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# CATALYTIC OXIDATION OF CYANOGEN CHLORIDE OVER A MONOLITHIC OXIDATION CATALYST

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GUILD ASSOCIATES, INC. Baltimore, MD 21236

April 1997



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AUTHORS Jeffrey M. Campbell and Joseph A. Rossin

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Increasing the concentration of	f water to slightly $> 0.25\%$ .	significantly increase	d the catalytic	c activity. Isothermal
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### PREFACE

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The work described in this report was performed under Contract No. DAAA15-93-C-0070, Design and Development of Regenerable Air Filtration Units and Integration into Host Applications. This work was started in September 1994 and completed in May 1995.

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## CATALYTIC OXIDATION OF CYANOGEN CHLORIDE OVER A MONOLITHIC OXIDATION CATALYST

#### 1. INTRODUCTION

Present air purification systems designed for removal of chemical warfare agents from air streams are based solely on activated, impregnated carbon, namely ASC whetlerite. While these filters function well against a wide range of chemical agents, they possess several shortcomings. First, the carbon filter has a limited capacity for agents which are removed by chemical reaction and those which are weakly adsorbed. Second, prolonged environmental exposure has been shown to reduce the capacity of these filters for agents which are removed by chemical reaction<sup>1</sup>. The result of these shortcomings is to impose change-out requirements which may present logistical as well as disposal burdens to the user. Catalytic oxidation is an alternative air purification technology which is being investigated as a means of alleviating the above mentioned burdens.

Catalytic oxidation has been proven to be a viable means of controlling industrial offgas emissions<sup>2,3</sup>. Offgas streams containing chlorinated organic molecules present an additional concern in that the chlorine associated with these compounds has the potential to poison/inhibit the catalyst<sup>2,4-9</sup>. Studies by Rossin et al.<sup>4,8,9</sup> report that the oxidation of chlorine-containing compounds over supported platinum catalysts will be significantly inhibited due to the adsorption of product HCl onto catalytic sites. The observed inhibition is; however, reversible. Deactivation resulting from the oxidation of chlorine-containing compounds has been observed by several investigators<sup>2,4-9</sup>. For example, Rossin and Farris<sup>8</sup> report a decrease in the conversion of 1,500 ppm chloroform over a 2% Pt/ $\alpha$ -Al<sub>2</sub>O<sub>3</sub> catalyst from 92 to 83% over the duration of a 65 hour experiment. Lester<sup>10</sup>; however, claims that a titania based catalyst is capable of oxidizing chlorine-containing compounds without deactivation.

The catalytic oxidation of nitrogen-containing compounds has the potential to form  $NO_x^{11-13}$ . Lester and Homeyer<sup>11</sup> studied the destruction of a number of amines over a monolithic catalyst which presumably contained platinum. In all cases, the  $NO_x$  selectivity was significant, especially at high reaction temperatures. Rossin<sup>12</sup>, in a study involving the oxidation of hydrogen cyanide over a platinum monolith, reported that while the catalyst could readily destroy hydrogen cyanide,  $NO_x$  selectivities were unacceptable. Later, Rossin and Campbell<sup>13</sup> reported a catalytic process for direct oxidation of nitrogen-containing compounds without the formation of  $NO_x$ . The catalyst employed by the process, whose details were not disclosed, represents a significant advantage in selectivity over conventional oxidation catalysts.

The objective of this study was to develop an experimentally based reactor design equation which would be capable of describing the oxidation of cyanogen chloride over a wide range of process conditions. The approach taken in meeting this objective was to first record isothermal reactor design data over a wide range of temperatures, concentrations and residence times, and then correlate the design data using a reactor design equation which takes into account all necessary transport resistances and kinetic effects. Also included in this effort was quantitatively identifying all reaction products formed during the oxidation of cyanogen chloride, assessing the effects of water on catalyst performance, and assessing the significance of catalyst deactivation.

#### 2. EXPERIMENTAL METHODS

*Materials*: Cyanogen chloride was obtained from Aberdeen Proving Ground. Analysis revealed the compound to be greater than 99% pure. Neat cyanogen chloride was metered into a 17 liter pressure vessel and diluted with either air or helium such that the concentration of cyanogen chloride in the vessel was 1.0% by volume. Calibration gases (namely CO, CO<sub>2</sub>, NO and NO<sub>2</sub>) were obtained from Matheson. The catalyst employed during this study was the Military Air Purification (MAP) catalyst and was obtained from Allied Signal. The catalyst was a platinum monolith with a cell density of 93 cells/cm<sup>2</sup>.

Catalyst Preparation: Catalyst cores were cut from a 30 cm diameter by 10.2 cm long monolith block using a diamond tip hole saw. The catalyst was a ceramic monolith possessing 93 channels per cm<sup>2</sup>. Once cut, the resulting catalyst core was approximately 2.2 cm diameter by 6.0 cm long. The center channels of the monolith were plugged with alundum cement so that only the channels near the outer circumference of the catalyst core (totaling 115 channels) remained open. The resulting catalyst volume was 7.42 cm<sup>3</sup>. Preparing the catalyst in this manner minimizes axial and radial temperature gradients, thereby allowing one to record reaction rate data under near isothermal conditions<sup>12,14</sup>. Following preparation, the catalyst core was wrapped with a thin layer of glass wool and loaded into the reactor. The glass wool was used to maintain a seal between the catalyst and the reactor wall, thereby preventing the gas stream from by-passing the catalyst. Once loaded into the reactor, seven type K fine-wire thermocouples were extended axially into unobstructed channels of the monolith. These thermocouples were located 0.5 cm above the monolith, and 0.5, 1.0, 2.0, 3.0, 4.0 and 5.0 cm from the inlet, and were used to assess the axial temperature gradient within the monolith.

*Equipment:* A schematic representation of the fixed bed reactor system is illustrated in Figure 1. Dry, oil-free air from a PSA air drier was metered to the reactor using either a 0-5 or 0-20 Nl/min mass flow controller. Nl is defined as one liter of dry air at 0°C and one atmosphere pressure. Dry air from the mass flow controller was then delivered to a water saturator, which was housed in a temperature controlled chamber. A back pressure regulator is located downstream of the water saturator. The concentration of water in the air stream may be controlled by controlling the temperature and pressure of the water saturator. In addition, the water saturator could be by-passed to conduct the experiments in dry air.



Figure 1: Schematic representation of fixed bed catalytic reactor system. BPR: back pressure regulator; GC: gas chromatograph; MFC: mass flow controller. Cyanogen chloride was delivered to the air stream from a compressed gas cylinder containing approximately 4.0% cyanogen chloride in helium. The reactant stream was metered into the humidified air stream via one of three mass flow controllers; 0-50 Nml/min, 0-200 Nml/min and 0-2,000 Nml/min. Nml refers to normal milliliter and is defined as one liter of air at 0°C, one atmosphere pressure. In some cases, the 0-2,000 Nml/min flow controller was replaced with a 0-5,000 Nml/min controller. Use of this flow controller bank allowed for achieving a wide range of cyanogen chloride concentrations over the desired air flow rate range.

Once blended, the feed stream was delivered to the catalytic reactor. The reactor consisted of a 66 cm long glass tube approximately 2.5 cm in diameter. The upper 40 cm of the glass reactor served as the pre-heat zone and was filled with 4 mm diameter glass beads. The glass beads provided further mixing of the feed gas as well as surface area for heating the incoming feed stream to reactor temperature. The reactor tube was housed in an 8.9 cm diameter by 60 cm long aluminum block. The aluminum block was electrically heated, and the temperature of the reactor was controlled by controlling the temperature of the aluminum block. The catalyst core was located approximately 7 to 10 cm below the pre-heat zone of the reactor.

Following the reactor, the effluent stream was delivered to a catalytic filter, which was designed to destroy any unreacted cyanogen chloride in the effluent stream. Following the catalytic filter, the air stream was delivered to two scrubber tanks in series. Each scrubber tank was filled approximately half way with a 10 molar sodium hydroxide solution and served to remove hydrochloric acid (a reaction product) from the effluent stream. Flow from the scrubber vessels was delivered to a back pressure regulator, which served to maintain a constant pressure on the reactor. Following the back pressure regulator, the air stream was vented into a fume hood. All lines in contact with cyanogen chloride were contained within the fume hood and were electrically heat traced.

Isothermal Reaction Rate Measurements: Reaction rate data were recorded at temperatures of 200, 250, 275 and 300°C at a pressure of 6±1 psig. The feed concentration of cyanogen chloride was varied between 100 and 10,000 ppm (274 and 27,400 mg/m<sup>3</sup>)<sup>1</sup> in humid ( $T_{dew} = 23 \pm 1^{\circ}$ C) air. The residence time, based on the reactor volume occupied by the unobstructed monolith channels and calculated at 0°C and 1 atm pressure, was adjusted in an effort to achieve conversions between 10 and 95%. Air flow rates were typically varied between 2 and 25 Nl/min. Reaction rate data were recorded Overnight, the catalyst was maintained at reaction only during day-time hours. The same catalyst core was used for all temperature under flowing, humid air. experiments conducted in this study. All process conditions were maintained for between 1 to 3 hours to ensure the achievement of steady-state. It should be noted that for feed concentrations greater than about 1,000 ppm, an increase in the catalyst temperature was observed. In these instances, the catalyst temperature was adjusted such that the average catalyst temperature (numerical average based on the thermocouple measurements) was  $\pm$ 2°C of the desired value. Both feed and effluent streams were analyzed continuously for

<sup>&</sup>lt;sup>1</sup> Concentrations reported in  $mg/m^3$  are referenced to 0°C, one atmosphere pressure.

 $\rm CO_2$  and cyanogen chloride during all runs. The conversion of cyanogen chloride was determined based on the concentration of cyanogen chloride in the feed and effluent streams.

*Effects of Water*: The effects of water concentration on the catalytic activity were determined by recording the conversion of cyanogen chloride as a function of reaction temperature for water concentrations of <0.03%, 0.1%, 0.25%, 0.60%, 1.14% and 2.71%. These concentrations correspond to dew point temperatures of  $<-35^{\circ}$ C,  $-22^{\circ}$ C,  $-11^{\circ}$ C,  $0^{\circ}$ C,  $10^{\circ}$ C and  $23^{\circ}$ C, respectively, at one atmosphere pressure. The dew point temperature of  $<-35^{\circ}$ C corresponds to air delivered from the PSA air drier. The run was initiated by heating the reactor to  $320^{\circ}$ C in flowing, humid air at a residence time of 0.25 seconds. Once at temperature, cyanogen chloride was introduced to the feed stream at a rate required to achieve a concentration of 1,000 ppm. These process conditions were maintained for approximately one hour, at which time, the catalyst temperature was decreased to  $170^{\circ}$ C at  $30^{\circ}$ C/hr, with the effluent stream sampled every 20 minutes ( $10^{\circ}$ C temperature intervals) for CO<sub>2</sub>, N<sub>2</sub>O and cyanogen chloride in the feed and effluent stream. For the run conducted with a dew point temperature of <-35, the run was initiated by heating the reactor to  $400^{\circ}$ C.

*Catalyst stability*: The stability of the catalyst was evaluated by exposing it to 1,000 ppm cyanogen chloride in humid ( $T_{dew} = 23\pm C$ ) air employing a residence time of 0.05 seconds at 275°C. These process conditions were maintained for 68 hours, at which time the run was terminated. During the run, the effluent stream was sampled for the concentration of CO<sub>2</sub>, N<sub>2</sub>O and cyanogen chloride every hour. Just prior to termination of the run, the effluent stream was sampled for the concentration of HCl and Cl<sub>2</sub> using Dräger tubes.

Upon termination of the run, an additional test was performed in dry air ( $T_{dew} < -35^{\circ}C$ ) employing a residence time of 0.50 seconds and a cyanogen chloride feed concentration of 1,000 ppm. This run was conducted at 400°C. Prior to the start of the run, the catalyst was exposed to the dry air stream overnight in order to remove all traces of water from the lines as well as equilibrate the catalyst with respect to the concentration of hydroxyls on the surface. The run was then initiated the next morning and discontinued following 8 hours of operation. Upon termination of the run, the catalyst was cooled to 275°C, and the previously conducted stability run was performed.

Sample Analysis: During the reaction rate studies, the concentration of cyanogen chloride in the feed and effluent stream, and the concentration of  $CO_2$  and  $N_2O$  in the effluent stream were monitored using a gas chromatograph. All analyses were performed using a Hewlett-Packard 5890 gas chromatograph (GC) equipped with both a flame ionization detector (FID) and a thermal conductivity detector (TCD) and automatic sampling valves. A sample of the GC trace is reported in Figure 2. Cyanogen chloride was analyzed using a 2 m by 3.2 mm diameter Hayesep Q column attached to the FID, while  $CO_2$  and  $N_2O$  were analyzed using a 2 m by 3.2 mm diameter Hayesep Q column attached to the TCD. Samples were analyzed first by sampling the effluent for  $CO_2$  and



Figure 2: Sample GC trace showing feed and effluent analysis. 1. Air, 2. CO<sub>2</sub>, 3. N<sub>2</sub>O, 4. H<sub>2</sub>O, 5. Cyanogen Chloride (effluent), 6. Cyanogen Chloride (feed).

 $N_2O$  at 60°C, then heating the oven to 120°C at 40°C/min and sampling the effluent for cyanogen chloride. Following this, the feed stream was sampled for cyanogen chloride. The above analysis sequence required 17 minutes to perform.

### 3. MODEL DEVELOPMENT AND FIT PARAMETER ESTIMATION

*Reaction Rate Model Development*: The monolithic oxidation catalyst employed in this study consisted of a series of straight, parallel channels. The monolith substrate is coated with a thin layer of catalyst support material, termed washcoat. The catalytic metals are impregnated onto the washcoat. The monolith employed in this study had square channels, with a channel density 93 cells/cm<sup>2</sup>. The thickness of the washcoat and substrate were not supplied by the manufacturer. For modeling purposes, the washcoat was assumed to be 0.0035 cm thick, and the substrate was assumed to be 0.0115 cm thick. A schematic representation of a monolith channel is provided below in Figure 3. Based on these dimensions, the monolith is 11.55% washcoat plus catalyst.



Figure 3: Schematic Representation of a Monolith Channel.

The oxidation of cyanogen chloride was modeled according to the reaction rate expression:

$$Rate = \frac{AC_{CK}}{1 + BC_{HCl} + DC_{CK}}$$
(1)

In the above reaction rate expression, A, B and D are fit parameters (s<sup>-1</sup>, cm<sup>3</sup>/mol and cm<sup>3</sup>/mol, respectively),  $C_{HCl}$  is the concentration of hydrochloric acid (mols/cm<sup>3</sup>),  $C_{CK}$  is the concentration of cyanogen chloride (mols/cm<sup>3</sup>) and Rate is the reaction rate (mols/s-cm<sup>3</sup> catalyst). The reaction rate expression may be derived by assuming the reaction occurs via adsorption of cyanogen chloride onto surface hydroxyls followed by decomposition of the adsorbed species. The concentration of oxygen was held constant during all experiments (21%) and was in great excess (meaning that regardless of the cyanogen chloride conversion, the concentration of oxygen was not taken into account. The rate expression also does not take into account the concentration of water.

As long as the concentration of water was greater than the concentration of cyanogen chloride, the concentration of water did not affect the reaction rate.

The above rate expression was incorporated into a fixed bed reactor design equation which took into account external mass transfer resistances and kinetic effects. The equations which govern mass transfer and chemical reaction rates within the channel of the monolith are<sup>15</sup>:

$$\overline{v}\frac{dC_{CK}^{g}}{dz} = k_{m}A[C_{CK}^{s} + C_{CK}^{g}]$$
<sup>(2)</sup>

$$Rate = k_m A' [C_{CK}^g - C_{CK}^s]$$
(3)

where  $\overline{v}$  is the average linear velocity within the channel at reaction conditions (cm/s),  $C_{CK}^g$  and  $C_{CK}^s$  refer to the gas and solid phase concentration of cyanogen chloride, respectively, calculated at reaction conditions (mols/cm<sup>3</sup>), dz is the differential reactor length (cm),  $k_m$  is the mass transfer coefficient (cm/s), A is the surface area of the monolith channel (cm<sup>2</sup>/cm<sup>3</sup> void), Rate is the reaction rate as defined by eq. 1 (mols/s-cm<sup>3</sup>-cat), and A' is the surface area of the catalyst (cm<sup>2</sup>/cm<sup>3</sup> catalyst). The above reaction rate expressions were numerically integrated using a multistep method. Data were correlated over the entire data set simultaneously, rather than by the standard technique of determining the fit parameters at each reaction temperature, followed by correlating the data using Arrhenius type equations. Therefore, both the activation energy/heats of adsorption and pre-exponential factors were determined simultaneously. The algorithm used to determine the fit parameters employed numerical derivatives and minimized the error between the predicted and experimental conversions. The mass transfer coefficient was determined from a limiting Sherwood number correlation<sup>15-16</sup>.

Sh = 
$$3.66(1 + 0.95 \text{ReSc} \frac{d}{z})^{0.45}$$
 (4)

$$k_{\rm m} = \frac{\rm ShD_{ab}}{2R_{\rm h}}$$
(5)

where Sh is the Sherwood number, Re is the Reynolds number, Sc is the Schmidt number, d is channel diameter (cm),  $D_{ab}$  is the diffusion coefficient for cyanogen chloride in air (cm<sup>2</sup>/s), and R<sub>h</sub> is the hydraulic radius of the channel (cm). The diffusion coefficient was estimated to be 0.067 cm<sup>2</sup>/s at 0°C and was allowed to increase proportional to temperature to the power of 1.5 (T<sup>1.5</sup>). Using the reactor design equations, the fit parameters were determined to be:

A = 
$$1.8631(08) \exp(-6.075/T) \text{ s}^{-1}$$
  
B =  $1.9446(04) \exp(+4.456/T) \text{ cm}^3/\text{mol}$   
D =  $3.0643(05) \exp(+2.695/T) \text{ cm}^3/\text{mol}$ 

#### 4. **RESULTS AND DISCUSSION**

Axial Temperature Deviation: Oxidation reactions are highly exothermic. When recording reaction rate data, it is desired to minimize heat transfer effects so that near isothermal operation of the catalyst is achieved. In this manner, kinetic effects will not be masked by heat transfer effects. For pelleted and granular catalysts, temperature excursions in the laboratory reactor can be and often are controlled by diluting the catalyst bed with inert particles of a similar mesh size. The monolithic oxidation catalyst consists of a single piece of substrate and thus cannot be diluted in the same manner as granular catalysts are. Therefore, recording reaction rates under near isothermal conditions presents a challenge to the investigator.

A differential mode of operation presents one option for minimizing the axial temperature gradients. Stock and Lowe<sup>17,18</sup> have measured quality reaction rate data for the oxidation of carbon monoxide by operating the monolith under differential conditions. However, a differential mode of operation was felt to be inappropriate for the present study. This is because the low residence times required to maintain differential conditions may result in entrance and exit effects dominating the fluid dynamics within the monolith channel, rather than the laminar flow that would be encountered by the application. Also, reaction rate inhibition due to the adsorption of reaction products onto catalytically reactive sites, as has been reported previously for the oxidation of chlorine-containing compounds<sup>8,9</sup>, may be difficult to assess, since the concentration of the reaction products will be minimal. Reaction products could always be added to the feed stream, but this would only add an additional order of complexity to an already complex problem.

The oxidation of cyanogen chloride in air is highly exothermic. The heat of reaction (at 298 K) was estimated to be 107.9 kcals/mol. The adiabatic temperature rise (corresponding to 100% destruction of CK) for cyanogen chloride concentrations of 1,000, 3,000 and 10,000 ppm were calculated to be 13.8, 41.4 and 138°C, respectively. Because of the heat generated by the reaction, one can anticipate difficulties in maintaining a uniform temperature along the catalyst length, especially as the concentration of reactant and reaction temperature are increased. The technique to minimize the axial temperature gradient, as described in the Experimental Methods section, involved limiting the heat generated within the volume of reactor occupied by the monolith core by obstructing the center channels of the monolith. Therefore, the reaction would only proceed in channels located near the reactor wall, and the ratio of the heat generated to surface area of the reactor tube is maximized.

Figure 4 reports the temperature deviation for the oxidation of cyanogen chloride at  $300^{\circ}$ C. Temperature deviation refers to the difference between the catalyst temperature recorded at discrete points within the monolith and the average of the temperature measurements. Data correspond to cyanogen chloride feed concentrations of 1,000, 3,000 and 10,000 ppm (2,740, 8,220 and 27,400 mg/m<sup>3</sup>) and were recorded under conditions where the conversion of cyanogen chloride was greater than 90%. As expected, the temperature uniformity of the catalyst was greatly improved at lower conversions and at

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Figure 4: Axial temperature deviation recorded during the oxidation of cyanogen chloride at 300°C.

lower reaction temperatures. For feed concentrations of 3,000 ppm (8,220 mg/m<sup>3</sup>) and below, the axial temperature deviation was typically less than  $\pm 5^{\circ}$ C. For feed concentrations of 10,000 ppm (27,400 mg/m<sup>3</sup>), the axial temperature deviation increased to almost  $\pm 10^{\circ}$ C. These results attest to the near isothermal operation achieved via modification of the monolith core.

*Effects of Water*: Figure 5 reports conversion as a function of temperature for the oxidation of 1,000 ppm (2,740 mg/m<sup>3</sup>) cyanogen chloride in air with different concentrations of water. Data reported in this figure were recorded at a residence time of 0.25 s. Results presented in this figure show that increasing the concentration of water from 0 to 0.25% ( $T_{dew} = <-35$  and  $-11^{\circ}$ C) has a significant effect on the conversion of cyanogen chloride; however, further increasing the concentration of water to 2.7% ( $T_{dew} = 23^{\circ}$ C) did not affect the conversion of cyanogen chloride. These results can be explained assuming the reaction is initiated by the hydrolysis of cyanogen chloride, rather than by an oxidative mechanism. A hydrolysis reaction involving cyanogen chloride would proceed via an interaction between a cyanogen chloride molecule and a hydroxyl group on the surface of the catalyst to form (possibly) an adsorbed -OCN species plus hydrochloric acid. The -OCN species may then decompose to form the reaction products.



Figure 5: Conversion as a function of temperature for the oxidation of 1,000 ppm cyanogen chloride in air as a function of water concentration.

One possible explanation for the effects of water on the catalyst performance stems from the chemisorption of water on the surface of the catalyst. At low water concentrations, the surface of the catalyst is not saturated with hydroxyls; i.e. there are sites available for adsorption of water. Under these conditions, increasing the concentration of water in the feed stream will increase the hydroxyl concentration at the surface of the catalyst, thereby increasing the catalytic activity. Upon further increasing the concentration of water in the feed stream, the catalyst surface becomes saturated with hydroxyls; i.e. there are no more sites available for water adsorption. As a result, the reaction rate is no longer a function of the concentration of water in the feed stream.

**Reaction Products:** The only carbon-containing reaction product identified in the reactor effluent stream was carbon dioxide. No carbon monoxide or products of partial oxidation were identified. Carbon balances were typically  $100\pm5\%$  for all runs conducted in this study. Figure 6 reports the product distribution of the nitrogen-containing reaction products as a function of temperature for the oxidation of 1,000 ppm cyanogen chloride at atmospheric pressure. This test was conducted by placing a small piece of monolith into a 0.95 cm o.d. glass tube and performing the test in the micro-reactor system. In this



Figure 6: Nitrogen-product concentration (A) and nitrogen-product selectivity (B) as a function of reaction temperature for the oxidation of 1,000 ppm cyanogen chloride in humid air.

manner, near isothermal operation could be achieved and the product distribution would not be skewed as a result of temperature effects. Nitrogen-containing reaction products consisted of N<sub>2</sub>, N<sub>2</sub>O, NO and NO<sub>2</sub>, with NO plus NO<sub>2</sub> being collectively referred to as NO<sub>x</sub> in Figure 6. The product distribution was found to vary as a function of temperature, with the formation of NO<sub>x</sub> favored at high temperatures, and the formation of the reduced products (N<sub>2</sub> and N<sub>2</sub>O) favored at the lower temperatures. At 400°C, approximately 65% of the nitrogen-containing products (on an atomic nitrogen basis) consist of NO<sub>x</sub>. Below 300°C, NO<sub>x</sub> decreases to less than 15%; however, under no conditions where the cyanogen chloride conversion was greater than 99% was NO<sub>x</sub> not measured in the effluent stream. Results presented in Figure 6 are important because NO<sub>x</sub> will be difficult to remove from a reactor effluent stream. Campbell and Rossin (1995) reported a catalyst capable of destroying nitrogen-containing compounds without the formation of NO<sub>x</sub>. Such a catalyst would be required to meet the demands associated military chemical defense applications.

Catalytic Deactivation: Figure 7 reports the conversion of 1,000 ppm (2,740 mg/m<sup>3</sup>) cyanogen chloride as a function of time-on-stream in humid air. Data were recorded at  $275^{\circ}$ C employing a residence time of 0.05 seconds. Results illustrate that the



Figure 7: Conversion as a function of time-on-stream for the oxidation of 1,000 ppm cyanogen chloride in humid air at 275°C.

conversion of cyanogen chloride remain constant at about 92% throughout the duration of the run. During the run, 7.4 cm<sup>3</sup> of monolith (about 0.85 g of washcoat) destroyed 99.6 g of cyanogen chloride. Results demonstrate that the catalyst is stable and will not be deactivated by exposure to cyanogen chloride.

Following completion of the run, the catalyst was exposed to dry air ( $T_{dew} = <$ -35°C) at 400°C overnight in order to remove all traces of moisture from the system. The stability of the catalyst was then evaluated under dry conditions by exposing the catalyst to 1,000 ppm cyanogen chloride at 400°C at a residence time of 0.5 s. The results of this test are shown in Figure 8. The conversion of cyanogen chloride rapidly decreased over the first hour of the run. Following this time, the conversion continued to decrease, but at a slower rate. A comparison of results presented in Figures 7 and 8 highlights the effects of water on the performance of the catalyst. In dry air, increasing the reaction temperature by 125°C and decreasing the residence time by a factor of ten was required in order to achieve a meaningful conversion. Following this run, the catalyst was cooled to 275°C overnight in flowing, humid air. In the morning, the catalyst was exposed to 1,000 ppm cyanogen chloride at 275°C at a residence time of 0.05 s in humid air. The conversion of cyanogen chloride was approximate 90%, consistent with the conversion obtained prior to the dry air exposure. Interestingly, exposure of the catalyst to cyanogen chloride under dry conditions did not irreversibly deactivate the catalyst.





Reaction Rate Measurements: Axial temperature gradients were assessed during each run from temperature measurements obtained using thermocouples located within channels near the outer circumference of the monolith. Typically, the axial temperature deviation was less than 3-5°C, with the deviation increasing to 10°C for runs conducted with a 10,000 ppm feed concentration. Even the 10°C temperature deviation was not deemed to be unacceptable, and therefore data recorded under these conditions were used in the fit parameter analysis.

Reaction rate data were recorded at temperatures of 200, 250, 275 and 300°C for cyanogen chloride feed concentrations between 100 and 10,000 ppm (274 and 27,400 mg/m<sup>3</sup>) at 6±1 psig in humid ( $T_{dew} = 23\pm1^{\circ}$ C) air. Figures 9 through 12 report the conversion of cyanogen chloride as a function of residence time at reaction temperatures of 200, 250, 275 and 300°C, respectively. Results presented in these figures show that increasing the concentration of cyanogen chloride has a significant effect on conversion, especially at 200°C. For example, increasing the feed concentration of cyanogen chloride from 300 ppm to 1,000 ppm at 200°C at a residence time of 0.1 seconds results in the conversion decreasing from about 75% to about 50%. At higher reaction temperatures, the effects of increasing the cyanogen chloride feed concentration on conversion are present but less pronounced. Results indicate that the reaction is being inhibited by the adsorption of reactant and/or reaction product (HCl) on catalytic sites.

Several different forms of the reaction rate expression were evaluated for their ability to correlate the experimental data. The form of the reaction rate expression which best described the experimental data is reported as eq. 1. This rate expression takes into account the adsorption of cyanogen chloride and product hydrochloric acid on the reaction sites. Failure to account for the adsorption of hydrochloric acid resulted in a poor data correlation. The solid lines in the Figures 9 through 12 represent the data correlated using the reactor design equation (eqs. 2 and 3). Figure 13 shows a parity plot of the predicted and experimental conversion of cyanogen chloride. Results presented in Figures 9 through 13 demonstrates the ability of the design equation to accurately describe the experimental data.

The form of the reaction rate expression and the effects of water concentration (Figure 5) suggest that the reaction is occurring via a catalyzed hydrolysis reaction, with the reaction being inhibited by the presence of product hydrochloric acid. The reaction rate expression may be derived by assuming gas phase cyanogen chloride adsorbs onto surface hydroxyls and is subsequently decomposed, with the reaction being inhibited by the presence of HCl. It should be noted that the reaction rate expression did not take into account the effects of water. This is because data used to derive the rate expression were recorded under conditions in which the concentration of water did not affect the catalytic activity.







Figure 10: Conversion of cyanogen chloride as a function of residence time in humid air at 250°C. Solid lines represent data as correlated using the reactor design equation.



Figure 11: Conversion of cyanogen chloride as a function of residence time in humid air at 275°C. Solid lines represent data as correlated using the reactor design equation.



Figure 12: Conversion of cyanogen chloride as a function of residence time in humid air at 300°C. Solid lines represent data as correlated using the reactor design equation.



Figure 13: Parity plot of predicted versus experimental conversion of cyanogen chloride.

Figure 14 reports the conversion of cyanogen chloride as a function of residence time at 300°C. Data were recorded for feed concentrations of 3,000 and 10,000 ppm (9,133 and 27,400 mg/m<sup>3</sup>) and are reported for the purpose of evaluating the ability the model to predict conversions greater than 99%. The high conversion scenario was evaluated because military air purification systems require that the challenge be reduced by several orders of magnitude. The solid lines in Figure 14 represent the cyanogen chloride conversion predicted using the reactor design equation. Results presented in this figure demonstrate that the reactor design equation can be used to accurately predict cyanogen chloride reduction ratios up to four orders of magnitude (which was the detection limits of our instruments) under isothermal conditions.

### 5. CONCLUSIONS

Cyanogen chloride may be readily oxidized over the monolithic oxidation catalyst at temperatures greater than about 250°C. Reaction products consisted of CO<sub>2</sub>, HCl, N<sub>2</sub>, N<sub>2</sub>O and NO<sub>x</sub>, with the nitrogen product selectivity being a strong function of catalyst temperature. Catalytic deactivation was not observed under humid conditions, indicating that the catalyst is very stable. In the absence of water, catalyst deactivation was



Figure 14: Reduction ratio,  $C_a / C_a^0$ , of cyanogen chloride as a function of residence time at 300°C.

observed. For water vapor concentrations less than 0.10%, the presence of water in the feed had a significant effect on the activity of the catalyst. In the absence of water, the oxidation of cyanogen chloride was very slow, requiring reaction temperatures greater than 400°C in order to proceed. The addition of a small amount of water to the feed stream greatly enhanced the catalytic activity. For water vapor concentrations greater than 0.25%, the concentration of water in the feed stream did not, to within experimental error, affect the catalytic activity. Under isothermal conditions, the oxidation of cyanogen chloride could be modeled employing a reaction mechanism which assumes the reaction occurs via adsorption and decomposition of cyanogen chloride onto surface hydroxyl groups. The reactor design equation was able to accurately describe the conversion of cyanogen chloride over a wide range of process conditions.

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# Appendix FORTRAN routine used to determine kinetic fit parameters

C-----C This routine, MONOLF, determines the kinetic fit parameters C for the oxidation of compound in air over a monolith C with a reaction rate expression: С С ACa С Rate = -----С (1 + BCp + DCa)С Where: С A, B and D are fit parameters С Ca is the reactant concentration С Cp is the concentration of reaction product С C The reaction is modeled according to: С C v dCa/dz = kA(Ca,g - Ca,s)С Rate = kA(Ca, s - Ca, g)С C Where v is the velocity, z is the reactor length, k is C the mass transfer coefficient, and A is the surface area C of the monolith. The subscripts s and g refer to the C gas and solid phase. C-----IMPLICIT REAL\*8 (A-H,O-Z) DIMENSION Y0(5), CONV(100), A1(6,7), P5(6) DIMENSION CONX(30), X(100) COMMON /SBCOM1/ XKM1, SA, SS, CA0, RHOB COMMON /SBCOM3/ X1, X2, X3 COMMON /SBCOM2/ WASHT, CELLR, TOUT, VEL common /sbcom3/ j, j5 DATA ONE /1.0D00/, TWO /2.0D00/, FOUR /4.0D00/, ZERO /0.0D00/ DATA RG /82.085D00/, KORD /2/, KORD1 /3/ C-----C N is the number of equations to be solved by EPISODE C-----N = 1OPEN(9,FILE='stat.out') OPEN(7,FILE='test.out') OPEN(5, FILE='monol.dat') REWIND(5) C-----C NDAT is the number of data points. C A0, B0 and D0 are initial guesses for the pre-exponential factors C corresponding to the fit parameters X1, X2 and X3. EA, EB and C ED are initial guesses for the corresponding activation energies. C-----READ(5,28) NDAT, A0, B0, D0, EA, EB, ED 28 FORMAT(I7,/,D18.8,/,D18.8,/,D18.8,/,D18.8,/,D18.8,/,D18.8) READ(5,26) RHOB, SAVEH, TOUT, EPS

READ(5,27) TREF, PRESS, CELLR, WASHT, SUBT, XKM READ(5,27) DELA, DELB, DELD, DEL1, DEL2, DEL4 26 FORMAT(3(D18.8,/),D18.8) 27 FORMAT(5(D18.8,/),D18.8) CLOSE(5) DELA = A0/DELADELB = B0/DELBDELD = D0/DELDDEL1 = EA/DEL1DEL2 = EB/DEL2DEL4 = ED/DEL4C------С SS surface area, cm\*\*2 catalyst/cm\*\*3 reactor void С SA surface area, cm\*\*2 catalyst/cm\*\*3 catalyst C-----CELLR = CELLR - SUBT CELLT = CELLR - TWO\*WASHT SS = FOUR/CELLTSA = FOUR\*CELLT/(CELLR\*CELLR - CELLT\*CELLT) VOID = (CELLT\*CELLT)/((CELLR+SUBT)\*(CELLR+SUBT)) C----c The DO 600 loop is used to re-evaluate the function with a c new set of fit parameters C-----DO 600 L = 1.50F1 = ZEROF2 = ZEROF3 = ZEROF4 = ZEROF5 = ZEROF6 = ZEROF1A = ZEROF1B = ZEROF1D = ZEROF11 = ZEROF12 = ZEROF14 = ZEROF2A = ZEROF2B = ZEROF2D = ZEROF21 = ZEROF22 = ZEROF24 = ZEROF3A = ZERO

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F3B = ZERO F3D = ZERO F31 = ZERO F32 = ZERO F34 = ZERO F4A = ZERO F4B = ZERO F4D = ZERO F41 = ZERO F42 = ZERO

F44 = ZERO F5A = ZERO F5B = ZERO F5D = ZERO F51 = ZERO F52 = ZERO F54 = ZERO F6A = ZERO F6B = ZERO F6D = ZERO F61 = ZERO F62 = ZERO F64 = ZERO OPEN(2,FILE='ck.dat') rewind(2)
c The DO 500 loop calculates conversions at a fixed set of process
c conditions, varying the fit parameters accordingly
C
DO 500 J = $1.NDAT$
READ(2,12) TAU, TEMP, CONC, CONV(J)
XKM1 = XKM*(TEMP/573.15D00)**1.5D00
VEL = TOUT*TEMP/(VOID*273.15D00*PRESS*TAU)
CA0 = CONC
TX = (ONE/TEMP - ONE/TREF)
12 FORMAT(4D12.6)
DO 400 J5 = $1.28$
IF(J5 EO 1) THEN
X1 = A0*DFXP(-FA*TX)
X2 = B0*DEXP(FB*?X)
$X3 = D0^* DEXP(ED^*TX)$
ELSE
ENDIF
F(15 EO 2) THEN
$X_{1} = (A_{0} + DEI A) * DEVD(EA * TV)$
X1 = (A0 + DEEA) + DEEA (-EA + 1A) $X2 = B0 + DEYD(EB + TY)$
X2 = D0 DEXI(ED TX) Y3 = D0*DEVP(ED*TY)
FI SE
ENDIE
IF(15 EO 2) THEN
$\frac{11}{10} = \frac{1}{10} = \frac{1}{10}$
$X1 = (A0 - DELA)^{*} DEAP(-EA^{*}IA)$ $Y2 = P0*DEVD(EP*TY)$
$X_2 = D0^* DEXP(ED^* IX)$
$AS = D0^* DEAP(ED^* IA)$
ELDE
IF(J) = A(t) DF(T) F(T) + tT T
$XI = AU^{+}DEXP(-EA^{+}IX)$
X2 = (B0 + DELB) * DEXP(EB * IX)
$X_{2} = DU^{*}DEXP(ED^{*}IX)$
Ir(JS.EQ.S) THEN
X1 = A0*DEXP(-EA*TX)
X2 = (B0 - DELB)*DEXP(EB*TX)

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X3 = D0*DEXP(ED*TX)
ELSE
ENDIF
IF(J5.EQ.6) THEN
 X1 = A0*DEXP(-EA*TX)
 X2 = B0*DEXP(EB*TX)
 X3 = (D0 + DELD)*DEXP(ED*TX)
ELSE
ENDIF
IF(J5.EQ.7) THEN
 X1 = A0*DEXP(-EA*TX)
 X2 = B0*DEXP(EB*TX)
 X3 = (D0 - DELD)*DEXP(ED*TX)
ELSE
ENDIF
IF(J5.EQ.8) THEN
 X1 = A0*DEXP(-(EA+DEL1)*TX)
 X2 = B0*DEXP(EB*TX)
 X3 = D0*DEXP(ED*TX)
ELSE
ENDIF
IF(J5.EQ.9) THEN
 X1 = A0*DEXP(-(EA-DEL1)*TX)
 X2 = B0*DEXP(EB*TX)
 X3 = D0*DEXP(ED*TX)
ELSE
ENDIF
IF(J5.EQ.10) THEN
 X1 = A0*DEXP(-EA*TX)
 X2 = B0*DEXP((EB+DEL2)*TX)
 X3 = D0*DEXP(ED*TX)
ELSE
ENDIF
IF(J5.EQ.11) THEN
 X1 = A0*DEXP(-EA*TX)
 X2 = B0*DEXP((EB-DEL2)*TX)
X3 = D0*DEXP(ED*TX)
ELSE
ENDIF
IF(J5.EQ.12) THEN
X1 = A0*DEXP(-EA*TX)
X2 = B0*DEXP(EB*TX)
X3 = D0*DEXP((ED+DEL4)*TX)
ELSE
ENDIF
IF(J5.EQ.13) THEN
X1 = A0*DEXP(-EA*TX)
X2 = B0*DEXP(EB*TX)
X3 = D0*DEXP((ED-DEL4)*TX)
ELSE
ENDIF
IF(J5.EQ.14) THEN
X1 = (A0 + DELA)*DEXP(-EA*TX)
X2 = (B0 + DELB)*DEXP(EB*TX)
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X3 = D0*DEXP(ED*TX)
ELSE
ENDIF
IF(J5.EO.15) THEN
 X1 = (A0 + DELA)*DEXP(-EA*TX)
 X2 = B0*DEXP(EB*TX)
 X3 = (D0 + DELD)*DEXP(ED*TX)
ELSE
ENDIF
IF(J5.EQ.16) THEN
 X1 = (A0 + DELA)*DEXP(-(EA + DEL1)*TX)
 X2 = B0*DEXP(EB*TX)
 X3 = D0*DEXP(ED*TX)
ELSE
ENDIF
IF(J5.EQ.17) THEN
 X1 = (A0 + DELA)*DEXP(-EA*TX)
 X2 = B0*DEXP((EB+DEL2)*TX)
 X3 = D0*DEXP(ED*TX)
ELSE
ENDIF
IF(J5.EQ.18) THEN
 X1 = (A0 + DELA)*DEXP(-EA*TX)
 X2 = B0*DEXP(EB*TX)
 X3 = D0*DEXP((ED+DEL4)*TX)
ELSE
ENDIF
IF(J5.EQ.19) THEN
 X1 = A0*DEXP(-EA*TX)
 X2 = (B0 + DELB)*DEXP(EB*TX)
 X3 = (D0 + DELD)*DEXP(ED*TX)
ELSE
ENDIF
IF(J5.EQ.20) THEN
 X1 = A0*DEXP(-(EA+DEL1)*TX)
 X2 = (B0 + DELB)*DEXP(EB*TX)
 X3 = D0*DEXP(ED*TX)
ELSE
ENDIF
IF(J5.EQ.21) THEN
 X1 = A0*DEXP(-EA*TX)
 X2 = (B0 + DELB)*DEXP((EB+DEL2)*TX)
 X3 = D0*DEXP(ED*TX)
ELSE
ENDIF
IF(J5.EQ.22) THEN
XI = A0*DEXP(-EA*TX)
 X2 = (B0 + DELB)*DEXP(EB*TX)
 X3 = D0*DEXP((ED+DEL4)*TX)
ELSE
ENDIF
IF(J5.EQ.23) THEN
X1 = A0*DEXP(-(EA+DEL1)*TX)
X2 = B0*DEXP(EB*TX)
```

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X3 = (D0 + DELD)*DEXP(ED*TX)
     ELSE
     ENDIF
     IF(J5.EQ.24) THEN
      X1 = A0*DEXP(-EA*TX)
      X2 = B0*DEXP((EB+DEL2)*TX)
      X3 = (D0 + DELD)*DEXP(ED*TX)
     ELSE
     ENDIF
     IF(J5.EQ.25) THEN
      X1 = A0*DEXP(-EA*TX)
      X2 = B0*DEXP(EB*TX)
      X3 = (D0 + DELD)*DEXP((ED+DEL4)*TX)
     ELSE
     ENDIF
     IF(J5.EQ.26) THEN
      X1 = A0*DEXP(-(EA+DEL1)*TX)
      X2 = B0*DEXP((EB+DEL2)*TX)
      X3 = D0*DEXP(ED*TX)
     ELSE
     ENDIF
     IF(J5.EQ.27) THEN
      X1 = A0*DEXP(-(EA+DEL1)*TX)
      X2 = B0*DEXP(EB*TX)
      X3 = D0*DEXP((ED+DEL4)*TX)
     ELSE
     ENDIF
     IF(J5.EQ.28) THEN
      X1 = A0*DEXP(-EA*TX)
      X2 = B0*DEXP((EB+DEL2)*TX)
      X3 = D0*DEXP((ED+DEL4)*TX)
     ELSE
     ENDIF
```

c setting initial conditions at reactor entrance

C-----

Y0(1) = 1.0D00

C-----

c EPISODE error control

c IERROR = 1 Absolute error control, Ymax(i) = 1

c IERROR = 2 Error relative to Y(i) is controlled.

c If Y(i) = 0, a divide error occurs

c IERROR = 3 Error relative to the largest value of

c Y(i) is controlled. c-----

> IERROR = 1 T0 = 0.0D00 MF = 22 INDEX = 1 H0 = SAVEH

CALL DRIVE(N, T0, H0, Y0, TOUT, EPS, IERROR, MF, INDEX)

CONX(J5) = (ONE - Y0(1))\*100.0D00

	IF(J5.EQ.1) THEN
	X(J) = CONX(1)
	WRITE(*,129) X(J), CONV(J)
	WRITE(9,129) $X(I)$ CONV(I)
	SIIM = SIIM + (X(I)-CONV(I))*(X(I)-CONV(I))
	SOM = SOM + (X(j) - CONV(j)) (X(j) - CONV(j))
	ELSE
	ENDIF
400	CONTINUE
	PA = (CONX(2) - CONX(1))/DELA
	PB = (CONX(4) - CONX(1))/DELB
	PD = (CONX(6) - CONX(1))/DELD
	P1 = (CONX(8) - CONX(1))/DEL1
	P2 = (CONX(10) - CONX(1))/DEL2
	P4 = (CONX(12) - CONX(1))/DEL4
	PAA = (CONY(2) - TWO*CONY(1) + CONY(3))/(DELA*DELA)
	PRP = (CONV(4) - TWO*CONV(1) + CONV(5))/(DELA DELA)
	PDD = (CONX(4) - TWO CONX(1) + CONX(3))/(DELD DELD)
	$PDD = (CONX(6) - 1WO^{+}CONX(1) + CONX(7))/(DELD^{+}DELD)$
	PII = (CONX(8) - 1WO*CONX(1) + CONX(9))/(DEL1*DEL1)
	P22 = (CONX(10) - TWO*CONX(1) + CONX(11))/(DEL2*DEL2)
	P44 = (CONX(12) - TWO*CONX(1) + CONX(13))/(DEL4*DEL4)
	PAB = (CONX(1) - CONX(2) - CONX(4) + CONX(14))/(DELA*DELB)
	PAD = (CONX(1) - CONX(2) - CONX(6) + CONX(15))/(DELA*DELD)
	PA1 = (CONX(1) - CONX(2) - CONX(8) + CONX(16))/(DELA*DEL1)
	PA2 = (CONX(1) - CONX(2) - CONX(10) + CONX(17))/(DELA*DEL2)
	PA4 = (CONX(1) - CONX(2) - CONX(12) + CONX(18))/(DELA*DEL4)
	PBD = (CONX(1) - CONX(4) - CONX(6) + CONX(19))/(DELB*DELD)
	PB1 = (CONY(1), CONY(4) = CONY(8) + CONY(20))/(DELBS DEED)
	PB1 = (CONX(1) - CONX(4) - CONX(0) + CONX(20))/(DELB DEE1) $PB2 = (CONY(1) - CONY(4) - CONY(10) + CONY(21))/(DELB *DE12)$
	PB2 = (CONX(1) - CONX(4) - CONX(10) + CONX(21))/(DELB*DEL2)
	PB4 = (CONX(1) - CONX(4) - CONX(12) + CONX(22))/(DELB*DEL4)
	PD1 = (CONX(1) - CONX(6) - CONX(8) + CONX(23))/(DELD*DEL1)
	PD2 = (CONX(1) - CONX(6) - CONX(10) + CONX(24))/(DELD*DEL2)
	PD4 = (CONX(1) - CONX(6) - CONX(12) + CONX(25))/(DELD*DEL4)
	P12 = (CONX(1) - CONX(8) - CONX(10) + CONX(26))/(DEL1*DEL2)
	P14 = (CONX(1) - CONX(8) - CONX(12) + CONX(27))/(DEL1*DEL4)
	P24 = (CONX(1) - CONX(10) - CONX(12) + CONX(28))/(DEL2*DEL4)
	F1 = F1 + TWO*CONV(J)*PA - TWO*CONX(1)*PA
	$F_2 = F_2 + TWO^*CONV(J)^*PB - TWO^*CONX(J)^*PB$
	$F_3 = F_3 + TWO*CONV(1)*PD - TWO*CONX(1)*PD$
	F4 = F4 + TWO*CONV(I)*P1 - TWO*CONX(I)*P1
	$F_{5} = F_{5} + TWO*CONV(D*P2 - TWO*CONV(1)*P2$
	$F_{2} = F_{2} + TWO^{*}CONV(J) + F_{2} + TWO^{*}CONV(J) + F_{4}$
	$F0 = F0 + 1 WO^{2} CONV(J)^{2}F4 - 1 WO^{2} CONX(1)^{2}F4$
	F1A = F1A + TWO*(CONV(I)*PAA - CONX(I)*PAA - PA*PA)
	FIB = FIB + TWO*(CONV(I)*PAB - CONY(I)*PAB - PA*PB)
	FID = FID + TWO (CONV(I) * PAD - CONV(I) * PAD - PA * PD)
	$\Gamma_{11} = \Gamma_{11} + \Gamma_{10} + (CONV(I) * DA = CONV(I) * DA = DA$
	$\Gamma_{11} = \Gamma_{11} + 1 \text{ WO}^{*}(\text{CONV}(J)^{*}\text{FA1} + \text{CONA}(1)^{*}\text{FA1} + \text{FA}^{*}\Gamma_{1})$ $\Gamma_{12} = \Gamma_{12} + TWO^{*}(\text{CONV}(J)^{*}\text{FA2} + OONV(J)^{*}\Gamma_{2} + $
	$\Gamma_{12} = \Gamma_{12} + 1 WU^{*}(UUNV(J)^{*}PA2 - UUNX(1)^{*}PA2 - PA^{*}P2)$
	$F14 = F14 + 1WO^{(CUNV(J)*PA4 - CUNX(1)*PA4 - PA*P4)}$
	$F2A = F2A + TWO^*(CONV(J)^*PAB - CONX(1)^*PAB - PB^*PA)$
	$F2B = F2B + TWO^*(CONV(J)^*PBB - CONX(J)^*PBB - PB^*PB)$
	$F2D = F2D + TWO^*(CONV(I)*PRD - CONX(I)*PRD - PR*PD)$
	F2I = F2I + TWO * (CONV(I) * PR1 - CONV(I) * PR1 - DR*P1)
	121 = 121 + 1000(0000000) 101 + 0000A(1) FD1 + FD + 1)

APPENDIX

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F22 = F22 + TWO\*(CONV(J)\*PB2 - CONX(1)\*PB2 - PB\*P2)F24 = F24 + TWO\*(CONV(J)\*PB4 - CONX(1)\*PB4 - PB\*P4)

 $\begin{array}{l} F3A = F3A + TWO*(CONV(J)*PAD - CONX(1)*PAD - PD*PA) \\ F3B = F3B + TWO*(CONV(J)*PBD - CONX(1)*PBD - PD*PB) \\ F3D = F3D + TWO*(CONV(J)*PDD - CONX(1)*PDD - PD*PD) \\ F31 = F31 + TWO*(CONV(J)*PD1 - CONX(1)*PD1 - PD*P1) \\ F32 = F32 + TWO*(CONV(J)*PD2 - CONX(1)*PD2 - PD*P2) \\ F34 = F34 + TWO*(CONV(J)*PD4 - CONX(1)*PD4 - PD*P4) \end{array}$ 

 $\begin{array}{l} F4A = F4A + TWO^*(CONV(J)^*PA1 - CONX(1)^*PA1 - P1^*PA) \\ F4B = F4B + TWO^*(CONV(J)^*PB1 - CONX(1)^*PB1 - P1^*PB) \\ F4D = F4D + TWO^*(CONV(J)^*PD1 - CONX(1)^*PD1 - P1^*PD) \\ F41 = F41 + TWO^*(CONV(J)^*P11 - CONX(1)^*P11 - P1^*P1) \\ F42 = F42 + TWO^*(CONV(J)^*P12 - CONX(1)^*P12 - P1^*P2) \\ F44 = F44 + TWO^*(CONV(J)^*P14 - CONX(1)^*P14 - P1^*P4) \end{array}$ 

F5A = F5A + TWO\*(CONV(J)\*PA2 - CONX(1)\*PA2 -P2\*PA) F5B = F5B + TWO\*(CONV(J)\*PB2 - CONX(1)\*PB2 -P2\*PB) F5D = F5D + TWO\*(CONV(J)\*PD2 - CONX(1)\*PD2 -P2\*PD) F51 = F51 + TWO\*(CONV(J)\*P12 - CONX(1)\*P12 -P2\*P1) F52 = F52 + TWO\*(CONV(J)\*P22 - CONX(1)\*P22 -P2\*P2) F54 = F54 + TWO\*(CONV(J)\*P24 - CONX(1)\*P24 -P2\*P4)

 $F6A = F6A + TWO^{*}(CONV(J)^{PA4} - CONX(1)^{PA4} - P4^{PA})$   $F6B = F6B + TWO^{*}(CONV(J)^{PB4} - CONX(1)^{PB4} - P4^{PB})$   $F6D = F6D + TWO^{*}(CONV(J)^{PD4} - CONX(1)^{PD4} - P4^{PD})$   $F61 = F61 + TWO^{*}(CONV(J)^{P14} - CONX(1)^{P14} - P4^{P1})$   $F62 = F62 + TWO^{*}(CONV(J)^{P24} - CONX(1)^{P24} - P4^{P2})$  $F64 = F64 + TWO^{*}(CONV(J)^{P44} - CONX(1)^{P44} - P4^{P4})$ 

500 CONTINUE

100	CO	NT	Π	JU.	E
F	AI(1,	7)	=	-F	1
A	A1(2,	7)	=	-F	2
F	A1(3,	7)	=	-F	3
A	A1(4,	7)	=	-F	4
Ā	1(5,	7)	=	-F	5
Ā	A1(6,	7)	=	-F	6
A	$\lambda 1(1)$	1)	=	F1	A
Æ	$\lambda 1(1)$	2)	=	Fl	В
F	1(1)	3)	=	Fl	D
A	$\lambda 1(1)$	4)	=	F1	1
A	$\lambda I(1)$	5)	=	F1	2
A	$\lambda 1(1)$	6)	=	Fl	4
ł	$\lambda 1(2)$	1)	=	F2	A
F	$\lambda_{1(2)}$	2)	=	F2	В
A	$\lambda 1(2)$	3)	=	F2	D
ŀ	$\lambda 1(2)$	(4)	=	F2	1
ł	$\lambda 1(2)$	5)	=	F2	2
A	1(2)	6)	=	F2	4
A	<u>1(3</u>	ń	=	F3	A
Ā	$\sqrt{3}$	2)	=	F3	В
Ā	1(3)	3)	=	F3	D

A1(3,4) = F31
A1(3,5) = F32
A1(3.6) = F34
A1(4,1) = F4A
$\Delta 1(4,2) = FAP$
$A_1(4,2) = F_4 B_1$
A1(4,3) = F4D
A1(4,4) = F41
A1(4,5) = F42
A1(4,6) = F44
A1(5,1) = F5A
A1(5,2) = F5B
A1(5,3) = F5D
A1(5.4) = F51
$A_1(5,5) = F52$
A1(5,6) = F54
A1(6,1) = E6A
A1(0,1) = FOA
A1(6,2) = F6B
A1(6,3) = F6D
A1(6,4) = F61
A1(6,5) = F62
A1(6,6) = F64
206 format(7d10.4)
CALL GAUSS(KORD, KORD1, A1, P5)
AOLD = A0
BOID = B0
DOLD = D0
O(D) = E
OLDI - EA
$OLD_2 = EB$
OLD4 = ED
A0 = A0 + P5(1)
B0 = B0 + P5(2)
D0 = D0 + P5(3)
EA = EA + P5(4)
EB = EB + P5(5)
ED = ED + P5(6)
EA0 = ((AOLD - A0)/A0)*((AOLD - A0)/A0)
EB0 = ((BOLD - B0)/B0)*((BOLD - B0)/B0)
ED0 = (DOLD - D0)/D0)*((DOLD - D0)/D0)
EE = ((OID) = E((OID) = E((OID)) = E((OID) = E((OID)) = E((OID) = E((OID)) = E((OID) = E((OID)) =
$EER = ((OLD1 - ER)/ER)^{((OLD1 - ER)/ER)}$
$EED = ((OLD2 - EB)/EB)^{*}((OLD2 - EB)/EB)$
$EED = ((OLD4 - ED)/ED)^*((OLD4 - ED)/ED)$
IF(A0.gt.2.0D00*AOLD) A0 = 2.0D00*AOLD
IF(A0.It.0.5D00*AOLD) A0 = 0.5D00*AOLD
IF(B0.gt.2.0D00*BOLD) B0 = 2.0D00*BOLD
IF(B0.lt.0.5D00*BOLD) B0 = 0.5D00*BOLD
IF(D0.gt.2.0D00*DOLD) D0 = 2.0D00*DOLD
IF(D0.lt.0.5D00*DOLD) D0 = 0.5D00*DOLD
IF(EA.gt.1.2D00*OLD1) EA = 1.2D00*OLD1
IF(EA.lt.0.8D00*OLD1) EA = 0.8D00*OLD1
IF(EB.gt, 1.2D00*OLD2) EB = 1.2D00*OLD2
IF(EB   t   0.8D00*OI   D2) FB = 0.8D00*OI D2
IE(ED  of  1.2D00*OLD2) ED = 0.0D00*OLD2
IE(ED + 0.0000000000000000000000000000000000
$I\Gamma(ED,II.0.8D00*0LD4) ED = 0.8D00*0LD4$

APPENDIX

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```
SUME = dsqrt(EA0+EB0+ED0+EEA+EEB+EED)/5.0D00
  ERROR = DSQRT(SUM)/DBLE(NDAT-1)
  WRITE(*, 551) ERROR
  WRITE(9, 551) ERROR
  WRITE(*, 547) A0, B0, D0, EA, EB, ED, SUME
  WRITE(9, 547) A0, B0, D0, EA, EB, ED, SUME
  SUM = 0.0D00
  IF(SUME.LT.1.0D-04) GOTO 699
600 CONTINUE
699 CONTINUE
800 CONTINUE
547 FORMAT(/,2X,'A0 = ',D12.6,2X,'B0 = ',D12.6,2x,'D0 = ',D12.6,/,
  1.
        2X,'EA = ',D12.6,2X,'EB = ',D12.6,2x,'ED = ',D12.6,/,
  1
        2x,'ERROR = ',d12.6,/)
551 FORMAT(/2X, 'XNORM = ', F8.5)
119 FORMAT(2x, 'Conversion: ', f10.4)
129 FORMAT(2x,'Pred. Conv: ',f10.4,2x,'Exp. Conv: ',f10.4)
999 CONTINUE
  close(9)
  STOP
  END
  SUBROUTINE GAUSS(KORD,KORD1,A,P5)
  IMPLICIT REAL*8 (A-H,O-Z)
  DIMENSION A(6,7), P5(7)
  KORD = 6
  KORD1 = KORD + 1
  KORD2 = KORD - 1
  DO 100 N1 = 1, KORD
    N3 = N1 + 1
    DO 101 N2 = N3,KORD
     IF(DABS(A(N2,N1)).GT.DABS(A(N1,N1))) GOTO 105
     GOTO 101
105
       DO 106 I = 1,KORD1
       SAVE1 = A(N2,I)
       SAVE2 = A(N1,I)
       A(N1,I) = SAVE1
106
        A(N2,I) = SAVE2
101
     CONTINUE
    DO 115 N = N1,KORD
     ASAVE = A(N,N1)
     IF(DABS(ASAVE).LE.1.0D-50) GOTO 115
     DO 116 M = N1,KORD1
       A(N,M) = A(N,M)/ASAVE
116
       CONTINUE
115
       CONTINUE
     IF(N1.EQ.KORD) GOTO 100
     NEQ = N1 + 1
     DO 120 N = NEQ,KORD
       IF(DABS(A(N,N1)).LE.1.0D-50) GOTO 120
       DO 121 N2 = N1, KORD1
        IF(DABS(A(N1,N)).LE.1.0D-50) GOTO 121
        A(N,N2) = A(N1,N2) - A(N,N2)
121
          CONTINUE
```

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APPENDIX
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120
      CONTINUE
100 CONTINUE
  P5(KORD) = A(KORD, KORD1)
  DO 200 N = 1, KORD2
    RHS = 0.0D00
    DO 210 I = 1,N
210
      RHS = P5(KORD1-I)*A(KORD-N,KORD1-I) + RHS
200 P5(KORD-N) = A(KORD-N,KORD1) - RHS
  RETURN
  END
  SUBROUTINE DIFFUN (N, T, Y, YDOT)
  IMPLICIT REAL*8 (A-H, O-Z)
  DIMENSION Y(N), YDOT(N)
  COMMON /SBCOM1/ XKM1, SA, SS, CA0, RHOB
  COMMON /SBCOM3/ X1, X2, X3
  COMMON /SBCOM2/ WASHT, CELLR, TOUT, VEL
  common /sbcom3/ j, j5
  DATA PT5 /0.5D00/, ONE /1.0D00/, TWO /2.0D00/, FOUR /4.0D00/
C-----
c The surface concentration of HCN, XAS, is determined using
c Newton's method of itteration, where
С
  0 = XKM1*SA*(Y(1) - XAS) - Rate == f(XAS)
С
c
   XAS = Y(1)
15 FORMAT(2X, 'XAS CONVERGENCE FAILURE, XAS = ',D12.6,/,
  1 2X,'ERROR = ',D12.6)
C-----
C SS surface area, cm**2 catalyst/cm**3 reactor void
C SA surface area, cm**2 catalyst/cm**3 catalyst
C-----
  A1 = -XKM1*SA*CA0*CA0*X3
  B1 = X3*CA0*CA0*XKM1*SA*Y(1) - XKM1*SA*CA0 -
  1 XKM1*SA*CA0*CA0*X2*(ONE - Y(1)) - X1*CA0
  C1 = XKM1*SA*CA0*Y(1) + XKM1*SA*CA0*CA0*X2*(Y(1) - Y(1)*Y(1))
  XAS = (-B1 - DSQRT(B1*B1 - FOUR*A1*C1))/(TWO*A1)
  IF(XAS.GT.ONE) WRITE(*,15) XAS
   YDOT(1) = -(ONE/VEL)*XKM1*SS*(Y(1) - XAS)
  RETURN
  END
   SUBROUTINE PEDERV(N,T,Y,PD,N0)
   IMPLICIT REAL*8 (A-H, O-Z)
   DIMENSION Y(N0), PD(N0,N0)
```

```
RETURN
END
```

C-----C File MONO1.DAT provides input parameters to run C Fit parameter routine, MONO1.F C-----

-	·
52	NUMBER OF DATA POINTS
.46429200E+04	~ <b>A</b> 0
.46250500E+08	B0
.33751100E+08	D0
.60752800E+04	EA/R
.44557700E+04	EB/R
.26948300E+04	ED/R
.1000000E+01	CATALYST BED DENSITY (G/CM**3)
.1000000E-04	INITIAL STEP SIZE (CM)
.5000000E+01	Catalyst Length, cm (TOUT)
.1000000E-05	ERROR TOLERANCE (EPS)
57315000E+03	Reference Temperature, K (TREF)
.13061000E+01	PRESSURE (ATM) (PRESS)
.10370000E+00	CELL RADIUS (CM)
.3500000E-02	WASHCOAT THICKNESS
.11500000E-01	SUBSTRATE THICKNESS
.77500000E+01	MASS TRNASFER COEFFICIENT, CM**2/S
.5000000E+04	DELA
.5000000E+04	DELB
.5000000E+04	DELD
.1000000E+02	DEL1
.1000000E+02	DEL2
.1000000E+02	DEL3

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