EMISSION MEASUREMENTS OF A J93 TURBOJET ENGINE

D. L. Davidson and A. F. Domal
ARO, Inc.

September 1973

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FOREWORD

The test program reported herein was conducted at the Arnold Engineering Development Center (AEDC), under sponsorship of the Department of Transportation (DOT), Office of the Secretary, under Interagency Agreement DOT-AS-20024, Program Element 921K. The test engine and engine support equipment were supplied by the Air Force Aero-Propulsion Laboratory. The program was accomplished with project management liaison by Mr. A. K. Forney of the DOT Federal Aviation Administration and technical liaison by Mr. A. J. Broderick and Mr. M. J. Scotto of the DOT Transportation Systems Center.

The test results were obtained by ARO, Inc. (a subsidiary of Sverdrup & Parcel and Associates, Inc.), contract operator of the AEDC, Air Force Systems Command, Arnold Air Force Station, Tennessee. The test was conducted in Propulsion Development Test Cell (J-2) of the Engine Test Facility under ARO Project RA109 between June 8 and July 31, 1972. The manuscript was submitted for publication on April 18, 1973.

The authors wish to express their appreciation to the General Electric Company (GE) for the use of essential test equipment during the test program, to Mr. R. S. Sowell of GE at Evendale, Ohio, for providing technical information and advice concerning operation of the test engine, and to Pratt & Whitney Aircraft at East Hartford, Connecticut, for providing information used in test systems design for the program. The authors also wish to express their appreciation to W. K. McGregor, J. D. Few, and B. L. Sieber of ARO, Inc., who obtained the spectroscopic data which are discussed in detail in Appendix III.

This report will also be published by the Department of Transportation as FAA-RD-73-66.

This technical report has been reviewed and is approved.

CHAUNCEY D. SMITH, JR.
Lt Colonel, USAF
Chief Air Force Test Director, ETF
Directorate of Test

FRANK J. PASSARELLO
Colonel, USAF
Director of Test
ABSTRACT

Exhaust gas emission measurements were made at the nozzle exit of a J93 turbojet engine at simulated flight conditions from sea-level static to Mach 2.0 at 75,000 ft and Mach 2.6 at 65,000 ft. Real time measurements of CO, CO$_2$, C$_x$H$_y$, NO, and NO$_x$ were taken over a range of afterburning and nonafterburning engine power settings using a gas sampling system designed to adhere to SAE ARP 1256 specifications. In addition, NO and OH were measured in situ by a narrow-line UV spectral absorption technique. Batch-type measurements of particulates and other trace constituents of the exhaust gas were also made. Major results of the test were that emissions vary significantly with combustor inlet pressure and temperature and, therefore, with Mach number and altitude. Specifically, NO$_x$ increased with the square root of the combustor inlet pressure and the 2.5 power of the combustor inlet temperature. Differences between the gas sampling and spectroscopic data on NO indicate the need for verification of sampling techniques.
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NOMENCLATURE

A/B Afterburning

A_x Emission index conversion constant for specie X

f/a Fuel to air mass ratio

h Altitude, ft

M Mach number

P Pressure, psia

r Jet radius, in.

R Physical radius, in.

T Temperature, °R

W_x Emission index of specie X, lbm per lbm of fuel

w Mass flow, lbm/sec

(X) Concentration of specie X, ppm or percent

a Molecular ratio of hydrogen to carbon in fuel
\( \phi \)  
Equivalence ratio or ratio of engine overall f/a to stoichiometric f/a

\( \phi_m \)  
Local equivalence ratio calculated from measured gas composition

**SUBSCRIPTS**

\( a \)  
Air

\( f \)  
Fuel

\( s \)  
Static

\( n \)  
Nozzle

\( t \)  
Total

\( 3,5,7,9 \)  
Engine stations (see Fig. 3b)
SECTION I
INTRODUCTION

1.1 BACKGROUND

According to the December 1968 report from the Secretary of Health, Education, and Welfare to the Congress, "Nature and Control of Aircraft Engine Exhaust Emissions," pursuant to PL 90-148, The Air Quality Act of 1967, commercial jet aircraft contribute only about 1 percent of the greater than 200 million tons of pollution dumped into the atmosphere over the United States each year. However, atmospheric pollution by the exhaust gases of turbojet engines is a problem because of the localized nature of their emission. Most of the engine-on time of a turbojet-powered aircraft is spent either at the airport during takeoff, climbout, approach, landing, and taxiing and at high altitudes during cruise. Therefore, two problem areas can be defined for potential turbojet pollution of the atmosphere. The first, the airport problem, has been the subject of several studies and will not be discussed in this report. The second, the high-altitude problem, is potentially more severe because of possible long-term residual effects in the stratosphere. Turbojet emission into the atmosphere at altitudes up to approximately 35,000 ft have a short mean residence time because of the motion induced by vertical convection in this region. However, in the stratosphere, a region from approximately 35,000 to 165,000 ft generally characterized by a positive temperature gradient, vertical motion is greatly reduced and no strong mechanism for removal of any emission in this region exists. As a result, there has been much speculation about the potential meteorological problems associated with aircraft operation in the stratosphere (Fig. 1, Appendix I). The need for evaluating this problem has been emphasized by the introduction of supersonic transports (SST's), which will operate almost exclusively in the stratosphere.

The Department of Transportation is aware of this problem and the need for its evaluation and, in 1971, instigated the Climatic Impact Assessment Program (CIAP). The objective of this program is to assess, by 1974, the impact of the environmental and meteorological changes due to the world wide high-altitude aircraft fleet as projected to 1990 (Ref. 1). The world aircraft fleet includes not only future SST's but also the subsonic jets that presently cruise in the low stratosphere and any other vehicle that would emit potential pollutants into the stratosphere.

The CIAP program is divided into the elements listed below:

1. Nature of stratosphere
2. Nature of propulsion effluents
3. Perturbed stratosphere
4. Perturbed troposphere
5. Physio-bio-botanical effects
In 1971, the Department of Transportation contracted, under an interagency funding agreement, with the Arnold Engineering Development Center (AEDC), Air Force Systems Command, to contribute, with two separate studies, to the second element of the CIAP program. The first study was the simulated altitude testing of a J93 turbojet engine to evaluate the generation of effluents during stratospheric flight and is the subject of this report. The second study was the testing of a J85 turbojet engine in the AEDC’s Propulsion Wind Tunnel to evaluate both the generation and the near-field reactions of engine effluents during supersonic stratospheric flight and will be the subject of a subsequent report.

1.2 SCOPE OF INVESTIGATION

The J93 test program was conducted during June and July 1972 in Propulsion Engine Development Test Cell (J-2) of the AEDC to determine the nature of effluents in the exhaust of an SST-type turbojet engine over a wide range of typical SST operating conditions from takeoff to high-altitude supersonic cruise. The engine was tested in a direct-connect installation, which allowed simulation of engine inlet pressures and temperatures and free-stream static pressures that are encountered in actual flight. Simulated flight conditions at which emission data were obtained varied from sea-level static to Mach 2.0 at 75,000 ft and Mach 2.6 at 65,000 ft and included both afterburning and nonafterburning engine power settings (Fig. 2). Emission measurements were taken at the nozzle exit using a gas sampling system designed to adhere to the Society of Automotive Engineer’s Aerospace Recommended Practice 1256 (Ref. 2). Real-time measurements of CO, CO$_2$, C$_x$H$_y$, NO, and NO$_x$ were made. In addition, NO and OH were measured in situ by a narrow-line UV spectral absorption technique. Batch-type measurements of particulates and other trace constituents of the exhaust gas were also made.

During the test program, some 169 data points were obtained at 8 flight conditions and are presented herein. To obtain these data, the engine was operated for a total of 65 hr including 12 hr in afterburning (Table I, Appendix II).

SECTION II
APPARATUS

2.1 J93 ENGINE

The General Electric YJ93-GE-3 engine (Refs. 3 and 4) was designed to power the XB-70, a Mach 3 bomber of the early 1960’s. Although it does not represent current technology in all respects (primarily maximum cycle temperature), it was an available engine which would operate over the wide range of altitude and Mach number for which data were desired and is quite similar, except for size, to the General Electric GE4/JP5 engine.
that was under development for the U.S. SST. It is representative of much of the engine technology that is flying today.

The engine (Fig. 3) is a single-spool turbojet engine with afterburner, having a maximum sea-level static thrust rating of 28,000 lb, maximum flight speed of Mach 3.2, and maximum altitude capability of 95,000 ft. The engine consists of the following major components: an 11-stage compressor section, a combustion section, a 2-stage turbine section, a high-augmentation afterburner section, a variable-area-ejector nozzle, and a control and accessory pod.

The engine compressor is an 11-stage unit having an 8.75:1 design pressure ratio at sea-level static. Engine airflow is 261 lb/sec at a 100-percent speed of 6825 rpm. The main combustor is a fully annular type, with 32 fuel nozzles and a dual energy ignition system; one for normal ground starts and the other for air starts. The turbine is a 2-stage, air-cooled unit with a conical drive shaft, supported both front and rear by bearings. The exhaust section consists of an afterburner and a guided expansion converging-diverging exhaust nozzle of fully variable area. The primary nozzle operates independently from the secondary nozzle. The fully throttleable afterburner (Fig. 3d) incorporates 32 radially oriented fuel nozzles and a circumferential flameholder containing 3 V-gutter rings and 6 catalytic igniters. Each of the fuel nozzles is supplied by two separate fuel supply systems. The local system is actuated first and is designed to promote smooth afterburner ignition; the fill system is actuated after the local system and is designed to provide as uniform afterburner fuel distribution as possible. The use of two systems also allows better control of fuel nozzle pressure for spray atomization than that possible with a single system.

Secondary airflow is directed aft between the engine shell and the air vehicle shroud (Fig. 3b), providing a layer of cooling air around the engine. The primary function of this air is to cool the engine compartment and the jet nozzle components. Flow equivalent to 7 percent of the engine primary airflow is required at maximum afterburner operation. At certain nonstandard operating conditions, the secondary airflow may exceed the 7 percent required for cooling. To provide aircraft inlet duct matching, the flow may reach as high as 20 percent of the primary flow.

The engine control system is hydro-mechanical, with an electrical temperature override. Engine controls and accessories are enclosed in an insulated pod below the compressor for environmental, assembly, and maintenance purposes.

2.2 TEST CELL

The test program was conducted in Propulsion Development Test Cell (J-2) of the AEDC's Engine Test Facility. The cell is 20 ft in diameter and 55 ft long. In it, propulsion devices may be tested over a wide range of simulated flight altitudes and velocities. The J93 was installed in the cell in a direct-connect type of installation (Fig. 4). This type of installation allows precise control of both the engine inlet total pressure and temperature associated with the flight altitude, Mach number, and inlet recovery and
the free-stream static pressure associated with the flight altitude being simulated. In this manner, the internal combustion and aerodynamic performance of the engine during flight, and therefore the exhaust gas composition at the nozzle exit, are reproduced in the test cell.

2.3 MEASUREMENTS

Instrumentation was provided to measure both the composition of the J93 exhaust gas and pertinent parameters controlling both the operation and the operational environments of the J93 engine. Gas composition measurements for engine emissions included both real-time sampling using process gas type instruments and in-situ absorption spectroscopy. Batch-type samples of engine exhaust gas and fuel were also taken for subsequent laboratory analysis.

2.3.1 Emission Measurements

The Society of Automotive Engineers E-31 Committee has established guidelines for the design of sample systems and testing approaches to the acquisition of emissions data on turbojet engines. These are contained in Aerospace Recommended Practice (ARP) 1256 (Ref. 2). Although this standard does not address some of the problems associated with the acquisition of data behind an afterburning engine, or from an engine in an altitude test cell, it was used for guidance in the design of the AEDC emission measurements system (Fig. 5).

Continuous flow samples of the exhaust were taken in a plane approximately 4 in. downstream of the nozzle exit using a single, movable probe. The sample probe was also used for pitot pressure measurements by closing the system sample valve (Fig. 5). A shielded iridium/iridium-rhodium thermocouple was used to determine exhaust gas temperature. The probe and thermocouple were mounted on a single vertical rake (Fig. 6). Remotely controlled rake actuators (Fig. 7) provided the capability to place the sample probe or the thermocouple mounted 2 in. above it at any point in the sample plane.

The gas was transported from the probe to an exhaust gas analyzer located outside the test cell approximately 60 ft from the probe tip. The gas was cooled to 300°F from total temperatures as high as 3100°F and maintained at that temperature in the gas transport line to prevent \( C_xH_y \) and \( H_2O \) condensation by a hot water thermal conditioning system. This system counterflowed in adjacent or coaxial lines the entire length of the sample transport line from the gas analyzer to the sample probe tip. Additional heating to overcome heat loss in certain areas was provided by electric heater tape. Transport time of the gas to the exhaust analyzer was 2.1 sec. All surfaces in the gas transport line that could contact the sample gas prior to analysis were made of either Teflon\textsuperscript{®} or stainless steel (primarily type 316).

The pressure of the exhaust flow at the sample probe tip varied from 3.5 to 37 psia. As the gas had to be delivered to the gas analyzer at 15 psia for proper operation
of the sensing instruments, four metal bellows pumps were arranged just upstream of the analyzer so that they could be operated in a variety of series and/or parallel configurations to boost the sample gas pressure as needed; control valves were installed to dump excess gas when the pressure was too high. With one exception, the portion of the sample gas to be analyzed for particulates bypasses these pumps to prevent possible particle trapping or agglomeration. The exception was the piezoelectric particulate density sensor which, like the gas sensing instruments, required essentially atmospheric pressure for proper operation.

The exhaust gas analyzer (Fig. 8) was designed to permit continuous sampling of 8 gases plus particulate density. This sampling required 13 different instruments (Table II). The analyzer had additional provisions both for taking bag and bottle samples of the gas and capturing exhaust gas particulates (Table IIb). One particulate capture method was the electrostatic capture of particulates on electron microscope grids using a device furnished by the DOT Transportation Systems Center (TSC). The other particulate capture method used standard, pre-weighed, gravimetric capture filters in conjunction with a positive displacement wet test meter to determine particulate density of the exhaust gas.

The analyzer's instruments were calibrated in place using certified calibration gases. A zero gas (dry nitrogen with zero concentrations of emission gases) and a minimum of two different span gases were used in the calibrations. Temperature conditioning was provided within the analyzer to prevent sample gas condensation; standard pressure and temperature instrumentation were used to ensure proper operation of both the analyzer and the rest of the emission measurement system. Ancillary measurements (Table IIc) of both exhaust gas state conditions and fuel properties were also made to afford a complete analysis of the exhaust gas composition. A more complete description of the emission measurement system and the performance of the system and its components is found in Ref. 5.

2.3.2 Spectroscopy

A new and promising technique for in-situ measurement of exhaust gas composition was used to measure both NO and OH in the J-93 exhaust. This technique consists of passing a light beam through the exhaust and measuring the optical absorptivity of the gas at specific optical wavelengths. The technique is very attractive because it avoids the flow disturbances of a sample rake and line. Additionally, it permits, by the proper optical wavelength selections, the measurement of a wide variety of exhaust gas constituents. Details of the operation and performance of this technique are presented in Appendix III.

2.3.3 Engine

Standard measurement techniques were used to determine engine and test cell pressures, temperatures, flows, speed, and vibrations to permit correlation with emission measurements and to allow safe and efficient conduct of the test program.
SECTION III
PROCEDURE

3.1 TEST

The instruments were calibrated in place by the use of standard zero and span gases before and after each test, and, additionally, every 2 to 3 hr during tests which were of four hours duration or longer. After calibration, the engine was started, and test conditions were set. The rake was moved into position, the gas pumps and pressure control valves were adjusted to establish the proper flow through the instruments, and the test cell, engine, and gas sensor instrumentation output levels were recorded by a digital data acquisition system. An analog display from each gas analysis instrument was monitored to ensure that the output levels were stabilized prior to recording the data point. After the data points were acquired, the probe was then moved to a new location in the stream, and the acquisition procedure was repeated. At each point, engine and test cell data were also recorded. Data were also acquired through a probe at the front of the engine during each test to ensure that no significant particulate, gaseous, or liquid contaminants were present in the air entering the engine.

ARP 1256 requires at least 12 samples be taken in the plane being surveyed. In the initial plans for the program, it was decided that 13 measurements would be made, one on the centerline and three on the centroid of equal areas in each of the four quadrants. After several tests it was determined that, within the objectives of this program, more useful data would be obtained with fewer points in a shorter time by obtaining more than three points along a single radius to better define the profiles.

At some times, it was desired to obtain data where something other than flight conditions was the variable. To permit rapid acquisition of data from such a parametric investigation, a single measurement was made by leaving the probe at a fixed point and varying the parameter under investigation.

Electrostatic particulate grids were obtained at each probe location. Single gravimetric filters, bag C_x H_y samples, and wet chemistry samples were obtained at each test condition because of the gas volume required for sufficient analysis resolution. At selected test conditions, sample bottles were filled with exhaust gas for subsequent analysis at the Bureau of Mines (BuMine) and the University of California at Los Angeles (UCLA). After test, the grids were shipped to the DOT Transportation System Center (TSC) for analysis using both scanning and transmission electron microscopes. The gravimetric, bag, and wet chemistry samples were analyzed at the AEDC chemical laboratory. Fuel samples were taken prior to each test and analyzed by the AEDC laboratory except for hydrogen/carbon ratio. This last analysis was accomplished by the Galbraith Laboratory in Knoxville, Tennessee.

Engine inlet pressure and temperature settings during the test program were determined from the geopotential altitude values from the U.S. 1962 Standard Atmosphere
and standard aerodynamic relationships. Engine inlet pressure at supersonic flight conditions was reduced to account for inlet recovery based on MIL-E5008C. JP-5 fuel which is quite similar to Jet A-1 was used throughout the test program.

After completion of the test program, a verification of the emission measurement system was made using a commercial calibration gas cross-reference service.

3.2 DATA REDUCTION

All digital data from test cell, engine, and gas sensor instrumentation recorded during a test were reduced to engineering units by a digital computer. All emission indices and the fuel/air ratio calculated from the gas composition were determined by the methods detailed in ARP 1256 (Ref. 2). These methods are based on a carbon mass balance of a $C_xH_{2x}$-air reaction and use the following general equations:

$$
\frac{W_X}{A_X(X)} = \frac{A_X(X)}{CO + CO_2 + \frac{C_xH_{2x}}{10^4}}
$$

and

$$
\frac{f/a}{10^4} = \frac{CO + CO_2 + \frac{C_xH_{2x}}{10^4}}{207 - 2\left(\frac{CO}{10^4}\right) - CO_2}
$$

where

$W_X$ = Mass emission rate of $X$, lbm'/1000 lbm of fuel  
$A_X$ = Constant, function of molecular weight of $X$  
$(X)$ = Concentration of $X$, ppm  
$CO, CO_2, C_xH_{2x}$ = Concentrations in ppm, percent, and ppm, respectively,  
$f/a$ = Fuel to air mass ratio

All exhaust components with the exception of NO were measured on a wet basis. The sample stream for NO measurement was dried prior to analysis by a column of indicating calcium sulfate. The NO measurement was converted to a wet basis by accounting for the absorbed water by the method detailed in ARP 1256.

All other calculated parameters were determined in accordance with accepted aero-thermodynamic and turbojet cycle calculation procedures. Engine net thrust and other predicted parameters were determined from J93 cycle performance tables (Ref. 6) provided by the General Electric Company. Batch analyses were conducted using accepted standard laboratory techniques.
SECTION IV
RESULTS AND DISCUSSION

Exhaust emission data were obtained at the nozzle exit of a J93 turbojet engine at 8 simulated flight conditions from takeoff to high altitude supersonic cruise (Fig. 2). Parametric variables investigated include flight Mach number, flight altitude, and engine power setting. Although thirteen continuous gas sensing instruments (Table II) were used to analyze the engine exhaust gas, some of these instruments provided unusable or only marginally usable data (Ref. 5). As a result, only the continuous sampling data from the instruments shown below for each gaseous constituent will be presented in this report.

- CO – Beckman 315B
- CO\textsubscript{2} – Beckman 315B
- C\textsubscript{x}H\textsubscript{y} – Beckman 402
- NO – Thermo-Electron 10A
- NO\textsubscript{x} – Thermo-Electron 10A

The piezoelectric particulate instrument provided no usable continuous data during the test. No significant concentration of any gaseous or particulate emission constituent was found in the engine inlet air during the test program.

4.1 EXHAUST JET PROFILE

An examination of the exhaust gas pressure and total temperature profiles (Fig. 9) at military or maximum nonafterburning power indicates that gas flow and gas state properties are fairly uniform across the nozzle exit until, at a radius ratio of approximately 0.8, mixing of primary nozzle flow with peripheral secondary air begins. This conclusion is substantiated by the fuel/air ratio, CO, C\textsubscript{x}H\textsubscript{y}, and NO\textsubscript{x} profiles. Both the fuel/air ratio and NO\textsubscript{x} profiles show the slightly increasing radial influence of exhaust gas temperature; the CO and C\textsubscript{x}H\textsubscript{y} data are flatter indicating little or no localized influence of gas temperature. Agreement of measured gas temperatures at the turbine discharge and at the nozzle exit is good. Agreement of measured and calculated fuel/air ratio is also good.

Exhaust gas profiles in afterburning (Fig. 10) are not as uniform because of the zone burning characteristic of the J93 afterburner and the lack of combustion time and mixing length between the afterburner flame holders and the nozzle exit. The pressure profile in afterburning, as expected, remains essentially flat, whereas the gas temperature has a strong radially increasing profile prior to mixing with the secondary air. Temperature at the centerline of the nozzle exit flow is only approximately 300°F higher than turbine exit temperature indicating a zone of poor fuel distribution and/or poor combustion about the centerline. This conclusion is substantiated by the fuel/air ratio, CO, and C\textsubscript{x}H\textsubscript{y} data with the fuel/air ratio being low and the CO and C\textsubscript{x}H\textsubscript{y} levels being very high near the centerline. The NO\textsubscript{x} profile shows the same dependence on exhaust gas temperature present in the military power profiles and, as a result, is low near the centerline.
An understanding of this strong radial profile in afterburning is related to the geometry of the afterburner fuel distribution. The turbine discharge flow of the J93, like most turbojets, is strongly annular in nature. To ensure adequate local fuel/air ratios in the afterburner combustion zone for proper ignition and continued burning, the fuel distribution is, as shown in Fig. 3d, also strongly annular. This arrangement results in a centerline zone behind the turbine cone in which local fuel/air ratios are dependent primarily on the limited radially inward mixing in the flow aft of the flameholder.

The strong radial exhaust gas profiles present in afterburning create a problem in comparative parametric analysis of the emission data. The military data are uniform enough to obtain, with use of the exhaust gas pressure and temperature data, an average emission index mass-weighted over the nozzle exit. However, in afterburning, a combination of the strong radial profile and a limited number of samples makes determining a meaningful mass-weighted average most difficult. Also, the suggestion of a significant circumferential profile near the center indicated in the data from four quadrants near a radius ratio of 0.4, together with the strong possibility that combustion near the centerline is not complete at the nozzle exit, further complicates this problem. Therefore, for the purpose of this report, data for the parametric comparisons that follow were taken from actual profile data points where the nearest calculated-to-measured fuel/air ratio match was obtained. This selection criterion generally meant that the data at radius ratios of 0.4 to 0.7 were chosen for the comparisons.

The data presented in Figs. 9 and 10 were from only one flight condition; however, these data are typical of the profiles from the remaining seven flight conditions.

4.2 CONTINUOUS EMISSION DATA

The emission data selected for parametric comparison by the method outlined above were converted to emission indices and equivalence ratios ($\phi$) and tabulated (Table III). Data were obtained at Mach numbers from 0 to 2.6, at altitudes from 0 to 75,000 ft, and at power settings from idle ($\phi = 0.12$) to military ($\phi = 0.22$ to 0.33) and to afterburning ($\phi = 0.4$ to 0.7). A comparison of these data (Fig. 11) shows that the concentrations of CO, $C_xH_y$, and NO$_x$ all vary significantly with all three external variables. For example, CO and $C_xH_y$ at military power vary about an order of magnitude, and NO$_x$ varies about 300 percent over the range of flight conditions investigated. Similar variances are present in the afterburning data. NO/NO$_x$ ratios vary approximately 20 percent except for the lower ratios present at sub-military power and near-minimum afterburning power settings. Average NO/NO$_x$ index ratio excluding these low values was approximately 0.9.

Other agencies have obtained data having similar or worse scatter (Ref. 7) and have concluded that there was no effect of either simulated flight altitude or Mach number on engine emissions. However, an understanding of the seemingly wide scatter of emission data is linked to the internal parameters that control the combustion processes inside a turbojet engine. Some of these parameters are combustion zone pressure, combustion zone inlet temperature, and overall equivalence ratio. Combustion pressure and temperature
are controlled both by flight conditions and engine power setting; combustion zone equivalence ratio is controlled by the engine power setting and, to a lesser extent, by flight velocity.

An afterburning turbojet such as the J93 has two combustion zones: the main combustor and the afterburner. Conditions at the inlet to the main combustor of the J93 varied from 670 to 1470°R and from 19 to 120 psia (Fig. 12); overall equivalence ratio in the main combustor varied from 0.12 to 0.33. Conditions at the inlet to the afterburner varied from 1920 to 2000°R and from 11 to 43 psia (Fig. 13); overall afterburner equivalence ratio varied from 0.45 to 0.71. It should be noted that, at military power (Fig. 12), emission data were obtained over a range of combustion zone pressures at constant inlet temperatures of approximately 1050 and 1220°R. Also, emission data were obtained at an essentially constant combustion zone pressure of 55 psia over a range of inlet temperature from 950 to 1460°R. These constant temperature and pressure lines made the analyses in Sections 4.2.2 and 4.2.3 of this report possible.

4.2.1 Effect of Engine Power Setting

Data obtained over a range of engine power settings at three flight conditions yielded consistently similar trends in emission indices versus engine power setting (Fig. 14). In nonafterburning, increasing power from idle (\( \phi = 0.12 \)) to cruise (\( \phi = 0.23 \)) and to military (\( \phi = 0.3 \)) decreased both the CO and \( \text{C}_x\text{H}_y \) emission indices and increased the NO\(_x\) index. Going from military to minimum afterburning increased CO and \( \text{C}_x\text{H}_y \) and decreased NO\(_x\). Increasing afterburning power decreased the CO and \( \text{C}_x\text{H}_y \) while the NO\(_x\) remained essentially constant. The increase in CO and \( \text{C}_x\text{H}_y \) going into afterburning can be explained by the fact that afterburning efficiencies, particularly at minimum afterburning, are lower than main combustor efficiencies. The subsequent decrease in these two indices is probably associated with both afterburning efficiency and the afterburning profile problem; as more fuel is used in afterburning, the local manifold is pressurized higher and flow is initiated to the fill manifold. As a result, exhaust gas profiles are more uniform, and overall afterburning efficiency is higher.

The decrease in NO\(_x\) going from military to afterburning can be explained by combustion zone dwell times. NO\(_x\) production is related to both the peak temperature in the combustion zone and the time gas flow spends at this peak temperature. The primary combustion zone of the main combustor (Fig. 3c) is a zone near the head of the combustor where both fuel flow and airflow are regulated to provide a near stoichiometric combustion at a relatively low velocity. The high temperature products of this combustion are cooled and diluted subsequently by controlled air bleeds in the side of the combustor. The overall flow arrangement of the main combustor provides a controlled high-efficiency burning to obtain the highest heat release possible without compromising thermal limits of engine components. This arrangement results in a low concentration of unreacted combustion products; however, because of the primary zone dwell time necessary for high burning efficiency, a significant amount of nitrogen is dissociated and reacts with available free oxygen to form nitrogen oxides. In the afterburner the combustion is much less controlled.
and occurs at a somewhat higher velocity than in the main combustor. The fuel is injected into engine total airflow, and as a result, the afterburner combustion zone fuel/air ratio is generally lower than stoichiometric. Also, dilution of the combustion products is not controlled and occurs almost instantaneously. Thus, both the peak temperature and the time at peak temperature are lower in the afterburner than in the primary combustor. Therefore, the primary effect of the afterburner combustion, although it does produce some NO\textsubscript{x}, is to dilute the NO\textsubscript{x} produced by the main combustor.

The NO/NO\textsubscript{x} ratio (Fig. 14b) generally increases from idle to cruise to military, decreases going into afterburning, and again increases as afterburning power is increased. This trend is essentially the opposite of the trend of the CO and C\textsubscript{x}H\textsubscript{y} indices with engine power setting; this apparent inverse correlation between either CO and C\textsubscript{x}H\textsubscript{y} and NO/NO\textsubscript{x} ratio is present in other data that will be discussed later.

The data in Fig. 14 are plotted versus equivalence ratio. Although this parameter is controlled directly by engine power setting, it should be realized that increasing power from idle to military increases both main combustor pressure and temperature. The data trends of Fig. 14 include the effect of these variables and not just the effect of equivalence ratio. Conversely, in afterburning, both the main combustor and the afterburner inlet conditions are essentially constant; the primary effects on data trends are those of equivalence ratio and overall afterburner efficiency as discussed above.

4.2.2 Effect of Combustion Zone Inlet Pressure

Military power data (Fig. 15) taken at essentially constant combustor inlet temperatures indicate that the C\textsubscript{x}H\textsubscript{y} emission indices decrease with increasing combustion zone pressure while the NO\textsubscript{x} indices increase. The CO indices increase with pressure at pressures below approximately 60 psia; no trend with pressure was found above approximately 80 psia.

Detailed evaluation of the NO\textsubscript{x} data indicates that the NO\textsubscript{x} index increases approximately with the square root of the combustion zone pressure. Also a consistent trend of decreasing NO/NO\textsubscript{x} index ratio with pressure was indicated by the data (Fig. 15b).

4.2.3 Effect of Combustion Zone Inlet Temperature

Military power data (Fig. 16) taken at essentially constant combustor inlet pressure indicate that the CO emission indices decrease with increasing combustion zone inlet temperature. Conversely, the NO\textsubscript{x} indices increase with increasing temperature. No significant trend of C\textsubscript{x}H\textsubscript{y} or NO/NO\textsubscript{x} ratio with temperature was found.

Detailed evaluation of the NO\textsubscript{x} data indicates that the NO\textsubscript{x} index increases approximately with the 2.5 power of the combustor inlet temperature.
4.2.4 Trend Evaluation

The trends of decreasing CO and C\(_x\)H\(_y\) emission indices with increasing combustion zone pressure and temperature are consistent with combustion efficiency trends predicted by currently accepted combustor technology. The increase and subsequent decrease of these indices in afterburning can be explained as previously discussed by overall afterburner efficiency.

The increase in NO\(_x\) with combustion zone inlet temperature is caused by the associated increase in peak flame temperatures that permit additional dissociation of nitrogen in the combustion flow. No definite cause of the increase of NO\(_x\) with combustion zone pressure is known.

An empirical relationship for the J93 NO\(_x\) data at military power was developed and is shown below:

\[
W_{NO_x} = \frac{5}{3} \times 10^{-6}(P_{in})^{0.5}(T_{in})^{2.5}
\]

Maximum deviation of the calculated values of the NO\(_x\) from measured values for the military power settings and two cruise settings at the eight flight conditions investigated was approximately 20 percent; average deviation was 10 percent. An attempt to use this relationship for idle power settings gave worse agreement, indicating that another variable, possibly dwell time at peak flame temperatures, should be included in the relationship. However, isolation of this variable was not possible.

A comparison of published NO\(_x\) emission data (Ref. 8) for higher pressure ratio engines with values predicted by the J93 empirical relationship gave fair agreement in both level and trend (Fig. 17). A deviation of 33 percent of the predicted value from the measured value at military power suggests again the presence of another variable. However, the probable variable, dwell time, is controlled directly by combustor geometry and it would be coincidental if engines of different sizes and cycles such as those involved in this comparison had the same dwell time.

No explanation for the decrease of NO(NO\(_x\)) emission indices ratio with increasing combustion zone pressure is known. However, the increase of the NO(NO\(_x\)) ratio with engine power setting discussed in Section 4.2.1 is probably caused by oxidation of the NO to NO\(_2\) or other nitrogen oxide compounds by either CO or C\(_x\)H\(_y\). This hypothesis is supported by the afterburning profile data (Fig. 18). Near the periphery of the primary exhaust flow (r/R\(_n\) = 0.7) where exhaust gas temperature and local fuel/air ratio is high (see Fig. 10), the NO(NO\(_x\)) ratio is high. At the same location, both CO and C\(_x\)H\(_y\) are low. As exhaust gas temperature and fuel/air ratio drop toward the center, both CO and C\(_x\)H\(_y\) rise and the NO index falls. On the centerline, CO and C\(_x\)H\(_y\) are quite high and the NO essentially disappears. Comparison of the NO, CO, and C\(_x\)H\(_y\) profiles suggest a stronger inverse NO/C\(_x\)H\(_y\) relationship, but this observation is not considered conclusive. The data in Fig. 18 are from only one test condition but are typical of all afterburning data obtained during the test program.
4.3 SPECTROSCOPIC DATA

Spectroscopic OH data were obtained at six flight conditions at both military and afterburning power settings. The OH values (Fig. 19) varied significantly with both flight conditions and engine power setting. However, comparison of the measured values with those predicted by equilibrium chemistry indicate that the variations can be explained by variations in state conditions of the engine exhaust gas (Fig. 20). The close agreement of the measured and calculated values also indicate that OH in the exhaust jet can be effectively calculated using equilibrium chemistry.

Spectroscopic NO data obtained at three flight conditions at both military and afterburning power settings (Fig. 21) also varied significantly with both flight condition and engine power setting in generally the same direction as the NO data from the emission measurement system. However, the spectroscopic NO data are significantly higher than the values obtained from the sampling system (Fig. 22). No explanation of this disagreement is known; the disagreement is greater than the estimated uncertainty, 28 percent (1 sigma), of the comparison. The measurement system accuracy (excluding transport line effects) is estimated to be 12 percent; the spectroscopic measurements are estimated to be accurate within 25 percent. If both the spectroscopic analysis and the sample system measurements are correct, there exists an inherent problem in the basic method of sampling by a probe and gas transport line to a sensing instrument. The possibility of continuing reactions in the transport line is raised, particularly in light of the NO/CxHy relationship suggested in the afterburning data. Certainly, the question raised is an important one, and one that needs early resolution.

Additional spectroscopic data and details of operation and performance of the spectroscopic technique used are given in Appendix III.

4.4 BATCH ANALYSIS DATA

Gravimetric filter data (Table IV) were obtained at all eight flight conditions investigated. No noticeable trend with flight condition was apparent in the data; however, the data scatter unavoidable in this type of analysis may have masked any trend. Also, the data at Mach 2 and 55,000 ft and above are not directly comparable with the lower altitude and higher Mach number data because of the large filter pore size necessitated by low engine exhaust pressures at these conditions. There is a suggestion of a trend for particulate density to increase from idle to cruise and to decrease from cruise to military engine power. Going into afterburning from military seems to increase particulate density; increasing afterburning power decreases particulate density. These possible trends are in general agreement with reported J85 smoke data (Ref. 9). Preliminary analysis of the particulate size data (Refs. 10 and 11) by TSC indicates that mean particle size is approximately 0.04 microns (Fig. 23). Agreement of the particle size data with previously reported TF30 data (Ref. 7) is not good. However, adjusting the J93 data to the resolution limit of the TF30 analysis yields better agreement. The J93 analysis is preliminary and incomplete at the time of this report; a detailed analysis for all flight conditions will be reported by TSC when completed.
The AEDC wet chemistry analysis of the oxides of sulfur and total aldehydes is shown in Table V. The concentrations of aldehydes are generally low except at engine idle. The concentration of the oxides of sulfur is higher and increases as expected with equivalence ratio. The data unexpectedly indicate that the predominant constituent of these oxides is SO$_3$. However, the analysis method used for SO$_x$ is also responsive to H$_2$SO$_4$, and an SO$_2$/H$_2$O reaction in the sample line may have occurred.

A preliminary analysis (Ref. 10) of the sample bottles of exhaust gas by the Bureau of Mines is shown in Table VI. Agreement of the gas concentrations with AEDC continuous and batch data is good with two exceptions. The BuMines NO$_x$ values are significantly lower than AEDC values. However, this disagreement is not unexpected because of the probability of reactions of the nitrogen compounds in the bottles prior to analysis; agreement of the summed total of the BuMines nitrogen compounds, including nitrites and nitrates, with AEDC's NO$_x$ values is quite good. The other exception is that the BuMines hydrocarbon levels at the Mach 2.6 conditions are two to three times higher than AEDC continuous values. Similar disagreement exists between AEDC's very limited batch samples (Table VII) and the voluminous continuous data obtained with the emission sample system. However, no confirmation of these disagreements is possible; the AEDC continuous data agreed well with the BuMines data at the Mach 2.0 conditions, and the continuous hydrocarbon analyzer performed well during posttest certifications (see Section 4.5).

The AEDC analysis of C$_x$H$_y$ distribution in the exhaust gas (Table VII) showed a trend of increasing carbon number with overall engine equivalence ratio. This trend, from very limited data, was unexpected and is somewhat contradictory to trends reported by the BuMines on sea-level static tests of a TF33 engine (Ref. 12).

The JP5 fuel used for the test program was within the MIL-T-5624G specifications in all significant respects (Table VIII). Sulfur content of the fuel was 0.054 percent by weight. Hydrogen/carbon ratio was approximately 1.9 as opposed to the 2.0 value assumed by ARP 1256. Trace metal concentration in the fuel (Table IX) was quite low with the expected exception of iron.

A comparison was made of the SO$_x$ emission index at military power determined by wet chemistry analysis of the exhaust gas and by fuel analysis. The exhaust gas value of 0.6 lbm/1000 lbm of fuel is lower than the fuel analysis value of 1.1 suggesting that a significant portion of the total exhaust sulfur content may be as sulfate or sulfur particulates.

The UCLA analysis of the exhaust gas sample bottles is not available at this time and will be reported by that organization when completed.

4.5 EMISSION MEASUREMENT SYSTEM VERIFICATION AND ACCURACY

A commercial calibration gas cross-reference service was used to verify overall operation of the emission measurement system. This service provided gases having the
same concentrations of CO, CO$_2$, NO, and C$_x$H$_y$ to each of its users; the certified values of the concentrations are not given with the gases. The users measure the concentrations using their respective measurement systems and calibrations gases and report the results to the supplier who statistically summarizes and reports the combined results. The results of the AEDC participation in this service (Table X) show that acceptable agreement with both the supplier and the remaining users was obtained.

The data shown in Table X allows a good assessment on total system accuracy of emission measurement systems of which the AEDC system is typical. Combined system accuracies (1 sigma) for individual concentration measurements are estimated to be 10 percent for NO and NO$_x$; CO, CO$_2$, and C$_x$H$_y$ concentrations accuracies are estimated to be 5 percent. Accuracies of the emission indices presented herein are estimated to be approximately 12 percent for the various oxides of nitrogen and 6 percent for the remaining gases. This evaluation does not include uncertainties introduced by interactions or reactions in the sample gas transport system, if any; such an evaluation is beyond the scope of this report.

4.6 MISCELLANEOUS

Water concentration in the exhaust gas was not measured because of unacceptable sensitivity shifts of the sensing instrument included in the emission measurement system. However, this concentration can be adequately determined on an overall basis using the equation

$$W_{H_2O} = \frac{9008\alpha}{12.01 + 1.008\alpha}$$

where $\alpha$ = the hydrogen/carbon ratio of the fuel. Values of the overall emission index from this equation are 1,284 and 1,234 lbm/1000 lbm of fuel based on the ARP 1256 value of 2.0 for $\alpha$ and the JP5 fuel average value of 1.9, respectively.

SECTION V
SUMMARY OF RESULTS

Exhaust emission data were obtained at the nozzle exit of an afterburning J93 turbojet engine at eight simulated flight conditions from takeoff to high-altitude supersonic cruise. Parametric variables investigated include flight Mach number, flight altitude, and engine power setting. The major results of the test program are summarized as follows:

1. Exhaust emission indices vary significantly with flight condition and engine power settings. The primary influencing parameters are the pressure and temperature at the entrance to the engine combustion zones.

2. The NO$_x$ production of the J93 at maximum nonafterburning power varied directly with the 0.5 power of the combustor inlet pressure and the 2.5 power of the combustor inlet temperature. Extrapolation of this relationship to operating conditions of higher pressure ratio engines suggests the presence of another variable, perhaps combustor dwell time.
3. Successful in-situ spectroscopic measurements of both NO and OH concentrations in the exhaust gases were made. However, spectrographic values of NO were significantly higher than values from conventional, continuous flow, gas capture analysis. Resolution of this difference, although possibly associated with continuing reaction in the gas transport line used in the conventional analysis, is necessary before the spectrographic technique, a potentially valuable and versatile measurement tool, can be fully accepted.

REFERENCES


APPENDIXES
I. ILLUSTRATIONS
II. TABLES
III. SPECTROSCOPIC DATA
Potential Meteorological Problems

- $H_2O$ (Cloud Cover; Precipitation)
- $CO_2$ (Greenhouse Effect)
- Particles (Blanket; Nucleus for Condensation)
- Depletion of Ozone (UV Shield)

Fig. 1 Environmental Interactions of Turbojet Emissions
a. External Configuration

Fig. 3 J93 Engine
b. Internal Configuration

Fig. 3 Continued
Primary Combustion Zone (Approximately)

Cooling Air Bleed (Typical)

Combustion Secondary Airflow
Compressor Discharge Airflow
Compressor Bleed Air, 8th Stage

Detail
1. First-Stage Turbine Nozzle Vane
2. First-Stage Turbine Rotor Blade
3. Second-Stage Turbine Nozzle Vane
4. Second-Stage Turbine Rotor Blade

c. Combustor and Turbine
Fig. 3 Continued
d. Afterburner

Fig. 3 Continued
e. Engine Operating Characteristics

Fig. 3 Continued
f. Engine Pumping Characteristics

Fig. 3 Concluded
Fig. 5 Emission Measurement System
Fig. 6 Sample Probe
Fig. 7 Sample Rake and Traversing System
Fig. 8 Exhaust Gas Analyzer
Fig. 9 Exhaust Jet Profile at Mach 2.0, 65,000 ft, and Military Power
Fig. 9 Continued

C. Fuel/Air Ratio

D. Carbon Monoxide

Fig. 9 Continued
e. Hydrocarbons

f. Oxides of Nitrogen

Fig. 9 Concluded
Fig. 10 exhaust jet profiles at Mach 2.0, 65,000 ft, and afterburning power.
c. Fuel/Air Ratio

d. Carbon Monoxide

Fig. 10 Continued
Fig. 10 Concluded

- Hydrocarbons
- Oxides of Nitrogen

Jet Radius Ratio, \( r/R_n \)
Carbon Monoxide Emission Index, lbm/1000 lbm fuel

Equivalence Ratio, $\phi$

solid symbols are at military power

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a. Carbon Monoxide

Fig. 11 J93 Exhaust Emissions
b. Hydrocarbons

Fig. 11 Continued

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Solid Symbols Are at Military Power
c. Oxides of Nitrogen

Fig. 11 Continued
Solid Symbols Are at Military Power

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NO/NO\textsubscript{x} Emission Index Ratio (as NO\textsubscript{2})

Equivalence Ratio, $\phi$

d. NO/NO\textsubscript{x} Ratio

Fig. 11 Concluded
Fig. 12 Main Combustor Conditions
Fig. 13 Afterburner Combustion Conditions
Fig. 14 Effect of Power Setting on J93 Exhaust Emission

a. CO, CₓHᵧ, and NOₓ
b. NO/NO\textsubscript{x} Ratio
Fig. 14 Concluded
Military Power
$\phi = 0.29$

Fig. 15 Effect of Combustion Zone Inlet Pressure on J93 Exhaust Emission a. CO, C$_x$H$_y$, and NO$_x$
Fig. 15 Concluded

Open Symbols: $T_{t3} = 1200$ to $1240^\circ$R
Closed Symbols: $T_{t3} = 1030$ to $1070^\circ$R

Military Power
$\dot{\gamma} = 0.29$
Fig. 16 Effect of Combustion Zone Inlet Temperature on J93 Exhaust Emission

a. CO, CₓHᵧ, and NOₓ
Military Power

$P_{s3} = 49$ to $58$ psia

$\phi = 0.22$ to $0.33$

Fig. 16 Concluded
\[ \text{NO}_x = \left[ \frac{5}{3} \times 10^{-8} \right] (p_{s3})^\frac{1}{2} (T_{t3})^\frac{5}{2} \]

**Sym**
- ○ JT3, JT8, JT9 Data (Ref. 7)
- △ Calculated Values

**Fig. 17** Empirical Prediction of NO\textsubscript{x} Production by High-Pressure Ratio Turbojets
Afterburning ($\phi = 0.46$)

Fig. 18 NO and NO$_x$ Exhaust Profiles in Afterburning
Fig. 19 Spectroscopic OH Data
Fig. 20 Comparison of Measured OH with Theoretical Equilibrium
Fig. 21 Spectroscopic NO Data

Solid Symbols Are at Military Power

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Equivalence Ratio, $\phi$
Fig. 22 Comparison of Spectroscopic NO with Emission Measurement System Values
Fig. 23 J93 Exhaust Particulates Size Distribution
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*Exhaust Gas Data Obtained
### TABLE II
**EMISSION MEASUREMENTS**

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<td>Chemiluminescence (CID)</td>
<td>Beckman 315BL</td>
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<td>Volume Concentration</td>
<td>Nondispersive Ultraviolet</td>
<td>Thermo-Electron</td>
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<td>Electrochemical (ECD)</td>
<td>Beckman 325BL</td>
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<td>CID</td>
<td>Dynasciences NR-210</td>
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<td>NOₓ</td>
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<td>ECD</td>
<td>Thermo-Electron 10A</td>
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<td>SOₓ</td>
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<td>Flame Photometry</td>
<td>Dynasciences NX-130</td>
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<td>Particulates</td>
<td>Mass/Volume</td>
<td>Electrostatic Capture with Piezoelectric Micro-Balance</td>
<td>Thermo-Systems 3210B</td>
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#### b. Batch

<table>
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<tr>
<th>Particulates</th>
<th>Mass/Volume</th>
<th>Filter with Gas Total Flowmeter</th>
<th>Gelman/Precision Scientific</th>
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<tbody>
<tr>
<td>NO, OH</td>
<td>Size Distribution</td>
<td>Electrostatic Capture with Electron Microscopy</td>
<td>DOT Laboratory</td>
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<tr>
<td>SO₂, SOₓ, Aldehydes</td>
<td>Remote Absorption Spectroscopy</td>
<td>Scanning UV</td>
<td>AEDC Laboratory</td>
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<tr>
<td>CₓHᵧ</td>
<td>Carbon Distribution</td>
<td>Wet Chemistry</td>
<td>AEDC Laboratory</td>
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<tr>
<td>Trace Gases</td>
<td>Volume Concentration</td>
<td>Bag Sample with Chromatography</td>
<td>AEDC Laboratory</td>
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<td></td>
<td></td>
<td>Bottle Sample with Wet Chemistry and Chromatography</td>
<td>BuMines Laboratory, UCLA Laboratory</td>
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#### c. Ancillary

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<tr>
<th>Fuel</th>
<th>H/C Ratio, Trace Metals, Aromatics, Olefins, Sulfur</th>
<th>Wet Chemistry and Spectroscopy</th>
<th>AEDC Laboratory, Galbraith Laboratory</th>
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<tr>
<td>T₁₉</td>
<td>°F</td>
<td>Ir/Ir-Rho Thermocouple</td>
<td>AEDC Laboratory</td>
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<tr>
<td>P₁₉</td>
<td>psia</td>
<td>Pitot Tube</td>
<td>AEDC Laboratory</td>
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## TABLE III
### J93 EXHAUST EMISSION DATA

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<th>φ</th>
<th>φm</th>
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*Military Power*
# TABLE IV
## GRAVIMETRIC PARTICULATE DATA

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<th>Nominal Filter Size, microns</th>
<th>Particulates, mg/M³</th>
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* M - Small pore membrane filter

P - Large pore polypropylene filter required because of low engine exhaust pressure, experienced weep-through

X - Broken filter required special analysis technique
<table>
<thead>
<tr>
<th>M</th>
<th>h, ft</th>
<th>φ</th>
<th>SO₂, ppm</th>
<th>SOₓ(As SO₂), ppm</th>
<th>Aldehydes(As CH₂O), ppm</th>
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<td>4.77</td>
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</tr>
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<td>0.331</td>
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<td>6.23</td>
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### TABLE VI
## BUMINES J93 EXHAUST GAS ANALYSIS

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<td>M</td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>h, ft</td>
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<td>75,000</td>
<td>75,000</td>
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<td>0.586</td>
<td>0.223</td>
<td>0.560</td>
</tr>
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</table>

**ANALYSIS** (Ref. 11):

- **$\text{O}_2$, percent**
  - 15.0
  - 8.6
  - 15.2
  - 10.8
  - 17.8
  - 9.8

- **$\text{CO}_2$, percent**
  - 3.6
  - 8.9
  - 3.7
  - 5.9
  - 2.7
  - 9.0

- **$\text{CO}$, ppmv**
  - 156
  - 673
  - 260
  - 1,860
  - 35
  - 220

- **$\text{H}_2$, ppmv**
  - 1.5
  - 42.7
  - 3.2
  - 114.6
  - 0.9
  - 7.7

- **$\text{CH}_4$, ppmv**
  - 0.2
  - 0.2
  - 0.5
  - 0.5
  - 0.5
  - 0.5

- **$\text{C}_x\text{H}_y$, ppmv**
  - 16.6
  - 53.4
  - 18.0
  - 20.0
  - 34.5
  - 43.9

- **Total Aldehyde, ppmv**
  - 0.56
  - 1.49
  - 1.96
  - 4.10
  - 0.24
  - 0.18

- **Formaldehyde, ppmv**
  - 0.53
  - 0.89
  - 0.25
  - 0.42
  - 0.02
  - 0.17

- **$\text{H}_2\text{O}_2$, ppmv**
  - ---
  - ---
  - ---
  - ---
  - ---
  - ---

- **$\text{N}_2\text{O}$, ppmv**
  - 1.98
  - 1.57
  - 1.45
  - 0.78
  - 1.23
  - 2.26

- **NO, ppmv**
  - 2.25
  - 2.20
  - 0.86
  - 2.13
  - 1.63
  - 2.33

- **$\text{NO}_x$, ppmv**
  - 4.88
  - 2.20
  - 1.70
  - 3.20
  - 3.00
  - 3.23

- **Nitrate, ppmv**
  - 6.7
  - 5.0
  - 3.2
  - 1.6
  - 7.5
  - 1.5

- **Nitrate, ppmv**
  - 62.4
  - 71.9
  - 44.8
  - 38.2
  - 72.2
  - 132.6

---

*Dash indicates not detectable, below 0.01 ppm*
# TABLE VII
AEDC ANALYSIS OF HYDROCARBONS IN J93 EXHAUST GAS

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<td>2.6</td>
<td>65,000</td>
<td>0.560</td>
<td>101</td>
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</table>

* Determined by analysis of batch sample
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<th>Item</th>
<th>Specified*</th>
<th>Actual</th>
<th>Deviation, One Sigma</th>
<th>No. of Samples</th>
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<td>18,300 min</td>
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<td>0.2</td>
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<tr>
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<td>14</td>
</tr>
<tr>
<td>at 100°F</td>
<td>---</td>
<td>0.796</td>
<td>0.001</td>
<td>16</td>
</tr>
<tr>
<td>Viscosity max, centistokes at -30°F</td>
<td>16.5</td>
<td>9.39</td>
<td>0.37</td>
<td>11</td>
</tr>
<tr>
<td>Sulfur, weight percent</td>
<td>0.4</td>
<td>0.059</td>
<td>0.018</td>
<td>11</td>
</tr>
<tr>
<td>Aromatics max, volume percent</td>
<td>25.0</td>
<td>13.51</td>
<td>0.74</td>
<td>11</td>
</tr>
<tr>
<td>Olefins max, volume percent</td>
<td>5.0</td>
<td>0.68</td>
<td>0.08</td>
<td>11</td>
</tr>
<tr>
<td>Smoke point min, mm</td>
<td>19.0</td>
<td>38.96</td>
<td>2.05</td>
<td>11</td>
</tr>
<tr>
<td>Flash point min, °F</td>
<td>140</td>
<td>122</td>
<td>13</td>
<td>14</td>
</tr>
<tr>
<td>Hydrogen/Carbon Ratio</td>
<td>---</td>
<td>1.909</td>
<td>0.023</td>
<td>13</td>
</tr>
</tbody>
</table>

*MIL-T-5624G
**TABLE IX**

**JP5 FUEL TRACE METAL ANALYSIS**

<table>
<thead>
<tr>
<th>Metal</th>
<th>Analysis* of 3 Samples, μg/ml (±10 to 30 percent)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
</tr>
<tr>
<td>Ag</td>
<td>0.003</td>
</tr>
<tr>
<td>Al</td>
<td>0.003</td>
</tr>
<tr>
<td>B</td>
<td>---</td>
</tr>
<tr>
<td>Ba</td>
<td>---</td>
</tr>
<tr>
<td>Be</td>
<td>---</td>
</tr>
<tr>
<td>Bi</td>
<td>---</td>
</tr>
<tr>
<td>Ca</td>
<td>0.003</td>
</tr>
<tr>
<td>Cb</td>
<td>---</td>
</tr>
<tr>
<td>Cd</td>
<td>---</td>
</tr>
<tr>
<td>Co</td>
<td>0.005</td>
</tr>
<tr>
<td>Cr</td>
<td>0.1</td>
</tr>
<tr>
<td>Cu</td>
<td>0.002</td>
</tr>
<tr>
<td>Fe</td>
<td>1.3</td>
</tr>
<tr>
<td>Mg</td>
<td>0.002</td>
</tr>
<tr>
<td>Mn</td>
<td>0.02</td>
</tr>
<tr>
<td>Mo</td>
<td>0.002</td>
</tr>
<tr>
<td>Na</td>
<td>---</td>
</tr>
<tr>
<td>Ni</td>
<td>0.05</td>
</tr>
<tr>
<td>P</td>
<td>---</td>
</tr>
<tr>
<td>Si</td>
<td>0.02</td>
</tr>
<tr>
<td>Sn</td>
<td>0.005</td>
</tr>
<tr>
<td>Ti</td>
<td>0.002</td>
</tr>
<tr>
<td>Zn</td>
<td>---</td>
</tr>
<tr>
<td>Pb</td>
<td>0.08</td>
</tr>
<tr>
<td>Total Ash (1 liter sample)</td>
<td>1.55</td>
</tr>
</tbody>
</table>

*Concentration values represented by dashes were not detectable.*


**TABLE X**

**EMISSION MEASUREMENT SYSTEM VERIFICATION DATA**

<table>
<thead>
<tr>
<th>Cross Reference Gas*</th>
<th>No. of Users</th>
<th>User Average Value</th>
<th>User Value Deviation (one sigma)</th>
<th>Certified Value</th>
<th>AEDC Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>CO, ppm</td>
<td>23</td>
<td>502</td>
<td>26</td>
<td>495</td>
<td>528</td>
</tr>
<tr>
<td>CO₂, percent</td>
<td>21</td>
<td>3.38</td>
<td>0.14</td>
<td>3.40</td>
<td>3.25</td>
</tr>
<tr>
<td>CₓHᵧ, ppm</td>
<td>17</td>
<td>325</td>
<td>15</td>
<td>334</td>
<td>325</td>
</tr>
<tr>
<td>NO, ppm</td>
<td>28</td>
<td>68</td>
<td>7</td>
<td>74</td>
<td>80</td>
</tr>
</tbody>
</table>

*References 13 and 14
APPENDIX III
SPECTROSCOPIC DATA

INTRODUCTION

A number of molecules produced during the combustion process within jet engines play roles in the chemical balance of the stratosphere (Ref. 15). Among these are nitric oxide (NO) and the hydroxyl radical (OH). To assess the impact of an additional source of these molecules, such as that from SST-like aircraft, on the chemical balance in the upper atmosphere, it is first necessary to determine their rate of emission from typical jet engines. The concentration of highly reactive species such as OH must be measured by an in situ technique, and it is also highly desirable to measure the more stable NO concentration in situ because of the uncertainty as to further chemical reactions within sampling probes and gas analyzer instruments. This appendix describes the application of a spectral absorption technique to the measurement of OH and NO concentrations in the exhaust plane of a YJ93-GE-3 engine operated at several simulated flight conditions.

The measurements were made as part of a program of exhaust emissions measurements on the YJ93-GE-3 engine at simulated flight conditions, conducted in Propulsion Development Test Cell (J-2) at the Arnold Engineering Development Center (AEDC). The engine was operated over a range of simulated Mach speeds from zero to Mach 2.6 and simulated altitudes from sea level to 75,000 ft at power settings from idle to maximum afterburning. Levels of emission of the constituents were primarily measured by sampling through a probe which could be positioned anywhere in the exhaust plane and by analyzing the gas using a variety of commercial process instruments. The results of the sampling measurements are described in the main body of this report.

The method used to make the in situ measurements of OH and NO concentrations is usually referred to as the "narrow line absorption" technique. In this method, the transmission from a narrow line source of radiation through a medium containing the species to be measured is determined and related, either through theoretical expressions or calibration, to the number of absorbing molecules along the absorbing path. The source of radiation is a discharge tube containing a fixed amount of the molecular species of interest. The essentials of the method are contained in the standard text by Mitchel and Zemansky (Ref. 16) and the application to OH in combustion gases has been described by many investigators, including Kaskan (Ref. 17) and Houghton and Jachimowski (Ref. 18). In the work described here, the resonance electronic bands of OH at 3064 Å and NO at 2269 Å were used. The details of the method used at AEDC are described by Davis, McGregor, and Mason (Ref. 19).

The main deterrent to such measurements is usually considered to be the harsh environment in which the laboratory-type instrumentation must be made to function. These hazards were overcome in this work, and a large amount of data were obtained.
DESCRIPTION OF THE METHOD

Theory

The narrow-line absorption technique is based on isolation of the measured absorption to the absorption at line center in order to take advantage of the relation:

$$\int_0^\infty k_\nu d\nu = \sqrt{\frac{\pi}{4 L_n}} \frac{k_o \Delta \nu_D}{\exp \left( \frac{a^2}{2} \right) \text{erfc}(a)}$$

(III-1)

where \( k_\nu \) is the absorption coefficient at wave number \( \nu \) defined by

$$k_\nu = \frac{1}{L} \ln \left( \frac{I_0}{I_\nu} \right),$$

(III-2)

with \( I_0 \) being the incident intensity, \( I \) the measured intensity, and \( L \) the path length in cm. The other symbols have the following definitions:

- \( k_\nu \) = the absorption coefficient at the line center wave number, \( \nu \).
- \( \Delta \nu_D = 2\nu_o \sqrt{2 \ln 2 \kappa T/M c^2} \), the Doppler width in cm^{-1}

where

\( \kappa = \text{Boltzmann's constant, } 1.38 \times 10^{-16} \text{ ergs/}^\circ\text{K} \)

\( T = \text{absolute temperature, } ^\circ\text{K} \)

\( M = \text{mass of absorbing molecule, } \text{gm} \)

\( c = \text{velocity of light, } 3 \times 10^{10} \text{ cm/sec} \)

\( a = \sqrt{\ln 2} \frac{\Delta \nu_N + \Delta \nu_c}{\Delta \nu_D} \), where \( \Delta \nu_N \) is the natural width and \( \Delta \nu_c \) is the collisional width of the line.

In almost all cases, \( \Delta \nu_N \ll \Delta \nu_c \) and may be neglected. For the OH molecule, several investigators have expressed "\( a \)" in terms of the pressure and temperature of the absorbing gas,

$$a = K(P/T)$$

(III-3)

where \( K \) is an experimentally determined constant, \( P \) is the pressure in atmospheres, and \( T \) the temperature in °K. Houghton and Jachimowski (Ref. 18) list several determinations of \( K \) by different investigators but the latest and most thorough work is by Nadler and Kaskan (Ref. 20), who show that \( K \) depends on which line of the (0,0) OH band is being used. Their value (\( K = 585 \)), appropriate for the \( R_2 \) (3) line, will be used in this work.
For the NO molecule, Thorson and Badger (Ref. 21) have shown that collisional broadening is much less than Doppler broadening so that "a" is small and

\[ \exp (a^2) \text{erfc} (a) \sim 1 \]

The absorption coefficient of a single line is expressed in terms of properties of the absorbing gas as follows:

\[
\int_{0}^{\infty} k_\nu d\nu = \frac{\pi e^2}{mc^2} N_{v''j''} f_{v'v''j'j''}
\]

where:

- \( e \) = Electron charge, \( 4.8 \times 10^{-10} \) esu
- \( m \) = Electron mass, \( 9.11 \times 10^{-28} \) gm
- \( N_{v''j''} \) = Number density of molecules in the rotational level (\( J'' \)) of the vibrational level (\( v'' \)) of the ground electronic state, cm\(^{-3} \)
- \( f_{v'v''j'j''} \) = Oscillator strength for the transition from the \( v'J' \) level of the upper electronic state to the \( v''J'' \) level of the ground electronic state.

Now, for Boltzmann equilibrium,

\[
N_{v''j''} = \sum_{v''j''} \frac{\exp \left[ -G(v'')/kT \right]}{\sum_{v''j''} \exp \left[ -G(v'')/kT \right]} \times \frac{(2J'' + 1)}{\sum_{j''} \exp \left[ -F(J'/\nu) / kT \right]} N_0
\]

where \( G(v'') \) is the vibrational energy of the \( v'' \) level, \( F(J'') \) is the rotational energy of the \( J'' \) rotational level, and the summations are the vibrational and rotational partition functions. The oscillator strength for OH, as given by Houghton and Jachimowski (Ref. 18) is \( 8.45 \times 10^{-4} \) and the rotational line strength as given by Dieke and Crosswhite (Ref. 22) with Learner's (Ref. 23) correction, will be used. By combining Eqs. (III-1), (III-4), and (III-5) and using the definitions given, the numerical equation for the number density of OH is given as follows:

\[
N_{OH} = \frac{Q_v Q_r}{W S_{\lambda}} \times \frac{Q_v Q_r}{\sqrt{\ln 2/\pi}} \exp (-\Delta \nu_D) k_\nu_0 \text{cm}^{-3}
\]

The partition functions \( (Q_v \text{ and } Q_r) \), as defined by the summations in Eq. (III-5), are easily calculable, and \( W \), defined as

\[
W = \frac{\sqrt{4 \ln 2/\pi}}{\Delta \nu_D} \exp (a^2) \text{erfc} (a)
\]
is also calculable for any given temperature and pressure. Values for $S_{J''}$ and $F(J'')$ are given for several $R_2$ lines in Table III-1.

In the case of NO, the oscillator strength as given by Perry-Thorne and Banfield (Ref. 24) should be used: $f_0 = 3.64 \times 10^{-4}$. The rotational line strengths are calculable from the formulas for $^{2}\Sigma \rightarrow ^{2}\Pi$ transitions given by Earls (Ref. 25), with the $^{2}\Pi$ state expressed as intermediate between Hund's cases "a" and "b". The resolvable lines are doublets arising from the same energy transition but different spin selection rule. Values for several useful lines arising from $Q_{22} + R_{12}$ transitions are given in Table III-2. The numerical equation becomes

$$N_{NO} = 2.59 \times 10^{16} \frac{\Delta \nu_D Q_y Q_{J'} k_{\nu_0}}{S_{J''} \exp[-F(J'')/\kappa T]} \text{ cm}^{-3}$$  \hspace{1cm} (III-8)

### TABLE III-1
LINE STRENGTH FACTORS AND TERM ENERGIES FOR SEVERAL LINES OF OH IN 3064 Å (0,0) BAND

| $R_2$ | $S(J''|^1$ | $S(J''|^2$ | $F(J'')$, cm$^{-1}$ |
|-------|------------|------------|---------------------|
| (1)   | 2.7        | 2.7        | 126.4               |
| (3)   | 8.9        | 8.8        | 289.0               |
| (4)   | 12.8       | 12.6       | 429.5               |
| (14)  | 53.2       | 45.7       | 3,841.5             |
| (15)  | 57.2       | 49.1       | 4,371.2             |
| (20)  | 77.4       | 57.3       | 10,762.3            |

$^1$Experimental value given by Dieke and Crosswhite (Ref. 22)
$^2$Value after applying Learner's correction (Ref. 23)

### TABLE III-2
LINE STRENGTH FACTORS AND TERM ENERGIES FOR SEVERAL LINES OF THE $Q_{22}$, $R_{12}$ BRANCH OF THE NO (0,0) $\gamma$-BAND AT 2269 Å

<table>
<thead>
<tr>
<th>$(J'')$</th>
<th>$S(J'')$</th>
<th>$F(J'')$, cm$^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>25/2</td>
<td>18.33</td>
<td>288.75</td>
</tr>
<tr>
<td>23/2</td>
<td>16.53</td>
<td>245.80</td>
</tr>
<tr>
<td>21/2</td>
<td>14.75</td>
<td>206.28</td>
</tr>
<tr>
<td>15/2</td>
<td>9.56</td>
<td>108.31</td>
</tr>
<tr>
<td>13/2</td>
<td>7.88</td>
<td>82.53</td>
</tr>
<tr>
<td>9/2</td>
<td>4.56</td>
<td>41.27</td>
</tr>
</tbody>
</table>

72
The lines of the NO gamma bands are much more closely spaced than those of OH so that they are not always resolvable with field-type instruments. This was the case for the measurements reported herein. In this case, one can write (Ref. 18):

\[
\left( \frac{I}{I_0} \right)_{\nu}^{\nu'} = \frac{I_1}{I_T} e^{-k_1 L} + \frac{I_2}{I_T} e^{-k_2 L} + \ldots + \frac{I_i}{I_T} e^{-k_i L}
\]  

(III-9)

where \((I/I_0)_{\nu}^{\nu'}\) is the transmissivity measured in the wave number interval \(\nu - \nu_a\), \(I_1, I_2, \ldots, I_i\) are the incident intensities of the individual lines within the interval, \(k_1, k_2, \ldots, k_i\) is the absorption coefficient of each of the lines, and \(I_T\) is the sum of all the incident individual line intensities from the source. In principle, Eq. (III-9) can be solved for the density of a species if the temperature is known, but it is generally easier to resort to a laboratory calibration. In the work reported here, a laboratory calibration was performed for the NO molecule, and the method of data reduction when using the calibration is described in the next section.

When the source line width is not much narrower than the absorption line width a correction is necessary which adjusts the measured transmissivity to line center transmissivity. Mitchel and Zemansky (Ref. 17) show that, for Doppler line shapes, the measured transmissivity \([I/I_0]_m\) of a line is related to the transmissivity at line center \([I/I_0]_0\) by the integral relation:

\[
\left( \frac{I}{I_0} \right)_m = \frac{\int_0^\infty \exp \left[ -\left( \frac{\omega - \omega_o}{a} \right)^2 \right] \exp \left[ -k_\nu_o \omega \exp \left( -\omega^2 \right) \right] d\omega}{\int_0^\infty \exp \left[ -\left( \frac{\omega - \omega_o}{a} \right)^2 \right] d\omega}
\]  

(III-10)

where

\[
k_\nu_o = -\frac{1}{L} \ln \left( \frac{I}{I_0} \right)_{\nu_o}
\]

\[
a = \frac{\Delta \nu_D \text{(source)}}{\Delta \nu_D \text{(absorber)}}
\]

\[
\omega = \frac{2(\nu - \nu_o)}{\Delta \nu_D} \sqrt{\ln 2}
\]

This correction is applicable for OH when "\(a\)" is small as it was in most cases under consideration in this work.

**Experimental**

The essential ingredients for application of the narrow-line absorption technique are a radiation source, optics for beaming the source radiation through the absorbing gas, a wavelength selective receiver (spectrometer), and a stable detector and signal conditioning.
system. The arrangement of this hardware is illustrated in Fig. III-1. The most unique requirement is that the source emit only radiation at the line center of the absorbing gas. The most convenient way at present of ensuring this condition is to create a gaseous discharge at low pressure and temperature in which the only radiation in the wavelength band of interest is the resonance radiation of the molecule to be studied. Several techniques are available for providing such a source, and the one used here will be described in the next section. The potential of tunable dye lasers in the ultraviolet is recognized, but stability and line width limitations do not currently make this device a contender.

![Fig. III-1 Physical Arrangement for Spectral Line Absorption Measurements](image)

It has been assumed in the development of the equation for the narrow-line absorption method that the absorbing path was uniform in both temperature and concentration. When this is not the case, inversion techniques can be employed for certain simple geometries. For axisymmetric streams such as jet engine exhausts, the Abel inversion can be applied. This follows by writing the fundamental absorption equation in the form:

$$\ln \left( \frac{I(y)}{I_o} \right)_{\nu_o} = -\int_{-\sqrt{R^2-y^2}}^{\sqrt{R^2-y^2}} k_{\nu_o}(x) \, dx$$  \hspace{1cm} (III-11)

where the appropriate geometry is illustrated in Fig. III-2. This is the form of an Abel integral transformation:

$$k_{\nu_o}(r) = \frac{1}{\pi} \int_0^R \frac{\ln \left( \frac{I(y)}{I_o} \right)_{\nu_o}}{\sqrt{y^2-r^2}} \, dy$$  \hspace{1cm} (III-12)

Many routines are available for performing this inversion, and it is possible to obtain both local temperature and concentration from an axisymmetric source by its use. Unfortunately, a very rigid traversing platform is required for either the instruments or the engine (or else, many instrument stations located in the plane of measurement) in order to obtain \( (I(y)/I_o) \) as a function of the distance from the centerline. Thus, the inversion could not be employed in this work, and it was necessary to assume a uniform path at some average temperature and concentration.
DESCRIPTION OF APPARATUS

Engine and Test Facility

The YJ93-GE-3 turbojet engine and the characteristics of the J-2 test cell are fully described in the main body of this report. It suffices to say here that the test was a direct-connect test in which ram air was supplied to the engine inlet at the temperature and pressure corresponding to a given airspeed and altitude and that the static pressure at the engine exit was maintained at the pressure corresponding to the given altitude. The engine could be operated at any power setting, but spectral absorption data were taken only at military power and several afterburner power settings.

The conversion of the spectral absorption measurements to an absolute average number density of the species in question requires knowledge of the static gas pressure and temperature along the path. The concentration in parts per million requires also the gas density, which is found from knowledge of the static pressure and temperature. In order to be consistent, since measurements of total pressure and temperature were not available for every condition that absorption data were obtained, the YJ93-GE-3 PFRT Cycle Review (Ref. 6) was used. The other engine variable required in the absorption measurements is the optical path length, or diameter of the engine exhaust stream. At the station 12 in. downstream of the nozzle exit where the optical path intercepted the gas stream, it was found that the total pressure survey provided the best indication of the diameter of the stream, and this value was used as the path length.

Radiation Source

The source of narrow-line radiation was provided by a gas discharge tube of special design shown schematically in Fig. III-3. A discharge is maintained through the capillary
tube by two electrodes at either end which are connected to a 2800-v power supply. Gas flows continually through the tube, and the pressure is adjusted to a value between 1 and 10 torr which appears to give the most stable operation. The tube is water cooled to remove the heat transferred to the Pyrex® walls and keep the gas as cool as possible in order to minimize Doppler broadening of the spectral lines. Fused silica windows are mounted at either end of the tube to allow transmission of ultraviolet radiation. A monitoring photomultiplier tube with an interference filter is used at one window to record any drift in the radiative output that might occur.

a. Diagram

b. Photograph

Fig. III-3 Discharge Tube Used to Produce Narrow-Line Radiation
Experience has led to the use of special gas mixtures to provide spectra which are intense and unperturbed by bands from other molecules. To produce intense, narrow-line OH resonance radiation, argon is bubbled through water at atmospheric pressure and then passes through a control valve, the discharge tube, and a vacuum pump. The argon evidently causes a selective excitation of the OH, producing the intense 3064 Å band shown in Fig. III-4, which was obtained with a 1/2-meter JACO spectrometer having a 2360 groove/mm grating.

Fig. III-4 Line Spectrum of the (0,0) OH Band at 3064 Å Produced by a Discharge through Flowing Argon with Entrained Water Vapor

The NO $\gamma$-band radiation was found to be most intense when a 12:3:1 mixture of Ar:N$_2$:O$_2$ at a pressure of about 5 torr was used. The (0,0) band of the $\gamma$-system of NO is shown in Fig. III-5 as obtained in second order using a 1-meter Jarrell-Ash instrument having a 1180 groove/mm grating. The lines of the $\gamma$-band are too closely spaced to be resolvable using the 1/2-meter instrument which had to be used in the test cell installation, so that an integrated band spectra as shown in Fig. III-6 had to be used. However, the principles of the narrow-line absorption method are still applicable since the source does emit narrow lines.

Fig. III-5 Line Spectrum of the (0,0) $\gamma$-Band of NO from a Discharge through a 12:3:1 Mixture of Ar:N$_2$:O$_2$
The receiver was a 1/2-meter Jarrell-Ash grating spectrometer having curved, adjustable slits, and a 2360 groove/mm grating blazed for maximum reflection at 3000 Å. The wavelength selector was operated remotely by a selsyn motor/servo pair. The wavelength position was transmitted by the signal from a 10-turn potentiometer, which served as the x-input for the x-y plotter used to record the data. The detector was a Hamamatsu type R-106 photomultiplier tube with S-19 spectral response. The source radiation was modulated at a frequency of 400 Hz, using an American Time Products Company tuning fork chopper. The ac signal from the detector passed through a Philbrick Mod 1003 preamplifier located within the spectrometer enclosure and to a Princeton Applied Research lock-in amplifier located in the instrument control room some 100 ft away. The data were recorded on both an x-y plotter and a strip chart recorder. The instrumentation system is shown in Fig. III-7.

For the OH measurements, a 30-micron slit width was used (resolution about 0.2 Å), whereas for the NO measurements, a 200-micron slit (1.6 Å bandpass) was used.
Installation

The orientation of the components of the measurement system with respect to the engine is diagrammed in Fig. III-8 and shown pictorially in Fig. III-9. In each case, the view is toward the rear of the engine. Two f-4.4 fused silica lenses of 10-cm focal length were placed at one focal distance from the source and spectrometer, respectively, so that a parallel beam of light was projected through a diameter of the engine exhaust at a point approximately 12 in. downstream of the nozzle exit plane. The distance from source unit to spectrometer was approximately 12 ft. The optical system was aligned by use of a small helium-neon laser placed at the centerline of the engine exhaust. The alignment did not have to be repeated during the 4-week test program.
The source unit and spectrometer were housed in insulated, water-cooled, aluminum enclosures mounted to the cell walls as shown in Fig. III-9. The enclosures were purged at atmospheric pressure with dry nitrogen. The source and spectrometer were attached to 1-in.-thick steel plates which were set on vibration isolation mounts attached to the base of the enclosures. No effect of vibration was detected during the testing. Thermocouples located inside the enclosures indicated that the temperature never exceeded approximately 100°F, and no adverse temperature effects were observed. The fused silica
windows through which the beam passed became coated with an oily-like substance during the tests even though a stream of nitrogen was aimed at them at all times. The degradation of the signal due to window-coating did not seriously affect the measured results although the data reduction procedure was made more difficult, as will be discussed in the following section.

DESCRIPTION OF DATA

OH Data

An example pair of OH spectra obtained during the test is shown in Fig. III-10. The upper spectra was obtained before the engine was started, and the lower spectra was obtained some 3 hours later during an afterburning data point. The magnitude of the absorption is notable as well as the reduction in the reference intensity as evidenced by the two spurious nitrogen bands and the lines from large rotational quantum number states (note particularly \( R_2(15) \)). The method of data reduction is outlined as follows:

1. Determine a reference value for the line to be measured using the \( R_2(15) \) line—\( I_0 \)

2. Determine the transmissivity of the line, \( (I/I_0)_m \)

![OH Spectra](image-url)
3. Correct the transmissivity to line center transmissivity using Eq. (III-10) and the given exhaust gas temperature. (A plot of measured transmissivity versus line center transmissivity for different values of $\alpha$ was determined from Eq. (III-10) and used for correction to line center transmissivity.)

4. Determine the number density using Eq. (III-6) at the appropriate gas pressure, temperature, and stream diameter and the constants of Table III-1.

The data reduction was carried out only for the $R_2(3)$ and $R_2(4)$ lines for this report.

NO Data

An example pair of NO spectra obtained during the test is shown in Fig. III-11. The upper trace is the reference spectra, and the lower trace is for an afterburner data point. In this case, the (1,1) and (2,2) bands from the source are also included in the spectral scans. The intensity of the (2,2) bandheads provides a convenient source of correction for source degradation since they experience very little absorption at the temperatures encountered in the exhaust stream.

![NO Spectra](image)

*Fig. III-11 NO Spectra Taken before YJ93-GE-3 Engine Ignition and during Afterburning*
A laboratory calibration was performed using an absorption tube of known length in which the pressure could be controlled. The spectra obtained in the laboratory at several different pressures are shown in Fig. III-12. The method of calibration selected was as follows:

1. Measure the transmissivity at the peak of the second bandhead.
2. Correct the measured transmissivity to line center, using Eq. (III-10).
3. Plot a calibration curve of N x L versus (I/I₀) (Fig. III-13).

The data reduction procedure was then straightforward and consisted of (1) correcting the reference intensity at the second bandhead using the (2,2) bandhead intensity, (2) measuring the transmissivity at the second bandhead, (3) correcting to line center transmissivity, (4) determining the density from Fig. III-13, using the path length given by the pitot pressure survey, and (5) correcting for the difference in temperature between the calibration cell and the engine exhaust.

This method of data reduction is subject to alteration to any number of schemes. The most scientific would be by use of Eq. (III-9), but difficulty arises in selecting a suitable wavelength increment in which the intensities of all the lines are measurable.
RESULTS AND DISCUSSION

Results

The results of the emissions measurements on the YJ93-GE-3 engine using the data reduction procedures just described are given in Tables III-3 and III-4 and Figs. III-14 and III-15. Examination of the tables leads to the following notable observations:

1. OH densities are strongly affected by afterburning, and values for the maximum afterburning power settings run in this series of tests are larger than values for the military power setting by about an order of magnitude.

2. OH densities are directly related to total fuel flow rate.

3. NO densities are not noticeably dependent on afterburning or total fuel flow rate, but remain about constant for all data points included.

4. NO concentrations measured by the in situ method are factors of from 1.5 to 5 larger than those measured by sampling and gas analyzers.
<table>
<thead>
<tr>
<th>Simulated Flight Conditions</th>
<th>Engine Power Setting</th>
<th>Total Fuel Flow Rate, lbm/hr</th>
<th>Average Total Density, cm(^{-3}) x 10(^{18})</th>
<th>Average Hydroxyl Density, cm(^{-3}) x 10(^{14})</th>
<th>Average Hydroxyl Concentration, ppm</th>
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<td>0</td>
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<td>4.45</td>
<td>6.41</td>
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<td>0.16</td>
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<tr>
<td>2.6</td>
<td>65</td>
<td>A/B</td>
<td>17,900</td>
<td>0.35</td>
<td>2.08</td>
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</table>
### TABLE III-4
CONCENTRATION OF NO IN EXHAUST OF YJ93-GE-3 ENGINE AT SIMULATED FLIGHT CONDITIONS

<table>
<thead>
<tr>
<th>Simulated Flight Conditions</th>
<th>Engine Power Setting</th>
<th>Total Fuel Flow Rate, lbm/hr</th>
<th>Average Total Density, cm⁻³ x 10¹⁸</th>
<th>Average Nitric Oxide Density, cm⁻³ x 10¹⁴</th>
<th>Average Nitric Oxide Concentration, ppm</th>
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<td>1.42</td>
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<tr>
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<td>55</td>
<td>A/B</td>
<td>18,950</td>
<td>0.55</td>
<td>1.53</td>
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<td>2.6</td>
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<td>7,100</td>
<td>0.70</td>
<td>2.26</td>
</tr>
<tr>
<td>2.6</td>
<td>65</td>
<td>A/B</td>
<td>17,900</td>
<td>0.35</td>
<td>2.16</td>
</tr>
</tbody>
</table>
The uncertainty for these measurements is estimated to be about ±10 percent based on several repeated data points. The error in the absolute number densities cannot be estimated with any reliability since it depends on many factors. Among these are the true path length, the nonuniform distribution of concentration and temperature, static temperature errors, and variations of source intensity during a spectral scan. However, for purposes of determining total emissions of OH and NO, the average molecular densities are believed to be accurate within ±20 percent.
Comparison of OH Results with Theoretical Predictions

In Fig. III-14, the density of OH is shown graphically for the flight conditions under which OH measurements were made and for the power settings of military, minimum afterburning, and the maximum afterburning power setting employed in these tests. At the right of the figure is shown the average measured concentration in parts per million and the equilibrium value calculated at the appropriate equivalence ratio and temperature corresponding to the afterburner combustion chamber. The theoretical value was obtained from the chemical equilibrium calculation procedure given by Osgerby and Rhodes (Ref. 26). It is to be noted that there is reasonable agreement between the calculated and measured values. However, the value measured in the exhaust stream from the afterburning runs is consistently greater than that calculated in the combustion chamber except for one point, the sea-level-static minimum afterburner run. The larger discrepancy between the measured and calculated values for the nonafterburning firings is not understood. The important point to be made is that the OH concentration is apparently governed by the finite rate chemistry in the flow and is not altered very much in the millisecond or so between the two stations. The equilibrium value of \([\text{OH}]\) at the static temperature and pressure existing at the measurement station is about two orders of magnitude less than the measured value. This observation is also supported by the modeling results reported by the Lockheed Research Group (Ref. 27).

Comparison of NO Results with Probe Measurements

Figure III-15 shows the NO molecular density results of Table III-4 in graphical form for the conditions run. Also, at the right of the figure are shown the concentrations determined by the absorption method and by the probe sampling method. The probe results were averaged [NO] data along the path taken from the data presented in the main body of this report. The total [NO\(_x\)] data from the sampling measurements are only slightly larger than the [NO] values. If both measurements are assumed correct, it is apparent that some of the NO must have been converted to some species other than an oxide of nitrogen along the path from sample probe to the analyzer instrument. Further investigation is required to resolve this discrepancy.

At this point, it should also be noted that the rather constant NO molecular density means that the total rate of emission of NO is about constant since the exhaust velocity and area do not vary much with engine flight speed and altitude. This fact is hidden when the measurement is expressed in terms of volume concentration (ppm) or emission index (weight of emissions/weight of fuel). Some absorption measurements were also attempted at the engine idle power setting, but the attenuation was much less than for the military power setting. It is, therefore, suggested that the total rate of emission is a function only of the primary combustor temperature which is controlled by the equivalence ratio in the primary combustor.

Effect of Particulates on Absorption Measurements

One problem in making the UV transmission measurements that was anticipated never arose. It was predicted that the submicron soot particles in the engine exhausts would
introduce scattering of the UV beam that would have to be distinguished from the absorption by the species of interest. However, no apparent reduction in transmitted intensity was detected at the idle engine condition where the particulate loading is probably highest (Ref. 9).

Further Possibilities for In Situ Measurements

The use of in situ techniques for measurement of the density of NO and OH has been demonstrated. It is possible to obtain concentrations of other minor species in the combustion gases such as NO, CH, SO, and CN using the same technique. Methods have also been developed at AEDC to determine concentration profiles of H₂O and CO₂ using measured infrared emission-absorption and band model parameters (Ref. 28). This iterative technique also provides perhaps the best method for obtaining static temperature profiles. The emission-absorption method may also be applied to CO in the infrared, although this has not been done to date. Also, progress has recently been made in the detection of NO₂ by absorption of continuum emission in the 2400 to 2600 Å range. It thus appears that the only species of interest which do not lend themselves to spectral techniques are the unburned hydrocarbons and the particulates.

SUMMARY

The results of the effort reported here on the in situ measurement of hydroxyl and nitric oxide concentrations in the exhaust of the YJ93-GE-3 jet engine by the resonance line absorption technique may be summarized as follows:

1. Spectral absorption techniques were successfully used to determine jet engine emissions of OH and NO.

2. OH emission densities of the YJ93-GE-3 engine were much larger for the afterburning engine than for the nonafterburning engine and, for the afterburner cases, corresponded to levels computed at the afterburner combustor fuel/air ratio, stagnation pressure and stagnation temperature. The values ranged from $0.15 \times 10^{14}$ cm⁻³ at Mach number 1.4, 35,000-ft altitude condition at military power, to a maximum of $10.4 \times 10^{14}$ cm⁻³ at the Mach number 1.4, 35,000-ft, maximum afterburner power setting.

3. NO emission densities for the YJ93-GE-3 engine were independent of whether the engine is afterburning or not afterburning and changed only slightly with flight conditions at the military primary combustor fuel flow. Values ranged from a minimum of about $1.4 \times 10^{14}$ cm⁻³ at the Mach number 2, 55,000-ft, minimum A/B condition to about $2.3 \times 10^{14}$ cm⁻³ at the Mach number 2.6, 66,000-ft, military condition.

4. NO emission levels in ppm for the YJ93-GE-3 engine determined in situ by spectral absorption were larger by factors of from about 1.5 to 5 than those measured by sampling through gas analyzers.
EMISSION MEASUREMENTS OF A J93 TURBOJET ENGINE

June 8 to July 31, 1972—Final Report

D. L. Davidson and A. F. Domal

Available in DDC

Exhaust gas emission measurements were made at the nozzle of a J93 turbojet engine at simulated flight conditions from sea-level static to Mach 2.0 at 75,000 ft and Mach 2.6 at 65,000 ft. Real time measurements of CO, CO₂, CₓHᵧ, NO, and NOₓ were taken over a range of afterburning and nonafterburning engine power settings using a gas sampling system designed to adhere to SAE ARP 1256 specifications. In addition, NO and OH were measured in situ by a narrow-line UV spectral absorption technique. Batch-type measurements of particulates and other trace constituents of the exhaust gas were also made. Major results of the test were that emissions vary significantly with combustor inlet pressure and temperature and, therefore, with Mach number and altitude. Specifically, NOₓ increased with the square root of the combustor inlet pressure and the 2.5 power of the combustor inlet temperature. Differences between the gas sampling and spectroscopic data on NO indicate the need for verification of sampling techniques.
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