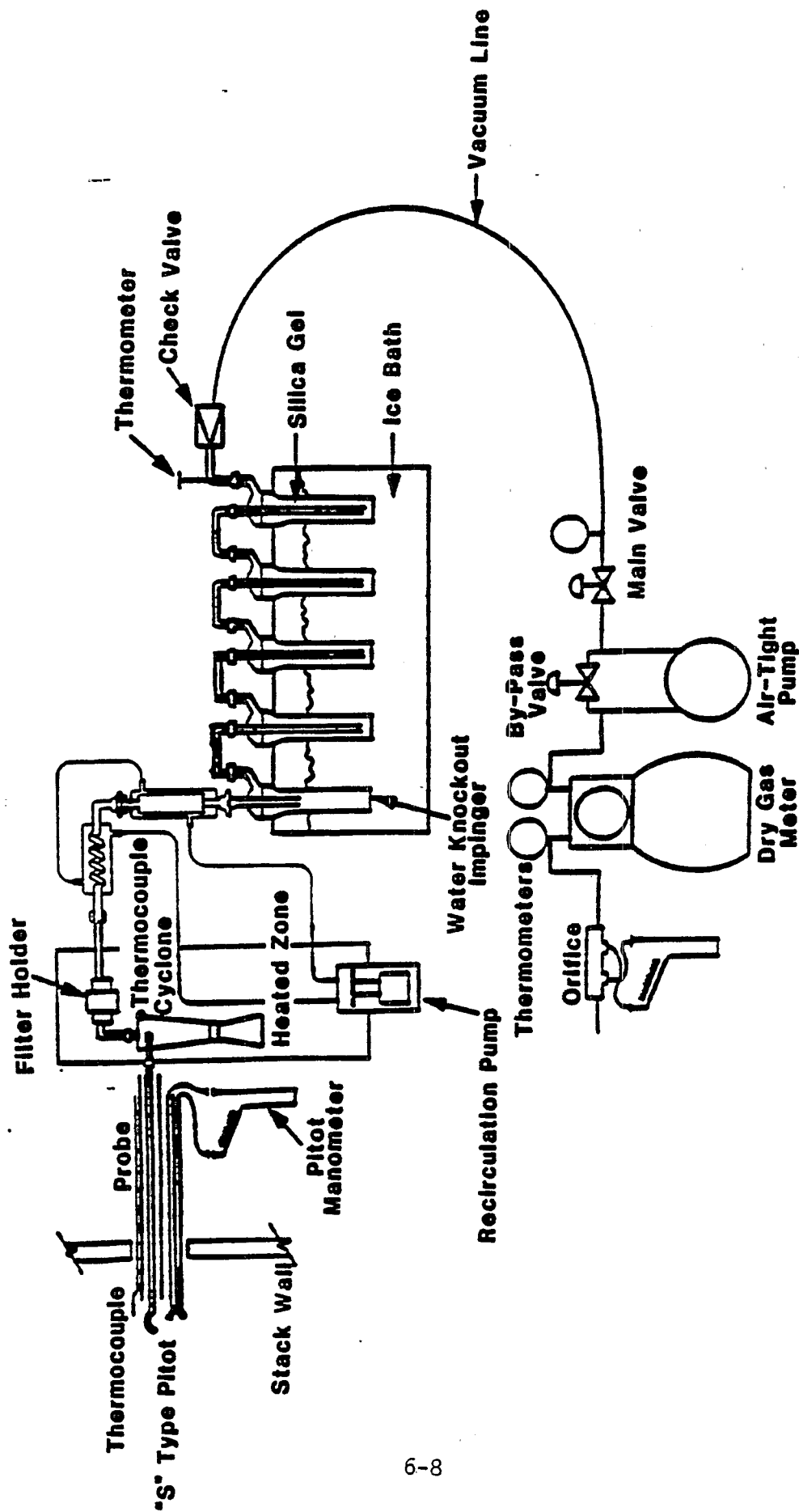


Figure 6-4 Modified Method 5 Train.

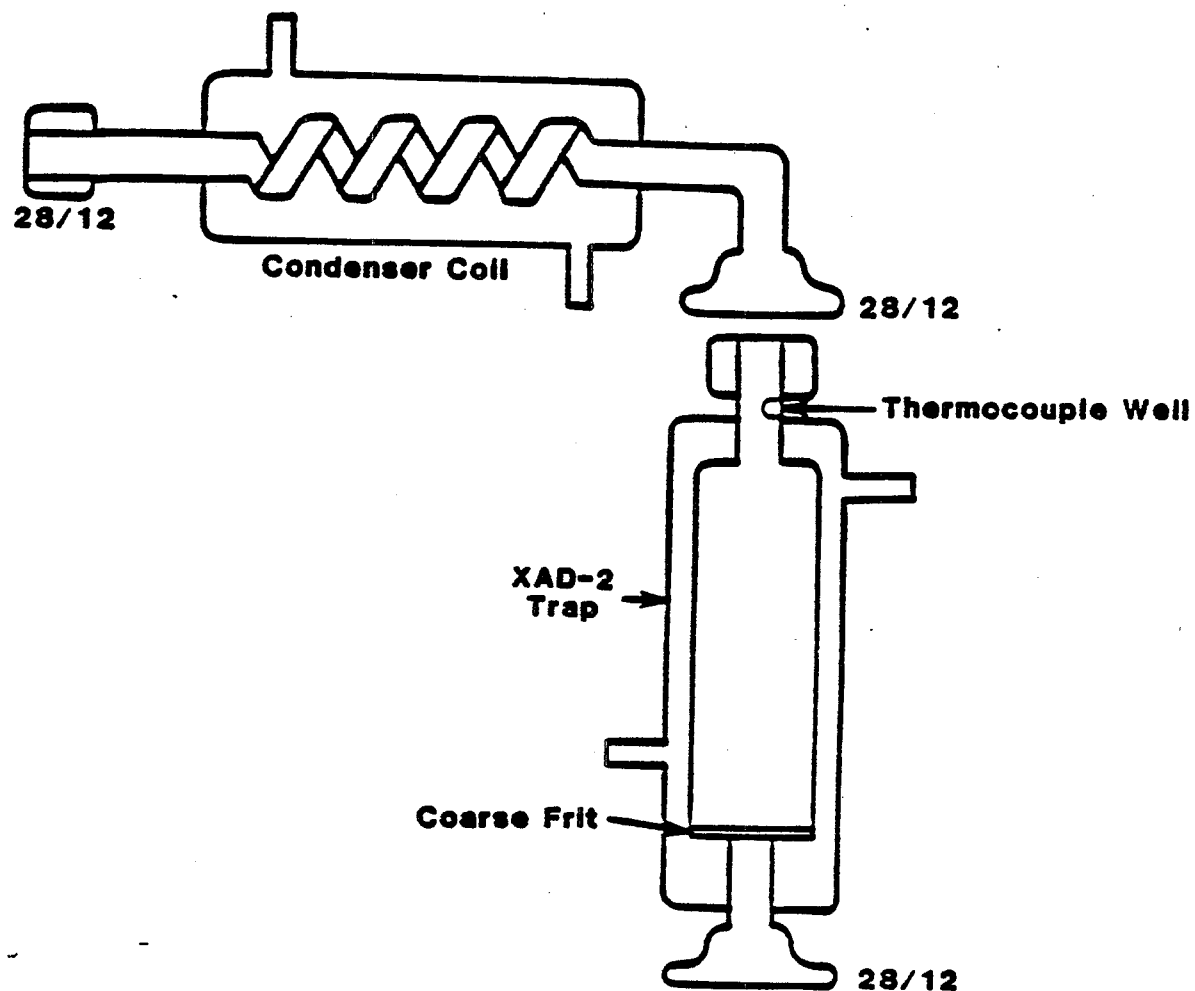


Figure 6-5 Adsorbent Sampling System.

Modifications to the ASME protocol that were instituted for this test site include the following:

1. Sample recovery was modified to include water in the sample train rinsing scheme. Water, acetone, and hexane were used in series to recover the probe, back-half/coil, and first impinger samples. Previous black liquor sampling experience has shown that water is necessary because the black liquor boiler particulate is soluble in water but insoluble in acetone.
2. The probe brush is specified in the ASME protocol as being inert material with a stainless steel handle. To ensure cleanliness, a separate nylon bristle brush attachable to a stainless steel handle was used for each probe cleaning.

6.1.2.2 HCl Determination.

HCl concentrations in the ESP outlet exhaust stack were determined using another modification of EPA Method 5. The sample train components and operation were identical to those of Method 5 with the following exceptions:

1. No knockout impinger was used.
2. During the first HCl train run, water in the first two impingers was replaced with 0.1 M NaOH. During the remaining HCl train (HCl runs 2 through 4), water in the first two impingers was replaced with 0.1M KOH. As discussed in Section 4.1, the substitution of KOH for NaOH in the impingers was performed at the suggestion of NCASI. The intent of the substitution was to eliminate NaCl interferences in the determination of HCl emissions by utilizing sodium, sulfate, and chloride material balances on the impinger solutions. Upon further study of this issue, it was determined that several additional chemical analysis would be needed to remove the potential NaCl interference. The uncertainty of the final result would be increased according to the uncertainty of each additional analysis. As a result, the plan to remove NaCl interferences was dropped and only the chloride analysis results were reported.
3. Sampling was single point isokinetic with the nozzle placed at points in the stack with approximate average velocity.
4. The moisture/NaOH or KOH in the impingers was saved for laboratory analysis by ion chromatography for total chlorides. The impinger catch was analyzed by Radian's Austin, Texas laboratory.

Recovery of the HCl train provided a sample consisting of three components: probe rinse, filter, and back-half rinse/impinger catch.

A total of four HCl train runs were performed at the ESP outlet stack sampling location. HCl train runs 1 and 2 were performed on sample days 1 and 2, while HCl train runs 3 and 4 were performed on sample day 3. The fourth HCl run was performed because a broken impinger may have invalidated data from the first HCl run. The HCl samples were collected over on-line sample times of 100 to 175 minutes at a sample flow rate of approximately 0.8 scfm.

6.1.2.3 Volumetric Gas Flow Rate Determination.

The volumetric gas flow rate was determined at the ESP inlet and outlet sampling locations using EPA Method 2. Based on this method, the volumetric gas flow rate was determined by measuring the average velocity of the flue gas and the cross-sectional area of the duct. The average flue gas velocity was calculated from the average gas velocity pressure (P) across a S-type pitot tube, the average flue gas temperature, the wet molecular weight, and the absolute static pressure.

6.1.2.4 Flue Gas Moisture Determination.

The moisture content of the flue gas was determined at the ESP inlet and outlet sampling locations using EPA Method 4. Based on this method, a measured volume of particulate-free gas was pulled through a chilled impinger train. The quantity of condensed water was determined gravimetrically and then related to the volume of gas sampled to determine the moisture content.

6.1.2.5 Flue Gas Molecular Weight Determination.

The integrated sampling technique described in EPA Method 3 was used at the ESP inlet and outlet sampling locations to obtain a composite flue gas sample for fixed gas (O_2 , CO_2 , N_2) analysis. The fixed gas analysis was used to determine the molecular weight of the gas stream. A small diaphragm pump and a stainless steel probe were used to extract single point flue gas samples. The samples were collected in a Tedlar^R bag. Moisture was removed from the gas sample by a water-cooled condenser so that the fixed gas analysis is on a dry basis.

The composition of the gas sample was determined using a Shimadzu Model 3BT analyzer instead of the Fyrite or Orsat analyzer prescribed in EPA Method 3. The Shimadzu instrument employs a gas chromatograph and a thermal conductivity detector to determine the fixed gas composition of the sample.

6.1.2.6 Continuous Monitors.

Continuous monitoring was performed at the boiler outlet sampling location for O_2 , CO_2 , CO , NO_x , SO_2 , and THC throughout the 4 to 6-hour period that MM5 dioxin sampling was being conducted each test day. The primary objectives of the continuous monitoring effort were to observe fluctuations in flue gas parameters and to provide an indication of combustion conditions. Sample acquisition was accomplished using an in-stack filter probe and a 150 ft length of heat-traced Teflon^R sample line connected to a mobile laboratory. The heat-traced sample line was maintained at a temperature of at least 120°C to prevent condensation in the sample line. The stack gas sample was drawn through the in-stack filter and sample line using pumps located in the mobile laboratory. Sample gas to be analyzed for CO , CO_2 , O_2 , SO_2 , and NO_x was pumped through a sample gas conditioner, which consisted of an ice bath and knockout trap. The sample gas conditioner removes moisture and thus provides a dry gas stream for analysis. A separate unconditioned gas stream was supplied to the THC analyzer for analysis on a wet basis.

An Anarad Model 412 nondispersive infrared (NDIR) analyzer was used to measure CO and CO_2 ; a Beckman Model 755 paramagnetic analyzer was used to measure O_2 ; a Teco Model 10 chemiluminescent analyzer was used to measure NO_x ; a Teco Model 40 pulsed fluorescence analyzer was used to measure SO_2 ; and a Beckman Model 402 flame ionization analyzer was used to measure THC.

6.2 LIQUID SAMPLING

Four types of liquid samples were obtained during this test program: strong black liquor, caustic (sodium hydroxide), white liquor, and weak black liquor. The corresponding sampling locations are shown on Figure 4-1 as A, D, E, F, and G, respectively.

6.2.1 Strong Black Liquor Sampling

Strong black liquor samples were taken from the mix tank where liquid sulfur is added to the concentrated black liquor from the evaporators. Black liquor from the mix tank is fed directly to the feed guns in the boiler. The host site performs daily strong black liquor sampling at this location using a dipper-type sampler.

Three identical composite strong black liquor samples were obtained during each of the three tests: a 1-liter composite was shipped to Troika for dioxin analysis, another 1-liter composite was returned to Radian/RTP for dioxin precursor analysis, and a 125 ml composite was sent to Radian/Austin for total chlorine analysis. The composite strong black liquor samples for each run were comprised of hourly grab samples from the mix tank. It was necessary to heat the running hourly sample composite to prevent the sample from solidifying prior to taking the final sample aliquots. This was accomplished by wrapping the sample composite bottle with rubber-coated heat tape.

6.2.2 Auxiliary Black Liquor Circuit Sampling

Samples of caustic, white liquor, and weak black liquor were obtained to indicate the relative amounts of chlorine entering the black liquor circuit through various input sources. One 125 ml composite sample of each stream was obtained during each test. Individual samples were taken twice during each test run, and the composite sample were prepared accordingly. The samples were analyzed for total chlorine content only. The chlorine-content data was used in conjunction with mass flow data to determine the relative amounts of chlorine associated with each potential source of chlorine input to the black liquor circuit. The chlorine analysis was performed by the Radian Analytical Services laboratory in Austin, Texas using ion chromatography.

Caustic samples were taken from a sample tap in the transfer line leading from caustic storage to the causticizing area in the pulping circuit. White liquor samples were taken from a sample tap on the white liquor storage tank because there was no sampling location available on the transfer line leading from the storage tank to the digesters. Weak black liquor samples were taken from a sample tap in the transfer line leading to the concentrators.

6.3 SOLID SAMPLING

Two types of solid samples were obtained at Site BLB-A: by-product salt cake from the ClO_2 generation process and soils from plant property. The sampling locations and methods are discussed below.

6.3.1 By-product Salt Cake Sampling

Samples of by-product salt cake from the ClO_2 generation system were obtained during Runs 01 and 02 at Site 04. This material was not being produced or fed to the black liquor circuit during Run 03, so no samples could be taken then. During Runs 01 and 02, approximately 300 grams of wet salt cake were obtained twice during the test run from a rotary filter used to separate precipitated salt cake from an aqueous by-product stream of the ClO_2 generation system. At the end of the test run a 125 g sample of the run composite was placed in a 125 ml bottle and sent to Radian's Austin Texas laboratory for total chloride analysis.

6.3.2 Soil Sampling

A single composite soil sample comprised of 10 individual soil samples was obtained at Site BLB-A. Soil sampling protocol for Tiers 3, 5, 6, and 7 of the National Dioxin Study are specified in the document, "Sampling Guidance Manual for the National Dioxin Study." A similar protocol was used for soil sampling at this test site. A total of 10 soil sampling locations were selected on plant property surrounding the black liquor boiler/power complex. The 10 individual soil sampling locations are shown in Figure 6-6 and are listed in Table 6-2. Soil samples were collected by forcing a bulb planter into the soil to a depth of 3 inches. The soil samples were then composited in a clean stainless steel bucket. Five hundred grams of the composite was placed in a 950 ml glass amber bottle and returned to Radian/RTP for archiving.

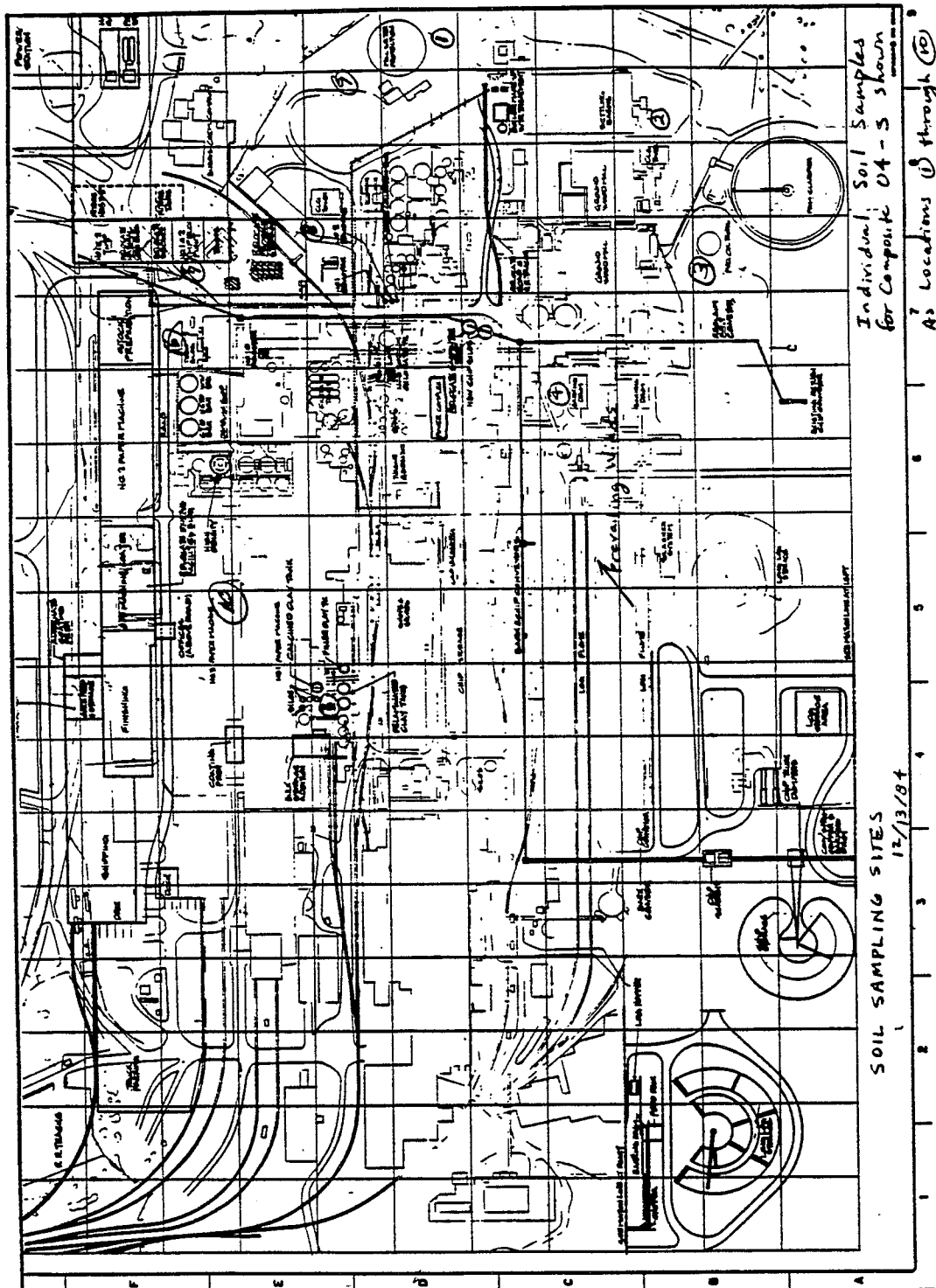


Figure 6-6 Soil Sampling Locations at Site BLB-A

TABLE 6-2. DESCRIPTION OF SOIL SAMPLING LOCATIONS AT SITE BLB-A

Individual Sample	Sampling Location Description ^a
1	Near "mill water protection" area (D9)
2	Near settling basins (B8)
3	Near fuel oil tanks (B7)
4	Near barking drum (C6)
5	Near clay tanks (E4)
6	Near technical building (F7)
7	Near TMP bleach plant (F7)
8	Near "CLG Tower" (E7)
9	Near "Mill water protection" area (D8)
10	Near No. 3 paper machine (E5)

^aSampling location coordinates for Figure 6-5 are listed in parenthesis.

7.0 ANALYTICAL PROCEDURES

Laboratory procedures used to quantify dioxins/furans and dioxin/furan precursors in the Tier 4 samples are described in this section. Analyses for dioxins/furans were performed by EPA's ECL-BSL and EMSL-RTP laboratories. The procedures used for these analyses are described in detail in the Analytical Procedures and QA Plan for the Analysis of Tetra through Octa CDD's and CDF's in Samples from Tier 4 Combustion and Incineration Processes (addendum to EPA/600/3-85-/019, April, 1985).

Black liquor boiler feed samples from Site BLB-A were analyzed to determine concentrations of chlorinated phenols (CP), chlorobenzenes (CB), polychlorinated biphenyls (PCB's), total organic halogen (TOX), and total chlorine. Procedures used for these analyses are detailed in Section 7.2.

7.1 DIOXINS/FURANS

The analytical procedures described in this section were used for the determination of PCDD and PCDF in stack effluent samples (MM5). Samples consisting of organic solvents, aqueous solutions and solids were prepared for analysis using slightly different procedures. The organic solvent samples consisted of rinses from the MM5 probe, nozzle, filter housing and condenser coil. The impinger catch was an aqueous solution and the solid samples included the filter and XAD^R resin. All aqueous and solid samples were extracted within 14 days of collection and then stored in this form until analysis. Isotopically-labeled surrogate compounds were added to all samples prior to extraction to allow determination of method efficiency and for quantification purposes.

Organic liquid samples (acetone and hexane or methylene chloride) were concentrated using a nitrogen blowdown apparatus. The residue, which contained particulates from the train probe and nozzle, was combined with the filter and handled as a solid sample. Solid samples were extracted with benzene in a Soxhlet apparatus for a period of at least 16 hours. The sample was then concentrated by nitrogen blowdown and subjected to chromatographic cleanup procedures.

Aqueous solutions were extracted with hexane by vigorous shaking for a three hour period. This extraction procedure was repeated three times, with the organic fractions ultimately being combined and concentrated for chromatographic cleanup.

The cleanup procedure involved using liquid chromatographic columns to separate the compounds of interest from other compounds present in the samples. Four different types of columns were used: a combination acid and base modified silica gel column, a basic alumina column, a PX-21 carbon/celite 545 column and a silica/diol micro column.. These were used in successive steps, with the last two being used only if necessary.

The cleaned samples were analyzed using high resolution gas chromatography/high resolution mass spectrometry (GC/MS). The conditions for the analyses were as follows:

Gas Chromatograph - Injector configured for capillary column, splitless injection, injector temperature 280°C, helium carrier gas at 1.2 ml/min, initial column temperature 100°C: final column temperature 240°C, interface temperature 270°C.

Mass Spectrometer - Varian/MAT Model 311A, electron energy 70ev, filament emission 1MA, mass resolution 8,000 to 10,000, ion source temperature 270°C.

7.2 DIOXIN/FURAN PRECURSORS

Feed samples for Site BLB-A were analyzed by Radian/RTP for chlorophenols (CP), chlorobenzenes (CB) and polychlorinated biphenyls (PCB's) by GC/MS, total organic halides (TOX) by GC/Hall detector, and total chlorine by Parr Bomb combustion followed by ion chromatography. Analytical procedures are discussed in the following sections.

7.2.1 GC/MS Analyses

The analytical procedures used for determining CP, CB, and PCB concentrations in feed samples are modified versions of procedures typically used for the analysis of MM5 train components.

7.2.1.1 Sample Preparation

A flow chart for the sample preparation procedure used for Site BLB-B feed samples is shown in Figure 7-1. The first step involved adding 200 mL of

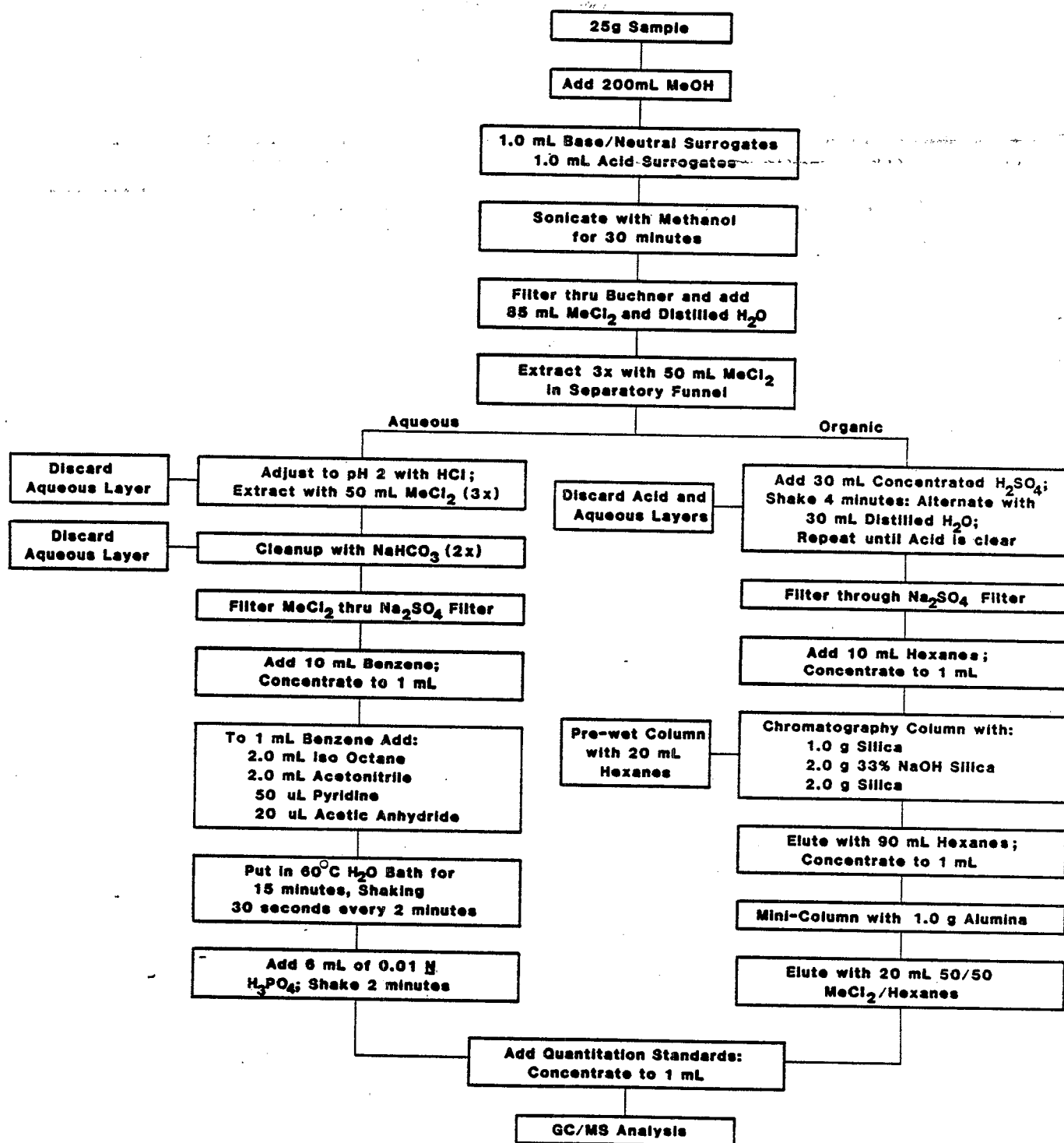


Figure 7-1. Sample preparation flow diagram for Site BLB-A precursor analyses.

methanol to the sample. The next step in the procedure involved adding labeled surrogate compounds to provide a measure of extraction method efficiency. The next step involved sonicating the sample for 30 minutes. The sonicated sample was filtered and rinsed with 85 mL methylene chloride (MeCl_2) and distilled H_2O . The filtrate was extracted three times with 50 mL MeCl_2 in a separatory funnel and the resulting aqueous and organic fractions saved for derivatization and/or further cleanup. These procedures involve initial extraction of the sample with an appropriate solvent, preliminary separation of the compounds of interest by solvent partitioning and liquid chromatography, and analysis of the processed fractions. Solutions containing CB and PCB are injected directly into the GC/MS, and solutions containing CP are derivatized prior to injection. Details on the procedures used for Site 05 samples are provided in the sections below.

The aqueous fraction (or acids portion) was acidified to pH 2 with HCl and then extracted three times with MeCl_2 , followed by two extractions with NaHCO_3 . The MeCl_2 from this extraction was dried with anhydrous Na_2SO_4 , exchanged to benzene, and concentrated using a nitrogen blowdown apparatus. Acetylation of any CP present in the sample involved the following steps:

1. Add 2.0 mL isooctane, 2.0 mL acetonitrile, 50 μL pyridine, and 20 μL acetic anhydride to the extract. Put the test tube containing the extract in a 60°C water bath for 15 minutes, shaking 30 seconds every 2 minutes.
2. Add 6 mL of 0.01 N H_3PO_4 to the test tube and agitate the sample for 2 minutes on a wrist action shaker.
3. Remove the organic layer and add the quantitation standard. Concentrate the sample in a Reacti-Vial at room temperature (using prepurified N_2) to 1 mL prior to GC/MS analysis.

Cleanup of the organic (or base/neutrals) layer from the initial MeCl_2 extraction involved successively washing the extract with concentrated H_2SO_4 and deionized water. The acid or water was added in a 30 mL portion and the sample was shaken for two minutes. After the aqueous and organic layers were completely separated, the aqueous or acid layer was discarded. The acid

washing procedure was repeated until the acid layer was colorless. The sample was then dried with anhydrous Na_2SO_4 , exchanged into hexanes and concentrated. Final cleanup of the sample by column chromatography involved the following procedure.

A glass macro-column, 20 mm o.d. x 230 mm in length, taper to 6 mm o.d. on one end was prepared. The column was packed with a plug of silanized glass wool, followed successively by 1.0 g silica, 2.0 g silica containing 33% (w/w) 1 M NaOH, and 2.0 g silica. The concentrate was quantitatively transferred to the column and eluted with 90 mL hexanes. The entire eluate was collected and concentrated to a volume of 1 mL in a centrifuge tube, as above.

A disposable liquid chromatography mini-column was constructed by cutting off a 5-mL Pyrex disposable pipette at the 2.0 mL mark and packing the lower portion of the tube with a small plug of siliconized glass wool, followed by 1 g of Woehlm basic alumina, which has been previously activated for at least 16 hours at 600°C in a muffle furnace and cooled in a desiccator for 30 minutes just before use. The concentrated eluate from above was quantitatively transferred onto the liquid chromatography column, and the centrifuge tube was rinsed consecutively with two 0.3-mL portions of a 3 percent MeCl_2 :hexanes solution, and the rinses were transferred to the chromatography column.

The column was eluted with 20 mL of a 50 percent (v/v) MeCl_2 :hexanes, and the elute was retained. The retained fraction was concentrated to a volume of approximately 1.0 mL by heating the tubes in a water bath while passing a stream of prepurified N_2 over the solutions. The quantitation standard was added and the final volume was adjusted to 1.0 mL prior to GC/MS analysis.

7.2.1.2 Analysis

Analyses for CP, CB and PCBs present in the feed sample extracts were performed using a Finnigan Model 5100 mass spectrometer using selected ion monitoring. A fused silica capillary column was used for chromatographic separation of the compounds of interest. Analytical conditions for the GC/MS analysis are shown in Table 7-1.

Tuning of the GC/MS was performed daily as specified in the Tier 4 QA Project Plan. An internal standard calibration procedure was used for sample quantitation. Compounds of interest were calibrated against a fixed concentration of either d_{12} -chrysene (CB, PCB) or d_8 -naphthalene (CP).

TABLE 7-1. INSTRUMENT CONDITIONS FOR GC/MS PRECURSOR ANALYSES

Parameter	Chlorobenzenes/ Polychlorinated Biphenyls	Chlorophenols
Column	30 m WB DB-5 (1.0 u film thickness) fused silica capillary	
Injector Temperature	290°C	290°C
Separator Oven Temperature	290°C	290°C
Column Head Pressure	9 psi	9 psi
He flow rate	1 mL/min	1 mL/min
GC program	40(4)-290°C, 10°/min & hold	40(1)-290°C, 12°/min & hold
Emission Current	0.50 ma	0.50 ma
Electron Energy	70 ev	70 ev
Injection Mode	Splitless 0.6 min, then 10:1 split	
Mode	Electron ionization, Selected Ion Monitoring	

Compounds of the calibration solution are shown in Table 7-2. For multi-point calibrations, this solution was injected at levels of 10, 50, 100, and 150 ng/mL.

Compound identification was confirmed by comparison of chromatographic retention times and mass spectra of unknowns with retention times and mass spectra for reference compounds. Since the selected ion monitoring technique was necessary for the types of samples analyzed, care was taken to monitor a sufficiently wide mass region to avoid the potential for reporting false positives.

The instrument detection limit was estimated to be approximately 500 picograms on column. For a 50 g sample and 100 percent recovery of the analyte, this corresponds to a feed sample detection limit of 10 ppb.

7.3 TOTAL CHLORINE ANALYSIS

Total chlorine concentrations in feed samples were determined by Parr Bomb combustion followed by ion chromatography (IC). A 0.5g sample was placed in the Parr Bomb with 10 mL of a 50 g/L Na_2CO_3 solution. After combustion of the samples according to standard procedures (ASTM 2015), the contents of the bomb were rinsed into a 100 mL flask and diluted to 100 mL. The resulting solution was analyzed for chloride concentration (Cl^-) by IC using standard anion conditions. For samples difficult to combust (such as sludges), 25 drops of paraffin oils were added to the bomb prior to combustion.

TABLE 7-2. COMPONENTS OF THE CALIBRATION SOLUTION

<u>Base/Neutrals</u>	<u>Acids</u>
4-chlorobiphenyl	2,5-dichlorophenol
3,3'-dichlorobiphenyl	2,3-dichlorophenol
2,4',5-trichlorobiphenyl	2,6-dichlorophenol
3,3',4,4'-tetrachlorobiphenyl	3,5-dichlorophenol
2,2',6,6'-tetrachlorobiphenyl	3,4-dichlorophenol
2,2,4,5,6-pentachlorobiphenyl	2,3,5-trichlorophenol
2,2',4,4',5,5'-hexachlorobiphenyl	2,3,6-trichlorophenol
2,2',3,4,4',5',6-heptachlorobiphenyl	3,4,5-trichlorophenol
2,2',3,3',4,4',5,5'-octachlorobiphenyl	2,4,5-trichlorophenol
2,2',3,3',4,4',5,6,6'-nonachlorobiphenyl	2,3,4-trichlorophenol
decachlorobiphenyl	2,3,5,6-tetrachlorophenol
p-dichlorobenzene	pentachlorophenol
1,2,4-trichlorobenzene	d ₆ -phenol (SS)
1,2,3,5-tetrachlorobenzene	d ₄ -2-chlorophenol (SS)
pentachlorobenzene	¹³ C ₆ -pentachlorophenol (SS)
hexachlorobenzene	d ₈ -naphthalene (QS)
d ₄ -1,4-dichlorobenzene (SS) ¹	2,4,6-tribromophenol (QS)
3-bromobiphenyl (SS)	d ₁₀ -phenanthrene (QS)
2,2',5,5'-tetrabromobiphenyl (SS)	d ₁₂ chrysene (QS)
2,2',4,4',6,6'-hexabromobiphenyl (SS)	
octachloronaphthalene (QS) ²	
d ₁₀ -phenanthrene (QS)	
d ₁₂ -chrysene (QS)	

¹Surrogate standard.²Quantitation standard.

8.0 QUALITY ASSURANCE/ QUALITY CONTROL (QA/QC)

This section summarizes the results of quality assurance and quality control (QA/QC) activities for Site BLB-A. The flue gas and ash dioxin/furan data for this site were generally within the QC specifications presented in the Tier 4 QAPP. All of the surrogate recoveries for labeled TCDDs were within the specified limits of 50 to 120 percent. The surrogate recoveries for the ESP outlet, Run 02 sample was the only run not within the QC limits of 40 to 120 percent for hepta- and octa-CDDs. The results of the analysis of the fortified laboratory QC sample were all within 33 percent of the true value which is well within the Tier 4 objective of ± 50 percent. These data indicate that the dioxin/furan results are within accuracy criteria specified for Tier 4.

The dioxin/furan precursor analysis of the feed samples was not as accurate as the dioxin/furan homologue analysis. Surrogate recoveries of the base neutrals fraction were generally within the specified QC limits of ± 50 percent; however, the surrogate acid fractions were generally below the specified limits. In spite of the low recoveries of the acid fractions, the dioxin/furan precursor results are considered a reasonable approximation of the true precursor concentration in the feed samples.

The following sections summarize the results of all Site BLB-A QA/QC activities. Manual gas sampling methods are considered in Section 8.1 and continuous emission monitoring and molecular weight determinations are considered in Section 8.2. The laboratory analysis QA/QC activities are summarized in Section 8.3.

8.1 MANUAL GAS SAMPLING

Manual gas sampling methods used at Site 04 included Modified Method 5 (MM5), the HCl acid train and EPA Methods 1 through 4. These methods are discussed in Section 6.0. Quality assurance and quality control (QA/QC) activities for the manual methods centered around 1) equipment calibration, 2) glassware precleaning, 3) procedural QC checks and 4) sample custody procedures. Key activities and QC results in each of these areas are

discussed in this section. Also discussed are problems encountered that may have affected data quality.

Pretest calibrations or inspections were conducted on pitot tubes, sampling nozzles, temperature sensors and analytical balances. Both pre- and post-test calibrations were also performed on dry gas meters. All of this equipment met the calibration criteria specified in the QAPP. Differences in pre- and post-test dry gas meter calibrations were less than 3.4 percent.

An extensive precleaning procedure was implemented for all sample train glassware and sample containers. This cleaning procedure, which is outlined in Table 8-1, was implemented to minimize the potential for sample contamination with substances that could potentially interfere with the analysis for dioxins and furans. To minimize the potential for contamination in the field, all sample train glassware was kept capped until use and a controlled environment was maintained in the recovery trailer during sample train assembly and recovery.

Procedural QC activities during manual gas sampling focused on:

- . inspecting equipment visually
- . collecting sampling train blanks
- . ensuring the proper location and number of traverse points
- . conducting pre-test, port change, and post-test sample train leak checks
- . maintaining proper temperatures at the filter housing, sorbent trap and impinger train
- . maintaining isokinetic sampling rates, and
- . recording all data on preformatted data sheets.

Results of isokinetic calculations for the MM5 test runs are shown in Table 8-2. As shown in Table 8-2, the average isokinetic sampling rate for the MM5 and sampling trains achieved the QA objective of ± 10 percent for all test runs.

Sample custody procedures used during this program emphasized careful documentation of the samples collected and the use of chain-of-custody records for samples to be transported. Steps taken to identify and document samples collected included labeling each sample with a unique alphanumeric

TABLE 8-1. GLASSWARE PRECLEANING PROCEDURE

NOTE: USE DISPOSABLE GLOVES AND ADEQUATE VENTILATION

1. Soak all glassware in hot soapy water (Alconox) 50° C or higher.
 2. Distilled/ deionized H₂O rinse (X3)^a
 3. Chromerge rinse if glass, otherwise skip to 6.
 4. High purity liquid chromatography grade H₂O rinse (X3).
 5. Acetone rinse (X3), (pesticide grade).
 6. Hexane rinse (X3), (pesticide grade).
 7. Cap glassware with clean glass plugs or hexane rinsed aluminum foil.
-

^a (X3) = three times.

TABLE 8-2. SUMMARY OF ISOKINETICS RESULTS
FOR MM5 SAMPLING TRAINS, SITE BLB-A

Run	ESP Outlets Isokinetics	Meets OC Objective?	ESP Inlet Isokinetics	Meets QC Objective?
01	91.7	Yes	92.5	Yes
02	106.5	Yes	98.7	Yes
03	108.8	Yes	104.7	Yes

NOTE: The quality assurance objective for MM5 sampling was isokinetics of 100 ± 10 percent.

code and logging the sample in a master sample logbook. All samples shipped to Troika or returned to Radian were also logged on chain-of-custody records that were signed by the sampler at shipment and then by the receiving laboratory when the samples arrived. Each sample container was also sealed with chain-of-custody seal so that the container could not be opened without tearing the seal.

8.2 CONTINUOUS MONITORING/ MOLECULAR WEIGHT DETERMINATION

Flue gas parameters monitored continuously during the MM5 test runs included CO, CO₂, O₂, total hydrocarbons (THC), SO₂ and NO_x. Concentrations of CO₂, O₂, and N₂ were also determined for integrated bag samples of stack gas. Quality control results for these analyses are discussed in this section.

Drift check and quality control standard analysis results for the continuously monitored flue gas parameters are summarized in Table 8-3. The acceptance criterion for drift checks was an instrument drift within ± 10 percent. All data reduction was performed by assuming a linear drift of instrument response over the test day. The largest calibration drifts were observed for CO and CO₂, both of which exceeded acceptance criterion for two of the three test runs. The instrument showing the smallest drift was the O₂ monitor.

The quality control standards for this program consisted of mid-range standards that were not used for instrument calibration but were analyzed immediately after calibration to provide data on day-to-day instrument variability. The acceptance criterion for each control standard was agreement within ± 10 percent of the running mean value. All of the instruments met this criterion on each test day except for the CO monitor. However, failure of the CO monitor to meet the acceptance criterion during test runs 2 and 3 is not entirely unexpected. The QC standard (2060 ppmv) was above the calibration range selected for the CO instrument during these test runs (0 to 520 ppmv) and, the CO instrument shows some non-linearity at low concentrations. The instrument was calibrated at a low range to maximize instrument accuracy near the CO levels present in the flue gas.

TABLE 8-3. SUMMARY OF DRIFT CHECK AND CONTROL STANDARD RESULTS

Test Date	Test Run	Parameter	Drift Check			Meets QC? ^b	QC Standard		Meets QC? ^c
			Input Concentration	Instrument Drift % ^a			Input Concentration	Output Concentration	
12/11/84	01	O ₂	21.2 %	- 0.77		yes	-	-	yes
12/12/84	02	O ₂	21.2 %	- 0.38		yes	3.12 %	3.16	yes
12/13/84	03	O ₂	21.2 %	ND ^e		ND ^e	3.12 %	3.19	yes
12/11/84	01	CO	5430 ppmv	0.12		yes	-	-	-
12/12/84	02	CO	5430 ppmv	1.92		yes	520 ppm	728 ppm	no
12/13/84	03	CO	5430 ppmv	e		e	520 ppm	480 ppm	no
12/11/84	01	CO ₂	17.6 %	6.01		yes	-	-	-
12/12/84	02	CO ₂	17.6 %	17.9		no	5.2 %	5.66 %	yes
12/13/84	03	CO ₂	17.6 %	ND ^e		ND ^e	5.2 %	7.0 %	no
12/11/84	01	SO ₂	958 ppmv	0.21		yes	83.1	86.9	yes ^d
12/12/84	02	SO ₂	958 ppmv	21.7 %		no	83.1	89.2	yes
12/12/84	03	SO ₂	958 ppmv	- 4.53 %		yes	83.1	90.1	yes
12/11/84	01	NO _x	816.1 ppmv	-		-	153	154	yes ^d
12/12/84	02	NO _x	84.6 ppmv	15.2 %		no	no QC	No QC	-
12/13/84	03	NO _x	84.6 ppmv	- 5.7 %		yes	21.0	23.1	yes ^d

TABLE 8-3. SUMMARY OF DRIFT CHECK AND CONTROL STANDARD RESULTS
(cont'd.)

Test Date	Test Run	Parameter	Drift Check		Meets QC? ^b	QC Standard		Meets QC?
			Input Concentration	Instrument Drift % ^a		Input Concentration	Output Concentration	
12/11/84	01	THC	90 ppmv	1.7 %	yes	19.7	20.1	yes
12/12/84	02	THC	90 ppmv	- 1.9 %	yes	19.7	20.0	yes
12/13/84	03	THC	90 ppmv			19.7	20.2	yes

^a Instrument drift is defined as the percent difference between the instrument response to the input concentration at the beginning and end of the test run.

^b QC criteria was instrument drift within ± 10 percent.

^c QC criteria was output concentration within ± 10 percent of the running mean concentration for this test site.

^d Output concentration within ± 10 percent of known value.

ND^e Not determined due to calibration gas leakage prior to completion of post test calibration procedure.

Molecular weight was determined by analyzing integrated bag samples of stack gas for CO₂, O₂, and N₂. Quality control for this analysis involved duplicate analyses of calibration gases immediately before and after sample analysis. Analysis of the calibration was to be repeated until two consecutive analyses within ± 5 percent were obtained. This same criterion of ± 5 percent applied to duplicate analyses required for each sample quantitation. These criteria were met for all molecular weight determinations where valid integrated bag samples were obtained.

8.3 LABORATORY ANALYSIS

QA/QC activities were carried out for dioxin/furan, precursor, and total chloride analyses performed on Site BLB-A samples. The dioxin/furan analyses of MM5 train samples performed by Troika are considered in Section 8.3.1; the precursor analyses of black liquor boiler feed samples performed by Radian/RTP are considered in Section 8.3.2; and the total chloride analyses of HCl train samples and process samples performed by Radian/Austin are considered in Section 8.3.3.

8.3.1 Dioxin/Furan Analyses

Two individual topics related to the dioxin/furan analyses at Site BLB-A are discussed in this section. Analytical recoveries of labeled surrogate compounds spiked onto MM5 train samples are reported in Section 8.3.1.1. Sample blank data are reported in Section 8.3.1.2.

8.3.1.1 Surrogate Recoveries of the Test Samples

Table 8-4 presents the analytical recovery data reported by Troika for four isotopically labeled surrogate compounds spiked onto the MM5 train samples. Samples were spiked with all four of the surrogates. The surrogate recovery results generally met the Tier 4 QA criteria of 50 to 120 percent recovery for the tetrachlorinated surrogates and 40 to 120 percent recovery for the hepta- and octa-chlorinated surrogates.

8.3.1.2 Sample Blanks

Table 8-5 summarizes the analytical results reported by Troika for internal laboratory blanks, laboratory fortified quality control (QC) samples, proof blank MM5 train samples, and field recovery blank MM5 train samples. In general, the data showed good surrogate recoveries, with values

TABLE 8-4. PERCENT SURROGATE RECOVERIES FOR SITE BLB-A
DIOXIN/FURAN ANALYSES

Sample	$^{37}\text{Cl}_4$ TCDD	$^{13}\text{C}_{12}$ TCDD	$^{37}\text{Cl}_4$ Hepta-CDD	$^{13}\text{C}_{12}$ Octa-CDD
<u>MM5 Train Samples</u>				
Inlet				
Run 01	90	90	55	76
Run 02	100	106	40	66
Run 03	96	62	53	42
Outlet				
Run 01	94	88	47	49
Run 02	85	91	34	51
Run 03	100	98	48	63
<u>Economizer Ash</u>				
a	-	89	-	93

^aA single economizer ash sample was taken during the pre-survey.

TABLE 8-5. ANALYSIS RESULTS FOR QUALITY CONTROL SAMPLES

Compound	Flue Gas Quality Control Samples					
	Laboratory Blank	Fortified Laboratory QC Sample		Proof Blank MW5 Train	Field Blank MW5 Train	
		Measured Value	True Value a,b		Inlet	Outlet
Amount Detected (Nanograms per Sample)						
Dioxins						
2378 TCDD	ND	0.4	0.4 (0)	ND	ND	ND
Other TCDD	ND	ND	ND (0)	ND	ND	ND
Penta CDD	ND	ND	ND (0)	ND	ND	ND
Hexa CDD	ND	1.4	1.6 (-14)	ND	ND	ND
Hepta CDD	ND	3.0	2.4 (20)	ND	ND	0.5
Octa CDD	0.9	3.6	3.2 (-11)	0.5	0.8	1.0
Furans						
2378 TCDF	ND	0.5	0.4 (20)	ND	ND	ND
Other TCDF	ND	ND	ND (0)	ND	0.2	ND
Penta CDF	ND	0.8	0.8 (0)	ND	ND	ND
Hexa CDF	ND	1.2	1.6 (-33)	ND	ND	ND
Hepta CDF	ND	3.1	2.4 (23)	ND	ND	ND
Octa CDF	0.9	3.7	3.2 (14)	ND	ND	ND
Surrogate Recoveries (Percent)						
³⁷ C ₁ -TCDD	94	100	NA	84	86	106
¹³ C ₁₂ -TCDD	94	92	NA	86	92	102
³⁷ C ₄ -Hepta CDD	54	57	NA	47	44	42
¹³ C ₁₂ -Octa CDD	65	72	NA	79	48	67

^aTrue values represent the amounts of each homologue spiked into the laboratory fortified QC samples.
^bValue shown in parenthesis is the percentage difference between the measured and the true value:

$$\% = \frac{\text{Measured Value} - \text{True Value}}{\text{True Value}} \times 100$$

ND = Not Detected

NA = Not Applicable

TCDD = Tetra-chlorinated dibenzo-p-dioxin

ranging from 42 to 106 percent. Comparison of the measured and spiked values for the laboratory fortified QC samples showed agreement well within the Tier 4 target of ± 50 percent. Dioxin and furan species were non-detectable in the proof blank MM5 train sample except the octa-CDD isomer for which 0.5 ng was detected. Small but detectable quantities of hepta-CDD, octa-CDD and tetra-CDF were found in the field blank MM5 trains. The octa-CDD isomer was detected in the inlet and outlet field blanks in quantities of 0.8 and 1.0 ng, respectively. The tetra-CDF isomer was detected at the inlet but was non-detectable in the outlet field blank. Finally, 0.5 ng of the hepta-CDD isomer was detected at the outlet. Table 8-6 gives a comparison of the dioxin/furan analytical results for the field blank and test run MM5 trains. For the inlet values, only the octa-CDD homologue had a detectable field blank value, equal to 50 percent of the minimum test run value. However, for the outlet data, the hepta-CDD, octa-CDD, and tetra-CDF isomers had reported field blank values equal to 100, 77, and 40 percent of the minimum test run value, respectively. This indicates that there might have been some blanking problems at Site BLB-A. Emissions data reported in Section 5.4 are not blank-corrected.

8.3.2 Precursor Analyses

Table 8-7 presents analytical recovery efficiencies for seven isotopically labeled compounds used as surrogates for the target precursor analytes in the Site BLB-A feed samples. The surrogate recovery values in Table 8-7 vary considerably by specific surrogate species but are fairly uniform between runs for the same species. Several of the recoveries are below the 50 percent objective stated in the Tier 4 QA Project Plan and are below those generally considered achievable when analyzing for similar compounds in water or from MM5 train components. In spite of the relatively low surrogate recovery values for some of the feed samples, the resulting analytical sensitivity for the target analytes was considered acceptable for the purpose of this study.

8.3.3 Total Chloride Analysis

Total chloride analyses were performed by Radian/Austin on the HCl train samples. QA/QC activities include total chloride analysis of field recovery blank HCl train samples and total chloride analysis of an aliquot of NaOH

TABLE 8-6. FIELD BLANK DIOXIN/FURAN DATA FOR SITE BLB-A MM5 SAMPLES

Isomer/Homologue	Amount Detected, Nanograms per Train					
	Field Blank Value		Minimum Test Run Value		Percentage ^a	
	Inlet	Outlet	Inlet	Outlet	Inlet	Outlet
<u>Dioxins</u>						
2378 TCDD	ND	ND	ND	ND	0	0
Other TCDD	ND	ND	0.05	0.3	0	0
Penta CDD	ND	ND	ND	ND	0	0
Hexa CDD	ND	ND	0.2	0.8	0	0
Hepta CDD	ND	0.5	0.5	0.5	0	100
Octa CDD	0.8	1.0	1.6	1.3	50	77
<u>Furans</u>						
2378 TCDF	ND	ND	ND	ND	0	0
Other TCDF	ND	0.2	ND	0.5	0	40
Penta CDF	ND	ND	ND	0.6	0	0
Hexa CDF	ND	ND	ND	1.0	0	0
Hepta CDF	ND	ND	ND	0.3	0	0
Octa CDF	ND	ND	ND	0.2	0	0

^aPercentage shown is the ratio of the field blank value to the minimum test run value, expressed as a percentage.

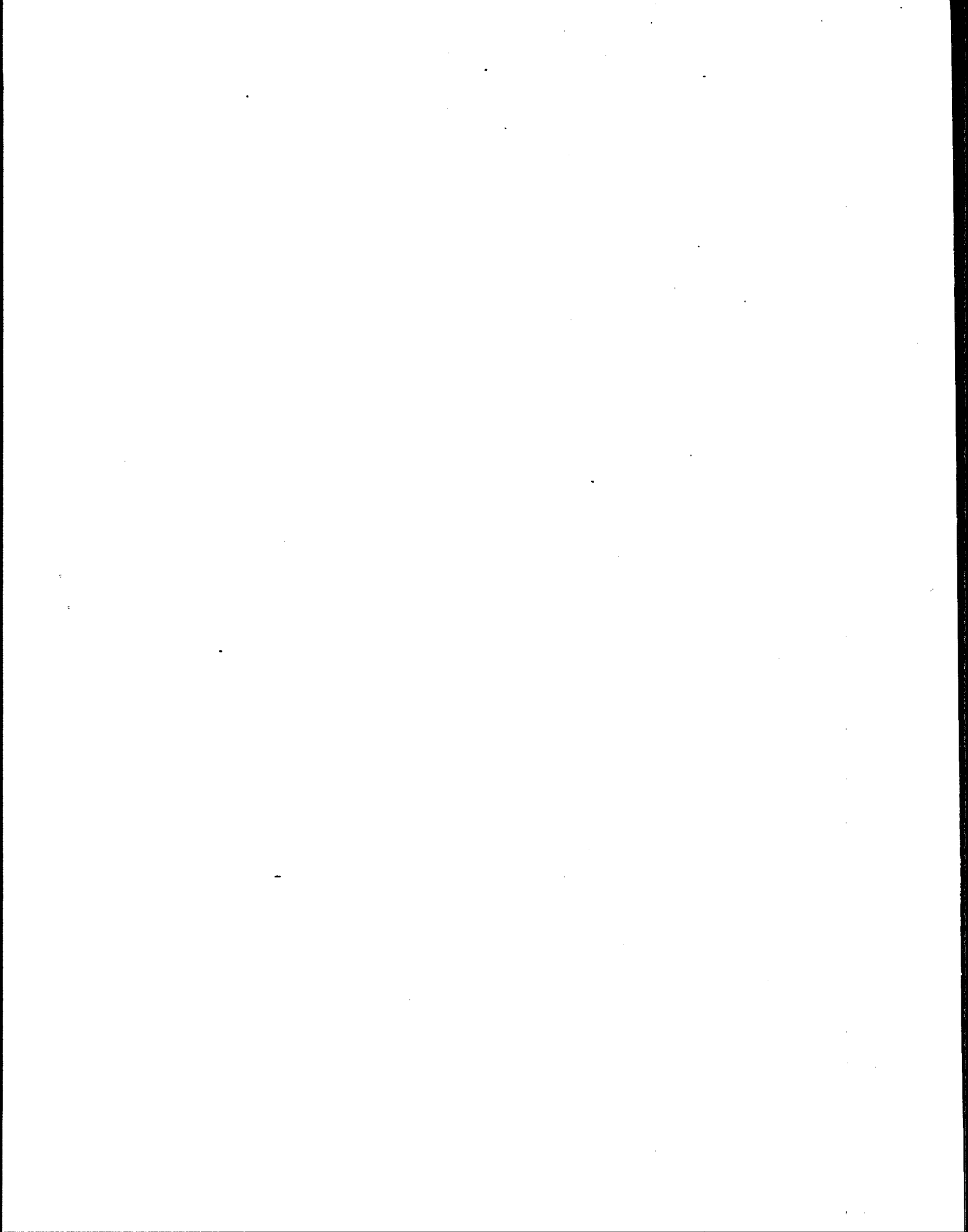
TABLE 8-7. PERCENT SURROGATE RECOVERIES FOR SITE BLB-A FEED SAMPLES

Surrogate Compound	Percent Surrogate Recovery			
	Black Liquor Feed			Average
	Run 01	Run 02	Run 03	
d ₄ -dichlorobenzene	35	100	85	73
bromobiphenyl	114	98	92	101
2',5,5' tetrabromobiphenyl	135	84	80	100
d ₆ -phenol	ND	12,78,77	25	38
d ₄ -2-chlorophenol	21	26,104,96	41	58
¹³ C ₆ -pentachlorophenol	92	35,79,68	40	63

solution used in the sample train impinger rinses. Less than 1 mg/l chloride was detected in the aliquot of NaOH solution analyzed, while chloride concentrations of 2 and 1 mg/l were detected in the front and back halves of the field recovery blank train samples, respectively. These values were well below the typical test run values for this site.

APPENDIX A-1

MODIFIED METHOD 5 AND
EPA METHODS 1-4 FIELD RESULTS



R A D I A N S O U R C E T E S T
E P A M E T H O D 2 - 5
(R A W D A T A)

PLANT : SITE 04
PLANT SITE :
SAMPLING LOCATION : LOCATION B-ESP INLET
TEST # : 04-MM5T-B1
DATE : 12/11/84
TEST PERIOD : 1420-2249

PARAMETER -----	VALUE -----
Sampling time (min.)	240
Barometric Pressure (in.Hg)	29.58
Sampling nozzle diameter (in.)	.312
Meter Volume (cu.ft.)	150.9299
Meter Pressure (in.H2O)	1.585208
Meter Temperature (F)	80.97916
Stack dimension (sq.in.)	15840
Stack Static Pressure (in.H2O)	-10
Stack Moisture Collected (gm)	1026.1
Absolute stack pressure(in Hg)	28.84471
Average stack temperature (F)	350.9375
Percent CO2	14
Percent O2	5
Percent N2	79
Delps Subroutine result	16.92305
DGM Factor	.9947
Pitot Constant	.84

R A D I A N S O U R C E T E S T
 E P A M E T H O D S 2 - 5
 F I N A L R E S U L T S
 PLANT : SITE 04
 PLANT SITE :
 SAMPLING LOCATION : LOCATION B-ESP INLET
 TEST # : 04-MM5T-B1
 DATE : 12/11/84
 TEST PERIOD : 1420-2249

PARAMETER -----	RESULT -----
Vm(dscf)	145.4338
Vm(dscm)	4.118685
Vw gas(scf)	48.38062
Vw gas (scm)	1.370139
% moisture	24.96234
Md	.7503766
MWd	29.88
MW	26.91447
Vs(fpm)	2616.968
Vs (mpm)	797.8562
Flow(acfm)	287866.5
Flow(acmm)	8152.379
Flow(dscfm)	135588.1
Flow(dscmm)	3839.854
% I	92.64972
% EA	31.53381

Program Revision:1/16/8

R A D I A N S O U R C E T E S T
E P A M E T H O D 2 - 5
(R A W D A T A)

PLANT : SITE 04
PLANT SITE :
SAMPLING LOCATION : LOCATION B-ESP INLET
TEST # : 04-MM5T-B2
DATE : 12/12/84
TEST PERIOD : 1100-1541

PARAMETER -----	VALUE -----
Sampling time (min.)	144
Barometric Pressure (in.Hg)	29.52
Sampling nozzle diameter (in.)	.312
Meter Volume (cu.ft.)	93.8
Meter Pressure (in.H2O)	1.632083
Meter Temperature (F)	88.98959
Stack dimension (sq.in.)	15840
Stack Static Pressure (in.H2O)	-10
Stack Moisture Collected (gm)	700.2
Absolute stack pressure(in Hg)	28.78471
Average stack temperature (F)	353.3542
Percent CO2	14
Percent O2	5
Percent N2	79
Delp's Subroutine result	16.67247
DGM Factor	.9973
Pitot Constant	.84

R A D I A N S O U R C E T E S T
 E P A M E T H O D S 2 - 5
 F I N A L R E S U L T S
 PLANT : SITE 04
 PLANT SITE :
 SAMPLING LOCATION : LOCATION B-ESP INLET
 TEST # : 04-MM5T-B2
 DATE : 12/12/84
 TEST PERIOD : 1100-1541

PARAMETER -----	RESULT -----
Vm(dscf)	89.12821
Vm(dscm)	2.524111
Vw gas(scf)	33.01443
Vw gas (scm)	.9349686
% moisture	27.02941
Md	.729706
MWd	29.88
MW	26.66891
Vs(fpm)	2592.76
Vs (mpm)	790.4756
Flow(acfm)	285203.6
Flow(acmm)	8076.967
Flow(dscfm)	129974.3
Flow(dscmm)	3680.871
% I	98.72038
% EA	31.53381

Program Revision:1/16/84

R A D I A N S O U R C E T E S T
E P A M E T H O D 2 - 5
(R A W D A T A)

PLANT : SITE 04
PLANT SITE :
SAMPLING LOCATION : LOCATION B-ESP INLET
TEST # : 04-MM5T-B3
DATE : 12/13/84
TEST PERIOD : 1230-1658

PARAMETER -----	VALUE -----
Sampling time (min.)	141
Barometric Pressure (in.Hg)	29.5
Sampling nozzle diameter (in.)	.312
Meter Volume (cu.ft.)	90.3589
Meter Pressure (in.H2O)	1.525532
Meter Temperature (F)	94.31914
Stack dimension (sq.in.)	15840
Stack Static Pressure (in.H2O)	-10
Stack Moisture Collected (gm)	830
Absolute stack pressure(in Hg)	28.76471
Average stack temperature (F)	353.1915
Percent CO2	14
Percent O2	5
Percent N2	79
Delp's Subroutine result	16.15068
DGM Factor	.9973
Pitot Constant	.84

R A D I A N S O U R C E T E S T
 E P A M E T H O D S 2 - 5
 F I N A L R E S U L T S
 PLANT : SITE 04
 PLANT SITE :
 SAMPLING LOCATION : LOCATION B-ESP INLET
 TEST # : 04-MM5T-B3
 DATE : 12/13/84
 TEST PERIOD : 1230-1658

PARAMETER -----	RESULT -----
Vm(dscf)	84.95314
Vm(dscm)	2.405873
Vw gas(scf)	39.1345
Vw gas (scm)	1.108289
% moisture	31.53779
Md	.6846221
MWd	29.88
MW	26.13331
Vs(fpm)	2538.104
Vs (mpm)	773.8123
Flow(acfm)	279191.5
Flow(acmm)	7906.702
Flow(dscfm)	119314.3
Flow(dscmm)	3378.981
% I	104.6838
% EA	31.53381

Program Revision:1/16/84

R A D I A N S O U R C E T E S T
E P A M E T H O D 2 - 5
(R A W D A T A)

PLANT : SITE 04
PLANT SITE :
SAMPLING LOCATION : LOCATION C-ESP OUTLET
TEST # : 04-MM5T-C1
DATE : 12/11/84
TEST PERIOD : 1420-2020

PARAMETER	VALUE -----
Sampling time (min.)	240
Barometric Pressure (in.Hg)	29.58
Sampling nozzle diameter (in.)	.309
Meter Volume (cu.ft.)	193.1
Meter Pressure (in.H2O)	2.264
Meter Temperature (F)	91.85714
Stack dimension (sq.in.)	12076.31
Stack Static Pressure (in.H2O)	-.85
Stack Moisture Collected (gm)	1287.2
Absolute stack pressure(in Hg)	29.5175
Average stack temperature (F)	346.72
Percent CO2	14
Percent O2	5
Percent N2	77
Delps Subroutine result	21.35512
DGM Factor	.9945
Pitot Constant	.84

R A D I A N S O U R C E T E S T
E P A M E T H O D S 2 - 5
F I N A L R E S U L T S

PLANT : SITE 04
PLANT SITE :
SAMPLING LOCATION : LOCATION C-FSP CUTLET
TEST # : 04-MM5T-C1
DATE : 12/11/84
TEST PERIOD : 1420-2020

PARAMETER -----	RESULT -----
Vm(dscf)	182.6704
Vm(dscm)	5.173226
Vw gas(scF)	60.69148
Vw gas (scm)	1.718783
% moisture	24.93878
Md	.7506123
MWd	29.32
MW	26.49693
Vs(fpm)	3290.109
Vs (mpm)	1003.082
Flow(acfm)	275919.3
Flow(acmm)	7814.033
Flow(dscfm)	133729.4
Flow(dscmm)	3787.215
% I	91.70922
% EA	32.62005

Program Revision:1/10/84

R A D I A N S O U R C E T E S T
E P A M E T H O D 2 - 5
(R A W D A T A)

PLANT : SITE 04
PLANT SITE :
SAMPLING LOCATION : LOCATION C-ESP OUTLET
TEST # : 04-MM5T-C2
DATE : 12/12/84
TEST PERIOD : 1100-1615

PARAMETER -----	VALUE -----
Sampling time (min.)	240
Barometric Pressure (in.Hg)	29.58
Sampling nozzle diameter (in.)	.309
Meter Volume (cu.ft.)	196.481
Meter Pressure (in.H2O)	2.483333
Meter Temperature (F)	30.09375
Stack dimension (sq.in.)	12076.31
Stack Static Pressure (in.H2O)	-.85
Stack Moisture Collected (gm)	1394.2
Absolute stack pressure(in Hg)	29.5175
Average stack temperature (F)	343.4167
Percent CO2	14.1
Percent O2	5.6
Percent N2	77
Delp's Subroutine result	20.82071
DGM Factor	.9945
Pitot Constant	.84

R A D I A N S O U R C E T E S T
 E P A M E T H O D S 2 - 5
 F I N A L R E S U L T S
 PLANT : SITE 04
 PLANT SITE :
 SAMPLING LOCATION : LOCATION C-ESP OUTLET
 TEST # : 04-MM5T-C2
 DATE : 12/12/84
 TEST PERIOD : 1100-1615

PARAMETER -----	RESULT -----
Vm(dscf)	209.4061
Vm(dscm)	5.930381
Vw gas(scf)	65.73653
Vw gas (scm)	1.861658
% moisture	23.8918
Md	.761082
MWd	29.556
MW	26.79506
Vs(fpm)	3189.879
Vs (mpm)	972.5241
Flow(acfm)	267513.6
Flow(acmm)	7575.986
Flow(dscfm)	132004.4
Flow(dscmm)	3738.365
% I	106.5056
% EA	38.02282

Program Revision:1/16/8

R A D I A N S O U R C E T E S T
E P A M E T H O D 2 - 5
(R A W D A T A)

PLANT : SITE 04
PLANT SITE :
SAMPLING LOCATION : LOCATION C-ESP OUTLET
TEST # : 04-MM5T-C3
DATE : 12/13/84
TEST PERIOD : 0945-1527

PARAMETER -----	VALUE -----
Sampling time (min.)	240
Barometric Pressure (in.Hg)	29.5
Sampling nozzle diameter (in.)	.309
Meter Volume (cu.ft.)	194.437
Meter Pressure (in.H2O)	2.4
Meter Temperature (F)	30.69792
Stack dimension (sq.in.)	12076.31
Stack Static Pressure (in.H2O)	-.85
Stack Moisture Collected (gm)	1511
Absolute stack pressure(in Hg)	29.4375
Average stack temperature (F)	341.625
Percent CO2	14.7
Percent O2	4.27
Percent N2	76.9
Delp's Subroutine result	20.41842
DGM Factor	.9945
Pitot Constant	.84

R A D I A N S O U R C E T E S T
 E P A M E T H O D S 2 - 5
 F I N A L R E S U L T S

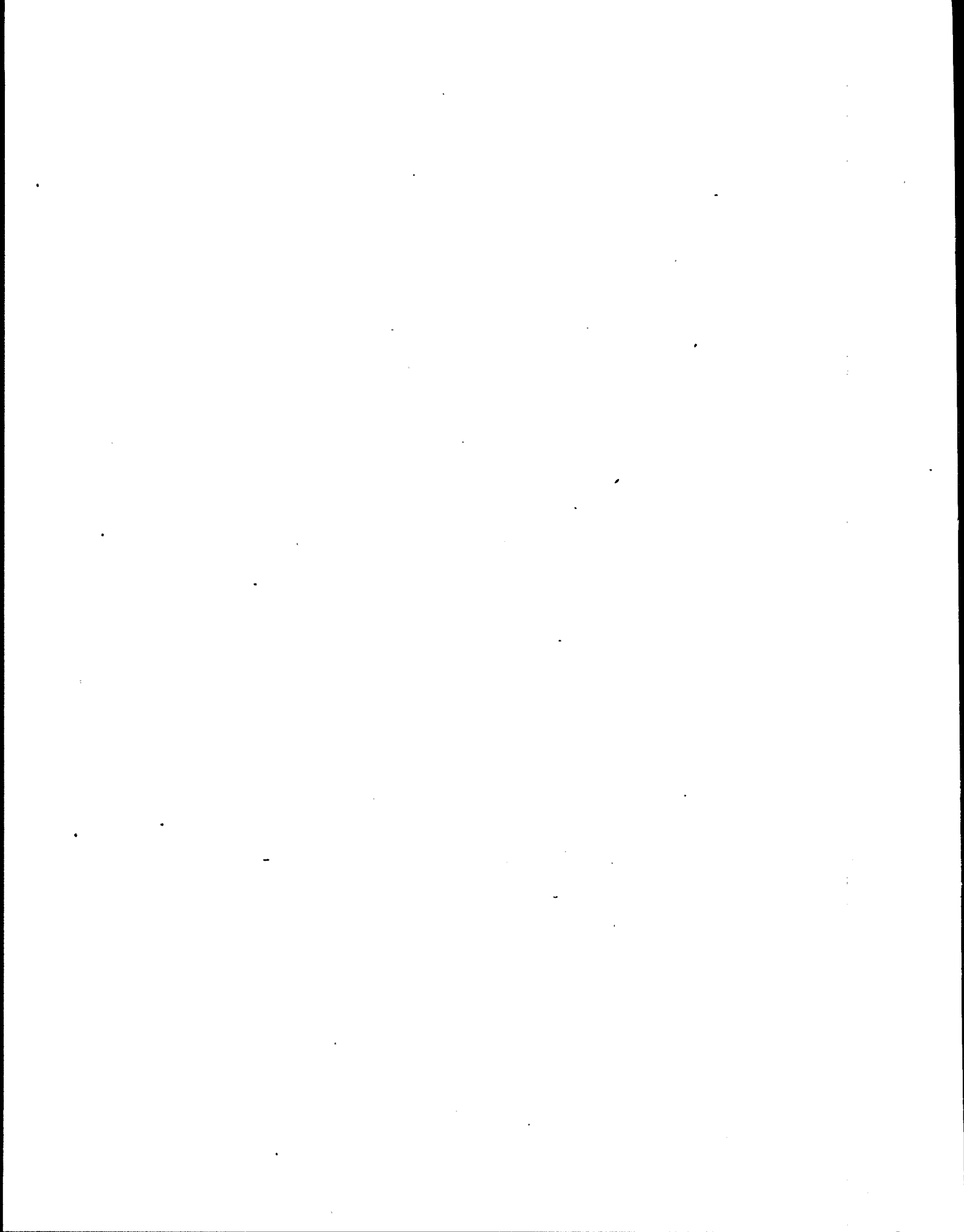
PLANT : SITE 04
 PLANT SITE :
 SAMPLING LOCATION : LOCATION C-ESP OUTLET
 TEST # : 04-MM5T-C3
 DATE : 12/13/84
 TEST PERIOD : 0945-1527

PARAMETER -----	RESULT -----
Vm(dscf)	206.3736
Vm(dscm)	5.844499
Vw gas(scf)	71.24365
Vw gas (scm)	2.01762
% moisture	25.66255
Md	.7433745
MWd	29.3664
MW	26.44949
Vs(fpm)	3152.89
Vs (mpm)	961.247
Flow(acfm)	264411.6
Flow(acmm)	7488.137
Flow(dscfm)	127376.8
Flow(dscmm)	3607.31
% I	108.7766
% EA	26.6349

Program Revision:1/16/84

APPENDIX A-2

CONTINUOUS EMISSION MONITORING RESULTS



ORPE TEST DATA - SITE 04 - TEST 1

FACTOR FOR 3S O2 NORMALIZATION OF OTHER PROCESS GASES			NORMALIZED / CORRECTED DATA - WITH ACTUAL O2						
			TIME	O2 (%)	CO (PPMV)	CO2 (%)	SO2 (PPMV)	NOX (PPMV)	THC (PPMV)
1.1683			1445	5.6	33.6	15.6	93.6	66.2	2.6
1.1765			1450	5.7		15.4	124.6	67.0	2.5
1.1831			1455	5.8	130.3				2.5
1.1726			1500	5.6	168.3	16.7	146.3	66.6	2.5
1.1748			1505	5.7	54.6	14.4	101.7	67.8	2.2
1.1699			1510	5.6		15.4	159.5	67.5	18.7
1.1680			1515	5.6	98.8	14.8	137.5	68.4	9.6
1.1661			1520	5.6		14.9	164.2	65.3	2.3
1.1505			1525	5.3		15.5	103.5	65.0	3.6
1.1862			1530	5.8		15.7	204.1	68.1	2.5
1.1547			1535	5.4		16.3	116.6	67.3	2.2
1.1651			1540	5.5	66.2	16.5	91.4	67.4	7.1
1.1788			1545	5.7	120.5	15.0	86.6	67.7	2.1
1.1860			1550	5.8	84.8	14.2	159.6	68.9	2.0
1.1744			1555	5.7	32.9	15.8	198.2	65.4	2.1
1.1711			1600	5.6	45.2	15.3	89.2	65.5	2.2
1.1809			1605	5.7	73.5	16.5	75.5	68.4	2.2
1.1802			1610	5.7	35.3	16.0	83.5	68.2	
1.1774			1615	5.7		16.9	88.8	69.5	
1.1848			1620	5.8		17.2	68.5	70.8	21.3
1.1823			1625	5.8	152.3	17.6	135.1	70.0	11.2
1.1772			1630	5.7		14.8	118.1	68.7	3.8
1.1894			1635	5.9	20.0	16.5	191.1	72.0	2.0
0.0176			1640						
1.1715			1645	5.6		16.3	262.6	68.8	2.0
1.1704			1650	5.6		16.3	281.2	68.8	1.9
1.1904			1655	5.9	137.0	15.8	371.5	72.6	2.4
1.1397			1700	5.2	84.7	16.6	90.7	67.2	1.5
1.1717			1705	5.6		16.6	101.9	68.5	2.9
1.1414			1710	5.2		15.9	24.2	71.4	3.4
1.1946			1715	5.9	218.4	16.8	63.8	73.2	2.0
1.2350			1720	6.4	33.0	16.9	104.9	77.7	1.7
1.2357			1725	6.4	78.6	14.8	94.6	76.0	1.6
1.2247			1730	6.3	261.4	17.2	123.9	75.8	2.4
1.2155			1735	6.2		16.3	73.9	72.8	1.6
1.2033			1740	6.0	11.4	16.7	83.0	72.1	1.5
1.2121			1745	6.1		16.2	131.3	72.7	2.3
1.2113			1750	6.1	248.8	16.5	54.5	75.2	1.6
1.2256			1755	6.3		17.4	257.3	73.5	1.8
1.2465			1800	6.5	36.6	16.4	323.5	75.3	1.8
1.2370			1805	6.4	10.4	14.3	366.4	76.6	1.7
1.1415			1815	5.2		16.5		72.5	7.6
1.1395			1820	5.2	39.9	15.9	18.4	71.1	16.3
1.1953			1830	5.9	157.3	15.7	19.1	76.5	9.4
1.1848			1835	5.8		15.6		75.0	2.0
1.1818			1840	5.8	29.2	16.7	3.2	71.9	1.7
1.1546			1845	5.4	129.8	16.4	0.3	75.7	1.7
1.1822			1850	5.8		16.5	8.2	76.7	
1.2243			1855	6.3		15.7	16.6	76.4	4.9
1.2162			1900	6.2	80.9	17.4	14.4	78.0	4.8
1.2240			1905	6.3	106.7	16.0	27.6	79.1	3.0
1.2277			1910	6.3	201.6	15.7	165.3	77.9	3.1
1.2029			1915	6.0	95.6	17.8	185.4	77.8	1.6
1.2102			1920	6.1	106.6	15.8	257.5	77.6	1.2
1.2152			1925	6.2	1.4	16.8	366.3	77.1	1.2
1.1918			1930	5.9	131.3	15.8	44.8	78.7	1.0
1.1864			1935	5.8	64.4	16.0	49.3	78.9	1.5
1.2090			1940	6.1		16.1	173.9	66.9	1.0
1.1979			1945	6.0		16.2	139.6	78.7	1.7
1.2076			1950	6.1	143.8	16.0	109.6	79.0	1.0
1.1953			1955	5.9		16.7	64.9	79.0	1.0
1.2168			2000	6.2	107.6	16.7	187.0	78.7	0.9
1.2017			2005	6.0		16.6	130.6	77.6	0.8
1.2117			2010	6.1	66.5	16.5	36.3	77.0	
1.1945			2015	5.9	33.5	16.4	15.7	76.3	2.8
1.2126			2020	6.1	78.2	16.7	40.4	78.9	1.9
1.2313			2025	6.4		16.8	27.9	82.2	3.6
1.2264			2030	6.3	197.9	16.3	28.6	82.1	2.2
1.2080			2035	6.1		16.1	20.3	81.6	3.4
1.2065			2040	6.1	6.7	16.3	47.9	37.9	2.6
1.2125			2045	6.1		16.8	326.5	78.5	3.2
1.2028			2050	6.0	3.6	16.7	93.0	81.3	4.9
1.1863			2055	5.8	11.3	17.0	128.6	80.6	1.6
1.1949			2100	5.9	74.8	17.1	219.7	81.6	1.0
1.2217			2105	6.2		16.6	389.6	83.3	1.0
1.2358			2110	6.4		16.8	232.7	83.1	0.9
1.2344			2115	6.4	227.8	17.1	117.2	82.7	0.8
1.2231			2120	6.3	15.8	16.7	87.6	82.6	1.5
1.2212			2125	6.2	15.7	16.8	437.0	80.7	0.9
1.2197			2130	6.2		17.1	100.3	84.8	0.6
1.2078			2135	6.1	92.5	17.1	86.2	85.2	1.0
1.2102			2140	6.1		16.3	19.0	81.2	0.7
1.2043			2145	6.0	2.5	17.4	10.0	85.4	0.8
1.2133			2150	6.1		16.6	8.0	83.3	0.7
1.1801			2155	5.7		16.5	6.2	79.7	1.1
1.1873			2200	5.8	28.5	16.5	9.0	84.2	0.8
1.2020			2205	6.0	119.0	17.8	6.7	85.7	1.5
1.2110			2210	6.1	89.4	17.1	14.1	83.4	2.2
1.2103			2215	6.1	19.9	16.6	0.5	85.8	1.2
1.2269			2220	6.3		16.8		86.5	5.5
1.2225			2225	6.3	142.5	17.4		87.1	1.2
1.2264			2230	6.3	65.3	16.5	9.1	85.5	0.7
1.2208			2235	6.2	6.4	17.2	10.2	84.9	0.7
1.2201			2240	6.2		17.3		72.9	0.7
1.2165			2245	6.2	55.9	16.2	12.7	84.1	2.9
1.2157			2250	6.2	75.6			63.5	3.0

NO. PTS. 95
MEAN 1.1976
STD. DEV. .0

NO. PTS. 95
MEAN 5.9
STD. DEV. 0.3

NO. PTS. 60
MEAN 84.4
STD. DEV. 65.2

NO. PTS. 93
MEAN 16.3
STD. DEV. 0.8

NO. PTS. 90
MEAN 112.1
STD. DEV. 102.1

NO. PTS. 94
MEAN 75.1
STD. DEV. 7.4

NO. PTS. 91
MEAN 2.5
STD. DEV. 3.6

CORRECTED DATA - SITE 04 - TEST 2

** FACTOR			NORMALIZED / CORRECTED DATA - WITH ACTUAL 02						
** FOR 3% O2									
** NORMALIZATION									
** OF									
** OTHER PROCESS									
** GASES			TIME	O2	CO	CO2	SO2	NOX	THC
**				(%)	(PPMV)	(%)	(PPMV)	(PPMV)	(PPMV)
**			*****	*****	*****	*****	*****	*****	*****
**	1.1316	**	1100	5.1	198.3	15.0	81.6	83.7	6.1
**	1.1799	**	1115	5.7	155.2	14.8	90.1	87.9	3.5
**	1.1729	**	1120	5.6	136.5	14.1	62.1	91.5	3.1
**	1.1655	**	1125	5.5	76.5	15.0	68.5	90.6	2.7
**	1.1455	**	1130	5.3		12.4	29.3	90.2	2.8
**	1.1481	**	1135	5.3	250.4	12.7	27.8	90.5	2.7
**	1.1612	**	1140	5.5	288.8	13.9	59.8	89.5	2.6
**	1.1430	**	1145	5.2	328.0	13.0	46.7	86.9	2.8
**	1.1661	**	1150	5.6	137.1	14.8	55.3	102.4	3.9
**	1.1582	**	1155	5.4	232.1	15.4	92.2	88.5	2.7
**	1.1447	**	1200	5.3	319.6	14.3	24.4	90.4	2.6
**	1.1462	**	1205	5.3	192.5	12.3	25.7	92.3	2.7
**	1.1508	**	1210	5.3	134.3	15.3	12.2	95.4	2.7
**	1.1594	**	1215	5.5	152.2	15.0	43.0	93.8	2.3
**	1.1619	**	1220	5.5	133.4	16.3	19.1	93.8	2.2
**	1.1639	**	1225	5.5	181.4	14.1	19.1	93.2	2.1
**	1.1437	**	1230	5.2	161.2	14.1	17.9	92.7	2.0
**	1.1637	**	1235	5.5	314.7	14.7	17.6	93.4	2.0
**	1.1540	**	1240	5.4	154.4	15.2	13.6	91.4	1.8
**	1.1651	**	1245	5.5	267.7	14.0	32.6	92.5	1.9
**	1.1634	**	1250	5.5	153.1	13.4	17.6	91.8	1.8
**	1.1633	**	1255	5.5	64.6	14.8	10.4	89.3	2.1
**	1.1678	**	1300	5.6	181.1	15.5	5.1	92.6	2.0
**	1.1759	**	1305	5.7	184.3	16.3	3.3	92.7	2.4
**	1.1631	**	1310	5.5	203.3	14.5	2.5	93.6	2.0
**	1.1724	**	1315	5.6	98.0	14.3	0.9	94.8	1.8
**	1.1830	**	1320	5.8		14.6	18.1	90.5	17.2
**	1.1840	**	1325	5.8		13.3	15.9	90.5	2.2
**	1.1900	**	1330	5.9	179.7	15.6	40.4	89.6	5.4
**	1.1876	**	1335	5.8	62.1	15.8	39.7	92.0	3.1
**	1.2138	**	1340	6.2	333.6	14.6	56.3	88.5	3.1
**	1.1638	**	1345	5.5		11.8	48.7	87.3	2.0
**	1.1947	**	1350	5.9	117.8	15.5	168.2	87.6	2.0
**	1.1864	**	1355	5.8		14.7	63.8	84.4	1.8
**	1.1927	**	1400	5.9	119.1	14.7	83.4	86.1	1.9
**	1.1633	**	1405	5.5	252.0	13.3	81.1	81.8	1.8
**	1.1947	**	1410	5.9	122.3	15.3	59.7	86.6	1.9
**	1.1900	**	1415	5.9		14.6	59.4	86.3	1.8
**	1.1781	**	1430	5.7		12.1	13.6	81.4	2.3
**	1.1739	**	1435	5.7	69.5	15.1	30.8	81.7	1.9
**	1.2887	**	1440	7.0	118.8	18.7	32.9	90.3	1.8
**	1.1725	**	1445	5.6	141.4	13.4	66.1	80.5	1.6
**	1.2071	**	1450	6.1		12.9	91.1	86.2	3.2
**	1.2049	**	1455	6.0	78.2	13.5	61.2	84.0	2.2
**	1.2191	**	1500	6.2		14.8	157.3	87.7	1.1
**	1.2131	**	1505	6.1	41.2	15.6	28.8	82.8	3.8
**	1.3780	**	1515	7.9		16.0	69.5	100.9	1.8
**	1.1770	**	1520	5.7	43.1	15.6	164.2	86.2	4.0
**	1.1886	**	1525	5.8	58.8	16.2	171.7	86.2	1.9
**	1.1940	**	1530	5.9	75.8	17.5	134.1	86.2	1.7
**	1.1894	**	1535	5.9	60.0	15.0	97.8	86.2	1.9
**	1.1834	**	1540	5.8	145.5	15.0	85.6	82.5	2.2
**	1.1846	**	1605	5.8	106.2	15.9	28.8	87.8	6.1
**	1.1740	**	1620	5.7	191.4	16.7	74.0	84.2	1.8

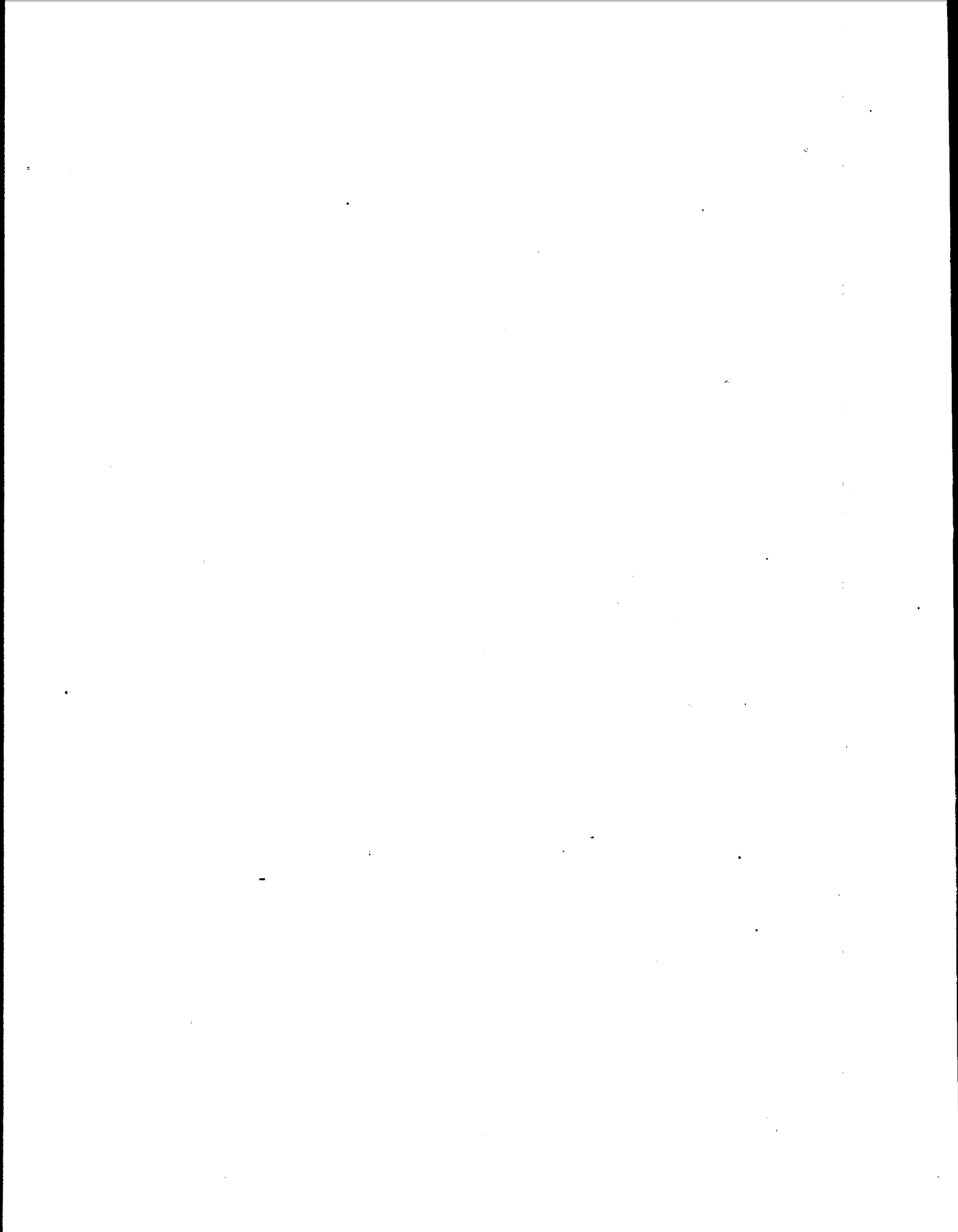
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STD. DEV.	.0	STD. DEV.	0.4	79.4	1.3	43.0	4.5	2.2	

CORRECTED DATA - SITE 04 - TEST 3

FACTOR FOR 3% O2 NORMALIZATION OF OTHER PROCESS GASES			NORMALIZED / CORRECTED DATA - WITH ACTUAL O2						
			TIME	O2 (%V)	CO (PPMV)	CO2 (%V)	SO2 (PPMV)	NOX (PPMV)	THC (PPMV)
*****			*****	*****	*****	*****	*****	*****	*****
**	1.2899	**	945	7.0	320.7	13.0	144.9	81.0	55.3
**	1.3077	**	950	7.2	3.4	14.9	167.9	81.0	12.5
**	1.2923	**	955	7.0		14.9	245.4	85.6	10.0
**	1.2788	**	1000	6.9	12.2	13.5	165.2	84.7	13.7
**	1.2728	**	1005	6.8	143.0	14.2	154.5	86.6	7.4
**	1.2651	**	1010	6.8	267.0	16.1	38.5	86.9	5.9
**	1.2615	**	1015	6.7	914.1	14.4	119.1	76.1	5.4
**	1.3276	**	1020	7.4	34.0	15.2	100.9	87.4	6.8
**	1.3182	**	1025	7.3	339.9	16.9	60.9	89.3	5.6
**	1.3105	**	1030	7.2	108.8	15.6	79.5	87.4	
**	1.2557	**	1055	6.6	51.9	15.6	131.7	84.6	3.9
**	1.2411	**	1110	6.5		15.0	208.9	86.2	2.5
**	1.2486	**	1115	6.6	370.4	13.9	46.9	84.2	2.7
**	1.2429	**	1120	6.5	399.6	15.8	41.9	81.9	8.4
**	1.2520	**	1125	6.6	261.0	15.4	56.6	82.7	4.8
**	1.2427	**	1130	6.5	235.0	16.1	27.3	86.5	4.5
**	1.2832	**	1135	6.9	222.8	14.2	60.4	85.8	3.6
**	1.2721	**	1140	6.8	7.5	14.0	132.3	88.0	3.4
**	1.2845	**	1155	7.0	41.2	15.8	126.5	89.3	3.5
**	1.4854	**	1230	8.8		17.4	84.6	107.5	5.2
**	1.3169	**	1235	7.3	114.7	15.5	54.3	97.4	3.5
**	1.3164	**	1240	7.3	174.5	15.1	43.5	96.4	3.3
**	1.2143	**	1355	6.2	58.4	13.0	122.3	82.7	5.4
**	1.2228	**	1400	6.3	202.4	15.4	146.7	86.4	4.3
**	1.2418	**	1410	6.5		16.1	120.3	86.2	3.5
**	1.2465	**	1415	6.5		16.0	128.2	85.5	3.6
**	1.2223	**	1420	6.3	158.4	17.7	61.9	83.6	3.7
**	1.2140	**	1425	6.2	111.8	15.3	82.2	85.6	8.5
**	1.2108	**	1430	6.1		14.9	80.8	88.0	5.2
**	1.2135	**	1435	6.1	198.0	17.5	84.8	85.5	3.9
**	1.2581	**	1440	6.7	41.4	16.2	35.7	86.5	3.8
**	1.2233	**	1445	6.3	162.0	16.2	153.0	83.6	3.5
**	1.2269	**	1450	6.3	165.9	16.1	176.2	82.8	3.7
**	1.2221	**	1455	6.3		14.1	139.1	80.3	3.2
**	1.2358	**	1500	6.4	121.3	16.4	110.5	84.3	3.0
**	1.2577	**	1505	6.7	27.8	15.1	151.7	85.0	2.9
**	1.2285	**	1510	6.3	104.1	16.3	257.4	83.9	2.8
**	1.2135	**	1515	6.1	59.9	18.6	271.5	82.1	2.8
**	1.2122	**	1520	6.1	66.0	15.6	298.1	83.4	2.9
**	1.2215	**	1525	6.2	150.4	13.6	196.2	82.3	2.9
**	1.2495	**	1530	6.6	318.0	14.7	100.5	81.8	3.2
**	1.2470	**	1535	6.5	118.8	14.1	136.5	83.7	3.1
**	1.2217	**	1540	6.2	163.6	18.5	255.5	80.2	4.0
**	1.2659	**	1545	6.8	20.0	17.1	272.3	85.1	3.2
**	1.2654	**	1550	6.8	67.1	16.2	307.7	86.0	3.2
**	1.2423	**	1555	6.5	42.2	18.1	227.9	84.6	3.9
**	1.2521	**	1600	6.6	130.6	18.4	162.2	86.7	3.7
**	1.2709	**	1605	6.8	38.9	19.3	334.9	88.4	3.1
**	1.2669	**	1610	6.8	216.6	14.4	388.2	89.6	2.9
**	1.2062	**	1615	6.1		16.9	153.0	81.6	2.8
**	1.2202	**	1620	6.2	107.9	16.5	68.3	82.6	2.7
**	1.2463	**	1625	6.5	293.5	17.0	141.6	87.0	2.8
**	1.2426	**	1630	6.5	34.1	16.2	87.3	85.4	2.8
**	1.2231	**	1635	6.3	316.8	15.3	47.7	86.5	2.9
**	1.2340	**	1640	6.4	42.1	14.4	119.7	88.8	3.2
**	1.2251	**	1645	6.3	156.0	19.2	98.7	88.2	3.0
**	1.2088	**	1650	6.1	96.4	16.5	95.0	84.6	3.0
**	1.2005	**	1655	6.0	27.5	13.8	63.9	83.2	2.9
**	1.1919	**	1700	5.9	66.8	16.2	55.7	84.9	3.1
**	1.2260	**	1705	6.3	58.4	16.7	83.3	84.9	3.0
**	1.2012	**	1710	6.0	366.9	18.0	16.4	83.3	2.7
**	1.2095	**	1715	6.1	151.9	16.2	75.5	83.1	2.6
**	1.2053	**	1720	6.0	435.9	16.4	18.8	83.3	2.3
**	1.2078	**	1725	6.1	47.8	16.7	63.8	84.5	3.0
**	1.2490	**	1730	6.6	33.0	15.2	61.0	90.8	3.0
**	1.2342	**	1735	6.4	171.2	17.1	12.6	88.3	2.6
**	1.2425	**	1740	6.5		14.4	24.4	88.8	2.3
**	1.2163	**	1745	6.2	400.2	15.5	4.5	83.9	2.2
**	1.2215	**	1750	6.2		16.5	6.5	82.8	9.6
**	1.2307	**	1755	6.4	270.5	18.3	3.9	83.0	3.0
**	1.2077	**	1800	6.1	143.9	16.0	8.6	78.4	3.0
**	1.2199	**	1805	6.2	277.9	17.9	20.9	80.9	0.7

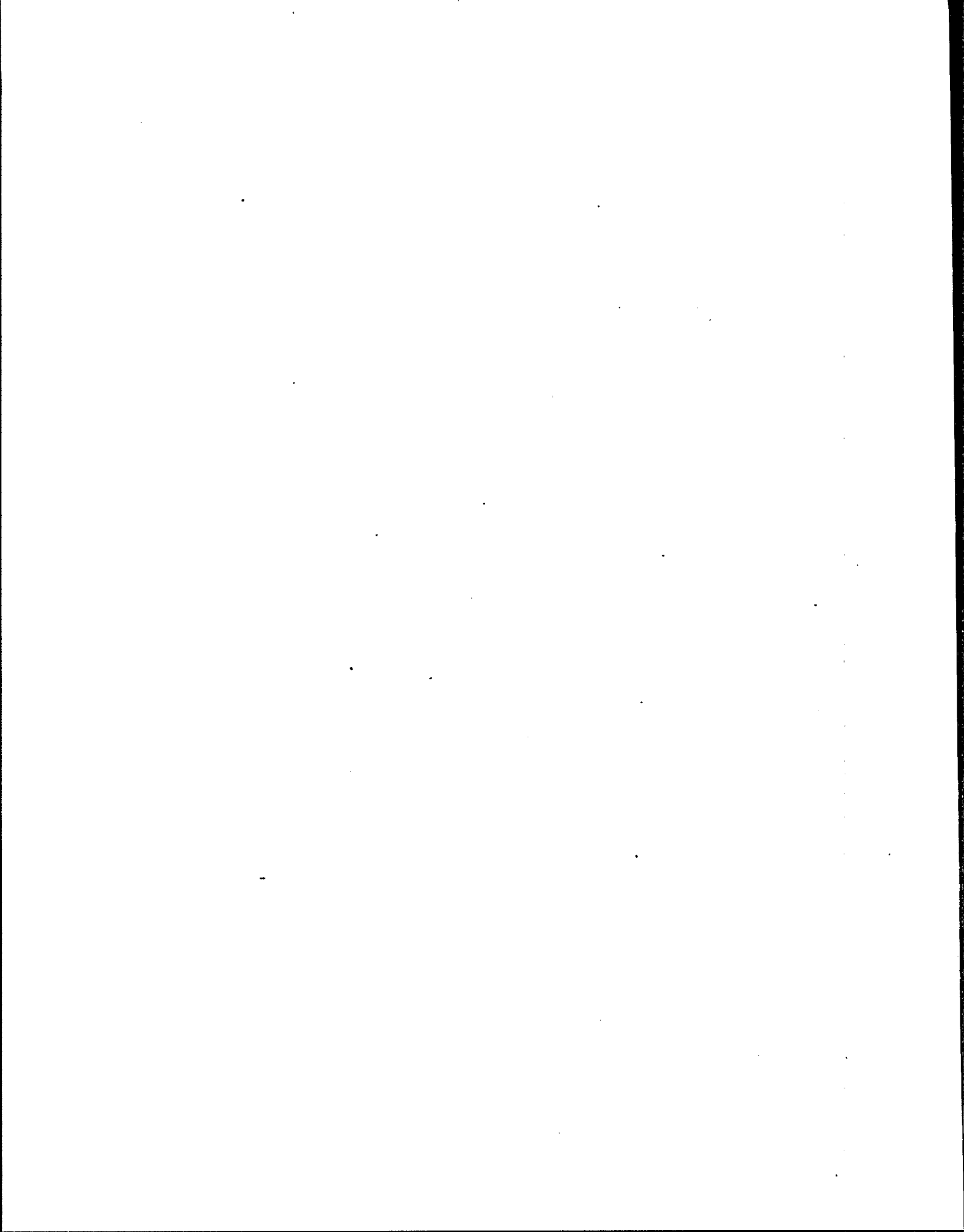
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APPENDIX A-3

HCl TRAIN RESULTS



R A D I A N S O U R C E T E S T
E P A M E T H O D 2 - 5
(R A W D A T A)

PLANT : SITE 04
PLANT SITE :
SAMPLING LOCATION : LOCATION C-ESP OUTLET
TEST # : 04-HCL-C-01
DATE : 12/11/84
TEST PERIOD : 1425-1719

PARAMETER -----	VALUE -----
Sampling time (min.)	174
Barometric Pressure (in.Hg)	29.58
Sampling nozzle diameter (in.)	.305
Meter Volume (cu.ft.)	144.175
Meter Pressure (in.H2O)	2.483333
Meter Temperature (F)	95.86111
Stack dimension (sq.in.)	12076.31
Stack Static Pressure (in.H2O)	-.85
Stack Moisture Collected (gm)	914.9
Absolute stack pressure(in Hg)	29.5175
Average stack temperature (F)	344.8333
Percent CO2	14.7
Percent O2	4.27
Percent N2	76.9
Delps Subroutine result	21.11785
DGM Factor	1.0037
Pitot Constant	.84

R A D I A N S O U R C E T E S T
E P A M E T H O D S 2 - 5
F I N A L R E S U L T S

PLANT : SITE 04
PLANT SITE :
SAMPLING LOCATION : LOCATION C-ESP OUTLET
TEST # : 04-HCL-C-01
DATE : 12/11/84
TEST PERIOD : 1425-1719

PARAMETER -----	RESULT -----
Vm(dscf)	136.7322
Vm(dscm)	3.872256
Vw gas(scf)	43.13754
Vw gas (scm)	1.221655
% moisture	23.98265
Md	.7601735
MWd	29.3664
MW	26.64044
Vs(fpm)	3244.78
Vs (mpm)	989.2621
Flow(acfm)	272117.8
Flow(acmm)	7706.377
Flow(dscfm)	133880
Flow(dscmm)	3791.481
% I	97.07468
% EA	26.6349

Program Revision:1/16/84

R A D I A N S O U R C E T E S T
E P A M E T H O D 2 - 5
(R A W D A T A)

PLANT : SITE 04
PLANT SITE :
SAMPLING LOCATION : LOCATION C-ESP OUTLET
TEST # : 04-HCL-C-02
DATE : 12/12/84
TEST PERIOD : 1535-1705

PARAMETER -----	VALUE -----
Sampling time (min.)	100
Barometric Pressure (in.Hg)	29.58
Sampling nozzle diameter (in.)	.305
Meter Volume (cu.ft.)	80.706
Meter Pressure (in.H2O)	2.31
Meter Temperature (F)	31.7
Stack dimension (sq.in.)	12076.31
Stack Static Pressure (in.H2O)	-.85
Stack Moisture Collected (gm)	547.4
Absolute stack pressure(in Hg)	29.5175
Average stack temperature (F)	340.9
Percent CO2	14.7
Percent O2	4.27
Percent N2	76.9
Delps Subroutine result	20.45646
DGM Factor	1.0037
Pitot Constant	.84

R A D I A N S O U R C E T E S T
 E P A M E T H O D S 2 - 5
 F I N A L R E S U L T S
 PLANT : SITE 04
 PLANT SITE :
 SAMPLING LOCATION : LOCATION C-ESP OUTLET
 TEST # : 04-HCL-C-02
 DATE : 12/12/84
 TEST PERIOD : 1535-1705

PARAMETER -----	RESULT -----
Vm(dscf)	86.49016
Vm(dscm)	2.449401
Vw gas(scf)	25.80991
Vw gas (scm)	.7309367
% moisture	22.98299
Md	.7701701
MWd	29.3664
MW	26.75406
Vs(fpm)	3136.474
Vs (mpm)	956.2419
Flow(acfm)	263034.9
Flow(acmm)	7449.148
Flow(dscfm)	131757
Flow(dscmm)	3731.357
% I	108.5658
% EA	26.6349

Program Revision:1/16/8

R A D I A N S O U R C E T E S T
E P A M E T H O D 2 - 5
(R A W D A T A)

PLANT : SITE 04
PLANT SITE :
SAMPLING LOCATION : LOCATION C-ESP OUTLET
TEST # : 04-HCL-C-03
DATE : 12/13/84
TEST PERIOD : 0945-1145

PARAMETER

VALUE

Sampling time (min.)	120
Barometric Pressure (in.Hg)	29.5
Sampling nozzle diameter (in.)	.305
Meter Volume (cu.ft.)	96.01196
Meter Pressure (in.H2O)	2.233333
Meter Temperature (F)	34.04167
Stack dimension (sq.in.)	12076.31
Stack Static Pressure (in.H2O)	-.85
Stack Moisture Collected (gm)	710.1
Absolute stack pressure(in Hg)	29.4375
Average stack temperature (F)	341.1667
Percent CO2	14.7
Percent O2	4.27
Percent N2	76.9
Delp's Subroutine result	20.1104
DGM Factor	1.0037
Pitot Constant	.84

R A D I A N S O U R C E T E S T
 E P A M E T H O D S 2 - 5
 F I N A L R E S U L T S
 PLANT : SITE 04
 PLANT SITE :
 SAMPLING LOCATION : LOCATION C-ESP OUTLET
 TEST # : 04-HCL-C-03
 DATE : 12/13/84
 TEST PERIOD : 0945-1145

PARAMETER -----	RESULT -----
Vm(dscf)	102.1106
Vm(dscm)	2.891773
Vw gas(scf)	33.48121
Vw gas (scm)	.9481879
% moisture	24.69265
Md	.7530735
MWd	29.3664
MW	26.55974
Vs(fpm)	3098.876
Vs (mpm)	944.7794
Flow(acfm)	259881.9
Flow(acmm)	7359.854
Flow(dscfm)	126900.6
Flow(dscmm)	3593.826
% I	110.8986
% EA	26.6349

Program Revision:1/16/8

R A D I A N S O U R C E T E S T
E P A M E T H O D 2 - 5
(R A W D A T A)

PLANT : SITE 04
PLANT SITE :
SAMPLING LOCATION : LOCATION C-ESP OUTLET
TEST # : 04-HCL-C-04
DATE : 12/13/84
TEST PERIOD : 1435-1628

PARAMETER -----	VALUE -----
Sampling time (min.)	113
Barometric Pressure (in.Hg)	29.5
Sampling nozzle diameter (in.)	.305
Meter Volume (cu.ft.)	96.01196
Meter Pressure (in.H2O)	2.258334
Meter Temperature (F)	33.66667
Stack dimension (sq.in.)	12076.31
Stack Static Pressure (in.H2O)	-.85
Stack Moisture Collected (gm)	667.3
Absolute stack pressure(in Hg)	29.4375
Average stack temperature (F)	340.5
Percent CO2	14.7
Percent O2	4.27
Percent N2	76.9
Delp's Subroutine result	20.2647
DGM Factor	1.0037
Pitot Constant	.84

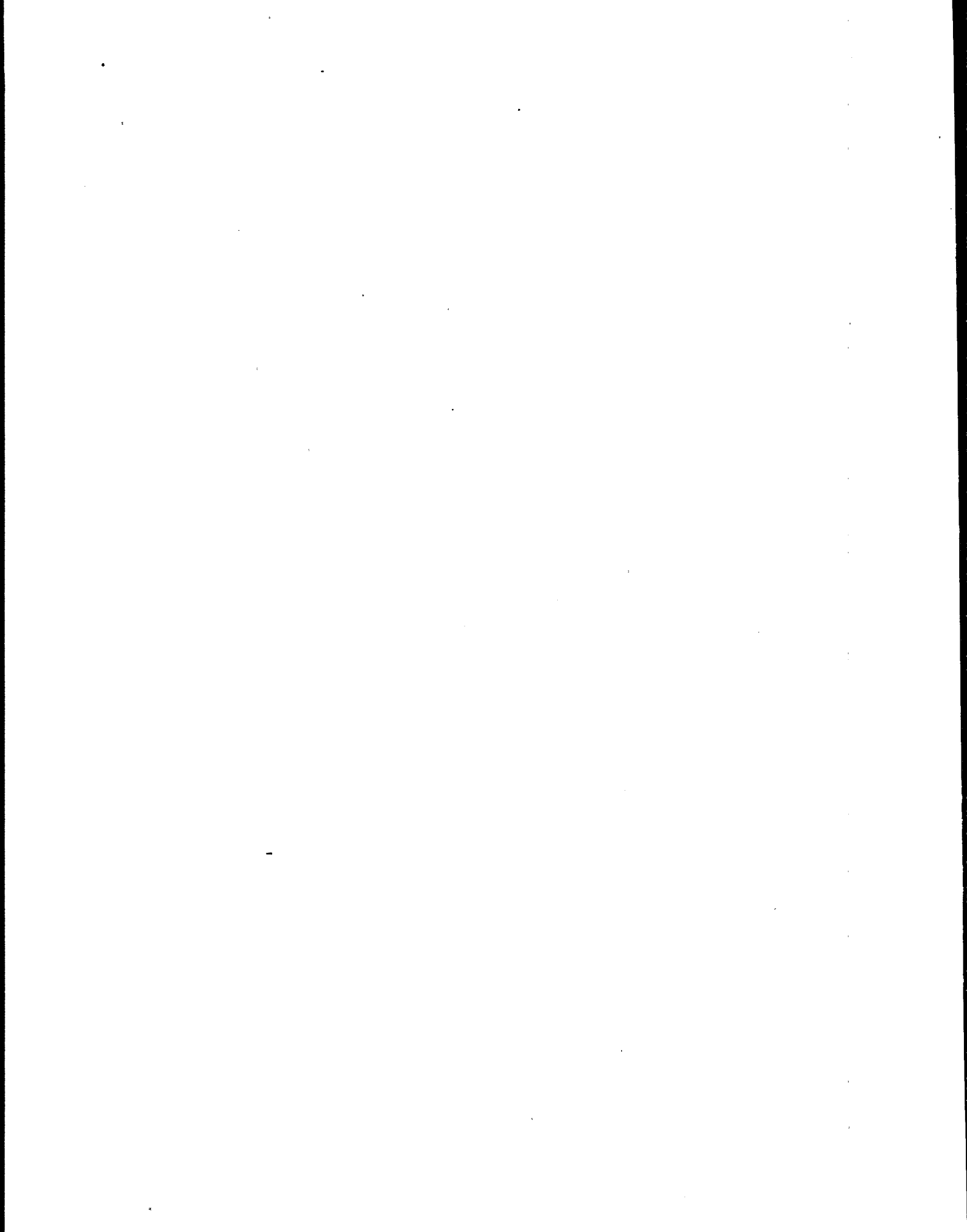
R A D I A N S O U R C E T E S T
 E P A M E T H O D S 2 - 5
 F I N A L R E S U L T S
 PLANT : SITE 04
 PLANT SITE :
 SAMPLING LOCATION : LOCATION C-ESP OUTLET
 TEST # : 04-HCL-C-04
 DATE : 12/13/84
 TEST PERIOD : 1435-1628

PARAMETER -----	RESULT -----
Vm(dscf)	102.1945
Vm(dscm)	2.894148
Vw gas(scf)	31.4632
Vw gas (scm)	.8910377
% moisture	23.54013
Md	.7645987
MWd	29.3664
MW	26.69074
Vs(fpm)	3114.981
Vs (mpm)	949.6891
Flow(acfm)	261232.4
Flow(acmm)	7398.102
Flow(dscfm)	129620.2
Flow(dscmm)	3670.843
% I	115.3922
% EA	26.6349

Program Revision:1/16/8

APPENDIX B

PROCESS MONITORING DATA



APPENDIX B-1

HOURLY AVERAGE VALUES
OF BOILER OPERATING PARAMETERS

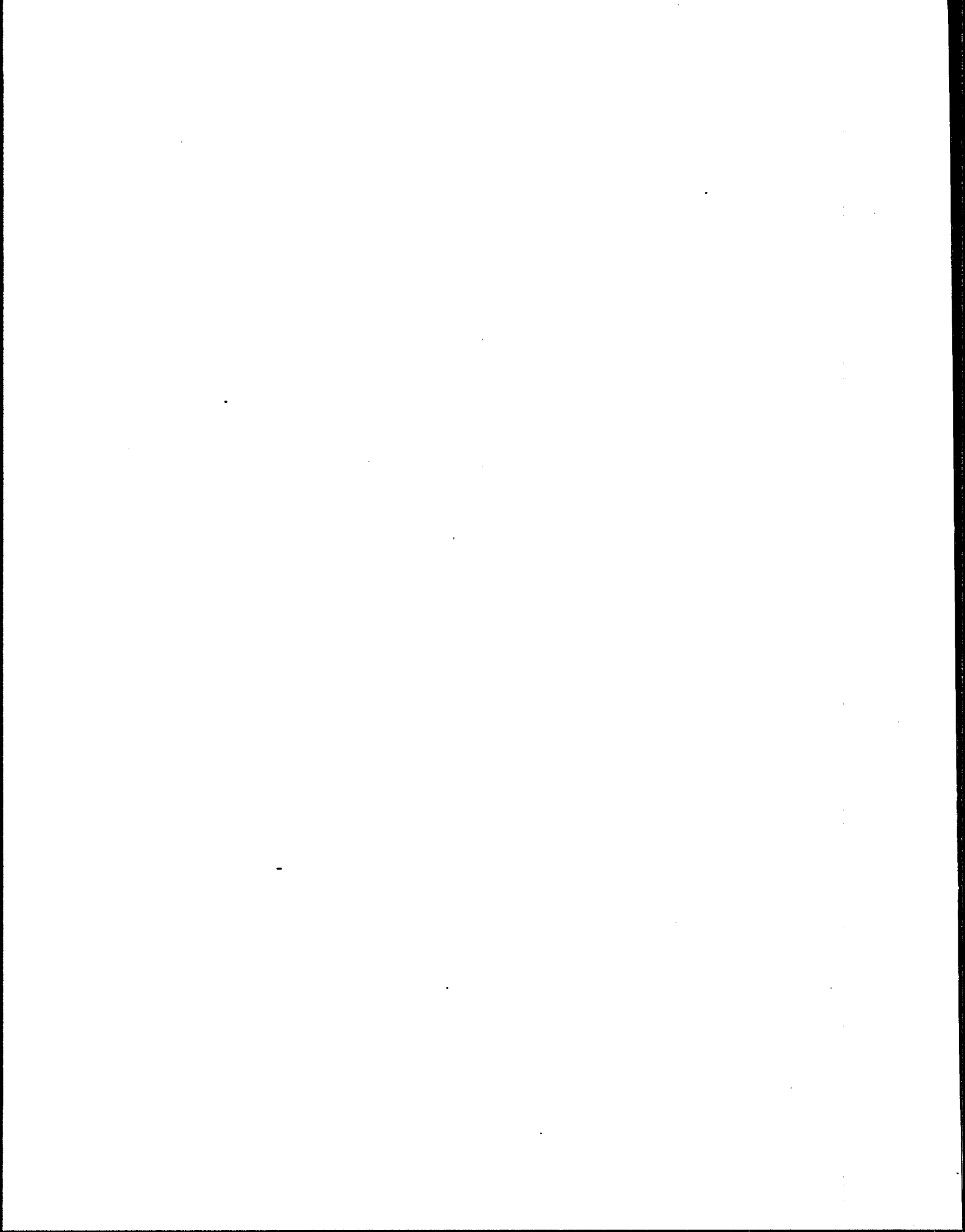


Table 1. Hourly Average Values^a of Various
Boiler Parameters, Run #1

Time ^b	Boiler Load 10 ³ lb/hr	Boiler Oxygen %O ₂ ^d	Stack Oxygen ^e %O ₂	Stack CO ^f ppmCO	Black Liq. Flow gpm	Black Liq. % Solids wt%	Stack TRS ^g ppm@8.70	Stack Opacity (%)
1500	396	3.6	5.9	1.6	258	64.9	3.8	15.0
1600	385	4.1	6.0	6.2	255	65.2	3.7	15.0
1700	388	4.0	6.3	0.7	253	65.5	4.2	15.1
1800	401	4.0	6.5	3.2	254	65.7	4.2	15.0
1900	382	4.4	6.5	1.8	259	65.8	4.0	15.0
2000	391	4.3	6.3	1.8	259	65.8	4.1	15.1
2100	387	4.3	6.9	2.5	257	65.9	4.1	15.0
2200	384	4.0	6.9	5.6	256	66.0	3.9	14.7
2300	390	3.8	6.7	9.6	255	66.0	4.2	14.9
AVG	389	4.1	6.4	3.7	256	65.6	4.0	15.0

a. Values shown in units used by host plant to convert

from:	to:	multiply by:
lb/hr	kg/hr	0.454
gpm	cumeter/min	0.00379

b. Value shown for time "t" is the average value for the one hour period preceding time "t"

c. Boiler load in 10³lb/hr steam @ 914.7 psi, 1360°R (1452 Btu/lb)

d. Boiler oxygen measured in superheater section

e. Stack oxygen measured at ESP outlet stack (wet basis)

f. Stack CO measured at ESP outlet stack (wet basis, O₂ as measured)

g. Stack TRS measured at ESP Outlet stack (dry basis, corrected to 8% O₂)

Table 2. Hourly Average Values^a of Various
Boiler Parameters, Run #2

Time ^b	Boiler Load ^c 10 ³ lb/hr	Boiler Oxygen ^d %O ₂	Black Liq. Flow gpm	Black Liq. % Solids wt%	Stack Oxygen ^e %O ₂	Stack CO ^f ppmv	Stack TRS ^g ppm ^v @8%O ₂	Stack Opacity (%)
1100	388	3.8	254	65.6	cal	33.5	cal	14.9
1200	391	3.3	256	65.6	cal	57.2	cal	15.1
1300	386	3.6	255	65.6	cal	24.1	cal	15.1
1400	391	3.6	254	65.5	5.9	10.9	4.2	14.6
1500	390	3.8	254	65.5	5.9	3.3	4.4	15.1
1600	380	3.4	253	65.6	6.0	3.3	4.7	14.9
1700	398	3.7	253	65.6	6.1	5.5	4.6	14.9
Avg	389	3.6	254	65.6	6.0	17.4	4.5	15.0

cal = no data available for O₂, TRS. Instrument calibrations in progress by host plant.

- Values shown in units used by host plant to convert
from: to: multiply by:
lb/hr kg/hr 0.454
gpm cumeter/min 0.00379
- Value shown for time "t" is the average value for the one hour period preceding time "t"
- Boiler load in 10³lb/hr steam @ 914.7 psi, 1360°R (1452 Btu/lb)
- Boiler oxygen measured in superheater section
- Stack oxygen measured at ESP outlet stack (wet basis)
- Stack CO measured at ESP outlet stack (wet basis, O₂ as measured)
- Stack TRS measured at ESP Outlet stack (dry basis, corrected to 8% O₂)

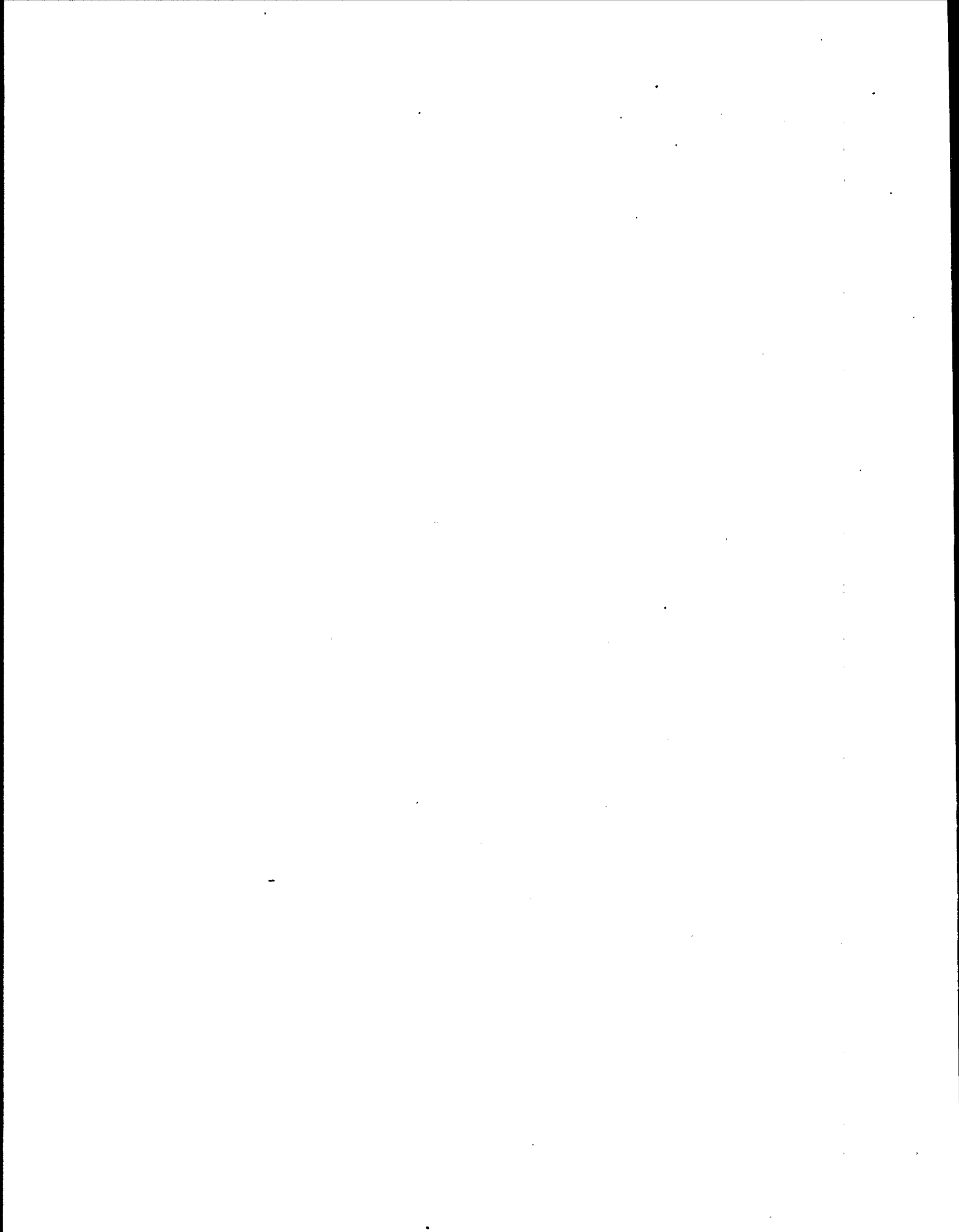
Table 3. Hourly Average Values^a of Various
Boiler Parameters, Run #3

Time ^b	Boiler Load ^c 10 ³ lb/hr	Boiler Oxygen ^d %O ₂	Black Liq. Flow gpm	Black Liq. % Solids wt%	Stack Oxygen ^e %O ₂	Stack CO ^f ppmv	Stack TRS ^g ppm ^v @8%O ₂	Stack Opacity (%)
1000	390	3.6	261	64.3	cal	53	cal	14.9
1100	382	3.1	263	64.0	cal	71	cal	15.0
1200	393	2.8	265	63.6	5.9	86	3.1	15.0
1300	384	2.9	265	63.5	5.4	96	3.3	15.1
1400	386	2.8	265	63.5	5.5	55	3.7	14.8
1500	387	2.9	264	63.5	5.4	79	3.6	14.9
1600	ND	ND	ND	ND	ND	ND	ND	ND
1700	ND	ND	ND	ND	ND	ND	ND	ND
1800	403	3.6	263	63.8	5.4	16	3.9	15.0
Mean	389	3.1	264	63.7	5.5	65	3.5	15.0

cal = no data available for O₂, TRS. Instrument calibrations in progress by plant personnel.

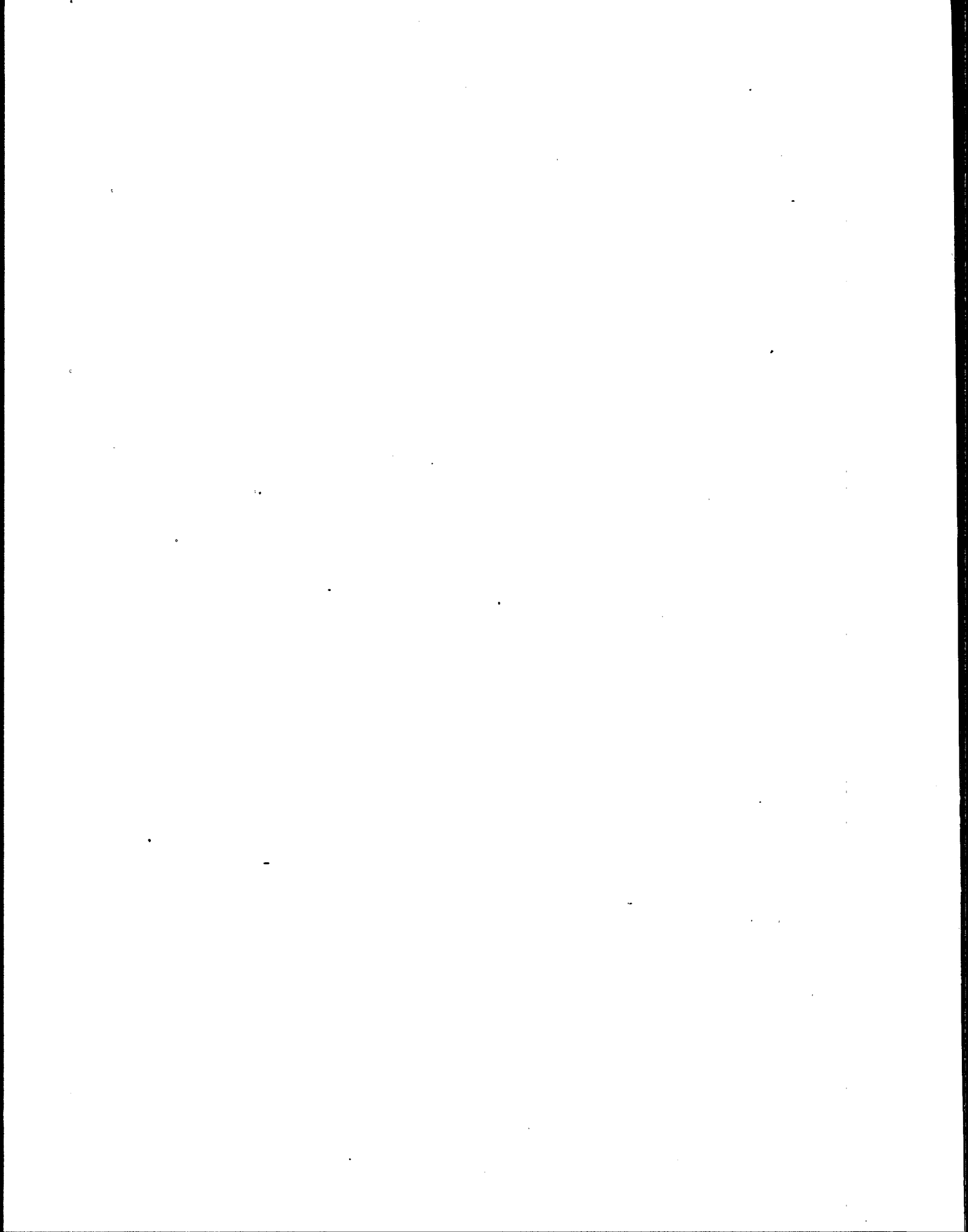
ND = No data available due to computer failure.

- Values shown in units used by host plant to convert
from: to: multiply by:
lb/hr kg/hr 0.454
gpm cumeter/min 0.00379
- Value shown for time "t" is the average value for the one hour period preceding time "t"
- Boiler load in 10³lb/hr steam @ 914.7 psi, 1360°R (1452 Btu/lb)
- Boiler oxygen measured in superheater section
- Stack oxygen measured at ESP outlet stack (wet basis)
- Stack CO measured at ESP outlet stack (wet basis, O₂ as measured)
- Stack TRS measured at ESP Outlet stack (dry basis, corrected to 8% O₂)



APPENDIX B-2

PLANT COMPUTER OUTPUT OF PROCESS DATA



POINT	DESCRIPTION	ENGR. UNITS	CURRENT VALUE	CURRENT AVERAGES		
				HOUR	SHIFT	DAY
RCF3264	CMPN STM OUTLET FLOW 3RB	MLB/HR	378.8	389.8	388.9	386.9
QAC3102	CNTRL BOILER OUTLET O2	%	4.161	3.776	4.099	3.722
RTI3069	PRI SHTR INLT. FLU GAS TP	DEG F	923.5	937.6	942.8	947.5
RTI3078	INDICATE PRI SCAM OUTLET	DEG F	303.5	303.7	305.1	306.0
REC3087	CONTROL PRIMARY AIR FLOW	MCFM	61.46	61.25	60.85	59.25
REC3089	CONTROL SECONDARY AIR FL	MCFM	73.55	73.70	72.44	70.54
REC3091	CONTROL TERTIARY AIR FLW	MCFM	42.04	43.29	42.53	42.21
REC3201	CONTROL TER BNRS AIR FLW	MCFM	6.737	6.870	7.146	7.233

RFI3741	PRMRY RLK LIQ NOZ FLOW	GPM	254.4	254.7	255.8	257.6
QOI3749	%SOLID NOZZLE DENSITY #1	%SOLID	66.02	66.02	65.81	65.02
RTC3746	CNTRL RLK LIQR NOZ TEMP	DEG F	216.9	216.7	216.9	217.0
RPC3751	CONTROL 9LK LIQR NOZ PRES	PSIG	40.01	39.97	39.99	39.77

RFI3234	INDCTE EPCP OUTLET FLOW	MCFM	116.9	110.4	109.0	108.6
RFI3231	INDCTE WPCP OUTLET FLOW	MCFM	117.9	113.4	110.9	110.7
RTI3228	EAST PCP OUTLET TEMP	DEG F	346.5	346.7	346.2	345.6
RTI3227	WEST PCP OUTLET TEMP	DEG F	347.0	346.7	347.4	346.6
RAI3233	INDCTE E PCP OTLET CAPAC	%	25.25	24.64	25.52	25.83
RAI3232	INDCTE W PCP OTLET CAPAC	%	13.22	13.56	13.01	12.90

RAI3245	INDICATE NO3RB STACK CO	PPM	18.11	9.620	3.607	23.46
R3TRSC	3RB CORRECTED TRS CONCEN	PPM	4.233	4.206	4.102	5.448
RAC3207AF	4 MIN OPACITY ALARM R3	PCT	15.31	10.92	14.97	14.96
RAI3235	#3 RR STACK O2 INDICATE	%	6.668	6.683	6.580	8.610

LOG IDENT - 455 TITLE - DIOXIN TEST TIME 1630 DATE 12-12-84

POINT	DESCRIPTION	ENGR. UNITS	CURRENT VALUE	HOUR	CURRENT AVERAGES SHIFT DAY
RFC3264	CMPN STM OUTLET FLOW 3RB MLR/HR		364.8	398.0	398.0
RAC3102	CNTRL BOILER OUTLET O2 %		3.833	3.727	3.727
RTI3069	PRI SHTR INLT FLU GAS TP DEG F		905.0	900.8	907.9
RTI3078	INDICATE PRI SCAM OUTLET DEG F		306.8	307.4	306.3
RFC3087	CONTROL PRIMARY AIR FLOW MCFM		58.99	58.31	58.67
RFC3089	CONTROL SECONDARY AIR FL MCFM		71.01	71.04	71.28
RFC3091	CONTROL TERTIARY AIR FLW MCFM		43.13	43.12	43.23
RFC3201	CONTROL TER RNRS AIR FLW MCFM		1.777	1.799	3.121
RFI3741	PRMRY BLK LIQ NOZ FLOW GPM		255.2	253.0	254.1
RTI3748	%SOLID NOZZLE DENSITY #1 %SOLID		65.51	65.59	65.57
RTC3746	CNTRL BLK LIQR NOZ TEMP DEG F		218.2	216.6	216.9
RPC3751	CNTRL BLK LIQR NOZ PRES PSIG		39.99	39.98	39.98
RFI3234	INDCTE EPCP OUTLET FLOW MCFM		108.7	105.0	109.1
RFI3231	INDCTE WPCP OUTLET FLOW MCFM		109.8	105.2	110.4
RTI3228	EAST PCP OUTLET TEMP DEG F		344.5	343.9	345.7
RTI3227	WEST PCP OUTLET TEMP DEG F		345.0	344.6	346.3
RAI3233	INDCTE E PCP OTLET CAPAC %		25.09	25.62	24.49
RAI3232	INDCTE W PCP OTLET CAPAC %		13.08	12.75	13.64
RAI3245	INDICATE NO3R8 STACK CO PPM		1.574	5.489	26.10
RTTRSC	3RB CORRECTED TRS CONCEN PPM		4.390	4.608	5.079
RAC3207AF	6 MIN OPACITY ALARM R3 PCT		14.86	14.91	14.95
RAI3235	#3 RB STACK O2 INDICATE %		5.683	6.129	12.56

12-13-84

DATE

TIME 1730

TIME

TITLE - DIOXIN TEST

455

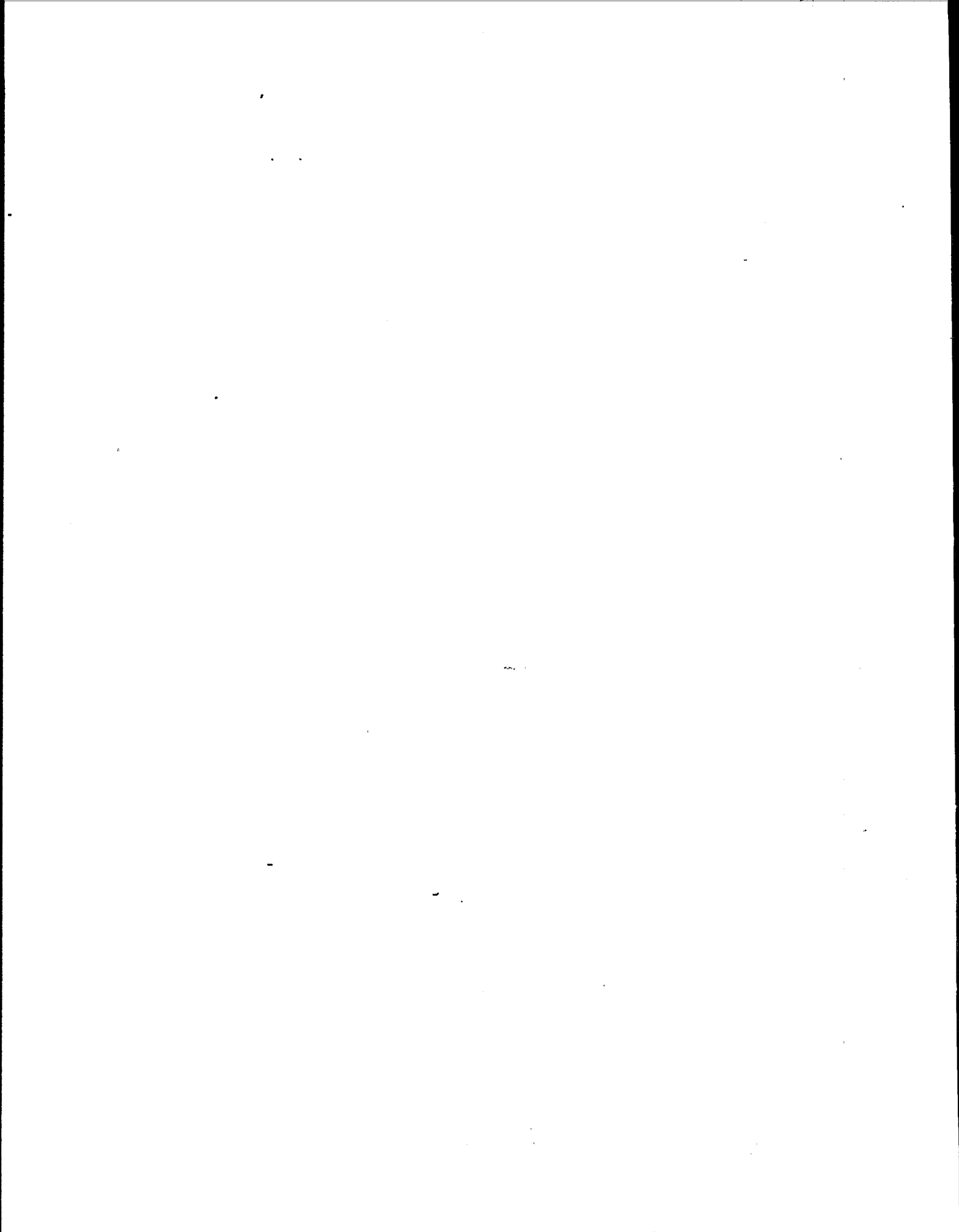
LOG IDENT -

POINT	DESCRIPTION	ENGR. CURRENT UNITS	CURRENT AVERAGES		
			VALUE	HOUR	SHIFT DAY
RCF3264	CMPN STM OUTLET FLOW 3RB	MLB/HR	402.7	384.0	387.3
RAC3102	CNTRL BOILER OUTLET O2	%	3.567	3.545	3.049
RTI3069	PRI SHTR INLT FLU GAS TP	DEG F	935.5	932.3	947.2
RTI3078	INDICATE PRI SCAH OUTLET	DEG F	309.8	309.8	308.5
RFC30A7	CONTROL PRIMARY AIR FLOW	MCFM	56.46	56.00	56.51
RFC3089	CONTROL SECONDARY AIR FL	MCFM	70.34	69.82	70.29
RFC3091	CONTROL TERTIARY AIR FLW	MCFM	41.68	41.31	41.61
RFC3201	CONTROL TER BNRS AIR FLW	MCFM	1.739	1.766	1.653

QFI3741	PRMRY BLK LIQ NOZ FLOW	GPM	263.3	263.2	263.1
RDI3748	XSOLID NOZZLE DENSITY #1	XSOLID	63.7A	63.55	63.77
RTC3746	CNTRL BLK LIQR NOZ TEMP	DEG F	217.2	217.2	217.2
RPC3751	CNTRL BLK LIQR NOZ PRES	PSIG	39.99	39.99	39.9A

RFI3234	INDCTE EPCP OUTLET FLOW	MCFM	100.0	102.2	107.1
RFI3231	INDCTE WPCP OUTLET FLOW	MCFM	103.2	103.1	108.7
RTI3228	EAST PCP OUTLET TEMP	DEG F	343.9	344.2	346.0
RTI3227	WEST PCP OUTLET TEMP	DEG F	342.7	343.2	346.0
RAI3233	INDCTE E PCP OTLET CAPAC	%	26.80	26.61	24.64
RAI3232	INDCTE W PCP OTLET CAPAC	%	12.63	13.41	13.66

RAI3245	INDICATE NO3RB STACK CO	PPM	16.00	27.33	67.09
R3TRSC	3RB CORRECTED TRS CONCEN	PPM	3.887	3.627	4.145
RAC3207AF	6 MIN OPACITY ALARM R3	PCT	14.96	14.89	14.94
RAI3235	#3 RH STACK O2 INDICATE	%	5.399	5.259	8.665



APPENDIX B-3

ELECTROSTATIC PRECIPITATOR ELECTRICAL DATA

1

Table 5. Electrostatic Precipitator Electrical Data for Site 04

Parameter ^{a,b}	Run 01		Run 02		Run 03	
	1445	2140	1310	1625	1020	-
West Inlet						
Voltage	36	38	37	36	35	ND
Current	175	150	150	175	175	ND
West Center						
Voltage	38	38	38	38	37	ND
Current	300	300	325	325	325	ND
West Outlet						
Voltage	35	35	35	35	34	ND
Current	275	250	250	325	300	ND
East Inlet						
Voltage	34	35	35	34	34	ND
Current	250	225	225	250	275	ND
East Center						
Voltage	38	37	37	36	37	ND
Current	175	175	175	200	225	ND
East Outlet						
Voltage	37	37	37	37	37	ND
Current	250	225	250	300	275	ND

- a. All voltage and current values are DC. Voltage in kv, current in mA.
b. Precipitator was set in opacity control mode during all test runs.

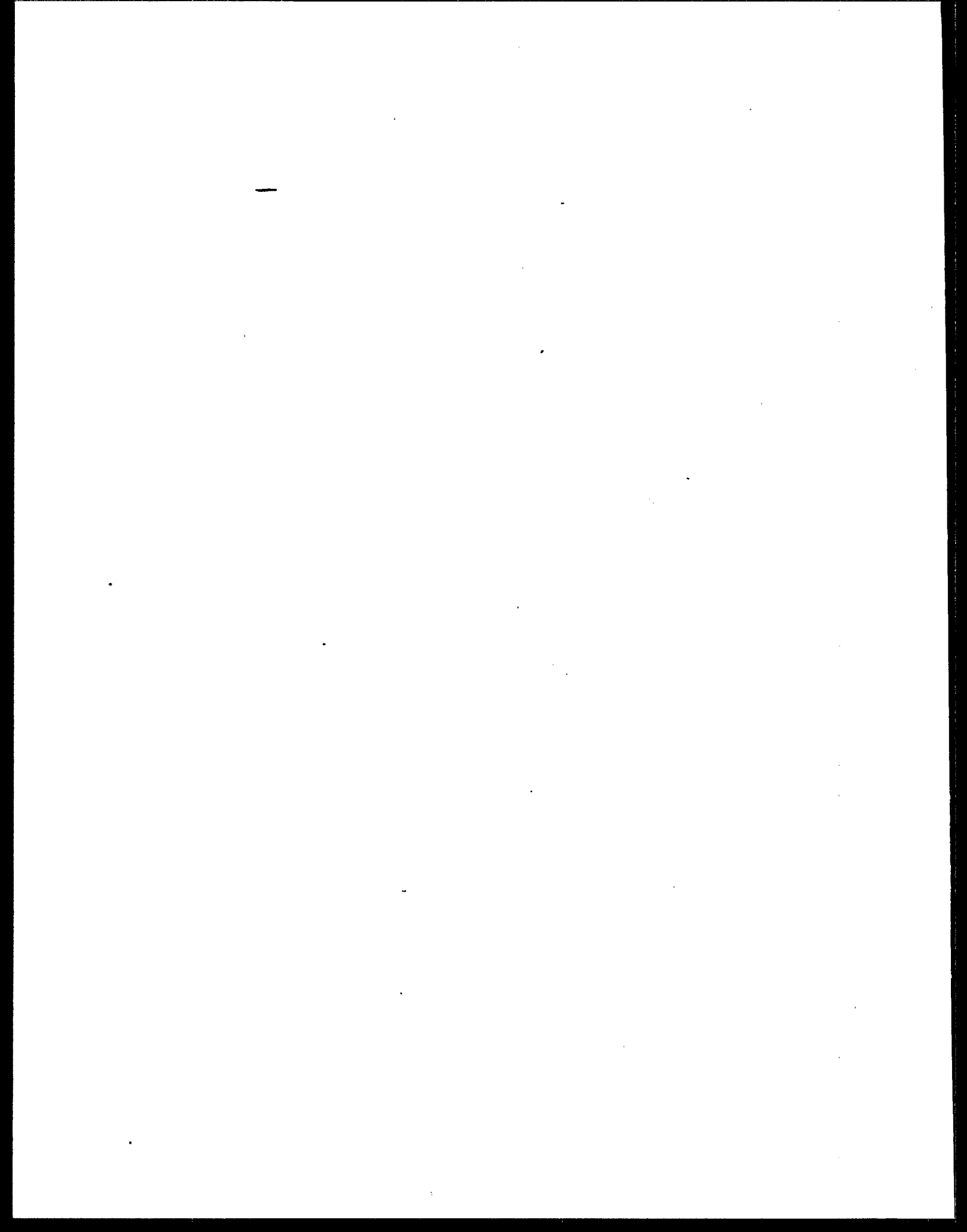
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APPENDIX B-4

DAILY CALIBRATION DATA FOR PLANT-MAINTAINED
TRS AND O₂ MONITORS



0954 12-11-84 PAGE PRINT CONSOL A 0954:40 12-11-84

RA13235 #3 RA STACK 02 INDICATE TIME 1 09:54

6 MIN. AVERAGE 09HR 09HR 07HR 06HR 05HR 04HR 03HR 02HR
57 21.07 21.03 21.06 5.656 5.439 5.846 5.526 5.751

51 21.07 21.03 21.01 5.711 5.429 5.846 7.410 6.455

45 21.07 21.03 21.01 5.880 5.422 5.560 5.344 5.910

39 21.06 21.05 20.92 5.974 5.539 5.518 5.291 5.935

33 21.05 21.06 6.959 6.786 5.896 5.616 5.353 5.780

27 21.02 21.06 5.674 6.118 5.609 7.630 5.543 5.720

21 21.02 21.07 5.722 5.910 5.600 5.810 6.737 5.998

15 21.02 21.07 5.779 6.008 5.603 5.939 5.664 5.919

09 21.02 21.07 7.276 6.032 5.652 6.014 5.536 5.741

03 21.03 21.07 5.608 5.883 5.673 5.374 5.651 5.695

40HR AVERAGE 09HR 08HR 07HR 06HR 05HR 04HR 03HR 02HR
20.74 21.04 13.13 5.975 5.596 5.942 5.776 5.872
01HR 00HR 23HR 22HR 21HR 20HR 19HR 18HR
5.874 6.368 6.255 5.935 5.963 6.231 6.116 5.857
17HR 16HR 15HR 14HR 13HR 12HR 11HR 10HR
6.201 6.428 6.089 6.491 6.573 6.821 6.443 10.73

DAY AVERAGE MON 10 SUN 09 SAT 08 FRI 07 THU 06 WED 05 TUE 04
7.275 6.713 7.164 9.093 9.249 8.438 7.586

TIME : 11:27

300 STACK TRS

RAT3236

6 MIN. AVERAGE 11HR 10HR 09HR 08HR 07HR 06HR 05HR 04HR

57 17.47 17.59 17.71 17.19 5.726 5.784 5.383

51 17.69 17.75 17.84 15.94 5.423 5.315 5.238

45 17.66 17.68 17.61 8.340 5.332 5.575 4.622

39 17.55 17.21 17.08 3.143 4.997 5.191 5.246

POW 33 17.85 17.76 17.76 5.434 5.226 5.058 4.562

27 17.64 17.53 17.68 17.70 5.394 5.747 5.831 5.159

21 17.81 17.75 17.86 17.31 5.484 5.098 5.800 5.309

15 17.53 17.62 17.60 17.57 5.343 5.885 5.332 5.073

09 17.61 17.42 17.16 17.16 5.093 5.397 5.239 5.107

03 17.97 17.97 17.60 17.60 5.484 4.845 4.430 4.777

400R AVERAGE 11HR 10HR 09HR 08HR 07HR 06HR 05HR 04HR

17.67 17.64 17.57 17.50 8.475 5.412 5.419 5.028

03HR 02HR 01HR 00HR 23HR 22HR 21HR 20HR

5.196 5.257 4.993 5.072 5.002 4.806 4.628 4.294

19HR 18HR 17HR 16HR 15HR 14HR 13HR 12HR

4.169 4.496 4.534 4.490 4.489 4.314 4.419 4.218

DAY AVERAGE TUE 11 MON 10 SUN 09 SAT 08 FRI 07 THR 06 WED 05

5.848 5.265 4.285 4.371 4.043 4.860 4.040

112A 12-12-84 PAGE PRINT CONSOLF 3 11:28:40 12-12-84

RA13235 #3 RR STACK 02 INDICATE TIME : 11:28

6 MIN. AVERAGE	11HR	10HR	09HR	08HR	07HR	06HR	05HR	04HR
57	21.10	21.11	21.0A	21.10	21.10	6.315	6.199	6.170
51	21.11	21.11	21.0A	21.09	21.09	6.305	6.301	6.01A
45	21.11	21.10	21.09	21.06	21.06	6.372	6.362	5.997
39	21.12	21.10	21.09	18.36	18.36	7.436	6.321	6.315
33	21.12	21.10	21.10	6.371	6.371	6.247	6.437	6.388
27	20.8A	21.11	21.10	21.10	6.440	6.384	6.199	7.2A7
21	21.11	21.11	21.10	21.10	6.419	6.60A	6.364	6.375
15	21.11	21.11	21.09	21.11	6.453	6.267	6.341	6.319
09	21.11	21.11	21.09	21.11	8.410	6.125	6.100	6.323
03	21.11	21.11	21.09	21.10	6.321	6.995	6.299	6.297
MINOR AVERAGE	11HR	10HR	09HR	08HR	07HR	06HR	05HR	04HR
	20.51	21.10	21.09	21.0A	13.1A	6.495	6.296	6.349
	03HR	02HR	01HR	00HR	23HR	22HR	21HR	20HR
	6.57A	6.455	6.407	6.7A0	6.652	6.696	6.6A3	6.947
	19HR	18HR	17HR	16HR	15HR	14HR	13HR	12HR
	6.864	6.335	6.505	6.470	6.261	5.984	5.853	6.232

DAY AVERAGE	TUE 11	MON 10	SUN 09	SAT 0A	FRI 07	THR 0A	WED 05
	7.826	7.275	6.713	7.164	9.093	9.249	8.43A

1149 12-13-84 PAGE PRINT CONSOLE 4 11:09:32 12-13-84

TIME : 11:44

388 STACK TRS

4 MIN. AVERAGE 11HR 10HR 09HR 08HR 07HR 06HR 05HR 04HR

57 3.997 3.444 17.24 .0023 5.109 4.340 4.909

51 3.193 2.982 17.07 .0135 5.035 4.293 5.473

45 4.324 3.838 3.809 17.18 .1090 4.354 4.250 4.891

39 4.203 3.450 3.730 17.42 3.706 4.750 4.140 4.770

PPM

33 3.663 3.490 3.874 16.80 4.829 4.734 4.582 4.575

27 3.492 3.400 12.53 16.97 4.443 4.533 4.354 4.700

21 3.794 2.702 17.24 17.03 4.504 4.510 4.841 4.765

15 4.304 4.303 17.18 16.55 3.620 4.297 4.467 5.051

09 4.157 4.043 17.44 16.37 5.069 4.400 5.050 4.565

03 3.833 3.934 17.15 14.00 5.042 4.943 5.279 4.321

470R AVERAGE

11HR 10HR 09HR 08HR 07HR 06HR 05HR 04HR

3.990 3.645 9.064 16.91 3.701 4.694 4.517 4.804

03HR 02HR 01HR 00HR 23HR 22HR 21HR 20HR

4.425 4.934 4.463 4.463 4.604 4.330 5.017 4.836

19HR 18HR 17HR 16HR 15HR 14HR 13HR 12HR

5.042 5.130 5.361 5.286 5.350 5.245 4.845 4.850

DAY AVERAGE WED 12 TUE 11 MON 10 SUN 09 SAT 08 FRI 07 THR 06

6.832 5.840 5.265 4.285 4.371 4.043 4.860

1150 12-13-84 PAGE PRINT CONSOLF 4 11:50:40 12-13-84

R413235 #3 RH STACK OP INDICATE TIME : 11:50

6 MIN. AVERAGE	11HR	10HR	09HR	08HR	07HR	06HR	05HR	04HR
57		5.579	5.358	20.91	21.08	5.462	5.556	5.354
51		7.306	5.405	20.91	21.24	5.551	5.619	5.342
45	5.311	5.578	7.040	20.90	21.21	5.596	5.522	5.565
39	5.241	5.452	5.612	20.91	15.90	7.049	5.566	5.584
33	5.216	5.587	5.776	20.91	5.524	5.514	7.587	5.859
27	5.310	5.665	13.40	20.91	5.552	5.659	5.458	5.584
21	5.838	7.660	20.91	20.90	5.709	5.708	5.502	5.656
15	5.394	5.541	20.91	20.90	5.739	5.876	5.577	5.690
09	5.521	5.354	20.91	20.90	5.773	8.039	5.570	5.781
03	5.403	5.319	20.90	20.91	5.614	5.941	6.073	5.586
4 HOUR AVERAGE	11HR	10HR	09HR	08HR	07HR	06HR	05HR	04HR
	5.388	5.908	11.58	20.89	12.35	6.018	5.763	5.652
	03HR	02HR	01HR	00HR	23HR	22HR	21HR	20HR
	5.759	6.141	5.797	5.690	6.114	7.681	5.601	5.809
	19HR	18HR	17HR	16HR	15HR	14HR	13HR	12HR
	6.029	6.126	5.755	5.745	6.129	6.002	5.908	5.870

DAY AVERAGE WED 12 TUE 11 MON 10 SUN 09 SAT 08 FRI 07 THR 06
 8.443 7.826 7.275 6.713 7.164 9.003 9.249

APPENDIX C
SAMPLE SHIPMENT LETTER

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RADIAN
CORPORATION

December 15, 1984

U. S. EPA ECC Toxicant Analysis Center
Building 1105
Bay St. Louis, MO 39529

Attention: Danny McDaniel

Subject: Tier 4 - Analysis Instructions

Dear Sir:

The objective of this letter is to clarify instructions and priorities for individual samples from specific Tier 4 combustion sites. This instruction letter is No. 4 and pertains to EPA Site No. 04 at

The Episode No. is 2549, and SCC numbers assigned to this site were numbers DQ000401 through DQ000499.

SCC numbers DQ000401 through DQ000406 have been assigned to Troika for internal QA/QC purposes. SCC numbers DQ000407 through DQ000418 have been assigned to samples included in this shipment and numbers DQ000419 have been assigned to samples being archived at Radian. All remaining SCC numbers are unused.

The sample shipment for EPA Site No. 04 consists of 4 boxes containing 57 samples in 67 containers. (Note-The Modified Method 5 samples are comprised of 6 components as listed below. Some MM5 sample runs have more than one container per component as indicated by asterisk.) The sample shipment was shipped air freight on December 15, 1984 by Federal Express under Airbill(s) No. 770332533 and No. 544549655

Instructions for extraction and analysis follow.

1. The following samples require immediate extraction and analysis (Priority #1 samples).

Radian Run #04-MM5-B-01
(Total of 6 train components)

SCC #	Component	Fraction
DQ000407	1	Filter
DQ000407	2* (3 containers)	Probe Rinse
DQ000407	3	Back Half/Coil Rinse
DQ000407	4* (2 containers)	Condensate
DQ000407	5	Impinger Solution
DQ000407	6	XAD Module

RADIAN

Radian Run # 04-MM5-BLANK
(Total of 6 train components)

SCC #	Components	Fraction
DQ000408	1	Filter
DQ000408	2	Probe Rinse
DQ000408	3	Back Half/Coil Rinse
DQ000408	4	Condensate
DQ000408	5	Impinger Solution
DQ000408	6	XAD Module

Radian Run # 04-MM5-C-01
(Total of 6 train components)

SCC #	Components	Fraction
DQ000409	1	Filter
DQ000409	2	Probe Rinse
DQ000409	3	Back Half/Coil Rinse
DQ000409	4*(2 container)	Condensate
DQ000409	5	Impinger Solution
DQ000409	6	XAD Module

Radian Run # 04-MM5-B-02
(Total of 6 train components)

SCC #	Components	Fraction
DQ000411	1	Filter
DQ000411	2*(3 containers)	Probe Rinse
DQ000411	3	Back Half/Coil Rinse
DQ000411	4	Condensate
DQ000411	5	Impinger Solution
DQ000411	6	XAD Module

Radian Run # 04-MM5-C-02
(Total of 6 train components)

SCC #	Components	Fraction
DQ000412	1	Filter
DQ000412	2	Probe Rinse
DQ000412	3	Back Half/Coil Rinse
DQ000412	4*(2 containers)	Condensate
DQ000412	5	Impinger Solution
DQ000412	6*(2 containers)	XAD Module

RADIAN
CORPORATION

Radian Run # 04-MM5-B-BLANK
(Total of 6 train components)

SCC #	Components	Fraction
<u>DQ000414</u>	1	<u>Filter</u>
<u>DQ000414</u>	2	<u>Probe Rinse</u>
<u>DQ000414</u>	3	<u>Back Half/Coil Rinse</u>
<u>DQ000414</u>	4	<u>Condensate</u>
<u>DQ000414</u>	5	<u>Impinger Solution</u>
<u>DQ000414</u>	6	<u>XAD Module</u>

Radian Run # 04-MM5-C-BLANK
(Total of 6 train components)

SCC #	Components	Fraction
DQ000415	1	Filter
DQ000415	2	Probe Rinse
DQ000415	3	Back Half/Coil Rinse
DQ000415	4	Condensate
DQ000415	5	Impinger Solution
DQ000415	6	XAD Module

Radian Run # 04-MM5-B-03
(Total of 6 train components)

SCC #	Components	Fraction
DQ000416	1	Filter
DQ000416	2*(2 containers)	Probe Rinse
DQ000416	3	Back Half/Coil Rinse
DQ000416	4	Condensate
DQ000416	5	Impinger Solution
DQ000416	6	XAD Module

Radian Run # 04-MM5-C-03
(Total of 6 train components)

SCC #	Components	Fraction
DQ000417	1	Filter
DQ000417	2*(2 containers)	Probe Rinse
DQ000417	3	Back Half/Coil Rinse
DQ000417	4	Condensate
DQ000417	5	Impinger Solution
DQ000417	6	XAD Module

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2. The strong black liquor samples are the only Priority #2 samples. They should be held for analysis by Troika pending the results of Priority #1 sample analysis.

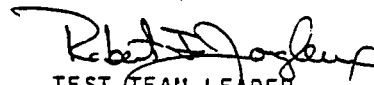
SCC #	Sample
DQ000410	SBL-01 Strong black liquor
DQ000413	SBL-02 Strong black liquor
DQ000418	SBL-03 Strong black liquor

3. The soil sample is the only Priority #3 sample. It will be held by Radian for analysis by Troika pending the results of Priority #1 and Priority #2 sample analyses.

SCC #	Sample
DQ000419	# 04-S-01 Soils

If there are any questions concerning this sample shipment, please contact either Bob Jongleux or Larry Keller at Radian Corporation (919) 541-9100.

Sincerely,


TEST TEAM LEADER

APPENDIX D

DIOXIN/FURAN ANALYTICAL DATA FOR MM5 SAMPLE TRAINS

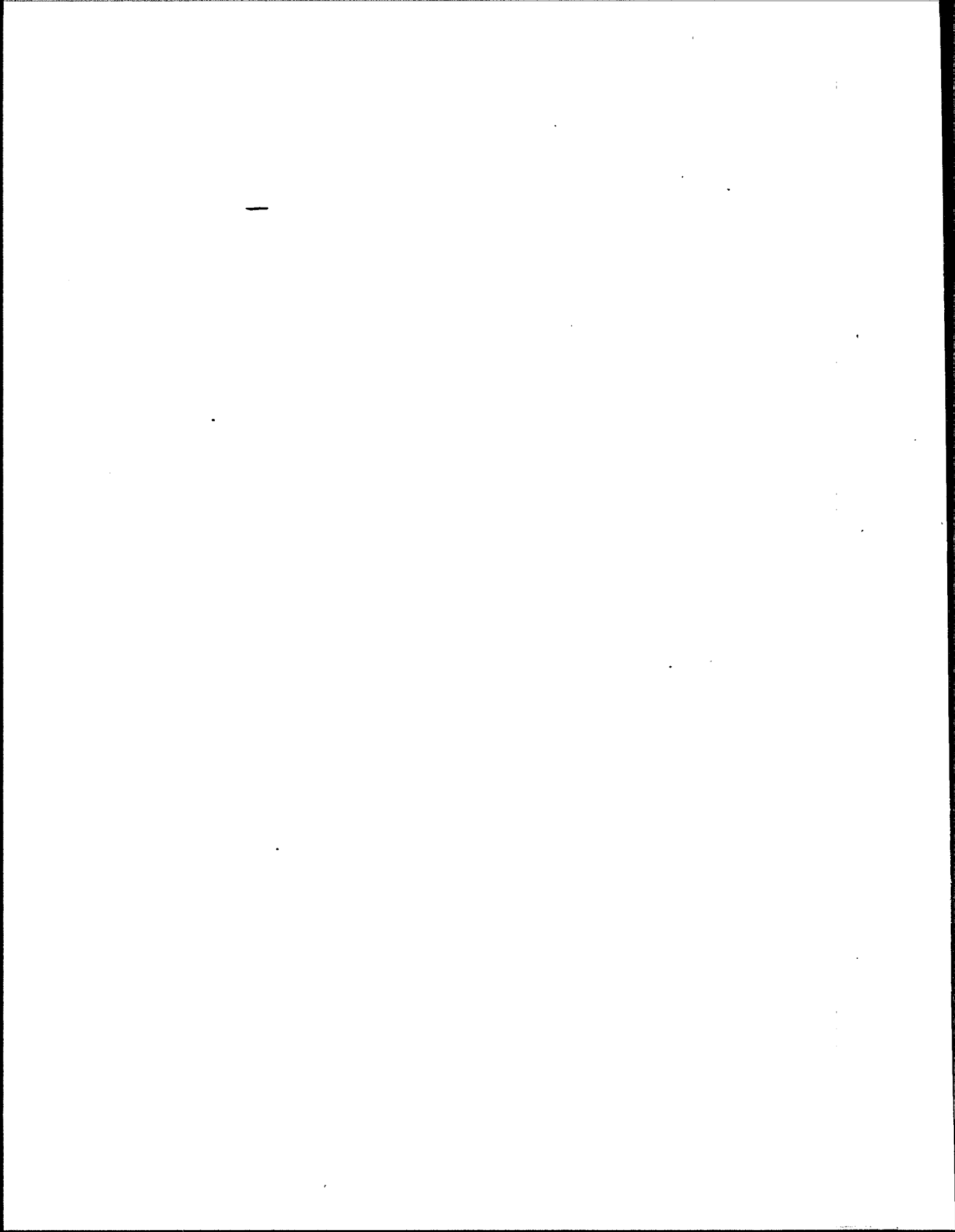


TABLE D-1. DIOXIN/FURAN ANALYTICAL DATA FOR MM5 TRAINS AT THE ESP INLET

Isomer/Homologue	Amount Detected Picograms Per Train		
	Run 01	Run 02	Run 03
<u>DIOXINS</u>			
2378 TCDD	ND (80)	ND (50)	ND (180)
Other TCDD	ND (60)	ND (200)	ND (400)
Penta-CDD	ND (310)	ND (60)	ND (240)
Hexa-CDD	500	200	ND (940)
Hepta-CDD	1200	500	1400
Octa-CDD	3300	1600	4900
TOTAL CDD	4000	2300	6300
<u>FURANS</u>			
2378 TCDF	ND (80)	50	ND (180)
Other TCDF	1000	500	1000
Penta-CDF	1300	600	ND (1480)
Hexa-CDF	1800	1000	1900
Hepta-CDF	1000	300	900
Octa-CDF	300	200	ND (630)
TOTAL CDF	5400	2650	3800

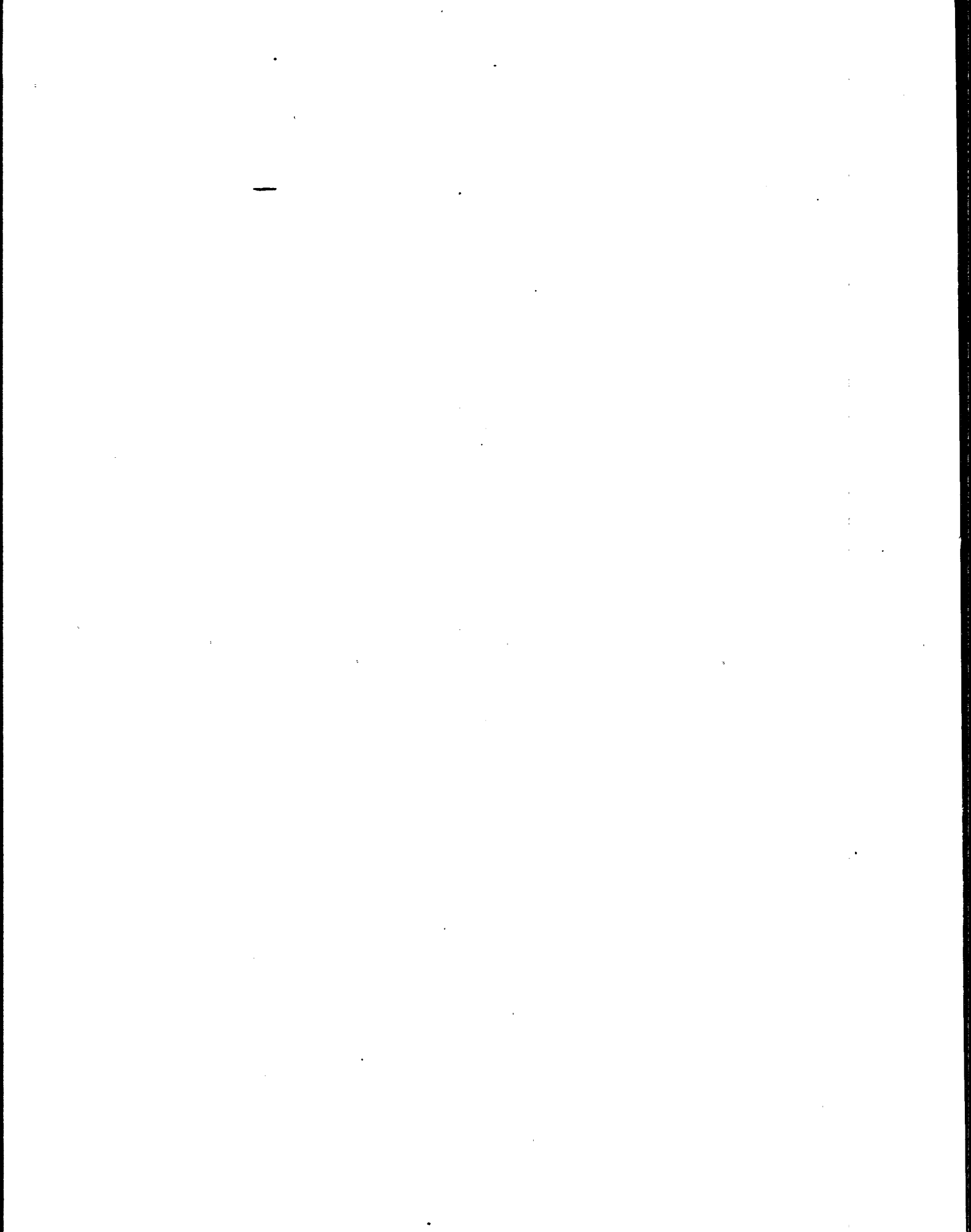
TABLE D-2. DIOXIN/FURAN ANALYTICAL DATA FOR MM5 TRAINS AT THE ESP OUTLET

Isomer/Homologue	Amount Detected Picograms Per Train		
	Run 01	Run 02	Run 03
<u>DIOXINS</u>			
2378 TCDD	ND (310)	ND (100)	ND (110)
Other TCDD	ND (310)	300	ND (110)
Penta-CDD	ND (130)	ND (120)	ND (220)
Hexa-CDD	800	ND (320)	ND (240)
Hepta-CDD	1600	500	500
Octa-CDD	3700	2200	1300
TOTAL PCDD	6100	3000	1800
<u>FURANS</u>			
2378 TCDF	ND (40)	ND (200)	ND (100)
Other TCDF	1700	300	200
Penta-CDD	800	ND (230)	ND (130)
Hexa-CDD	700	300	ND (510)
Hepta-CDD	1000	1000	300
Octa-CDD	400	1500	100
TOTAL PCDD	4600	3100	600

APPENDIX E

RUN-SPECIFIC DIOXIN/FURAN EMISSIONS DATA

- E-1 Run-specific Dioxin/Furan Emissions Data at the ESP Inlet
(As-measured Concentrations)
- E-2 Run-specific Dioxin/Furan Emissions Data at the ESP Outlet
(As-measured Concentrations)
- E-3 Run-specific Dioxin/Furan Emissions Data at the ESP Inlet
(Concentrations Corrected to 3 Percent Oxygen)
- E-4 Run-specific Dioxin/Furan Emissions Data at the ESP Outlet
(Concentrations Corrected to 3 Percent Oxygen)



APPENDIX E-1

RUN-SPECIFIC DIOXIN/FURAN EMISSIONS DATA AT THE ESP INLET

(As-measured Concentrations)

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TABLE E-1. DIOXIN/FURAN EMISSIONS DATA AT THE ESP INLET FOR
RUN 01, SITE BLB-A (AS-MEASURED CONCENTRATIONS)

Dioxin/Furan Isomer	Isomer Concentration In Flue Gas (ng/dscm)	Isomer Concentration In Flue Gas (ppt)	Isomer Hourly Emissions Rate (ug/hr)
DIOXINS			
2378 TCDD	ND (1.94E-02)	ND (1.45E-03)	ND (4.47E+00)
Other TCDD	ND (1.94E-02)	ND (1.45E-03)	ND (4.47E+00)
Penta-CDD	ND (7.52E-02)	ND (5.08E-03)	ND (1.73E+01)
Hexa-CDD	1.21E-01(N/A)	7.47E-03(N/A)	2.80E+01
Hepta-CDD	2.91E-01(N/A)	1.65E-02(N/A)	6.71E+01
Octa-CDD	8.01E-01(N/A)	4.19E-02(N/A)	1.85E+02
Total PCDD	1.21E+00	6.58E-02	2.80E+02
FURANS			
2378 TCDF	ND (1.46E-02)	ND (1.14E-03)	ND (3.36E+00)
Other TCDF	2.43E-01(N/A)	1.91E-02(N/A)	5.59E+01
Penta-CDF	3.16E-01(N/A)	2.23E-02(N/A)	7.27E+01
Hexa-CDF	4.37E-01(N/A)	2.80E-02(N/A)	1.01E+02
Hepta-CDF	2.43E-01(N/A)	1.43E-02(N/A)	5.59E+01
Octa-CDF	7.28E-02(N/A)	3.95E-03(N/A)	1.68E+01
Total PCDF	1.31E+00	8.77E-02	3.02E+02

NOTE: Isomer concentrations shown are at as-measured oxygen conditions.

ND = not detected (detection limit in parenthesis).

N/A = detection limit not applicable. QA samples indicate the detection limit for positive samples.

ng = 1.0E-09g

ug = 1.0E-06g

ppt = parts per trillion, dry volume basis

8760 operating hours per year

TABLE E-2. DIOXIN/FURAN EMISSIONS DATA AT THE ESP INLET FOR
RUN 02, SITE BLB-A (AS-MEASURED CONCENTRATIONS)

Dioxin/Furan Isomer	Isomer Concentration In Flue Gas (ng/dscm)	Isomer Concentration In Flue Gas (ppt)	Isomer Hourly Emissions Rate (ug/hr)
DIOXINS			
2378 TCDD	ND (1.98E-02)	ND (1.48E-03)	ND (4.38E+00)
Other TCDD	1.98E-02(N/A)	1.48E-03(N/A)	4.38E+00
Penta-CDD	ND (2.38E-02)	ND (1.61E-03)	ND (5.26E+00)
Hexa-CDD	7.94E-02(N/A)	4.88E-03(N/A)	1.75E+01
Hepta-CDD	1.98E-01(N/A)	1.12E-02(N/A)	4.38E+01
Octa-CDD	6.35E-01(N/A)	3.32E-02(N/A)	1.40E+02
Total PCDD	9.33E-01	5.08E-02	2.06E+02
FURANS			
2378 TCDF	ND (7.94E-02)	ND (6.24E-03)	ND (1.75E+01)
Other TCDF	1.98E-01(N/A)	1.56E-02(N/A)	4.38E+01
Penta-CDF	2.38E-01(N/A)	1.68E-02(N/A)	5.26E+01
Hexa-CDF	3.97E-01(N/A)	2.55E-02(N/A)	8.76E+01
Hepta-CDF	1.19E-01(N/A)	7.00E-03(N/A)	2.63E+01
Octa-CDF	7.94E-02(N/A)	4.30E-03(N/A)	1.75E+01
Total PCDF	1.03E+00	6.92E-02	2.28E+02

NOTE: Isomer concentrations shown are at as-measured oxygen conditions.

ND = not detected (detection limit in parenthesis).

N/A = detection limit not applicable. QA samples indicate the detection limit for positive samples.

ng = 1.0E-09g

ug = 1.0E-06g

ppt = parts per trillion, dry volume basis

8760 operating hours per year

TABLE E-3. DIOXIN/FURAN EMISSIONS DATA AT THE ESP INLET FOR RUN 03
 — SITE BLB-A (AS-MEASURED CONCENTRATIONS)

Dioxin/Furan Isomer	Isomer Concentration In Flue Gas (ng/dscm)	Isomer Concentration In Flue Gas (ppt)	Isomer Hourly Emissions Rate (ug/hr)
DIOXINS			
2378 TCDD	ND (7.47E-02)	ND (5.58E-03)	ND (1.51E+01)
Other TCDD	ND (7.47E-02)	ND (5.58E-03)	ND (1.51E+01)
Penta-CDD	ND (9.96E-02)	ND (6.73E-03)	ND (2.02E+01)
Hexa-CDD	ND (3.90E-01)	ND (2.40E-02)	ND (7.91E+01)
Hepta-CDD	5.81E-01(N/A)	3.29E-02(N/A)	1.18E+02
Octa-CDD	2.03E+00(N/A)	1.06E-01(N/A)	4.12E+02
Total PCDD	2.61E+00	1.39E-01	5.30E+02
FURANS			
2378 TCDF	ND (1.66E-01)	ND (1.30E-02)	ND (3.36E+01)
Other TCDF	4.15E-01(N/A)	3.26E-02(N/A)	8.41E+01
Penta-CDF	ND (6.14E-01)	ND (4.34E-02)	ND (1.25E+02)
Hexa-CDF	7.88E-01(N/A)	5.06E-02(N/A)	1.60E+02
Hepta-CDF	3.73E-01(N/A)	2.20E-02(N/A)	7.57E+01
Octa-CDF	ND (2.61E-01)	ND (1.42E-02)	ND (5.30E+01)
Total PCDF	1.58E+00	1.05E-01	3.20E+02

NOTE: Isomer concentrations shown are at as-measured oxygen conditions.

ND = not detected (detection limit in parenthesis).

N/A = detection limit not applicable. QA samples indicate the detection limit for positive samples.

ng = 1.0E-09g

ug = 1.0E-06g

ppt = parts per trillion, dry volume basis

8760 operating hours per year

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APPENDIX E-2

RUN-SPECIFIC DIOXIN/FURAN EMISSIONS DATA AT THE ESP OUTLET

(As-measured Concentrations)

1

TABLE E-4. DIOXIN/FURAN EMISSIONS DATA AT THE ESP OUTLET FOR
RUN 01, SITE BLB-A (AS-MEASURED CONCENTRATIONS)

Dioxin/Furan Isomer	Isomer Concentration In Flue Gas (ng/dscm)	Isomer Concentration In Flue Gas (ppt)	Isomer Hourly Emissions Rate (ug/hr)
DIOXINS			
2378 TCDD	ND (6.00E-02)	ND (4.48E-03)	ND (1.36E+01)
Other TCDD	ND (6.00E-02)	ND (4.48E-03)	ND (1.36E+01)
Penta-CDD	ND (2.51E-02)	ND (1.70E-03)	ND (5.71E+00)
Hexa-CDD	1.55E-01(N/A)	9.52E-03(N/A)	3.52E+01
Hepta-CDD	3.09E-01(N/A)	1.75E-02(N/A)	7.03E+01
Octa-CDD	7.16E-01(N/A)	3.74E-02(N/A)	1.63E+02
Total PCDD	1.18E+00	6.45E-02	2.68E+02
FURANS			
2378 TCDF	ND (7.74E-03)	ND (6.08E-04)	ND (1.76E+00)
Other TCDF	3.29E-01(N/A)	2.58E-02(N/A)	7.47E+01
Penta-CDF	1.55E-01(N/A)	1.09E-02(N/A)	3.52E+01
Hexa-CDF	1.35E-01(N/A)	8.69E-03(N/A)	3.08E+01
Hepta-CDF	1.93E-01(N/A)	1.14E-02(N/A)	4.40E+01
Octa-CDF	7.74E-02(N/A)	4.19E-03(N/A)	1.76E+01
Total PCDF	8.90E-01	6.11E-02	2.02E+02

NOTE: Isomer concentrations shown are at as-measured oxygen conditions.

ND = not detected (detection limit in parenthesis).

N/A = detection limit not applicable. QA samples indicate the detection limit for positive samples.

ng = 1.0E-09g

ug = 1.0E-06g

ppt = parts per trillion, dry volume basis

8760 operating hours per year

TABLE E-5. DIOXIN/FURAN EMISSIONS DATA AT THE ESP OUTLET FOR
RUN 02, SITE BLB-A (AS-MEASURED CONCENTRATIONS)

Dioxin/Furan Isomer	Isomer Concentration In Flue Gas (ng/dscm)	Isomer Concentration In Flue Gas (ppt)	Isomer Hourly Emissions Rate (ug/hr)
DIOXINS			
2378 TCDD	ND (1.69E-02)	ND (1.26E-03)	ND (3.78E+00)
Other TCDD	5.06E-02 (N/A)	3.78E-03 (N/A)	1.13E+01
Penta-CDD	ND (2.02E-02)	ND (1.37E-03)	ND (4.54E+00)
Hexa-CDD	ND (5.40E-02)	ND (3.32E-03)	ND (1.21E+01)
Hepta-CDD	8.43E-02 (N/A)	4.77E-03 (N/A)	1.89E+01
Octa-CDD	3.71E-01 (N/A)	1.94E-02 (N/A)	8.32E+01
Total PCDD	5.06E-01	2.80E-02	1.13E+02
FURANS			
2378 TCDF	ND (3.37E-02)	ND (2.65E-03)	ND (7.56E+00)
Other TCDF	5.06E-02 (N/A)	3.98E-03 (N/A)	1.13E+01
Penta-CDF	ND (3.88E-02)	ND (2.74E-03)	ND (8.70E+00)
Hexa-CDF	5.06E-02 (N/A)	3.25E-03 (N/A)	1.13E+01
Hepta-CDF	1.69E-01 (N/A)	9.92E-03 (N/A)	3.78E+01
Octa-CDF	2.53E-01 (N/A)	1.37E-02 (N/A)	5.67E+01
Total PCDF	5.23E-01	3.08E-02	1.17E+02

NOTE: Isomer concentrations shown are at as-measured oxygen conditions.

ND = not detected (detection limit in parenthesis).

N/A = detection limit not applicable. QA samples indicate the detection limit for positive samples.

ng = 1.0E-09g

ug = 1.0E-06g

ppt = parts per trillion, dry volume basis

8760 operating hours per year

TABLE E-6. DIOXIN/FURAN EMISSIONS DATA AT THE ESP OUTLET FOR
RUN 03, SITE BLB-A (AS-MEASURED CONCENTRATIONS)

Dioxin/Furan Isomer	Isomer Concentration In Flue Gas (ng/dscm)	Isomer Concentration In Flue Gas (ppt)	Isomer Hourly Emissions Rate (ug/hr)
DIOXINS			
2378 TCDD	ND (1.88E-02)	ND (1.41E-03)	ND (4.08E+00)
Other TCDD	ND (1.88E-02)	ND (1.41E-03)	ND (4.08E+00)
Penta-CDD	ND (3.77E-02)	ND (2.55E-03)	ND (8.15E+00)
Hexa-CDD	ND (4.11E-02)	ND (2.53E-03)	ND (8.89E+00)
Hepta-CDD	8.56E-02(N/A)	4.85E-03(N/A)	1.85E+01
Octa-CDD	2.23E-01(N/A)	1.16E-02(N/A)	4.82E+01
Total PCDD	3.08E-01	1.65E-02	6.67E+01
FURANS			
2378 TCDF	ND (1.71E-02)	ND (1.35E-03)	ND (3.71E+00)
Other TCDF	3.42E-02(N/A)	2.69E-03(N/A)	7.41E+00
Penta-CDF	ND (2.23E-02)	ND (1.57E-03)	ND (4.82E+00)
Hexa-CDF	ND (8.73E-02)	ND (5.60E-03)	ND (1.89E+01)
Hepta-CDF	5.14E-02(N/A)	3.02E-03(N/A)	1.11E+01
Octa-CDF	1.71E-02(N/A)	9.28E-04(N/A)	3.71E+00
Total PCDF	1.03E-01	6.64E-03	2.22E+01

NOTE: Isomer concentrations shown are at as-measured oxygen conditions.

ND = not detected (detection limit in parenthesis).

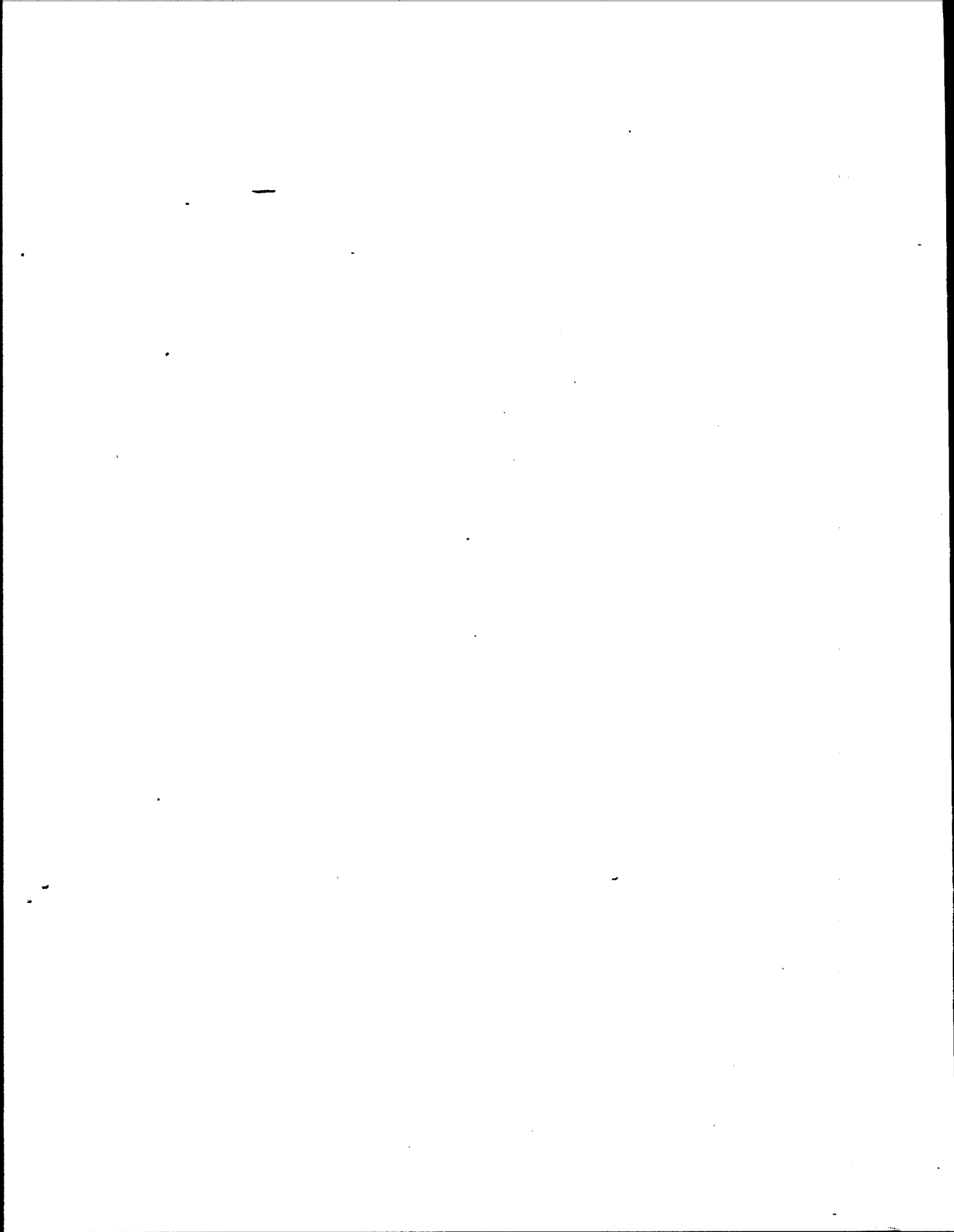
N/A = detection limit not applicable. QA samples indicate the detection limit for positive samples.

ng = 1.0E-09g

ug = 1.0E-06g

ppt = parts per trillion, dry volume basis

8760 operating hours per year



APPENDIX E-3

RUN-SPECIFIC DIOXIN/FURAN EMISSIONS DATA AT THE ESP INLET

(Concentrations Corrected to 3 Percent Oxygen)

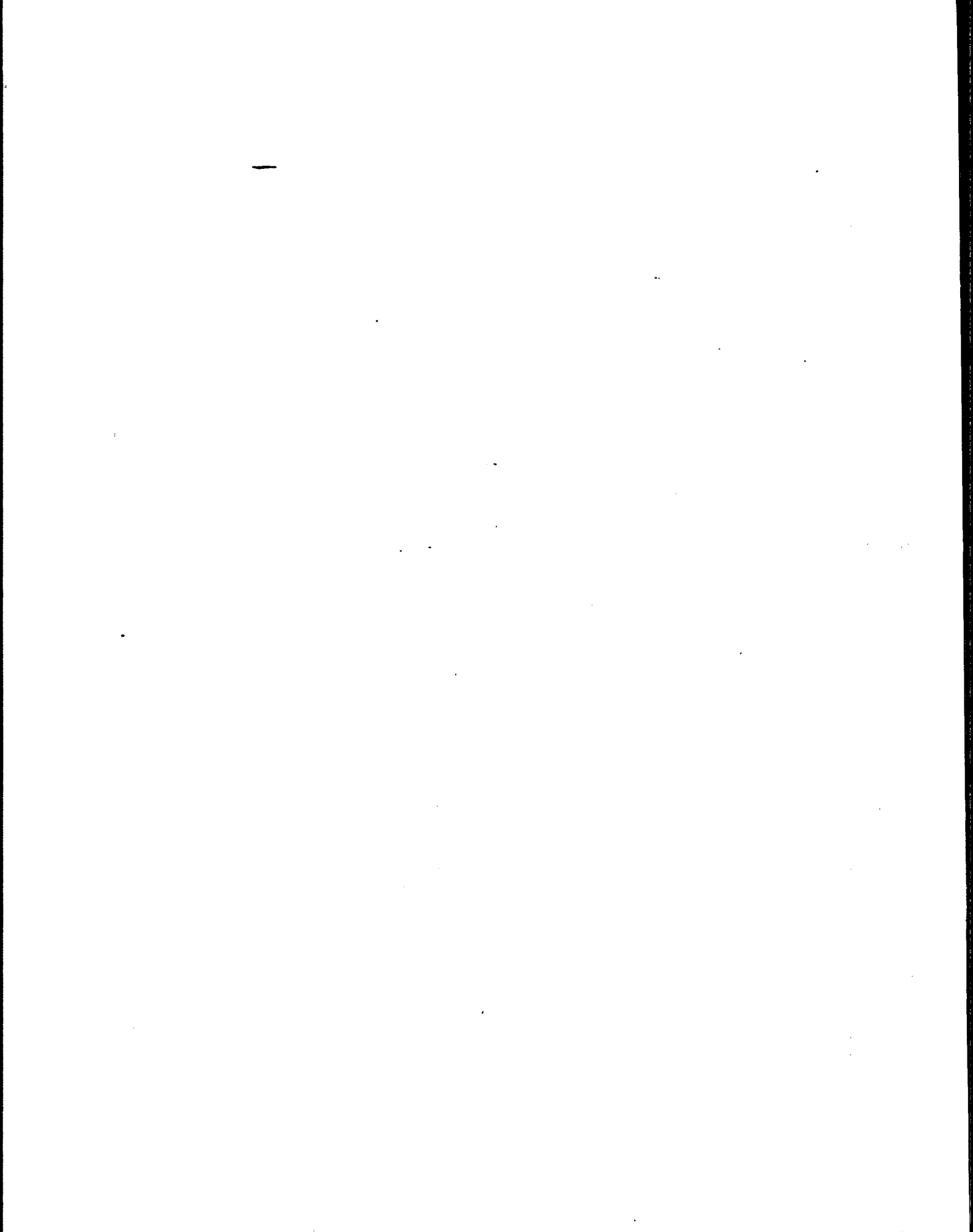


TABLE E-7. DIOXIN/FURAN EMISSIONS DATA AT THE ESP INLET FOR RUN 01,
 SITE BLB-A (CONCENTRATIONS CORRECTED TO 3 PERCENT OXYGEN)

Dioxin/Furan Isomer	Isomer Concentration In Flue Gas (ng/dscm @ 3% oxygen)	Isomer Concentration In Flue Gas (ppt @ 3% oxygen)	Isomer Hourly Emissions Rate (ug/hr)
DIOXINS			
2378 TCDD	ND (2.18E-02)	ND (1.63E-03)	ND (4.47E+00)
Other TCDD	ND (2.18E-02)	ND (1.63E-03)	ND (4.47E+00)
Penta-CDD	ND (8.46E-02)	ND (5.72E-03)	ND (1.73E+01)
Hexa-CDD	1.37E-01(N/A)	8.40E-03(N/A)	2.80E+01
Hepta-CDD	3.28E-01(N/A)	1.85E-02(N/A)	6.71E+01
Octa-CDD	9.01E-01(N/A)	4.71E-02(N/A)	1.85E+02
Total PCDD	1.37E+00	7.41E-02	2.80E+02
FURANS			
2378 TCDF	ND (1.64E-02)	ND (1.29E-03)	ND (3.36E+00)
Other TCDF	2.73E-01(N/A)	2.15E-02(N/A)	5.59E+01
Penta-CDF	3.55E-01(N/A)	2.51E-02(N/A)	7.27E+01
Hexa-CDF	4.92E-01(N/A)	3.15E-02(N/A)	1.01E+02
Hepta-CDF	2.73E-01(N/A)	1.61E-02(N/A)	5.59E+01
Octa-CDF	8.19E-02(N/A)	4.44E-03(N/A)	1.68E+01
Total PCDF	1.47E+00	9.86E-02	3.02E+02

NOTE: Isomer concentrations shown are corrected to 3% oxygen.

ND = not detected (detection limit in parenthesis).

N/A = detection limit not applicable. QA samples indicate the detection limit for positive samples.

ng = 1.0E-09g

ug = 1.0E-06g

ppt = parts per trillion, dry volume basis

8760 operating hours per year

TABLE E-8. DIOXIN/FURAN EMISSIONS DATA AT THE ESP INLET FOR RUN 02,
—SITE BLB-A (CONCENTRATIONS CORRECTED TO 3 PERCENT OXYGEN)

Dioxin/Furan Isomer	Isomer Concentration In Flue Gas (ng/dscm @ 3% oxygen)	Isomer Concentration In Flue Gas (ppt @ 3% oxygen)	Isomer Hourly Emissions Rate (ug/hr)
DIOXINS			
2378 TCDD	ND (2.23E-02)	ND (1.67E-03)	ND (4.38E+00)
Other TCDD	2.23E-02 (N/A)	1.67E-03 (N/A)	4.38E+00
Penta-CDD	ND (2.68E-02)	ND (1.81E-03)	ND (5.26E+00)
Hexa-CDD	8.93E-02 (N/A)	5.49E-03 (N/A)	1.75E+01
Hepta-CDD	2.23E-01 (N/A)	1.26E-02 (N/A)	4.38E+01
Octa-CDD	7.14E-01 (N/A)	3.74E-02 (N/A)	1.40E+02
Total PCDD	1.05E+00	5.71E-02	2.06E+02
FURANS			
2378 TCDF	ND (8.93E-02)	ND (7.02E-03)	ND (1.75E+01)
Other TCDF	2.23E-01 (N/A)	1.75E-02 (N/A)	4.38E+01
Penta-CDF	2.68E-01 (N/A)	1.90E-02 (N/A)	5.26E+01
Hexa-CDF	4.46E-01 (N/A)	2.86E-02 (N/A)	8.76E+01
Hepta-CDF	1.34E-01 (N/A)	7.88E-03 (N/A)	2.63E+01
Octa-CDF	8.93E-02 (N/A)	4.84E-03 (N/A)	1.75E+01
Total PCDF	1.16E+00	7.79E-02	2.28E+02

NOTE: Isomer concentrations shown are corrected to 3% oxygen.

ND = not detected (detection limit in parenthesis).

N/A = detection limit not applicable. QA samples indicate the detection limit for positive samples.

ng = 1.0E-09g

ug = 1.0E-06g

ppt = parts per trillion, dry volume basis

8760 operating hours per year

TABLE E-9. DIOXIN/FURAN EMISSIONS DATA AT THE ESP INLET FOR RUN 03,
SITE BLB-A (CONCENTRATIONS CORRECTED TO 3 PERCENT OXYGEN)

Dioxin/Furan Isomer	Isomer Concentration In Flue Gas (ng/dscm @ 3% oxygen)	Isomer Concentration In Flue Gas (ppt @ 3% oxygen)	Isomer Hourly Emissions Rate (ug/hr)
DIOXINS			
2378 TCDD	ND (8.40E-02)	ND (6.28E-03)	ND (1.51E+01)
Other TCDD	ND (8.40E-02)	ND (6.28E-03)	ND (1.51E+01)
Penta-CDD	ND (1.12E-01)	ND (7.57E-03)	ND (2.02E+01)
Hexa-CDD	ND (4.39E-01)	ND (2.70E-02)	ND (7.91E+01)
Hepta-CDD	6.54E-01(N/A)	3.70E-02(N/A)	1.18E+02
Octa-CDD	2.29E+00(N/A)	1.20E-01(N/A)	4.12E+02
Total PCDD	2.94E+00	1.57E-01	5.30E+02
FURANS			
2378 TCDF	ND (1.87E-01)	ND (1.47E-02)	ND (3.36E+01)
Other TCDF	4.67E-01(N/A)	3.67E-02(N/A)	8.41E+01
Penta-CDF	ND (6.91E-01)	ND (4.89E-02)	ND (1.25E+02)
Hexa-CDF	8.87E-01(N/A)	5.69E-02(N/A)	1.60E+02
Hepta-CDF	4.20E-01(N/A)	2.47E-02(N/A)	7.57E+01
Octa-CDF	ND (2.94E-01)	ND (1.59E-02)	ND (5.30E+01)
Total PCDF	1.77E+00	1.18E-01	3.20E+02

NOTE: Isomer concentrations shown are corrected to 3% oxygen.

ND = not detected (detection limit in parenthesis).

N/A = detection limit not applicable. QA samples indicate the detection limit for positive samples.

ng = 1.0E-09g

ug = 1.0E-06g

ppt = parts per trillion, dry volume basis

8760 operating hours per year

1

APPENDIX E-4

RUN-SPECIFIC DIOXIN/FURAN EMISSIONS DATA AT THE ESP OUTLET

(Concentrations Corrected to 3 Percent Oxygen)

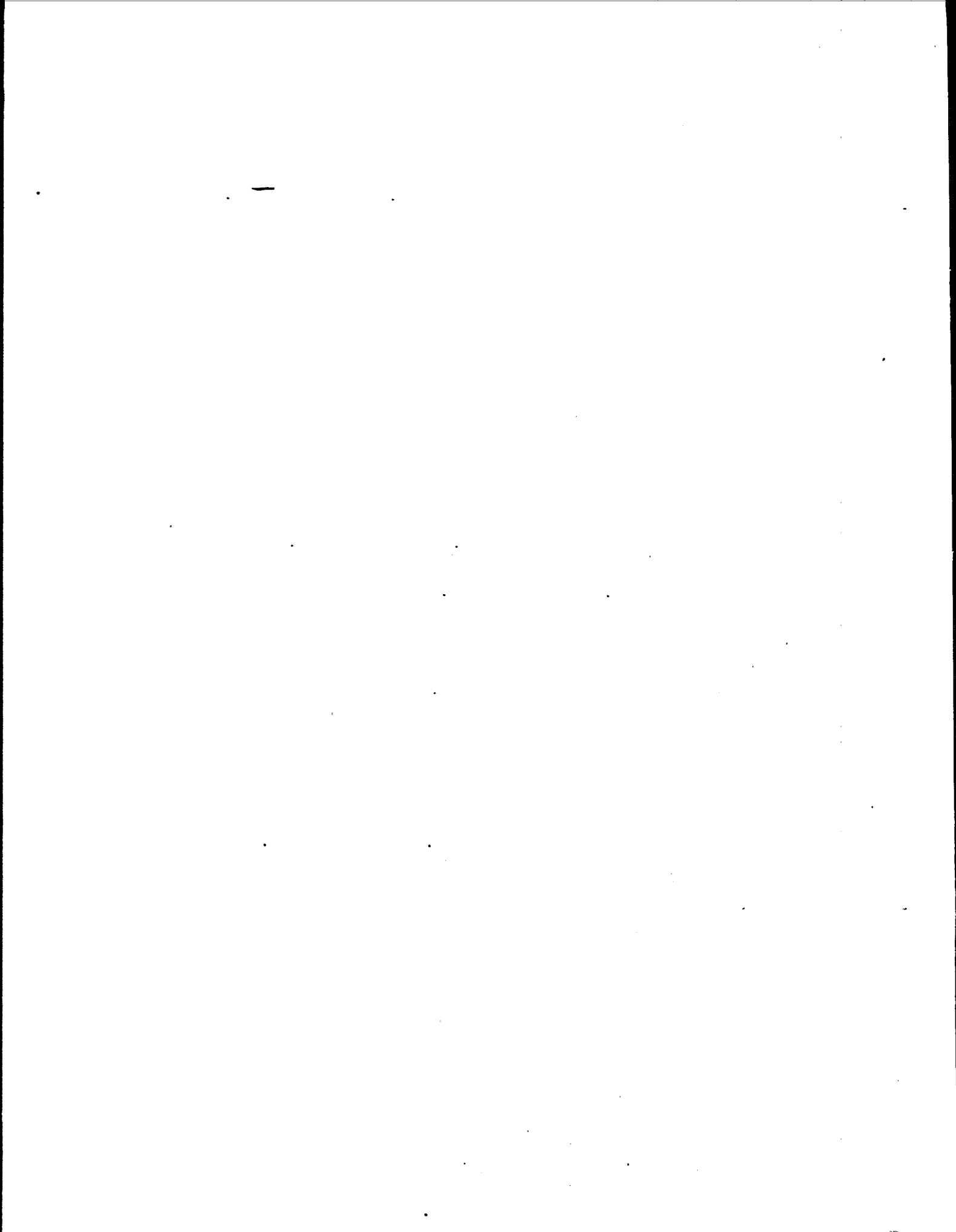


TABLE E-10. DIOXIN/FURAN EMISSIONS DATA AT THE ESP OUTLET FOR RUN 01,
SITE BLB-A (CONCENTRATIONS CORRECTED TO 3 PERCENT OXYGEN)

Dioxin/Furan Isomer	Isomer Concentration In Flue Gas (ng/dscm @ 3% oxygen)	Isomer Concentration In Flue Gas (ppt @ 3% oxygen)	Isomer Hourly Emissions Rate (ug/hr)
DIOXINS			
2378 TCDD	ND (6.75E-02)	ND (5.04E-03)	ND (1.36E+01)
Other TCDD	ND (6.75E-02)	ND (5.04E-03)	ND (1.36E+01)
Penta-CDD	ND (2.83E-02)	ND (1.91E-03)	ND (5.71E+00)
Hexa-CDD	1.74E-01(N/A)	1.07E-02(N/A)	3.52E+01
Hepta-CDD	3.48E-01(N/A)	1.97E-02(N/A)	7.03E+01
Octa-CDD	8.05E-01(N/A)	4.21E-02(N/A)	1.63E+02
Total PCDD	1.33E+00	7.25E-02	2.68E+02
FURANS			
2378 TCDF	ND (8.70E-03)	ND (6.84E-04)	ND (1.76E+00)
Other TCDF	3.70E-01(N/A)	2.91E-02(N/A)	7.47E+01
Penta-CDF	1.74E-01(N/A)	1.23E-02(N/A)	3.52E+01
Hexa-CDF	1.52E-01(N/A)	9.77E-03(N/A)	3.08E+01
Hepta-CDF	2.18E-01(N/A)	1.28E-02(N/A)	4.40E+01
Octa-CDF	8.70E-02(N/A)	4.72E-03(N/A)	1.76E+01
Total PCDF	1.00E+00	6.87E-02	2.02E+02

NOTE: Isomer concentrations shown are corrected to 3% oxygen.

ND = not detected (detection limit in parenthesis).

N/A = detection limit not applicable. QA samples indicate the detection limit for positive samples.

ng = 1.0E-09g

ug = 1.0E-06g

ppt = parts per trillion, dry volume basis

8760 operating hours per year

TABLE E-11. DIOXIN/FURAN EMISSIONS DATA AT THE ESP OUTLET FOR RUN 02,
SITE BLB-A (CONCENTRATIONS CORRECTED TO 3 PERCENT OXYGEN)

Dioxin/Furan Isomer	Isomer Concentration In Flue Gas (ng/dscm @ 3% oxygen)	Isomer Concentration In Flue Gas (ppt @ 3% oxygen)	Isomer Hourly Emissions Rate (ug/hr)
DIOXINS			
2378 TCDD	ND (1.97E-02)	ND (1.47E-03)	ND (3.78E+00)
Other TCDD	5.91E-02 (N/A)	4.42E-03 (N/A)	1.13E+01
Penta-CDD	ND (2.37E-02)	ND (1.60E-03)	ND (4.54E+00)
Hexa-CDD	ND (6.31E-02)	ND (3.88E-03)	ND (1.21E+01)
Hepta-CDD	9.86E-02 (N/A)	5.58E-03 (N/A)	1.89E+01
Octa-CDD	4.34E-01 (N/A)	2.27E-02 (N/A)	8.32E+01
Total PCDD	5.91E-01	3.27E-02	1.13E+02
FURANS			
2378 TCDF	ND (3.94E-02)	ND (3.10E-03)	ND (7.56E+00)
Other TCDF	5.91E-02 (N/A)	4.65E-03 (N/A)	1.13E+01
Penta-CDF	ND (4.53E-02)	ND (3.21E-03)	ND (8.70E+00)
Hexa-CDF	5.91E-02 (N/A)	3.79E-03 (N/A)	1.13E+01
Hepta-CDF	1.97E-01 (N/A)	1.16E-02 (N/A)	3.78E+01
Octa-CDF	2.96E-01 (N/A)	1.60E-02 (N/A)	5.67E+01
Total PCDF	6.11E-01	3.61E-02	1.17E+02

NOTE: Isomer concentrations shown are corrected to 3% oxygen.

ND = not detected (detection limit in parenthesis).

N/A = detection limit not applicable. QA samples indicate the detection limit for positive samples.

ng = 1.0E-09g

ug = 1.0E-06g

ppt = parts per trillion, dry volume basis
8760 operating hours per year

TABLE E-12. DIOXIN/FURAN EMISSIONS DATA AT THE ESP OUTLET FOR RUN 03,
SITE BLB-A (CONCENTRATIONS CORRECTED TO 3 PERCENT OXYGEN)

Dioxin/Furan Isomer	Isomer Concentration In Flue Gas (ng/dscm @ 3% oxygen)	Isomer Concentration In Flue Gas (ppt @ 3% oxygen)	Isomer Hourly Emissions Rate (ug/hr)
DIOXINS			
2378 TCDD	ND (2.03E-02)	ND (1.51E-03)	ND (4.08E+00)
Other TCDD	ND (2.03E-02)	ND (1.51E-03)	ND (4.08E+00)
Penta-CDD	ND (4.05E-02)	ND (2.74E-03)	ND (8.15E+00)
Hexa-CDD	ND (4.42E-02)	ND (2.72E-03)	ND (8.89E+00)
Hepta-CDD	9.21E-02(N/A)	5.21E-03(N/A)	1.85E+01
Octa-CDD	2.40E-01(N/A)	1.25E-02(N/A)	4.82E+01
Total PCDD	3.32E-01	1.77E-02	6.67E+01
FURANS			
2378 TCDF	ND (1.84E-02)	ND (1.45E-03)	ND (3.71E+00)
Other TCDF	3.68E-02(N/A)	2.90E-03(N/A)	7.41E+00
Penta-CDF	ND (2.40E-02)	ND (1.69E-03)	ND (4.82E+00)
Hexa-CDF	ND (9.40E-02)	ND (6.03E-03)	ND (1.89E+01)
Hepta-CDF	5.53E-02(N/A)	3.25E-03(N/A)	1.11E+01
Octa-CDF	1.84E-02(N/A)	9.98E-04(N/A)	3.71E+00
Total PCDF	1.11E-01	7.15E-03	2.22E+01

NOTE: Isomer concentrations shown are corrected to 3% oxygen.

ND = not detected (detection limit in parenthesis).

N/A = detection limit not applicable. QA samples indicate the detection limit for positive samples.

ng = 1.0E-09g

ug = 1.0E-06g

ppt = parts per trillion, dry volume basis

8760 operating hours per year

1. 1

1. 1

1. 1

APPENDIX F
RUN-SPECIFIC RISK MODELING INPUT DATA

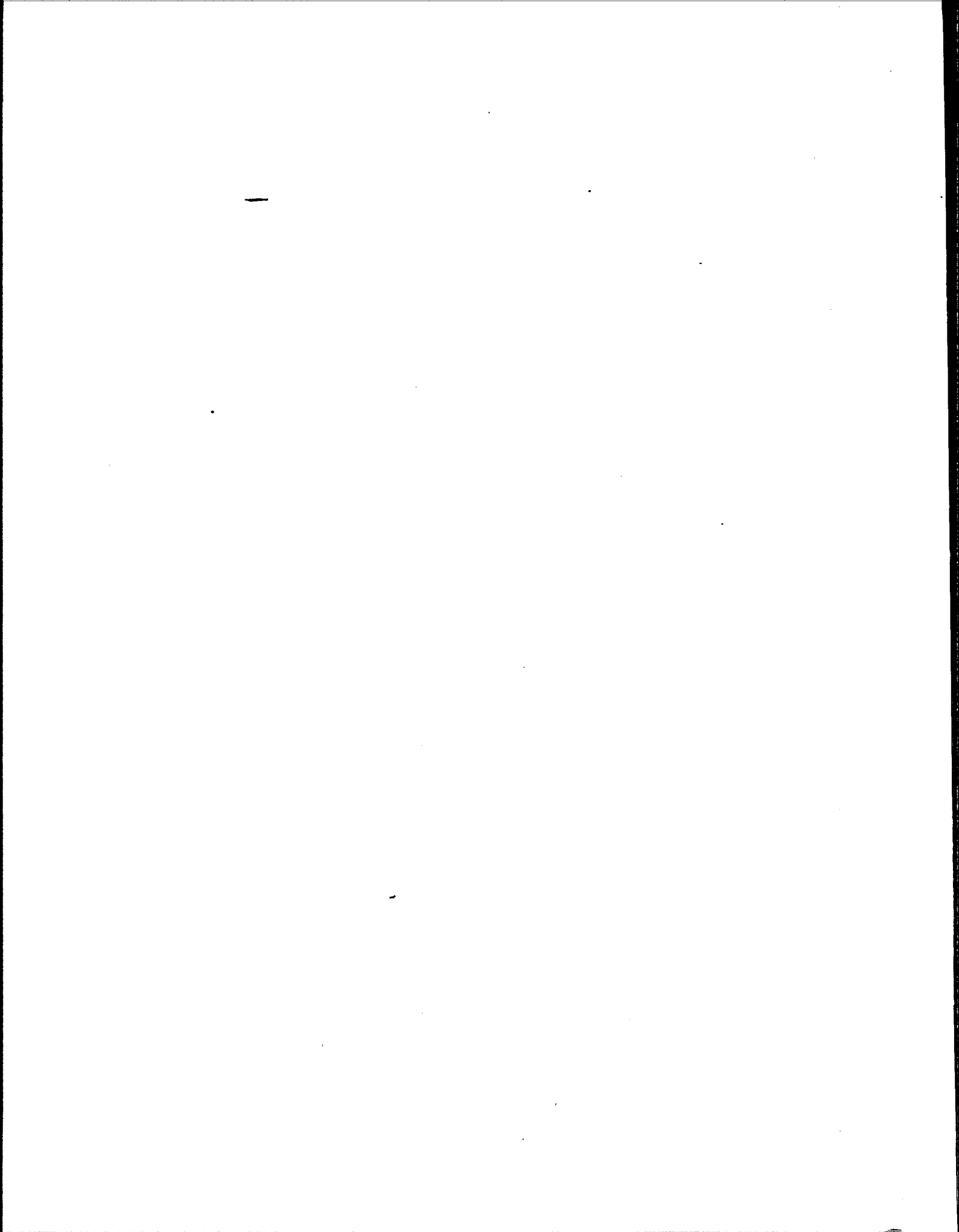


TABLE F-1. RISK MODELING PARAMETERS FOR RUN 01, SITE BLB-A

Latitude = 34 50 39
 Longitude = 80 53 22
 Stack Height (From Grade Level) = 68.6 m
 Stack Diameter (ID) = 3.2 m
 Flue Gas Flow Rate (Dry Standard) = 3787.2 dscmm
 Flue Gas Exit Temperature = 448.0 K
 Flue Gas Exit Velocity (Actual) = 1003.1 mpm

Dioxin/Furan Isomer	Isomer Concentration In Flue Gas (ng/dscm)	Isomer Hourly Emissions Rate (ug/hr)	Relative Potency Factor	2,3,7,8 - TCDD Equivalent Emissions (mg/yr)
2378 TCDD	ND (6.00E-02)	ND (1.36E+01)	1.000	ND (1.19E+02)
Other TCDD	ND (6.00E-02)	ND (1.36E+01)	.010	ND (1.19E+00)
2378 TCDF	ND (7.74E-03)	ND (1.76E+00)	.100	ND (1.54E+00)
Other TCDF	3.29E-01	7.47E+01	.001	6.55E-01
Penta-CDD	ND (2.51E-02)	ND (5.71E+00)	.500	ND (2.50E+01)
Penta-CDF	1.55E-01	3.52E+01	.100	3.08E+01
Hexa-CDD	1.55E-01	3.52E+01	.040	1.23E+01
Hexa-CDF	1.35E-01	3.08E+01	.010	2.70E+00
Hepta-CDD	3.09E-01	7.03E+01	.001	6.16E-01
Hepta-CDF	1.93E-01	4.40E+01	.001	3.85E-01
Octa-CDD	7.16E-01	1.63E+02	.000	.00E+00
Octa-CDF	7.74E-02	1.76E+01	.000	.00E+00
Net 2378 TCDD Equivalent Atmospheric Loading				4.75E+01

ND = not detected (detection limit in parentheses).

N/A = detection limit not available

ng = 1.0E-09g

ug = 1.0E-06g

mg = 1.0E-03g

Standard conditions: 293 K (20 C) temperature and 1 atmosphere pressure.
 8760 operating hours per year

TABLE F-2. RISK MODELING PARAMETERS FOR RUN 02, SITE BLB-A

Latitude = 34 50 39
 Longitude = 80 53 22
 Stack Height (From Grade Level) = 68.6 m
 Stack Diameter (ID) = 3.2 m
 Flue Gas Flow Rate (Dry Standard) = 3738.4 dscmm
 Flue Gas Exit Temperature = 446.2 K
 Flue Gas Exit Velocity (Actual) = 972.5 mpm

Dioxin/Furan Isomer	Isomer Concentration In Flue Gas (ng/dscm)	Isomer Hourly Emissions Rate (ug/hr)	Relative Potency Factor	2,3,7,8 - TCDD Equivalent Emissions (mg/yr)
2378 TCDD	ND (1.69E-02)	ND (3.78E+00)	1.000	ND (3.31E+01)
Other TCDD	5.06E-02	1.13E+01	.010	9.94E-01
2378 TCDF	ND (3.37E-02)	ND (7.56E+00)	.100	ND (6.63E+00)
Other TCDF	5.06E-02	1.13E+01	.001	9.94E-02
Penta-CDD	ND (2.02E-02)	ND (4.54E+00)	.500	ND (1.99E+01)
Penta-CDF	ND (3.88E-02)	ND (8.70E+00)	.100	ND (7.62E+00)
Hexa-CDD	ND (5.40E-02)	ND (1.21E+01)	.040	ND (4.24E+00)
Hexa-CDF	5.06E-02	1.13E+01	.010	9.94E-01
Hepta-CDD	8.43E-02	1.89E+01	.001	1.66E-01
Hepta-CDF	1.69E-01	3.78E+01	.001	3.31E-01
Octa-CDD	3.71E-01	8.32E+01	.000	.00E+00
Octa-CDF	2.53E-01	5.67E+01	.000	.00E+00
Net 2378 TCDD Equivalent Atmospheric Loading				2.58E+00

ND = not detected (detection limit in parentheses).

N/A = detection limit not available

ng = 1.0E-09g

ug = 1.0E-06g

mg = 1.0E-03g

Standard conditions: 293 K (20 C) temperature and 1 atmosphere pressure.
 8760 operating hours per year

TABLE F-3. RISK MODELING PARAMETERS FOR RUN 03, SITE BLB-A

Latitude = 34 50 39
 Longitude = 80 53 22
 Stack Height (From Grade Level) = 68.6 m
 Stack Diameter (ID) = 3.2 m
 Flue Gas Flow Rate (Dry Standard) = 3607.3 dscmm
 Flue Gas Exit Temperature = 445.2 K
 Flue Gas Exit Velocity (Actual) = 961.2 mpm

Dioxin/Furan Isomer	Isomer Concentration In Flue Gas (ng/dscm)	Isomer Hourly Emissions Rate (ug/hr)	Relative Potency Factor	2,3,7,8 - TCDD Equivalent Emissions (mg/yr)
2378 TCDD	ND (1.88E-02)	ND (4.08E+00)	1.000	ND (3.57E+01)
Other TCDD	ND (1.88E-02)	ND (4.08E+00)	.010	ND (3.57E-01)
2378 TCDF	ND (1.71E-02)	ND (3.71E+00)	.100	ND (3.25E+00)
Other TCDF	3.42E-02	7.41E+00	.001	6.49E-02
Penta-CDD	ND (3.77E-02)	ND (8.15E+00)	.500	ND (3.57E+01)
Penta-CDF	ND (2.23E-02)	ND (4.82E+00)	.100	ND (4.22E+00)
Hexa-CDD	ND (4.11E-02)	ND (8.89E+00)	.040	ND (3.12E+00)
Hexa-CDF	ND (8.73E-02)	ND (1.89E+01)	.010	ND (1.66E+00)
Hepta-CDD	8.56E-02	1.85E+01	.001	1.62E-01
Hepta-CDF	5.14E-02	1.11E+01	.001	9.74E-02
Octa-CDD	2.23E-01	4.82E+01	.000	.00E+00
Octa-CDF	1.71E-02	3.71E+00	.000	.00E+00
Net 2378 TCDD Equivalent Atmospheric Loading				3.25E-01

ND = not detected (detection limit in parentheses).

N/A = detection limit not available

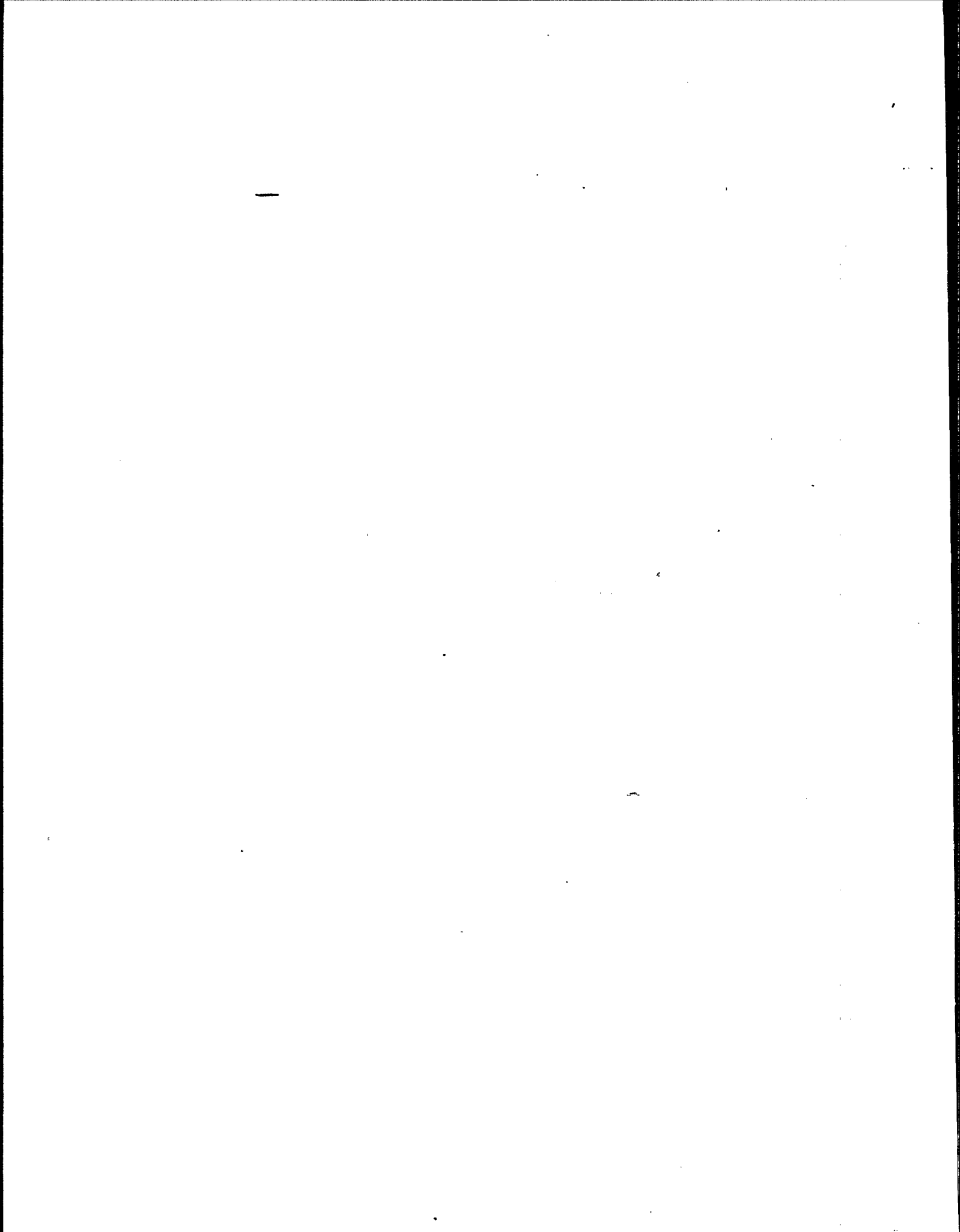
ng = 1.0E-09g

ug = 1.0E-06g

mg = 1.0E-03g

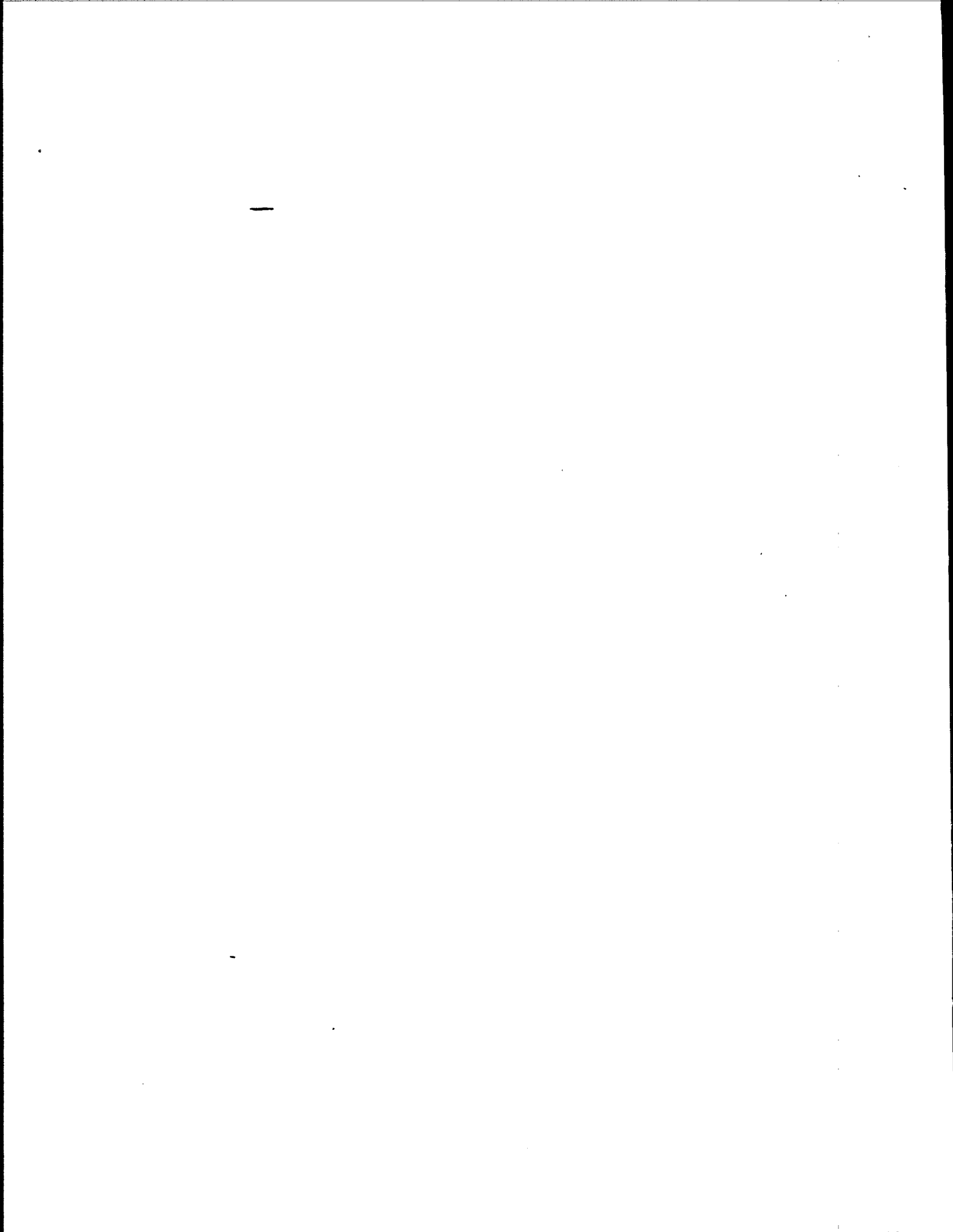
Standard conditions: 293 K (20 C) temperature and 1 atmosphere pressure.

8760 operating hours per year



APPENDIX G

ERROR ANALYSIS OF CONTROL DEVICE EFFICIENCY CALCULATIONS



APPENDIX G

ERROR ANALYSIS OF CONTROL DEVICE EFFICIENCY CALCULATIONS

Objective: Given the analytical uncertainty of the dioxin/furan analyses ($\pm 50\%$ accuracy), estimate the uncertainty of the control device efficiency calculations.

Let: $C_{out,meas}$ = the measured concentration of a given dioxin/furan homologue at the outlet location.

$C_{in,meas}$ = the measured concentration of a given dioxin/furan homologue at the inlet location.

$C_{out,max}$ = the maximum possible concentration of the dioxin/furan homologue given the measured value $C_{out,meas}$.

$C_{out,min}$ = the minimum possible concentration of the dioxin/furan homologue given the measured value $C_{out,meas}$.

$C_{in,max}$ = the maximum possible concentration of the dioxin/furan homologue, given the measured value $C_{in,meas}$.

$C_{in,min}$ = the minimum possible concentration of the dioxin/furan homologue, given the measured value $C_{in,meas}$.

E = the removal efficiency of the control device

Assuming ± 50 percent analytical accuracy:

$$C_{min} = C_{meas} - 0.5 C_{meas} = 0.5 C_{meas}$$

$$C_{max} = C_{meas} + 0.5 C_{meas} = 1.5 C_{meas}$$

Note that:
$$E_{max} = \frac{C_{in,max} - C_{out,min}}{C_{in,max}} = 1 - \frac{C_{out,min}}{C_{in,max}}$$

$$\begin{aligned} E_{max} &= 1 - \frac{0.5 C_{out,meas}}{1.5 C_{in,meas}} = 1 - \frac{1}{3} (1 - E_{meas}) \\ &= \frac{2}{3} + \frac{1}{3} E_{meas} \end{aligned}$$

and:

$$\begin{aligned} E_{\min} &= \frac{C_{\text{in,min}} - C_{\text{out,max}}}{C_{\text{in,min}}} = 1 - \frac{C_{\text{out,max}}}{C_{\text{in,min}}} \\ &= 1 - \frac{1.5 C_{\text{out,meas}}}{0.5 C_{\text{in,meas}}} \\ &= 1 - 3 (1 - E_{\text{meas}}) \end{aligned}$$

$$E_{\min} = 3 E_{\text{meas}} - 2$$

Now, $E_{\min} > 0 \implies$ positive control (i.e., emissions reduction across the control device)

$$(3E_{\text{meas}} - 2) > 0$$

$$E_{\text{meas}} > 2/3$$

Therefore, if E_{meas} is larger than 66.7 percent, the true removal efficiency can safely be assumed to be greater than zero.

And, $E_{\max} < 0 \implies$ negative control (i.e., emissions increase across the control device)

$$2/3 + 1/3 E_{\text{meas}} < 0$$

$$E_{\text{meas}} < -2$$

Therefore, if E_{meas} is less than -200 percent, the true efficiency can safely be assumed to be less than zero.

To summarize:

$E_{\text{meas}} > 66.7 \text{ percent} \implies$ positive control

$-200 < E_{\text{meas}} < 66.7 \text{ percent} \implies$ no definitive conclusions can be drawn

$E_{\text{meas}} < -200 \text{ percent} \implies$ negative control

TECHNICAL REPORT DATA (Please read Instructions on the reverse before completing)		
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7. AUTHOR(S) Lawrence E. Keller, Dennis R. Kinsley Robert F. Jongleux	10. PROGRAM ELEMENT NO.	
9. PERFORMING ORGANIZATION NAME AND ADDRESS Radian Corporation Post Office Box 13000 Research Triangle Park, NC 27709	11. CONTRACT/GRANT NO. 68-03-3148	
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	15. SUPPLEMENTARY NOTES EPA Project Officers: Donald Oberacker, ORD William B. Kuykendal, OAQPS	
16. ABSTRACT This report summarizes the results of a dioxin/furan emissions test of a black liquor recovery boiler equipped with a wet-bottom electrostatic precipitator for particulate matter emissions control. Black liquor recovery boilers are used at kraft pulp mills to produce process steam and to reclaim inorganic chemicals from spent wood pulping liquors. The test is the forth in a series of several dioxin/furan emissions tests being conducted under Tier 4 of the National Dioxin Study. The primary objective of Tier 4 is to determine if various combustion sources are sources of dioxin and/or furan emissions. If any of the combustion sources are found to emit dioxin or furan, the secondary objective of Tier 4 is to quantify these emissions. Black liquor recovery boilers are one of 8 combustion source categories tested in the Tier 4 program. The test black liquor boiler, hereafter referred to as boiler BLB-A, was selected for this test after an initial information screening and a one-day pretest survey visit. Data presented in the report include dioxin (tetra through octa homologue + 2378 TCDD) and furan (tetra through octa homologue + 2378 TCDF) results for stack samples. In addition, process data collected during sampling are also presented.		
17. KEY WORDS AND DOCUMENT ANALYSIS		
a. DESCRIPTORS Air Emissions Combustion Sources Dioxin Furans 2,3,7,8 Tetrachlorodibenzo-p-dioxin Black Liquor Boiler Pulp and Paper	b. IDENTIFIERS/OPEN ENDED TERMS Air Pollution Emissions Data	c. COSATI Field/Group
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