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ENVIRONMENTAL ASSESSMENT OF
AN ENHANCED OIL RECOVERY
STEAM GENERATOR EQUIPPED
WITH A LOW-NOX BURNER
Volume I. Technical Results

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ENVIRONMENTAL ASSESSMENT OF AN ENHANCED OIL RECOVERY STEAM GENERATOR EQUIPPED WITH A LOW-NO_x BURNER

Volume I Technical Results

By

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SECTION 1

INTRODUCTION

This report describes and presents results for a set of environmental assessment tests performed for the Environmental Protection Agency's Air and Energy Engineering Research Laboratory (EPA/AEERL) under the Combustion Modification Environmental Assessment (CMEA) program, EPA Contract No. 68-02-3188. The CMEA started in 1976 with a 3-year study, the NO $_{\rm X}$ Control Technology Environmental Assessment (NO $_{\rm X}$ EA, EPA Contract No. 68-02-2160), having the following four objectives:

- Identify potential multimedia environmental effects of stationary combustion sources and combustion modification technology
- Develop and document control application guidelines to minimize these effects
- Identify stationary source and combustion modification R&D priorities
- Disseminate program results to intended users

During the first year of the NO_X EA, data for the environmental assessment were compiled and methodologies were developed. Furthermore, priorities for the schedule and level of effort to be devoted to evaluating the various source/fuel/control combinations were identified. This effort revealed major data gaps, particularly for noncriteria pollutants (organic emissions and trace elements) for virtually all combinations of stationary

combustion sources and combustion modification techniques. Consequently, a series of seven environmental field test programs was undertaken to fill these data gaps. The results of these tests are documented in seven individual reports (References 1-1 through 1-7) and in the NO_X EA final report summarizing the entire 3-year effort (Reference 1-8).

The current CMEA program has, as major objectives, the continuation of multimedia environmental field tests initiated in the original NO_X EA program. These new tests, using standardized Level 1 sampling and analytical procedures (Reference 1-9) are aimed at filling the remaining data gaps and addressing the following priority needs:

- Advanced NO_x controls
- Alternate fuels
- Secondary sources
- EPA program data needs
 - -- Residential oil combustion
 - -- Wood firing in residential, commercial, and industrial sources
 - -- High interest emissions determination (e.g., listed and candidate hazardous air pollutant species)
- Nonsteady-state operation

The petroleum reserves which can be recovered through primary production methods have been essentially exhausted in the oil fields in Kern County, California. These fields still contain significant reserves, although the remaining crude is too viscous to be produced by normal means. This crude is currently being produced using what has been termed enhanced oil recovery (EOR). In one popular process, near saturated (80 to 90 percent quality)

steam is injected into a field. This steam heats the oil, thereby decreasing its viscosity and allowing it to be pumped.

The steam for injection is raised by crude oil-fired steam generators (often termed steamers) which have uncontrolled NO $_{\rm X}$ emissions in the 300 ppm range. Since Kern County is only in borderline attainment of the NO $_{\rm Z}$ ambient air quality standard, EOR steamers have received close regulatory attention with respect to reducing NO $_{\rm X}$ emissions.

One approach to reducing NO_X emissions from these steamers incorporates a $low-NO_X$ emission burner design. One such burner was developed in Japan by Mitsubishi Heavy Industries (MHI) and is currently marketed in the United States by CE-Natco (a steamer manufacturer).

A steamer equipped with an MHI low- NO_X burner was tested in the current CMEA program. These tests, described in this report, were conducted to quantify a broad emissions spectrum from the burner and to compare selected species emissions to those from a steamer equipped with a "standard" burner. Thus, a similar unit with a standard burner was also tested (in less depth, however) in this program.

In addition to the tests described in this report, another EOR steamer, this one equipped with a $low-NO_X$ burner developed under EPA contract by the Energy & Environmental Research Corporation, was also tested. Results from these tests are documented in Reference 1-10.

Table 1-1 lists all the tests performed in the CMEA program, outlining the source tested, fuel used, combustion modifications implemented and the level of sampling and analysis performed in each case. Results of these test programs are discussed in separate reports.

TABLE 1-1. COMPLETED TESTS DURING THE CURRENT PROGRAMa

Source	Description	Test points unit operation	Sampling protocol	Test collaborator		
Spark-ignited, natural- gas-fired reciprocating internal combustion engine	Large bore, 6-cylinder, opposed piston, 186-kk (250 Bhp)/cyl, 900-rpm Model 38TDS8-1/8	Baseline (pre-NSPS) Increased air-fuel ratio aimed at meeting proposed NSPS of 700 ppm corrected to 15 percent 02 and standard atmospheric conditions	Engine exhaust: SASS Method 5 Gas sample (C ₁ -C ₆ HC) Continuous NO, NO _x , CO, CO ₂ , O ₂ , CH ₄ , TUHC Fuel Lube of	Fairbanks Morse Division of Colt Industries		
Compression ignition, diesel-fired, reciprocating internal combustion engine	Large bore, 6-cylinder opposed piston, 261-kW (350 Bhp)/cyl, 900-rpm Model 38TDD8-1/8	Baseline (pre-NSPS) Fuel injection retard aimed at meeting pro-posed NSPS of 600 ppm corrected to 15 percent 02 and standard atmospheric conditions	Engine exhaust: SASS Method 8 Method 5 Gas sample (C ₁ -C ₆ HC) Continuous NO, NO _x , CO, CO ₂ , O ₂ , CH ₄ , TUHC Fuel Lube oil	Fairbanks Morse Division of Colt Industries		
Low-NO _X , residential, condensing-heating system furnished by Karlsons Blueburner Systems Ltd. of Canada	Residential hot water heater equipped with M.A.N. low-MO, burner, 0.55 ml/s (0.5 gal/hr) firing capacity, condensing flue gas	Low-NO _x burner design by M.A.H.	Furnace exhaust: SASS Method 8 Method 5 Gas sample (C1-C6 HC) Continuous NO, NO, CO, CO2, O2, CH4, TUHC Fuel Waste water	New test		
Rocketdyne/EPA low-NO _x residential forced warm air furnace	Residential warm air furnace with modified high-pressure burner and firebox, 0.83 ml/s (0.75 gal/hr) firing capacity	Low-NO _x burner design and integrated furnace system	Furnace exhaust: SASS Method 8 Controlled condensation Method 5 Gas sample (C ₁ -C ₆ HC) Continuous NO, NO, CO, CO ₂ , O ₂ , CH ₄ , TUHĈ	New test		
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TABLE 1-1. (continued)

Description	Test points unit operation	Sampling protocol	Test collaborator
400-MW tangentially fired; new MSPS design aimed at meeting 301 ng/J NO _X limit	ESP inlet and outlet, one test	ESP inlet and outlet SASS Method 5 Controlled condensation Gas sample (C ₁ -C ₆ HC) Continuous NO, NO _x , CO, CO ₂ , O ₂ Coal Bottom ash ESP ash	Exxon Research and Engineering (ER&E) conducting cor- rosion tests
1.14 kg/s steam (9,000 lb/hr) firetube fired with a mixture of coal-oil-water (COW)	Baseline (COW) Controlled SO ₂ emissions with limestone addition	Boiler outlet SASS Method 5 Method 8 Controlled condensation Gas sample (C ₁ -C ₆ HC) Continuous O ₂ , CO ₂ , CO, NO _x	Envirocon per- formed particulate and sulfur emission tests
1.89 kg/s steam (15,000 lb/hr) hot water firetube fired with a mixture of coal-oil- water (COW)	Baseline (COW) Controlled SO ₂ emissions with soda ash (Na ₂ CO ₃) addition	Boiler outlet SASS Method 5 Method 8 Controlled condensation Gas Sample (C ₁ -C ₆ HC) Continuous O ₂ , CO ₂ , NO _X , SO ₂ , CO	Adelphi University
3.03 kg/s steam (24,000 lb/hr) watertube fired with a mixture of coal-oil (COM)	Baseline test only with COM	Boiler outlet SASS Method 5 Controlled condensation Continuous O ₂ , CO ₂ , NO _x , TUHC, CO Fuel	PETC and General Electric (GE)
	400-MW tangentially fired; new NSPS design aimed at meeting 301 ng/J NO _X limit 1.14 kg/s steam (9,000 lb/hr) firetube fired with a mixture of coal-oil-water (COW) 1.89 kg/s steam (15,000 lb/hr) hot water firetube fired with a mixture of coal-oil-water (COW) 3.03 kg/s steam (24,000 lb/hr) watertube fired with a mixture of	400-MW tangentially fired; new NSPS design aimed at meeting 301 ng/J NO _X limit 1.14 kg/s steam (9,000 lb/hr) firetube fired with a mixture of coal-oil-water (COW) 1.89 kg/s steam (15,000 lb/hr) hot water firetube fired with a mixture of coal-oil-water (COW) 1.89 kg/s steam (15,000 lb/hr)	400-MM tangentially fired; new MSPS design aimed at meeting 301 ng/J NO _X limit 1.14 kg/s steam (9,000 lb/hr) firetube fired with a mixture of coal-oil (COM) 1.89 kg/s steam (IS,000 lb/hr) attertube fired with a mixture of coal-oil (COM) 3.03 kg/s steam (24,000 lb/hr) watertube fired with a mixture of coal-oil (COM) 3.03 kg/s steam (24,000 lb/hr) watertube fired with a mixture of coal-oil (COM) 3.03 kg/s steam (24,000 lb/hr) watertube fired with a mixture of coal-oil (COM) 3.03 kg/s steam (24,000 lb/hr) watertube fired with a mixture of coal-oil (COM) 3.03 kg/s steam (24,000 lb/hr) watertube fired with a mixture of coal-oil (COM) 3.03 kg/s steam (24,000 lb/hr) watertube fired with a mixture of coal-oil (COM) 3.03 kg/s steam (24,000 lb/hr) watertube fired with a mixture of coal-oil (COM) 3.03 kg/s steam (24,000 lb/hr) watertube fired with a mixture of coal-oil (COM) 3.03 kg/s steam (24,000 lb/hr) watertube fired with a mixture of coal-oil (COM) 3.03 kg/s coal let and outlet SASS Method 5 Method 5 Method 8 Controlled condensation SO2, CO2, NO2, SO2, CO2, NO3, SO2, CO3, NO3, SO3, CO3, NO3, SO3, CO3, CO3, NO3, SO3, CO3, NO3, TUHC, CO

TABLE 1-1. (continued)

Source	Description	Test points unit operation	Sampling protocol	Test collaborator
TOSCO Refinery vertical crude oil heater	2.54 Ml/day (16,000 bbl/day) natural draft process heater burning oil/refinery gas	Baseline Staged combustion using air injection lances	Heater outlet SASS Method 5 Controlled condensation Gas sample (C ₁ -C ₆ HC) Continuous O ₂ , NO _X , CO, CO ₂ , HC N ₂ O grab sample Fuel of 1 Refinery gas	KYB coordinating the staged com- bustion operation and continuous emission monitoring
Mohawk-Getty 011 industrial boiler	8.21 kg/s steam (65,000 1b/hr) watertube burning mixture of refinery gas and residual oil	Baseline Ammonia injection using the noncatalytic Thermal DeNO _X Process	Economizer outlet SASS Method 5, 17 Controlled condensation Gas Sample (C ₁ -C ₆ HC) Ammonia emissions H ₂ O grab sample Continuous O ₂ , NO _x , CO, CO ₂ Fuels (refinery gas and residual oil)	New test
Industrial boiler	2.52 kg/s steam (20,000 lb/hr) watertube burning wood waste	Baseline (dry wood) Wet (green) wood	Boiler outlet SASS Method 5 Controlled condensation Gas sample (C ₁ -C ₅ HC) Continuous O ₂ , NO _x , CO Fuel Flyash	North Carolina Department of Natural Resources, EPA IERL-RTP
industrial boiler	3.16 kg/s steam (29,000 lb/hr) firetube with refractory firebox burning wood waste	Baseline (dry wood)	Outlet of cyclone particulate collector SASS Method 5 Controlled condensation Gas sample (C ₁ -C ₆ HC) Continuous O ₂ , NO _x , CO Fuel Bottom ash	North Carolina Department of Natural Resources, EPA IERL-RTP

TABLE 1-1. (continued)

Source	Description	Test points unit operation	Sampling protocol	Test callaborator
Enhanced oil recovery steam generator	15 MM (50 million Btu/hr) steam generator burning crude ofl equipped with MHI low-HO _X burner	Performance mapping Low NO _X operation	Steamer outlet: SASS Hethod 5 Method 8 Andersen impactors Gas sample (C ₁ - C ₆ HC) Continuous O2, NO _X , CO, CO ₂ N ₂ O grab sample Fuel	Getty Oil Company, CE-Natco
Pittsburgh Energy Technology Center (PETC) industrial boiler	3.03 kg/s steam (24,000 lb/hr) watertube fired with a coal-water slurry (CWS)	Baseline test only with CWS	Boiler outlet: SASS Method 5 Method 8 Gas sample (C ₁ - C ₆ HC) Continuous O ₂ , NO _X , CO, CO ₂ , TUHC N ₂ O grab sample Fuel Bottom ash Collector hopper ash	PETC and General Electric
Spark-ignited, natural gas-fuel reciprocating internal combustion engine nonselective NO _X reduction catalyst	610 kH (818 hp) Waukesha rich-burn engine equipped with DuPont NSCR system	Low NO _x (with catalyst) 15-day emissions monitoring	Catalyst inlet and outlet SASS NH3 HCN N ₂ O grab sample Continuous O ₂ , CO ₂ , NO _X TUHC Lube oil	Southern California Gas Company
Industrial boiler	180 kg/hr steam (400 lb/hr) stoker fired with a mixture of coal and waste plastic beverage containers	Baseline (coal) Coal and plastic waste	Boiler outlet SASS YOST Method 5 Hctlod 8 HCl Continuous O ₂ , NO _X , CO, CO ₂ , TUHC H ₂ O grab sample Fuel Bottom ash Cyclone ash	Vermont Agency of Environmental Conservation

TABLE 1-1. (concluded)

Source	Description	Test points unit operation	Sampling protocol	Test callaborator
Industrial boiler	7.6 kg/s steam (60,000 lb/hr) watertube retrofit for coal-water-slurry (CWS) firing	Baseline test with CWS 30-day emissions monitoring	Boiler outlet SASS VOST Method 5 Method 8 Gas sample (C ₁ -C ₆ HC) N ₂ O grab sample Continuous NO _x , CO, CO ₂ , O ₂ , TUHC, SO ₂	EPRI, OuPont
Enhanced oil recovery steam generator	15 MM (50 million Btu/hr) steam generator burning crude oil, equipped with the EPA/EER low NO _X burner	Low NO _X (with burner) 30-day emission monitoring	Steamer outlet SASS YOST Method 5 Method 8 Controlled condensation Andersen impactors Grab sample (C ₁ -C ₆ HC) N ₂ O grab sample Continuous NO _X , CO, CO ₂ , O ₂ , SO ₂ Fuel	Chevron U.S.A., EERC
Spark-ignited, natural gas-fired reciprocating internal combustion engine selective NO _X reduction catalyst	1490 kW (2000 hp) Ingersoll-Rand lean burn engine equipped with Englehard SCR system	Low NO _X (with catalyst) 15-day emissions monitoring	Catalyst inlet and outlet SASS YOST NH3 HCN N ₂ O grab sample Continuous O ₂ , CO ₂ , CO, NO, NO _x , NO _x + NH ₃	Southern California Gas Company

^aAcronymns used in the table: EERC, The Energy and Environmental Research Corporation; EPA IERL-RTP. The Environmental Protection Agency's Industrial Environmental Research Laboratory-Research Triangle Park; EPRI, The Electric Power Research Institute; HC, hydrocarbons; NSCR, nonselective catalytic reduction; NSPS, new source performance standard; SASS, source assessment sampling system; SCR, selective catalytic reduction; TUHC, total unburned hydrocarbon; VOST, volatile organic sampling train

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SECTION 2

SOURCE DESCRIPTION

Tests were performed on two CE-Natco model STOF steam generators rated at 50 million Btu/hr heat output. One unit was equipped with a standard North American burner; the other had been retrofitted with the PM low-NO $_{\rm X}$ burner manufactured by Mitsubishi Heavy Industries (MHI). The primary objective of the tests was to measure the NO $_{\rm X}$ reduction performance of this burner as a function of its operational parameter settings (when compared to a standard burner) and to obtain data on emissions of noncriteria pollutant categories and species at a nominal low-NO $_{\rm X}$ setting.

Figure 2-1 illustrates the physical design of the MHI PM burner. As shown, the rectangular burner throat is divided into five nozzles. Typically about 30 percent of the total combustion air is delivered through the central primary air nozzle. This air is mixed with a centralized oil spray comprising approximately half the total fuel fired, forming an oxygen deficient diffusion flame. A premixed flame is obtained by mixing the remaining fuel with about 60 percent of the total air, evenly delivered through each of the upper and lower nozzles. This mixing takes place in a zone offset from the burner which delays ignition until the fuel and air have mixed.

The remaining combustion air (about 10 percent) is delivered through an overfire air (OFA) injection system which injects this air approximately

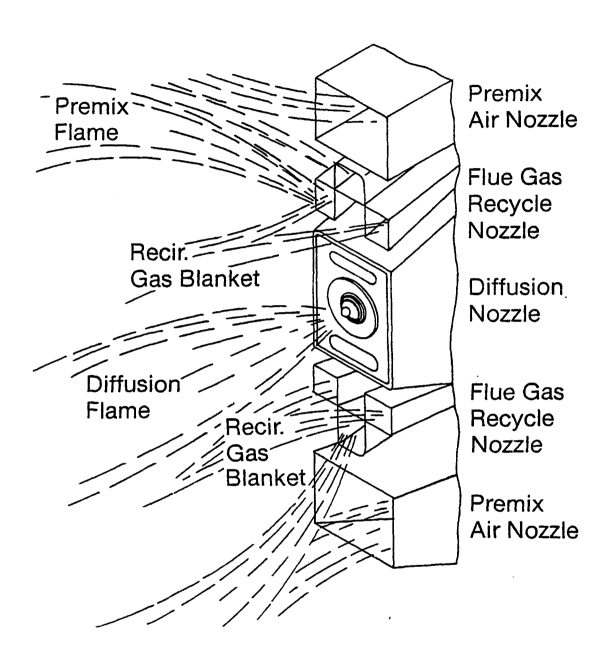


Figure 2-1. The MHI PM burner nozzle.

halfway down the length of the cylindrical furnace through three sets of three ports equally spaced around the furnace circumference. This OFA is designed to ensure that sufficient excess air and mixing are achieved before combustion gas leaves the furnace.

Recirculated flue gas is delivered to the nozzles separating the central (diffusion) and outer (premix) air nozzles. This gas is used to shape the diffusion flame and to maintain separation between the diffusion and premixed flames. Typically about 15 percent of the total combustion product gas is recirculated.

Figure 2-2 shows a sketch of the steamer retrofitted with the burner system. The additional flue gas recirculation (FGR) and OFA systems along with the burner are shown.

In the test program performed, one day of flue gas emission testing was performed on the steamer equipped with the conventional burner. In these tests, flue gas NO_{X} emissions were measured at two steamer loads while varying the excess air fuel. The steamer equipped with the $\mathrm{low-NO}_{\mathrm{X}}$ burner was then subjected to two days of performance/emissions mapping tests in which flue gas composition (NO_{X} , CO , CO_{2} , O_{2} and smoke) was characterized while varying burner operation at full steamer load. In these tests, the following were varied: the FGR rate; the relative distribution of combustion air among the premixed flame nozzles, the diffusion flame nozzles, and the OFA ports; and the overall excess air level. Finally, comprehensive emissions testing (flue gas organics, particulate load, particle-size distribution, and SO_{2} and SO_{3} emissions) was performed on the $\mathrm{low-NO}_{\mathrm{X}}$ burner-equipped steamer with the burner set at a nominal $\mathrm{low-NO}_{\mathrm{X}}$ condition.

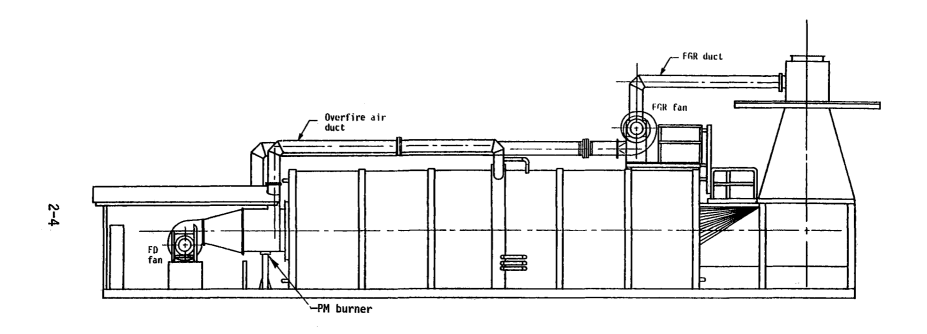


Figure 2-2. Schematic of test steamer.

The fuel fired in both steamers for all tests was local Kern County crude. The fuel ultimate analysis is given in Table 2-1.

The concentrations of 72 trace elements in the fuel were also obtained using spark source mass spectrometry (SSMS) supplemented by atomic absorption spectrometry. Results of these analyses are given in Table 2-2.

TABLE 2-1. FUEL ULTIMATE ANALYSIS

Component (wt percent)	
Carbon Hydrogen Sulfur Nitrogen Oxygen Ash	86.88 10.84 1.06 0.76 0.43 0.03
Higher heating, MJ/kg value (Btu/lb)	43.2 (18,560)
API gravity	13.3

TABLE 2-2. FUEL TRACE ELEMENT CONCENTRATIONS

Element	Concentration (µg/g)	Element	Concentration (µg/g)	Element	Concentration (µg/g)
Aluminum	4.0	Holmium		Samarium	
Antimony	0.07	Iodine		Scandium	0.08
Arsenic	0.4	Iridium		Selenium	0.2
Barium	0.5	Iron	95	Silicon	34
Beryllium	<0.01	Lanthanum	0.4	Silver	0.03
Bismuth	a	Lead	0.3	Sodium	22
Boron	0.2	Lithium	0.1	Strontium	0.5
Bromine	0.3	Lutetium		Tantalum	2
Cadmium	~~	Magnesium	14	Tellurium	0.06
Calcium	48	Manganese	0.4	Terbium	
Cerium	0.2	Mercury	<0.01	Thallium	
Cesium		Molybdenum	1	Thorium	0.8
Chlorine	4	Neodymium	<0.2	Thulium	
Chromium	0.8	Nickel	90	Tin	0.07
Cobalt	2	Niobium	<0.03	Titanium	3
Copper	5	Osmium	~	Tungsten	
Dysprosium		Palladium .		Uranium	
Erbium		Phosphorus	0.8	Vanadium	80
Europium		Platinum		Ytterbium	
Fluorine	27	Potassium	4	Yttrium	0.6
Gadolinium		Praseodymium	0.06	Zinc	1
Gallium	0.5	Rhenium		Zirconium	0.3
Germanium	0.2	Rhodium			
Gold		Rubidium	0.02		
Hafnium		Ruthenium			

 $^{^{\}rm a}$ -- denotes present at less than the detection limit of 0.02 µg/g.

SECTION 3

EMISSION RESULTS

As noted in Section 2, the objectives of these tests were to evaluate the NO_{X} emission reduction performance of the MHI PM low NO_{X} burner system retrofitted to an enhanced oil recovery steam generator and to quantitate emissions of noncriteria pollutant species from the retrofit steamer. To satisfy these objectives a brief series of flue gas emission measurement tests was performed on an identical unit equipped with a conventional burner. A relatively comprehensive series of performance/emission mapping tests was performed next on the unit equipped with the low NO_{X} burner. Finally, a set of comprehensive environmental assessment flue gas characterization tests was performed on the low NO_{X} burner equipped-steamer with burner operation set at a nominal low NO_{X} setting.

Section 3.1 summarizes results of the tests of the conventional burner-equipped steamer and the performance/emission mapping tests on the low ${\rm NO}_{\rm X}$ burner-equipped steamer. Results of the comprehensive emission testing of the low ${\rm NO}_{\rm X}$ burner-equipped steamer are discussed in Section 3.2.

3.1 PERFORMANCE/EMISSION MAPPING TESTS

Flue gas emissions of NO_X , CO, CO_2 , O_2 , and stack gas smoke readings were measured on the steamer equipped with a conventional North American burner at two loads (full load and about 75 percent of rated capacity) and several excess air settings. These measurements were performed at the stack

using a continuous flue gas monitoring system as described in Appendix A. To supplement these measurements, Getty Oil Company personnel performed complementary monitoring of the combustion gas at the steamer furnace exit.

Results of these tests are summarized in Table 3-1. The data in the table clearly suggest that there was some combustion gas dilution through air inleakage between the furnace outlet sampling location and the stack sampling location. Stack 0_2 levels are consistently higher and $C0_2$ consistently lower than corresponding furnace outlet levels. CO levels (corrected to 3 percent 0_2) are comparable at the two locations at full load. However, $N0_X$ levels at full load were generally about 40 to 50 ppm higher at the stack location than at the furnace outlet. At 75 percent load, $N0_X$ levels at both locations were comparable; however, CO levels were apparently increased. Reasons for both these apparent increases (if they were indeed real) can only be speculated.

The stack location NO_X emissions data are plotted in Figure 3-1 as NO_X versus stack gas O_2 . The figure shows a steady decrease in NO_X emissions as excess air is reduced until flue gas O_2 falls below about 3 percent. Below this O_2 level, the rate of NO_X emissions increases. However, referring to Table 3-1, as flue gas O_2 is decreased below 3 percent, the smoke number increases to unacceptable levels. For practical operation, then, the conventional burner appears capable of attaining full load NO_X emissions of about 300 ppm (3 percent O_2) with flue gas O_2 about 3.7 percent at acceptable CO emissions and smoke number. At 75 percent load, NO_X emissions are reduced to about 250 ppm (3 percent O_2) at flue gas O_2 of 4.0 percent and acceptable CO and smoke number.

Following the conventional burner-equipped unit testing, a relatively comprehensive series of performance emission mapping tests was performed on

TABLE 3-1. FLUE GAS EMISSIONS SUMMARY: CONVENTIONAL BURNER

	Fuel	flow	Hea	t input	Stack ^a Furnace outle				utle t ^b	tle t ^b				
Test no.	(1/s)	(BPO)	(MW)	(million Btu/hr)	0 ₂ (%)	CO ₂	(ppm)c	(ppm)c	Smoke	0 ₂ (%)	CO ₂	(ppm)c	NO _X	SO ₂
Full load	•••													
1 2 3 4 5 6	0.39 0.39 0.39 0.39 0.39 0.39	210 210 210 210 210 210	16.3 16.3 16.3 16.3 16.3	55.5 55.5 55.5 55.5 55.5 56.3	3.7 5.1 6.6 3.7 2.9 2.3	12.8 12.0 10.8 13.1 12.2 14.4	23 34 31 42 54 46	302 340 365 305 278 197	3.0 4.0 4.0 8.0 8.5	3.0 4.4 6.2 2.8 1.9 1.3	14.1 12.9 11.5 14.1 14.8 15.3	42 48 49 42 46 54	263 288 305 256 227 206	580 580 584 593 598 603
75% load		٠												
7 8 9	0.29 0.29 0.29	160 160 160	12.4 12.4 12.4	42.3 42.3 42.3	4.2 6.4 6.1	13.3 11.7 12.7	78 96 133	246 296 290	4.0 2.5 3.0	2.8 5.3 4.0	14.1 12.1 13.2	40 46 42	254 300 277	584 608 578

aEmission measurements by Acurex.
bEmission measurements by Getty Oil Company.
cOry at 3 percent O₂.

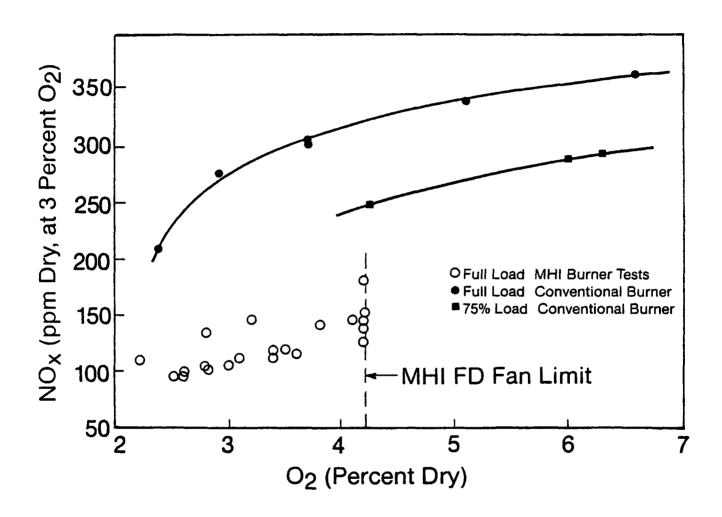


Figure 3-1. NO_X emissions versus flue gas O_2 .

the steamer equipped with the low NO_X burner. Recall from Section 2 that this burner directs combustion air to a central diffusion flame nozzle, outboard premixed flame nozzles, and overfire air injection ports. Thus, the distribution of combustion air among these streams, as well as the total air fired, are adjustable burner operating parameters. In addition, the rate at which flue gas is recirculated to the burner (to separate the diffusion and premixed air flames) is a further adjustable parameter. All of these were varied in the testing performed.

Results of the performance/emission mapping tests are summarized in Table 3-2. Again, flue gas composition both at the steamer stack and at the furnace outlet (measurements performed by Getty Oil Company personnel) are shown. For these tests there was general agreement between the stack O_2 and CO_2 readings and those at the furnace outlet. This suggests that negligible air inleakage occurred between these locations for the low NO_X burner-equipped steamer, in contrast to the apparent case for the steamer with the conventional burner. Corresponding NO_X and CO levels were, in general, similarly comparable (although for a few test points stack CO levels were significantly higher than furnace outlet levels).

The data in Table 3-2 show that NO_X emissions from the unit varied from 95 to 180 ppm (corrected to 3 percent O_2) with changes in the parameters investigated. Certain conditions resulted in NO_X emissions at the stack below 100 ppm (3 percent O_2 , dry), but these were, in general, accompanied by high CO emissions and high smoke spot. Conditions which resulted in NO_X in the 110 ppm range with moderate CO are also noted.

The variation in NO_X emissions with overall excess air (stack gas O_2) for this unit was shown in Figure 3-1. The scatter in the figure results

TABLE 3-2. MHI BURNER PERFORMANCE TEST RESULTS

				Air dist	tribution	Fuel	ra te	Hea	t input		Stackb					Furnace outlet ^c			
Test no.	FGR rate (1)	OFA (%)	Premixa flame air (%)	Diffusiona flame air (%)	(1/s)	(BPD)	(MW)	(Million Btu/hr)	02	CO ₂	(ppm)d	(Pbw)q	TUHC (ppm)d	Smoke	02	CO (ppm)d	(1) CO5	NO _x	50 ₂ (ppm)d
1	8.9	19	48	33	0.383	208	16,1	55.0	3.5	13.4	99	119	3.2	>10	4.3	70	12.9	124	597
2	9.4	18	52	30	0.385	209	16.2	55.2	2.7	13.9	266	102	4.4	10	3.3	80	13.9	113	594
3	13,4	19	52	29	0.381	207	16.0	54.7	2.6	13.9	215	99	3.2	9.5	3.1	80	13.9	108	582
4	9.9	19	53	29	0.390	212	16.4	56.0	2.5	14.0	236	95	5.3	10	3.1	73	14.0	109	592
5	9.9	13	52	35	0.379	206	16.0	54.5	2.6	14.0	. 269	97	3.9	8	3.0	76	13,9	105	576
6	9.4	15	51	34	0.377	205	15,9	54.2	3.4	13.9	60	119	2.2	6	4.0	59	13.0	126	573
7	8.8	15	51	34	0.388	211	16.3	55.8	3.8	13.3	60	140	1.4	3.5	4.4	44	12.9	140	581
8	9.1	11	56	33	0,386	210	16.3	55.5	3.2	13.2	51	145	1.1	3.5	3.3	45	13.9	144	568
9	9.8	9	57	34	0.386	210	16.3	55.5	2.2	14.6	79	110	1.0	8	2.3	58	14.5	112	585
10	9.3	В	57	35	0.388	211	16.3	55.8	3.1	13.3	141	111	4.5	10	2.2	104	14.7	102	616
11	8.8	8	57	35	0.386	210	16.3	55.5	4.1	12.5	70	145	8.5	6	3.2	54	13.8	126	595
12	8.2	8	58	34	0.396	215	16.7	56.8	4.2	12.5	64	180	8.6	3.5	4.2	46	12.9	174	570
13	8.4	8	55	37	0.388	211	16.3	55.8	4.2	12.4	51	126	1.8	4	4.6	55	12.5	125	587
34	8.8	8	55	37	0.390	212	16.4	56.0	3,4	13.0	85	111	1.1	8	3.6	67	13.5	114	573
15	6.6	8	55	37	0.390	212	16.4	56.0	4.2	12.3	66	131	1.4	6	4.5	58	12.7	131	572
16	2,6	8	55	37	0.390	212	16.4	56.0	4.2	12.3	54	152	1.1	2.5	4.6	53	12.6	149	582
17	8.4	7	59	34	0.405	220	17.0	58.2	4.2	12.6	60	143	1.4	4	4.1	50	13.1	144	556
18	9.1	6	58	36	0.377	205	15.9	54.2	3.6	12.9	62	116	0.5	6	3.3	64	13.8	113	558
19	9.8	7	58	35	0.386	210	16.3	55.5	2.8	13.5	80	106	0	8	2.3	87	14.5	98	574
20	9.8	3	62	35	0.388	211	16.3	55.8	2.8	13.6	64	133	0	6	2.5	55	14.5	126	583
21 (SASS)	9.5	10	54	36	0.386	210	16.3	55,5	3.0	13.3	93	106	0	8	2.5	88	14.5	108	586

^aPremix and diffusion nozzle combustion air flows were not measured. Values shown here were estimated based on blower discharge pressure and static pressure readings in the windbox for diffusion and premix zones.

^bEmission measurements by Acurex.

^cEmission measurements by Getty Oil Company.

dDry at 3 percent O₂.

from changes in NO_{X} emissions with the split in air flowrates among the diffusion and premixed flames and the OFA ports, and FGR rates at constant overall excess air. In general, though, NO_{X} emissions with the low- NO_{X} burner at full load were roughly half those of the conventional burner at a given flue gas O_2 .

Figure 3-2 shows steamer stack gas CO emissions versus stack gas 0_2 for both burners at full load. Again the scatter in the data for the low-NO $_{\rm X}$ burner results from variations in air distribution and FGR rate at constant stack 0_2 . The data in Figure 3-2 show that CO emissions from the low-NO $_{\rm X}$ burner increased steeply at flue gas 0_2 below 2.5 to 3.0 percent. This contrasts with conventional burner behavior where CO emissions were still low at flue gas 0_2 down to 2.5 percent. The higher CO levels from the low-NO $_{\rm X}$ burner, which were accompanied by high smoke spot (see Table 3-2) are attributed to flame impingement which was observed at the 4 and 8 o'clock positions at virtually all burner settings. Higher CO levels are attributed to increased flame impingement and excessively low diffusion zone stochiometries during low 0_2 and high OFA tests.

The effect of OFA flowrate on both CO and NO_X levels for the low- NO_X burner is illustrated in Figure 3-3. CO levels decrease sharply at OFA rates below 10 percent. At 3 percent OFA, CO levels are nearly those of the conventional burner (see Figure 3-2). NO_X emissions at minimum OFA, however, are not significantly higher than those at high OFA rates.

The effect of FGR on NO_X and CO emissions from the low- NO_X burner is shown in Figure 3-4. FGR had a greater effect at a higher O_2 and lower OFA levels (4 percent and 8 percent OFA) than it did at lower O_2 and higher OFA levels (O_2 of 2.6 percent and 19 percent OFA). CO responded in an opposite

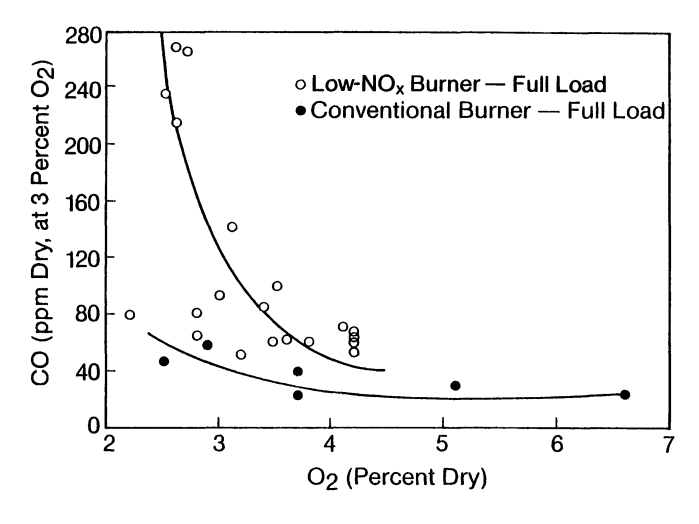
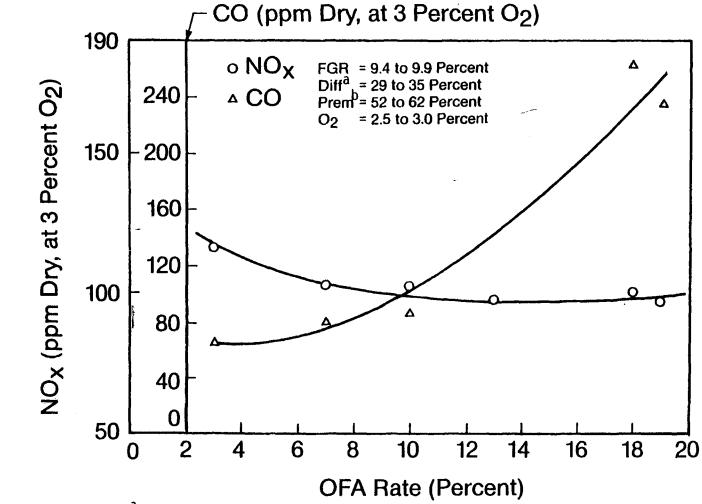
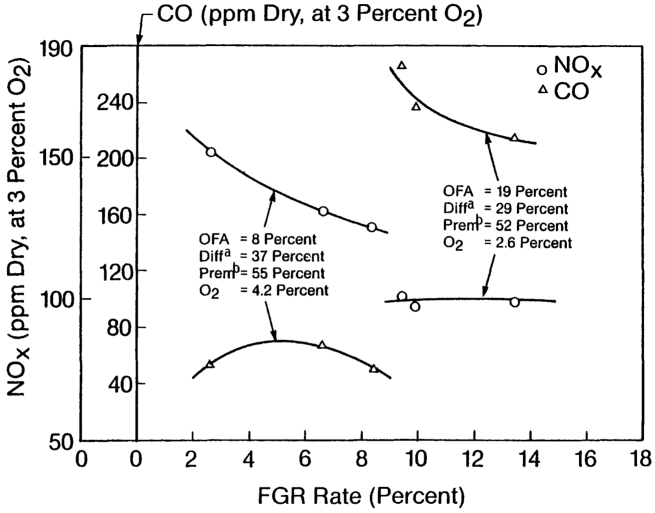


Figure 3-2. CO emissions versus flue gas 0_2 .



^aDiff: Diffusion flame air Prem: Premixed flame air

Figure 3-3. Effect of OFA rate on NO_X and CO emissions from the low- NO_X burner.



a b Prem: Diffusion flame air Premixed flame air

Figure 3-4. Effect of FGR rate on NO_X and CO emissions from the low NO_X burner.

manner. This can be explained in part by the greater mixing occurring at higher burner stoichiometries combined with lower FGR rates. This mixing tended to partly cancel the low- NO_X properties of the split flame. Conversely, the higher FGR rates combined with lower burner stoichiometry, while keeping the flames separate, will tend to cause greater impingement of the premix flame, which increased the CO levels.

Figure 3-5 shows a crossplot of the $\mathrm{NO_X/CO}$ emission data for the $\mathrm{low-NO_X}$ burner. This figure shows that, as the burner is adjusted to give $\mathrm{NO_X}$ emissions below about 110 ppm (3 percent $\mathrm{O_2}$), CO emissions (and smoke, see Table 3-2) increase significantly. Thus, for this burner/steamer combination, minimum $\mathrm{NO_X}$ emissions at acceptable operation appear to be 110 ppm.

3.2 ENVIRONMENTAL ASSESSMENT TESTING

Following the performance/emission mapping tests discussed in Section 3.2, a set of burner operating conditions was selected for comprehensive emissions testing. The sampling protocol for these comprehensive tests included:

- o Continuous monitoring for NO_X , O_2 , CO_2 , and total unburned hydrocarbons (TUHC)
- Source assessment sampling system (SASS) for particulate size fractionation, and organic emissions
- ullet EPA Method 5/8 for particulate mass emissions, and SO₂ and SO₃ emissions
- Andersen impactor train sampling for particle size distribution determination



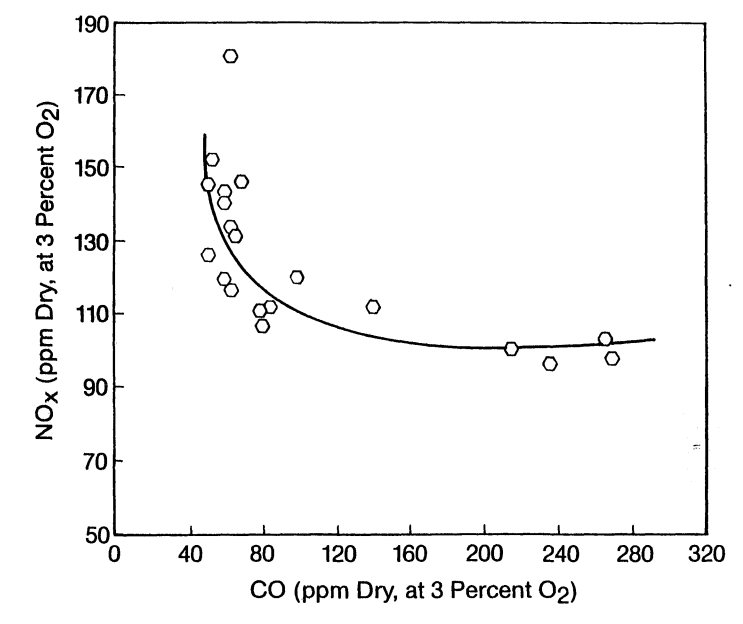


Figure 3-5. NO_{X} emissions versus CO for the MHI low NO_{X} burner.

- \circ Grab sample for onsite analysis of C_1 to C_6 hydrocarbons by gas chromatography (GC)
- Grab sample for laboratory analysis of N₂O

All flue gas sampling for these tests was performed at the steamer stack. In addition, as for other testing performed in this program, Getty Oil Company personnel performed continuous flue gas monitoring at the steamer furnace outlet.

The analysis protocol for SASS samples included:

- Analyzing SASS train samples for total organic content in two boiling point ranges: 100° to 300°C by total chromatographable organics (TCO) analysis and greater than 300°C by gravimetry (GRAV)
- Analyzing the SASS train sorbent module and particulate extracts for the 58 semivolatile organic priority pollutant species including many of the PAH compounds
- Performing infrared (IR) spectrometry analysis of the GRAV residue of organic sample extracts
- Performing direct insertion probe low resolution mass spectrometry
 (LRMS) analysis of selected sample extracts

All aspects of the sampling and analysis protocols conformed to a modified EPA Level 1 protocol (Reference 3-1). Details of the procedures used are discussed in Appendix A.

Bioassay testing of SASS samples, a normal part of the comprehensive testing performed in this project, was not done in these tests due to the limited amount of particulate sample obtained and the very low organic content of the sorbent module extract.

Results of these comprehensive tests are discussed in the following subsections. Section 3.2.1 further details the steamer operating condition during the tests performed; Section 3.2.2 presents the criteria pollutant and other gas phase species emission results; and Section 3.3.3 summarizes organic category and species emission results.

3.2.1 Burner and Steamer Operation

The burner operating conditions selected for comprehensive testing matched those noted for Test 21 in Table 3-2. This operating point was selected since it represented about a minimum NO_X condition with CO emissions below 100 ppm (see Figure 3-5). The specific steamer and burner operating conditions for these tests are summarized in Table 3-3.

The steamer efficiency noted in Table 3-3 was calculated based on the ASME heat loss method (ASME PTC 4.1). The relative contributions to the calculations are summarized in Table 3-4. As shown in this table, most of the efficiency loss was through dry gas loss and moisture loss from the fuel hydrogen. The overall efficiency noted (82.8 percent) compares favorably to the efficiency of conventional burner-equipped units.

3.2.2 Criteria Pollutant and Other Gas Phase Species Emissions

Table 3-5 summarizes the gaseous and particulate emission levels measured during the comprehensive tests. Continuous emission monitor measurements from both the stack location and the furnace outlet location (obtained by Getty Oil personnel) are noted in the table. As shown in the table, steamer stack NO_X emissions were just below 110 ppm (3 percent O_2) with CO emissions of 93 ppm (3 percent O_2), negligible TUHC emissions, and a smoke reading of 8.

TABLE 3-3. STEAMER/BURNER OPERATING CONDITIONS: COMPREHENSIVE TESTS

Fuel flow, 1/s (BPD) Heat input, MW (million Btu/hr) Feedwater flow, 1/s (BPD) Steam pressure, MPa (psig)	0.386 (210) 16.3 (55.5) 6.72 (3,650) 4.55 (660)
Air flows (percent) Diffusion Premix Overfire	36 54 10
FGR rate (percent)	9.5
Steamer efficiency (percent)	82.8

TABLE 3-4. STEAMER THERMAL EFFICIENCY

Heat loss efficiency (percent)	
Dry gas loss	6.8
Loss due to fuel moisture	
Loss due to water from the combustion of fuel hydrogen	6.3
Loss due to combustibles in the flyash	0.6
Radiation loss	2.0
Unmeasured loss	1.5
Total loss	17.2
Efficiency (percent)	82.8

TABLE 3-5. FLUE GAS EMISSIONS

	Stack ^a		Furnace ou		tlet ^b	
Pollutant	Ra	ange	Average	Rar	ige	Average
As measured: 02, percent dry C02, percent dry N0x, ppm dry N20, ppm dry C0, ppm dry TUHC, ppm dry S02, ppm dry Continuous monitor Method 8 S03, ppm Method 8 Bacharach smoke number	2.7 to 3.3 13.1 to 13.5 108 to 115 12.9 to 20.5c 45 to 135 <1dee 8		3.0 13.3 106 17.0 93 <1 c 594 3.1	2.4 to 2.7 14.4 to 14.5 110 to 112d 68 to 75d 550 to 610ddd		2.5 14.4 111 d 71 d 600 d d
	ppm	ng/J ^f	lb/million Btuf	ppm	ng/J ^f	1b/million Btuf
Corrected to 3% 0 ₂ NO _X (as NO ₂) N ₂ 0 CO TUHC (as CH ₄) SO ₂ Continuous monitor Method 8 SO ₃ (as H ₂ SO ₄) Method 8	106 17 93 <1 d 594	73.7 11 39 <0.2 d 574 4.5	0.171 0.026 0.091 <0.001 d 1.34	108 d 69 d 584 d	77.2 d 29 d 565 d	0.179 d 0.069 d 1.31 d
Particulate Method 5 SASS Andersen	mg/dscn 39 118 579	14 30 219	0.033 0.071 0.0489	d d d	d d d	d d d

^aEmission measurements by Acurex.

^bEmission measurements by Getty Oil Company.

CRange over duplicate analysis of 6 separate gas samples.

dMeasurement not performed at this location.

eExtractive sampling procedure, range not applicable.

fHeat input basis.

gAverage of two trains run.

The data in Table 3-5 show relatively good agreement between the monitor measurements at the stack and furnace outlet locations. In addition, there was good agreement between the flue gas SO_2 levels measured with a continuous monitor at the furnace outlet and by the Method 8 train run at the stack.

The Method 8 results suggest that SO₃ represents about 0.5 percent of the total sulfur oxides present in the flue gas. This ratio is significantly lower than the 5 to 10 percent range typical for residual oil-fired utility and industrial boilers (Reference 3-2). The SO₂ and SO₃ levels measured by Method 8 in the flue gas would be as expected for complete conversion of all the sulfur in a 1.2 weight percent sulfur fuel oil with heating value as noted in Table 2-1. This compares favorably to the 1.06 percent sulfur content measured in the fuel.

Particulate emissions were measured at 39 mg/dscm by Method 5, 118 mg/dscm by SASS, and 57 mg/dscm as an average of two Andersen impactor trains. These are in fair agreement. The Method 5 result is the most trustworthy, since this reference method involves a multipoint (traverse) isokinetic sampling procedure.

Particle size distribution results from the two Andersen impactor trains run are shown in Figure 3-6. Results from the two runs are similar. The mean particle diameter of emitted particulate was in the 3 to 4 µm range, for runs 1 and 2, respectively.

Emissions of nitrous oxide were also measured in these tests. The level noted, at 17 ppm, is about 16 percent of the NO_X emission level. Tests of several other fossil fuel combustion sources have shown that N_2O emissions are generally in the range of 20 percent of the NO_X emission level. These

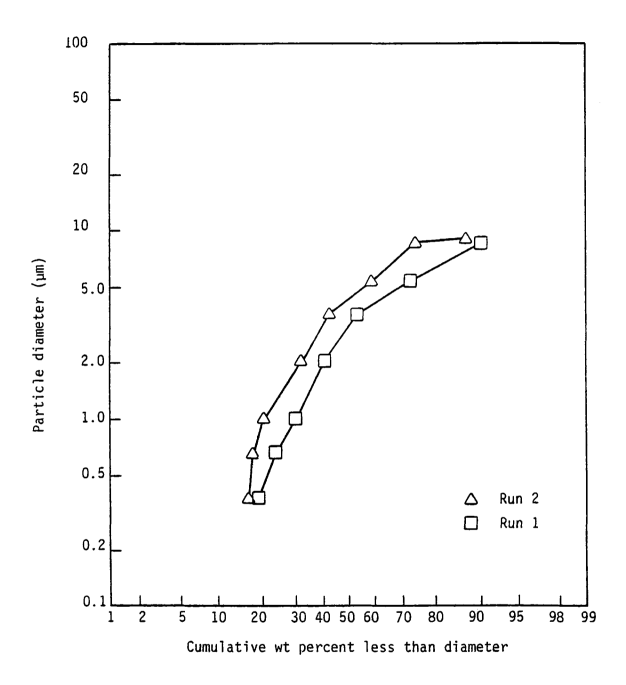


Figure 3-6. Emitted particle size distribution.

data are summarized in Figure 3-7. The point noted for this study falls on the curve corresponding to other data. The curve noted in Figure 3-7 was obtained from a least squares fit of all the data points shown in the figure, with the constraint that the curve pass through the origin. The relationship shown, $N_2O = 0.22 \ NO_X$, had a correlation coefficient (r^2) of 0.88.

3.2.3 Organic Species Emissions

Organic analyses were performed on specified flue gas samples according to EPA Level 1 protocol (Reference 3-1) as outlined in Appendix A. Volatile organics having boiling points in the C_1 to C_6 range of less than $100\,^{\circ}\text{C}$ (212°F) were determined by analysis of flue gas grab samples by onsite gas chromatography. The SASS train particulate, organic module sorbent (XAD-2), and organic module condensate (OMC) samples were extracted with methylene chloride in a Soxhlet apparatus. The extracts (XAD-2 and OMC extracts were combined) were then subjected to total chromatographable organic (TCO) and gravimetric (GRAV) analyses to determine species within the $100\,^{\circ}$ to $300\,^{\circ}$ C (212° to $572\,^{\circ}$ F), and greater than $300\,^{\circ}$ C ($572\,^{\circ}$ F) boiling point ranges, respectively. Infrared (IR) spectra of the GRAV residue of the extracts were also obtained.

The extracts were also analyzed via gas chromatography/mass spectrometry (GC/MS) for the semivolatile organic priority pollutant species (including many polynuclear aromatic hydrocarbons (PAH's)). Other major chromatogram peaks were identified and approximately quantitated.

Since the total organic contents (TCO and GRAV) of the extract samples were less than 15 mg, liquid chromatographic separations were not performed. However, low resolution mass spectrometry (LRMS) analysis of the particulate extract was performed.

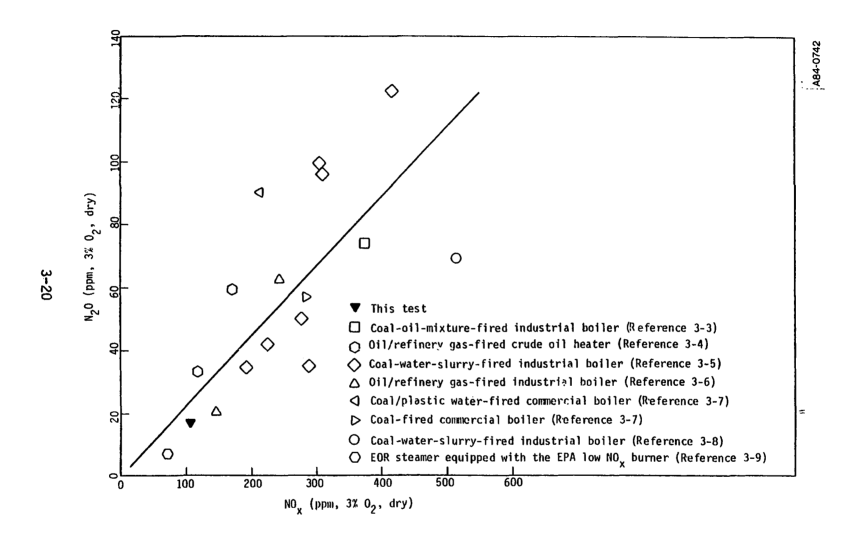


Figure 3-7. N_2O versus NO_X emissions for external combustion sources.

3.2.3.1 Total Organic Analysis

Table 3-6 summarizes measured organic emissions from the low NO_{X} burner-equipped steamer by organic boiling point range. The organic emissions are dominated by the volatile (C_1 to C_6) fraction, which is further composed primarily of compounds in the C_3 and C_4 boiling range. No semivolatile organics were detected. Nonvolatile organics (nominally C_{16+}) were found in the particulate, though not in the sorbent module. This confirms the high smoke emissions for the tests and suggests soot formation was occurring.

The C_1 , C_3 , and C_4 volatile hydrocarbon levels noted in Table 3-6 correspond to 0.3, 4.6, and 0.9 ppm, respectively, as measured. The total hydrocarbon monitor (which was unheated) read <1 ppm (see Table 3-5) for the tests. The two measurements are in fair agreement; most C_4 would not reach the total hydrocarbon monitor, and the response factor for C_3 hydrocarbon would be less than 1 ppm (as methane).

3.2.3.2 Infrared Spectra of Total Sample Extracts

The results of the IR spectrometry analysis of total sample extracts are summarized in Table 3-7. The SASS particulate spectrum suggest only the presence of aliphatic hydrocarbons in the organic fraction. The XAD-2 extract spectrum is consistent with the presence of oxygenated species such as carboxylic acids and alcohols. An interpretable spectrum for this sample was obtained despite its low organic content as noted in Table 3-6.

3.2.3.3 Low Resolution Mass Spectrometry Analysis of Total Sample Extracts

The SASS particulate extract was subjected to LRMS analysis via direct insertion probe to obtain compound category composition information. The compound categories searched for with the characteristic ions used to

TABLE 3-6. TOTAL ORGANIC EMISSIONS SUMMARY

Organic category	mg/dscm	ng/J
Volatile organics analyzed in the field by gas chromatography C1 C2 C3 C4 C5 C6 Total C1-C6	0.2 0 8.4 2.2 0 0	0.07 0 3.0 0.80 0
Semivolatile organics analyzed by TCO Filter XAD-2 Total C7-C16	<0.004 <0.004	<0.001 <0.001
Nonvolatile organics analyzed by gravimetry Filter XAD-2 Total C ₁₆₊	0.3 <0.1 0.3	0.11 <0.04 1.1
Total organics	11.1	4.0

TABLE 3-7. SUMMARY OF INFRARED SPECTRA OF TOTAL SAMPLE EXTRACTS

Sample	Wave number (cm ⁻¹)	Intensity	Possible assignment	Possible compound categories present
Particulate extract	2980-2910	Strong	C-H stretch	Aliphatic hydrocarbons
XAD-2	3500-3020 1660 1270-1130	Strong Strong Strong	O-H stretch C=O stretch C-O stretch	Oxygenated hydrocarbons such as carboxylic acids or alcohols

identify them are listed in Table 3-8. Table 3-9 notes compound categories found and their relative abundance (intensity). As noted, aliphatic hydrocarbons was the major organic category present in the sample. Minor categories present were ketones and heterocyclic nitrogen compounds. Specific compounds detected suggested that the ketones were chiefly fluoren-9-one, and the nitrogen heterocyclics were chiefly ethyl carbazole. The LRMS results suggest that fluoren-9-one and ethyl carbazole were present at levels in the 250 µg/g of particulate range. Confirmation of this by GC/MS is discussed in Section 3.2.3.4.

3.2.3.4 Gas Chromatography/Mass Spectrometry of Total Sample Extracts

Capillary column GC/MS analyses for the semivolatile organic priority pollutant species, a category which includes several polynuclear aromatic hydrocarbons (PAH's), were performed on the SASS particulate and XAD-2 extracts. The compounds sought in the analyses and their respective detection limits are listed in Table 3-10. In addition, major peaks in the chromatogram, other than these compounds, were identified and quantitated. Results of the analyses are summarized in Table 3-11.

Of the PAH's, only naphthalene, phenanthrene, and pyrene were found, and, except for the naphthalene, which was present at the highest concentration, these were found only in the particulate. The other species detected were generally oxygenated aromatics and fused aromatics. Benzoic acid was present at relatively high levels, followed by fluoren-9-one and ethyl carbazole. The levels of the fluorenone and ethyl carbazole, at 180 and 110 µg/g particulate respectively, confirm the qualitative results of the LRMS analyses discussed in Section 3.2.3.3.

TABLE 3-8. COMPOUND CLASSES AND FRAGMENT IONS SEARCHED FOR BY DIRECT INSERTION PROBE LRMS

Compound class	Fragment ions (m/e ⁻)
Polynuclear aromatic hydrocarbons	178,202,216,228,252,276
Aliphatic hydrocarbons	57,71
Halogenated aliphatics	49,63,79,81,93,95,107,109
Aromatic hydrocarbons	50,51,77,78,79,91
Ethers	45,59,73
Alcohols	45,59,61,73,75
Pheno1s	51,77,94
Nitriles	54,68,82
Phthalate esters	149,167
Amines	44,58
Ketones	51,71
N-heterocyclics	117,129,167,179
Mercaptans, sulfides	47,61,75
Benzothiophenes	57,58,59,69,70,85,97,111,125
Carboxylic acids	60,73,149
Amides	58,72,86,100

TABLE 3-9. SASS PARTICULATE EXTRACT LRMS RESULTSª

Intensity ^b	Ca tegory	MW range		
Major categories				
100 10 10 1	Aliphatic hydrocarbons Hetercyclic nitrogen compounds Ketones Polynuclear aromatic hydrocarbons	150 to 250 150 to 200 150 to 200 150 to 250		
Specific compounds				
10 10	Fluoren-9-one Ethyl carbazole	180 195		

aTotal organic content of this sample is 3.0 mg/g particulate, GRAV compounds. b100: major component; 10: minor component; 1: trace component.

TABLE 3-10. COMPOUNDS SOUGHT IN THE GC/MS ANALYSIS AND THEIR DETECTION LIMITS (ng/ μ 1 INJECTED)

Acid	i comp	pounds	
2,4,6-trichlorophenol p-chloro-m-cresol 2-chlorophenol 2,4-dichlorophenol 2,4-dimethylphenol	5 5 5 5 5 5	2-nitrophenol 4-nitrophenol 2,4-dinitrophenol 4,6-dinitro-o-cresol pentachlorophenol phenol	5 20 20 20 5 1
Base ner	ıtral	compounds	
1,2,4-trichlorobenzene 1,2-dichlorobenzene 1,2-diphenylhydrazine (as azobenzene) 1,3-dichlorobenzene 1,4-dichlorobenzene 2,4-dinitrotoluene 2,6-dinitrotoluene 2-chloronaphthalene 3,3'-dichlorobenzidine 3-methyl cholanthrene 4-bromophenyl phenyl ether 4-chlorophenyl phenyl ether 7,12-dimethyl benz(a)anthracene N-nitrosodimethylamine N-nitrosodimethylamine N-nitrosodiphenylamine acenaphthene acenaphthene acenaphythylene anthracene benzo(ghi)perylene benzo(b)fluoranthene benzo(k)fluoranthene benzo(a)anthracene benzo(a)pyrene	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	benzo(c)phenathrene bis(2-chloroethoxy)methane bis(2-chloroethyl)ether bis(2-chloroisopropyl)ether bis(2-ethylhexyl)phthalate butyl benzyl phthalate chrysene di-n-butyl phthalate di-n-octyl phthalate dibenzo(a,h)anthracene dibenzo(c,g)carbazole diethyl phthalate dimethyl phthalate dimethyl phthalate fluoranthene fluorene hexachlorobenzene hexachlorobutadiene hexachlorocyclopentadiene hexachloroethane indeno(1,2,3-cd)pyrene isophorone naphthalene nitrobenzene perylene phenanthrene pyrene	40 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1

TABLE 3-11. COMPOUNDS DETECTED IN THE GC/MS ANALYSES

Species		oarticulate ^a (µg/dscm)	XAD-2 extract ^a , ^b (μg/dscm)	Total flue gas ^c (μg/dscm)
Semivolatile organic priority pollutants				
Naphthalene Phenanthrene Phenol Pyrene	1.6 2.6 1.9 0.97	0.19 0.30 0.22 0.11	1.2 <0.04 <0.04 <0.04	1.4 0.30 0.22 0.11
Other compounds identified				
Benzofurandione Benzoic acid Benzothiazole Dichlorodibenzosulfone Ethyl benzoate Ethyl carbazole Fluoren-9-one Terphenyl	28 110 180 45	3.3 13 20 5.2	0.44 34 0.52 0.40 	0.44 34 3.3 0.52 0.40 13 20 5.2

^a27.0 dscm sampled, 3.11g particulate on filter.

^bAverage of duplicate injections.

^cSum of average of duplicate XAD-2 result plus filter result.

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- 3-4. DeRosier, R., "Environmental Assessment of a Crude-Oil Heater Using Staged Air Lances for NO_X Reduction," Acurex Report TR-82-94/EE, November 1983.
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- 3-8. VanBuren, D., and L. R. Waterland, "Environmental Assessment of a Coal-Water-Slurry-Fired Industrial Boiler," Acurex Draft Report TR-84-155/EE, March 1985.
- 3-9. Castaldini, C., et al., "Environmental Assessment of an Enhanced Oil Recovery Steam Generator Equipped with the EPA Low NO_X Burner," Acurex Draft Report TR-85-174/EE, January 1985.

SECTION 4

OUALITY ASSURANCE ACTIVITIES

Specific quality assurance (QA) activities performed to determine the accuracy and precision of the laboratory analyses performed on samples collected in this test program included:

- Spiking a sample of cleaned XAD-2 resin from the lot used in this test with TCO, GRAV, and semivolatile priority pollutant compounds and analyzing the spiked resin to determine the accuracy (recovery) of the resin extraction and subsequent analyses
- Analyzing NBS flyash for mercury to determine the accuracy of the atomic absorption technique used
- Performing duplicate TCO and GC/MS injections on the SASS train
 XAD-2 extract to determine the precision of these measurements

The following paragraphs discuss results of these QA activities.

4.1 ACCURACY DETERMINATIONS

A sample of XAD-2 resin from the cleaned lot used for these tests was spiked with 3.0 mg bis(2-ethylhexyl)phthalate, 200 μ g tetradecane, and 100 μ g each of d₈-naphthalene, phenanthrene, and pyrene. Thus, this resin contained 0.3 mg TCO compounds (tetradecane and naphthalene) 3.2 mg GRAV compounds (the phthalate, phenanthrene, and pyrene) and 100 μ g each of the three polynuclear aromatics for the semivolatile organic priority pollutant analysis.

Results of the analyses of this spiked resin are shown in Table 4-1. As noted, the recovery of the TCO analysis was 125 percent, of the GRAV analysis was 81 percent, and averaged 77 percent for the GC/MS analyses. If these are interpreted to be the accuracy of these measurements, all fall within the project accuracy objective (Reference 4-1) also noted in Table 4-1.

A sample of NBS 1633a flyash was analyzed by the cold vapor atomic absorption technique used for sample determinations in this project. The analysis result was 0.18 ppm Hg in the sample; the NBS certified value is 0.16 ppm. Thus, the accuracy of this measurement was within 11 percent, again within the QA objective of ± 20 percent for this measurement.

4.2 PRECISION DETERMINATIONS

The XAD-2 extract samples from the SASS train for this test were analyzed in duplicate for TCO content, and for the semivolatile organic priority pollutants and other major peaks by GC/MS. The two TCO measures were 0.087 and 0.094 ng/injection, giving a relative standard deviation of 5.5 percent. This is within the precision objective of this measurement of 10 percent (Reference 4-1).

Results of the duplicate GC/MS injections are summarized in Table 4-2. The relative standard deviations for all compounds quantitated were well within the project precision objective of 50 percent for this measurement.

TABLE 4-1. XAD-2 RESIN SPIKE AND RECOVERY RESULTS

Measurement	Spiked amount (mg)	Recovered amount (mg)	Percent recovery	Implied accuracy	Accuracy objective ^a
Total chromatographable organics (TCO)	0.3	0.4	125	+25	±50
Gravimetric organics (GRAV)	3.2	2.6	81	-19	±50
Semivolatile organic priority pollutants: dg-Naphthalene Phenanthrene Pyrene	0.1 0.1 0.1	0.077 0.077 0.077	77 77 77	-23 -23 -23	
Average			77	-23	-50 +100

aReference 4-1.

TABLE 4-2. DUPLICATE GC/MS ANALYSIS RESULTS FOR THE XAD-2 EXTRACT

Compound	Run 1 µg/train	Run 2 µg/train	Relative standard deviation (%)
Phenol Benzofurandione Benzoic acid Dichlorodibenzosulfone Ethyl benzoate	26	37	24.7
	10	14	23.6
	960	900	4.6
	13	15	10.1
	10	12	12.9

REFERENCE FOR SECTION 4

4-1. "Quality Assurance Plan for the Combustion Modification Environmental Assessment," Acurex Corporation for EPA Contract 68-02-2160, September 10, 1982.

SECTION 5

SUMMARY

A comprehensive emissions testing program was performed on an enhanced oil recovery steam generator (EOR steamer) equipped with an MHI PM low NO_X burner, with less detailed comparison testing performed on an identical unit equipped with a conventional North American burner.

Full load NO $_{\rm X}$ emissions from the conventional burner-equipped boiler varied from 365 ppm (corrected to 3 percent 0 $_{\rm 2}$) with stack 0 $_{\rm 2}$ of 6.6 percent to 197 ppm with stack 0 $_{\rm 2}$ of 2.3 percent. However, smoke emission levels were unacceptably high at the lower 0 $_{\rm 2}$, lower NO $_{\rm X}$ levels. A practical NO $_{\rm X}$ emission limit (acceptable CO and smoke emissions) of about 300 ppm (corrected to 3 percent 0 $_{\rm 2}$) with flue gas 0 $_{\rm 2}$ of 3.7 percent could be maintained. At 75 percent load NO $_{\rm X}$ emissions were reduced to about 250 ppm with stack 0 $_{\rm 2}$ of 4.0 percent and acceptable CO and smoke emissions.

Full load NO_X emissions from the low- NO_X burner-equipped steamer varied from 95 to 180 ppm (corrected to 3 percent O_2) with variations in the overall excess air level (as measured by stack O_2), the distribution of combustion air among the burner's premixed flame nozzle, diffusion flame nozzle, and overfire (OFA) air ports, and the rate of flue gas recirculation (FGR) to the burner to separate premixed and diffusion flames. Again, unacceptably high CO and smoke emissions existed at the lower NO_X conditions. A NO_X emission

level of about 110 ppm (3 percent 0_2) could be maintained with acceptable CO and smoke.

Comprehensive emissions testing of the low-NO $_{\rm X}$ burner-equipped boiler was performed with the burner operation at a nominal low-NO $_{\rm X}$ setting. With 54 percent of the combustion air supplied to the burner's premix flame, 36 percent to the diffusion flame, and 10 percent to the OFA ports, and with 9.5 percent FGR and 3.0 percent stack O_2 , $NO_{\rm X}$ emissions were 106 ppm, CO emissions were 93 ppm, and Bacharach smoke number was 8. At this condition SO_2 and SO_3 emissions were 594 ppm and 3.1 ppm respectively, and particulate emissions were 39 mg/dscm. The mean particle size of the particulate was in the 3 to 4 μ m range (two separate impactor train runs).

Total organic emissions were 11.1 mg/dscm, 97 percent of which was in the volatile (C_1 to C_6) boiling point range; the remainder was in the nonvolatile (C_{16}) boiling point range. The nonvolatiles were condensed on flue gas particulate and consisted largely of aliphatic hydrocarbons, heterocyclic nitrogen compounds (ethyl carbazole), and ketones (fluorenone).

Of the polynuclear aromatic hydrocarbons specifically analyzed for in flue gas emissions, only naphthalene (1.4 $\mu g/dscm$), phenanthrene (0.3 $\mu g/dscm$), and pyrene (0.11 $\mu g/dscm$) were detected. Other compounds identified as comprising the flue gas organics included benzoic acid, ethyl carbazole, and fluoren-9-one with emission levels in the 13 to 34 $\mu g/dscm$ range, and phenol, benzofurandione, benzothiazole, ethyl benzoate, and terphenyl with emission levels in the 0.1 to 5.2 $\mu g/dscm$ range.

APPENDIX A

SAMPLING AND ANALYSIS METHODS

Emission test equipment was provided primarily by Acurex Corporation. Onsite equipment included a continuous flue gas monitoring system; the source assessment sampling system (SASS) for particulate mass, semivolatile, and nonvolatile organic emissions measurement; a combined EPA Method 5 and 8 train for measuring particulate, SO₂ and SO₃ emissions; an Andersen cascade impactor train for measuring emitted particle size distribution; gas grab sampling equipment and an onsite gas chromatograph equipped with a flame ionization detector (GC/FID) for determining flue gas C₁ to C₆ hydrocarbon emissions; and gas grab sampling equipment for laboratory determination of N₂O emissions by gas chromatography using an electron capture detector (GC/ECD). All the above flue gas emission sampling was performed at the steam generator stack.

In addition, Getty Oil Company provided flue gas monitoring of O_2 , CO_2 , CO_3 , and CO_4 , and CO_5 at the steam generator furnace outlet location.

The following sections summarize the equipment sampling and analysis procedures used by Acurex in the evaluation of the steam generator/low NO_X burner.

A.1 CONTINUOUS MONITORING SYSTEM

Rack-mounted monitors and recorders located in a mobile emission laboratory were used for continuous measurement of NO_X , CO, total unburned



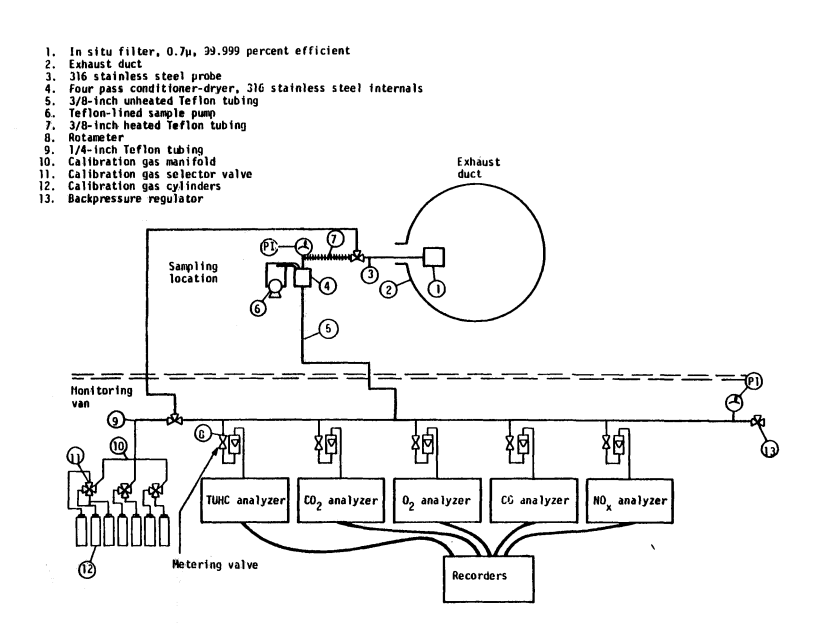


Figure A-1. Continuous monitoring system.

hydrocarbon (TUHC), CO₂, and O₂. Figure A-1 illustrates the continuous flue gas extractive sampling system and monitors arrangement. Flue gas was drawn through an in-stack filter and a heated stainless steel probe to a gas conditioning and refrigeration system designed to remove water. An unheated line was then used to bring the conditioned gas to the monitors. Calibration gases were used to monitor and correct the drift in the instruments. The calibration gases follow the same path as the flue gas being monitored in that both are conditioned at the stack prior to analysis. Table A-1 lists the instrumentation constituting the continuous monitoring and flue gas extractive sampling system used in this test program.

A.2 PARTICULATE AND SULFUR OXIDE EMISSIONS

Particulate mass emissions were measured in accordance with EPA Reference Method 5 and SO₂ and SO₃ emissions were measured in accordance with EPA Reference Method 8. A combined Method 5/8 train employing the Acurex High Volume Stack Sampler (HVSS), illustrated schematically in Figure A-2, was used in this program. A glass-lined stainless-steel probe was used to isokinetically extract the gas sample from the stack.

Particulate was removed by a heated 142 mm (5.6 in.) diameter glass fiber filter. Both the filter and the sampling probes were maintained at 120°C (250°F) as specified by Method 5.

The impinger train consisted of four glass impingers with a fritted glass filter placed between the first and second impingers as specified by Method 8. The first impinger contained 100 ml of 80 percent isopropanol (20 percent water); the second and third impingers contained 100 ml of 3 percent $\rm H_{2}O_{2}$ in water; and the fourth impinger contained 200g of silica gel.

TABLE A-1. CONTINUOUS MONITORING EQUIPMENT IN THE MOBILE LABORATORY

Instrument	Principle of operation	Manufacturer	Instrument model	Range
NO NO _X	Chemiluminescence	Thermo Electron	10 AR	0-100 ppm 0-500 ppm 0-1,000 ppm 0-5,000 ppm
со	Nondispersive infrared (NDIR)	ANARAD	500R	0-1,000 ppm
TUHC	Flame ionization detector	Beckman	400	0-10 ppm 0-100 ppm 0-1000 ppm
C0 ₂	Nondispersive infrared (NDIR)	ANARAD	AR500	0-20 percent
02	Fuel cell	Teledyne		0-5 percent 0-25 percent
Sample gas conditioner	Refrigerant dryer-condenser	Hankinson	E-4G-SS	10 scfm
Strip chart recorder	Dual pen analog	Linear	400	0-10 mV 0-100 mV 0-1V 0-10V

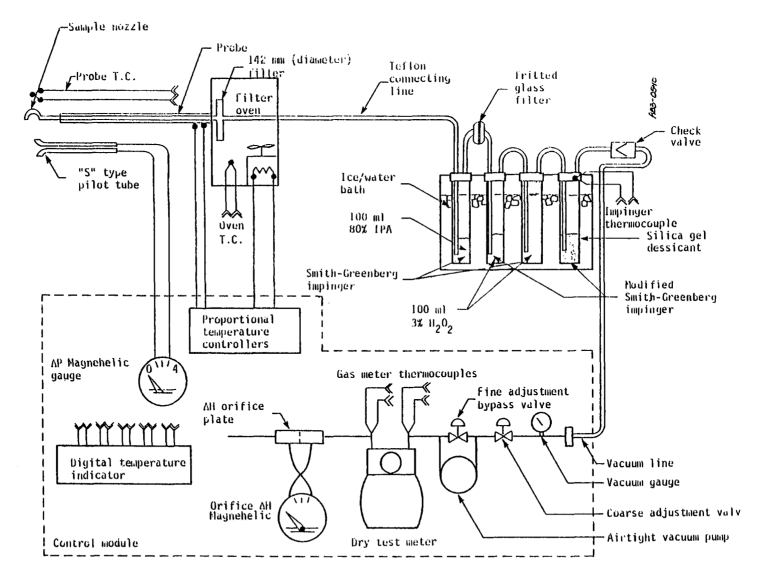


Figure A-2. Schematic of Method 5/8 sampling train.

 SO_3 (H₂SO₄ mist) is collected in the first impinger and SO_2 (oxided to SO_4) in the second and third. These were determined in the laboratory by titration with 0.01N barium perchlorate using thorin indicator.

A.3 ORGANIC EMISSIONS

Emissions of organic compounds and compound categories were sampled using the source assessment sampling system (SASS). Designed for Level 1 environmental assessment (Reference A-1), the SASS collects large quantities of gas and solid samples required for subsequent analyses of inorganic and organic emissions.

The SASS, illustrated in Figure A-3, is generally similar to the system utilized for total particulate mass emission tests (a high volume Method 5 train) with the exception of:

- Particulate cyclones heated in the oven with the filter to 230°C
 (450°F)
- The addition of a gas cooler and organic sampling module
- The addition of necessary vacuum pumps to allow a sampling rate of
 2 1/s (4 cfm)

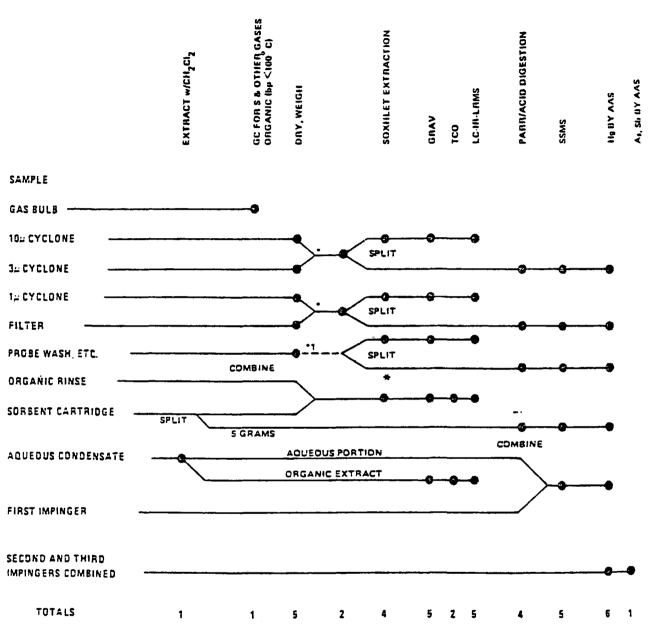
The particulate cyclones shown were not used for these tests because of the low particulate loading in the flue gas.

Schematics outlining the standard sampling and analytical procedures using the SASS equipment are presented in Figures A-4 and A-5. The inorganic analyses of SASS train samples noted in the figures were not performed for these tests.

The SASS train particulate, XAD-2 resin, and organic module condensate (OMC) were extracted with methylene chloride in a Soxhlet apparatus. The

Note: T.C. = Thermocouple

Figure A-3. Source assessment sampling system schematic.



^{*} if required, sample should be set aside for biological analysis at this point,

Figure A-4. Flue gas analysis protocol for SASS samples.

This step is required to define the total mass of particulate catch, If the sample exceeds 10% of the total cyclone and filter sample weight, proceed to analysis. If the sample is less than 10% of the catch, hold in reserve,

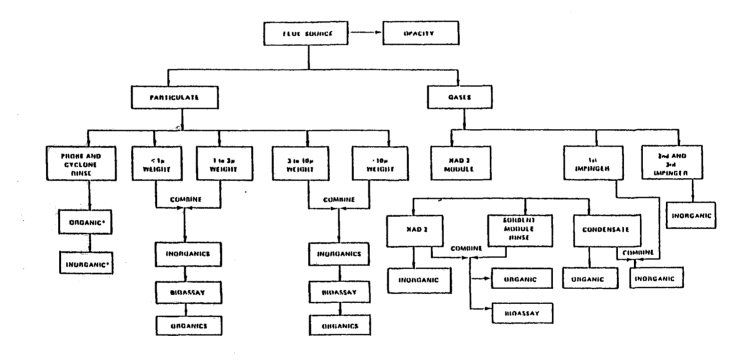


Figure A-5. Flue gas sample analysis protocol.

XAD-2 and OMC organic extracts were combined for analysis. The extracts were analyzed for total organic content in two boiling point ranges: 100° to 300° C (nominally C_7 to C_{16} organics) by GC/FID for total chromatographable organics (TCO) and >300°C (nominally > C_{16} organics) by gravimetry (GRAV). Infrared (IR) spectra were obtained of the GRAV residue of extracts. GC/mass spectrometry (MS) in accordance with EPA Method 625 for the semivolatile organic priority pollutant species was also performed on extract samples. Extract samples containing total organic content corresponding to emissions of >0.5 mg/dscm were analyzed by low resolution mass spectrometry to identify the major compound categories present. Figure A-6 illustrates the organic analysis methodology generally followed.

A.4 PARTICLE SIZE DISTRIBUTION

An Andersen 2000 Mark III in-stack cascade impactor was used to measure particle size distribution. The impactor was preheated inside the stack for 30 minutes prior to the start of sampling. Sampling was performed isokinetically at a point of average stack gas velocity.

The Mark III impactor consists of multiple stages which collect different particle sizes. Each stage consists of orifices of a specific diameter above a collection plate containing a glass fiber substrate. The orifice sizes of each stage are different and are arranged in descending order, the largest being stage 0.

For sampling, the stack gas was drawn in through the stainless-steel nozzle into the heated preseparator and impactor. The gas flowed through a stainless-steel probe and a Teflon line into the condensor train consisting of a series of 3 Lexan impingers. The gas was then pulled through a carbon

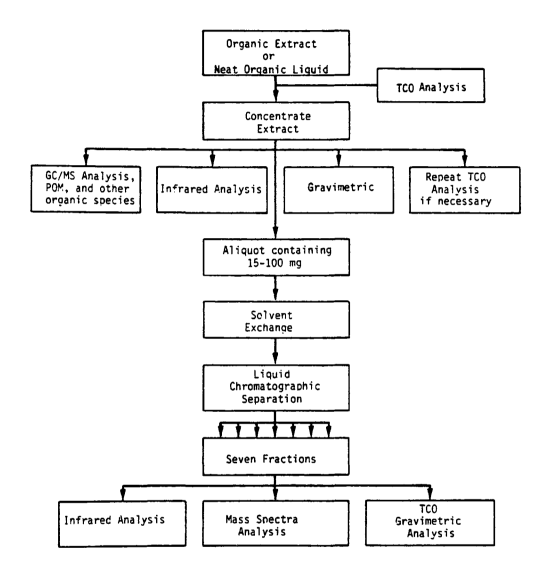


Figure A-6. Organic analysis methodology.

vane pump, dry gas meter, and calibrated orifice. The temperature of the gas leaving the impactor and the impactor temperature were measured during each test with type K thermocouples.

After a test, the Mark III impactor was carefully disassembled and the glass fiber substrates returned to their original foil containers. Any particulate matter which adhered to the impaction plates was brushed onto the appropriate filter. The samples were desiccated for 24 hours and weighed to the nearest 0.01 mg. The nozzle and Mark III inlet cone were rinsed thoroughly with acetone into a labeled amber jar. These washings were transferred to tared aluminum pans, evaporated, then desiccated for 24 hour and weighed to the nearest 0.1 mg.

A.5 C1 TO C6 HYDROCARBON SAMPLING AND ANALYSIS

Samples of flue gas for C_1 to C_6 hydrocarbon analysis were collected using a grab sampling procedure employing the apparatus illustrated in Figure A-7. The equipment consisted of a heated, 0.64-cm (1/4-in.) OD pyrex-lined, stainless-steel probe fitted with a 7- μ m sintered stainless-steel filter at the probe inlet. The outlet of the probe was directly attached to a diaphragm vacuum pump which was in turn attached to a 500-ml stainless-steel heated sampling cylinder. The sampling cylinder was insulated with heat tape powered by a varying voltage controller. The heating jacket kept the sample gas above the dew point to minimize sample loss due to water condensation.

Prior to sampling, the gas cylinder was purged with stack gas for 3 minutes and then sealed. The trapped flue gas was then analyzed onsite with a Varian Model 3700 gas chromatograph (GC) equipped with a flame ionization detector.

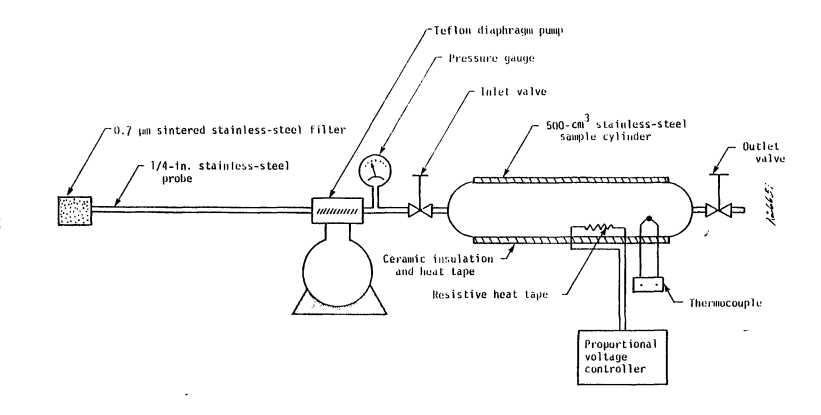


Figure A-7. C_1 to C_6 hydrocarbon sampling system.

Table A-2 lists the design specifications of the Varian GC. A 1.85m (6 ft) long, 0.32 cm (1/8 in.) diameter stainless-steel column packed with Poropak Q 60/80 mesh was used to separate the hydrocarbons into their respective components (C_1 to C_6). The GC was calibrated with repeated injections of a Scott Speciality standard gas containing C_1 to C_6 hydrocarbons (each having a concentration of 15 ppm). The chromatographic responses for the standards and the samples were recorded on a Hewlett Packard Model 3390A reporting integrator.

A.6 NoO EMISSIONS

Stack gas grab samples were extracted into stainless-steel cylinders similar to those used for C_1 to C_6 hydrocarbon sampling (Section A.5) for laboratory analysis for N_2O . For the analysis each sample cylinder was externally heated to $120\,^{\circ}\text{C}$ ($250\,^{\circ}\text{F}$); then a 1-ml sample was withdrawn with a gas-tight syringe for injection into a gas chromatograph. The analytical equipment consisted of a Varian 3700 gas chromatograph equipped with a ^{63}Ni electron capture detector and a 3.65m (12 ft) stainless-steel column packed with Poropak Super Q, 80/100 mesh. The injector temperature was kept at ^{30}C , the detector at 350°C, and the column temperature at 33°C. Elution time for N_2O was approximately 5 minutes, with a flowrate of 20 ml/min of nitrogen.

TABLE A-2. GAS CHROMATOGRAPH SPECIFICATIONS

Varian Model 3700	Gas	Chroma tograph:		
Sensitivity		1 x 10 ⁻¹² and range	A/mV at attenuation 10 ⁻¹² A/mV	1

 -10^{-11} to 10^{-9} A (reversible with internal switch) Zero range

 5×10^{-15} A; 0.5 μ V peak to peak Noise (inputs capped)

220 ms on all ranges (approximate 1 sec response to 99 percent Time constant

of peak)

Carrier gas (helium), combustion air, fuel gas (hydrogen) Gas required

TECHNICAL REPORT DATA (Please read Instructions on the reverse before completing)				
3. RECIPIENT'S ACCESSION NO.				
5. REPORT DATE February 1986				
6. PERFORMING ORGANIZATION CODE				
8. PERFORMING ORGANIZATION REPORT NO.				
TR-84-161/EE				
10. PROGRAM ELEMENT NO.				
11. CONTRACT/GRANT NO.				
68-02-3188				
13. TYPE OF REPORT AND PERIOD COVERED Final; 1/84 - 1/85				
14. SPONSORING AGENCY CODE				
EPA/600/13				

15. SUPPLEMENTARY NOTES AEERL project officer is Robert E. Hall, Mail Drop 65, 919/541-2477. Volume II is a data supplement.

16. ABSTRACT The report discusses results from sampling flue gas from an enhanced oil recovery steam generator (EOR steamer) equipped with an MHI PM low-NOx burner. The tests included burner performance/emission mapping tests, comparative testing of an identical steamer steamer equipped with a conventional burner, and comprehensive testing of the low-NOx-burner-equipped steamer. Comprehensive test measurements included continuous flue gas monitoring; source assessment sampling system testing with subsequent laboratory analysis to give total flue gas organics in two boiling point ranges and specific quantitation on the semivolatile organic priority pollutants; Cl to C6 hydrocarbon sampling; Methods 5/8 sampling for particulate and SO2 and SO3 emissions; and emitted particle size distribution tests using Andersen impactors. Full-load NOx emissions of 110 ppm (3% O2) could be maintained from the low-NCx burner at acceptable CO and smoke emissions, compared to about 300 ppm (3% O2) from the conventional-burner-equipped steamer. At this low-NOx condition, CO, SO2, and SO3 emissions were 93, 594, and 3.1 ppm, respectively. Particulate emissions were 39 mg/dscm with a mean particle diameter of 3 to 4 micrometers. Total organic emissions were 11.1 mg/dscm, almost exclusively volatile (C1 to C6) organics. Three PAHs were detected at from 0.1 to 1.4 micrograms/dscm.

17. KEY WORDS AND DOCUMENT ANALYSIS				
a. DESCRIPTORS	b. IDENTIFIERS/OPEN ENDED TERMS	c. COSATI Field/Group		
Pollution	Pollution Control	13B		
Boilers Oil Burners	Stationary Sources Low-NOx Burners	13A		
Crude Oil Oil Recovery	Environmental Assess- ment	11H,08G 081		
Assessments		14B		
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