

December 7, 1990

RECEIVED

DEC 10 1990

DER-BAOM

Mr. C. H. Fancy, P.E., Chief Bureau of Air Regulation Florida Department of Environmental Regulation 2600 Blair Stone Road Tallahassee, Florida 32399-2400

RE:

Sulfur Storage Permit Modification AC29-187327 Sulfuric Acid Plants "C" and "D" Permit Modification AC29-186931

Dear Mr. Fancy:

In reference to your letter dated October 25, 1990 stating incompleteness of the application for modification of the construction permit, for molten sulfur storage and handling, the following is offered:

1. Please submit the calculation sheet showing the derivation/assumptions of the revised emission estimates for PM/PM₁₀, SO₂, TRS/H₂S and VOC emissions.

Derivation/assumptions for emissions estimates were supplied in the original permit application dated June 28, 1989. The revised emission calculations are based on the emissions stated in the construction permit number AC29-167204 at specific condition 8. The increase in through-put will result in a 10% increase in the number of trucks being unloaded. Therefore, the expected emissions from the increased trucks should result in a 10% increase in total emissions. The revised table for specific condition 8 was derived by multiplying the expected emissions by a factor of 1.10.

Mr. C.H. Fancy - 12/07/90 Page 2

Why does this application state that the sulfuric acid production for the facility is increasing from 6900 TPD to 7600 TPD, when the application for permit modifications of sulfuric acid plants "C" and "D" requests a facility increase from 6900 TPD to 7300 TPD?

Applications for permit modifications for "C" and "D" sulfuric acid plants were submitted to increase the production rate for each plant from 2400 tons/day to 2600 tons/day.

It is anticipated that applications to increase the production rates for "A" and "B" sulfuric acid plants from 1050 tons/day to 1200 tons/day will be submitted within the next year. The modification to the sulfur handling permit was calculated to allow this increase before the fact, to avoid having to modify the permit again in less than a year's time.

Answers to questions 1 through 6 concerning the "C" and "D" Sulfuric Acid Plant permit modifications are attached, as provided by Dr. John B. Koogler.

A reply to your November 13 letter on SO₂ modeling for these plants will be provided by mid-December.

Should you have additional questions, please call Jim Martin at (813) 782-1591.

Sincerely,

J. E. Parsons General Manager

JEP/CJM/tjj

Attachment

cc: P.R. Roberts/T.A. Edwards

C.J. Martin/Env. File

C. Fred Deuel

Jerry Campbell (HCEPC)

Harry Kerns (DER SW District)

o Phillips

DETERMINATION OF NOx CONCENTRATIONS IN SULFURIC ACID PLANT STACK EMISSIONS

Alan A. Pratt CF Industries, Inc.

 NO_{\times} concentrations in "C" and "D" Sulfuric Acid Production plant stack emissions were determined through manual sampling techniques and analyzed using a chemiluminescence detector (NO_{\times} Box) at the University of South Florida, Tampa.

INTRODUCTION

The chemiluminescence detector-based method for trace NO_{\times} in air samples is used and approved by the Environmental Protection Agency. (1)

Concentrations of nitrogen oxides in ambient air are determined by photometrically measuring the light intensity resulting from the chemiluminescent reaction of nitric oxide (NO) with ozone (O3). NO2 and NO3 are also measured by conversion to NO.

Normally, $NO_{\rm x}$ analyzers are used for continuous operation. Since CF Industries, Inc. does not currently own a chemiluminescence analyzer, consultation of remote sampling techniques was done with Dr. Robert S. Braman of the University of South Florida (USF) and analyses were acquired using their $NO_{\rm x}$ analyzer.

EXPERIMENTAL

Apparatus. A Thermal Electron Corporation Model 14 B/E Chemiluminescent NO_{\times} analyzer donated to USF by the DER (#17149) was used for the analysis. Inboard flow rate was controlled by a mass flow meter set to approximately 250 mL./min. The output signal was recorded by means of a Linear Instruments, Inc. Model 252A integrating recorder. The apparatus arrangement is shown in figure 1.

Procedure - Sampling. Sampling tubes were constructed of $^{1}/_{4}$ " glass tubing to which a coating of cobalt oxide (CoO) had been deposited. CoO absorbs NO, NO₂, and NO₃ gas and releases as NO upon heating. Sampling tubes were "blanked" before use by heating with a heating coil while carrier gas was passed through and into the NO_x analyzer. After cooling and capping, the tubes were taken to CFII for sample collection.

Sampling was achieved by purging inlet lines to the continuous SO₂ monitors with a vacuum pump and pulling a

50mL. sample from a "T" through the sample tube with a Hamilton Gastight air syringe. Sampling apparatus is shown in figure 2.

Several ambient air samples were taken for background data in 200 mL. volumes. All sample tubes were capped immediately after sampling for transport to USF for analysis.

Procedure - Analysis. Sample tubes were connected inline to the NO_{\times} analyzer and heated with a heating coil wrapped around the tube with 40V a/c current. Response times averaged 2 minutes. A typical response is shown in figure 3.

Procedure - Response Calibration. A vanadium oxide packed tube was blanked. Triplicate injections into the cool packed tube of standard KNO3 solution was done at each of four different volumes. Heating the tube at 15V produced response. A calibration regression was established for sample comparison (figure 4).

RESULTS

Table I lists the results of each analysis. Results were obtained in units of nanograms N per 50 mL. sample and reported as ppm (uL/L) NO. A sample calculation is as follows:

 $\frac{\text{ng. N}}{\text{N}} \times \frac{1.000 \text{ ml.}}{\text{1 L}} \times \frac{1 \text{ ug}}{\text{10}^3 \text{ ng}} \times \frac{30 \text{ ug NO/mole}}{\text{14 ug N/mole}} \times \frac{22.4 \text{ uL}}{\text{30 ug.}}$

 $\times 298 = u1/L$ 273

LITERATURE CITED

(1) Federal Register. Vol. 54, No. 211, 1989, 40 CFR Part 60. Method 7E.

Figure 1. Apparatus Arrangement

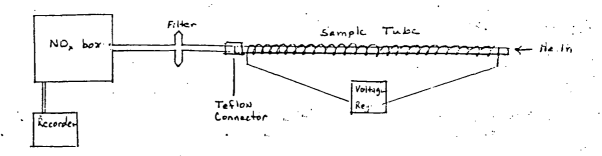


Figure 2. Sampling Apparatus

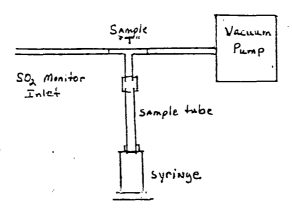


Figure 3. Typical Response of Chemiluminescence Detector

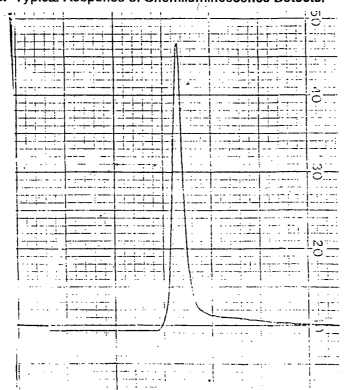
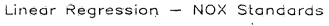


Figure 4. Standard Regression Curve



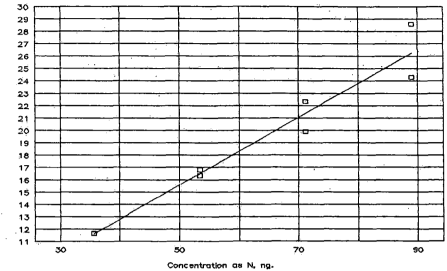


Table I. Concentration of NOX - Results

CSAP			
Sample #	Units	N, ng	NO, ppm.
1	39.88	138.50	4.84
2	41.93	145.96	5.10

DISAP			
Sample #	Units	N, ng	NO, ppm.
1.	43.75	152.58	5.33
2	22.85	76.52	2.67
3	19.09	62.84	2.20

Ambient Air	 * -	4	a the
Sample #	Units	N, ng	NO, ppm.
1	6.29	16.26	0.14

1. Is the 99.7% efficiency (listed on page 5 of the application) of the dual absorption towers based on current test data or is that the proposed efficiency of the towers after they are repacked?

Since the towers have already been repacked (see Question 2, below), this efficiency represents current and proposed operation efficiency. The 99.7 percent efficiency for sulfur dioxide is based on the sulfur recovery necessary to comply with the NSPS emission limit of 4.0 pounds of sulfur dioxide per ton of 100 percent sulfuric acid. The sulfur (M.W. = 32) required to produce 1.0 ton of $\rm H_2SO_4$ (M.W. = 98) is 653.1 pounds. Sulfur loss in the stack gas is 2.0 pounds per ton (2.0 lbs. sulfur per 4.0 lbs $\rm SO_2$). The efficiency of sulfur recovery is (653.1-2.0) x 100/(653.1) = 99.7 percent.

2. How are the absorption towers to be repacked? Is the packing material to be a new type?

There is to be no change in the configuration of the absorption towers. The changes that were made pursuant to the 1988 permitting, in concert with a recent maintenance re-packing of the towers, have resulted in greater capacity than anticipated at the time of permitting.

3. The application states that NO_{X} emissions will be minimized by operating the burners of the sulfuric acid plants within the limits established by the designer. What, specifically, are these limits?

No specific operating parameters are specified by the designer. Operation of the sulfur furnace is based on conditions established by CF and operating practices that are consistent throughout the industry. The sulfur feed rate to the sulfur furnace will be 35.4 tons per hour (corresponding to a production rate of 2,600 tons per day of 100 percent H_2SO_4). The oxygen and SO_2 at the furnace exit will be maintained in the range of 9.2 - 9.4 percent and 11.8 - 11.6 percent, respectively. These operating conditions will result in a furnace temperature of 2000° - 2100°F.

Another factor related to $\mathrm{NO}_{\mathbf{x}}$ that also needs updating is the $\mathrm{NO}_{\mathbf{x}}$ concentration in the stack gas from the CF sulfuric acid plants. In the original application, a "typical" NO_x concentration of 2.1 x 10^{-6} lb NO $_{\rm x}$ per cubic foot (18 ppm ${
m NO}_{
m X}$ by volume) was used. This concentration was measured at a 2250 ton per day sulfuric acid plant in Polk County by Koogler & Associates personnel in about 1976 (using EPA Method 7) and has been used in several sulfuric acid plant permitting projects since that time. Actual measurements recently made by CF personnel on the subject sulfuric acid a chemiluminescent NO_x analyzer plants with attached) showed $NO_{\mathbf{x}}$ concentrations in the stack gas in the range of 5 ppm by volume. Using this site-specific measurement, the predicted NO_x increment is deminimis. amended application includes these data.

4. Please show the correlation between the gas flow rate of 146,162 dscfm (listed on page 6 of the application) and the gas flow rate of 67,500 dscf/ton of acid (listed on page 7b).

The gas flow of 67,500 dscf/ton of 100 percent acid is a "typical" gas volume for double absorption sulfuric acid plants. The gas flow factor specific to the CF Industries "C" and "D" plants averages 64,500 dscf/ton (based on stack test data). This latter factor should have been used in the calculations referenced on page 7b of the application as should have a NO_X concentration of 5 ppm. Amended sections of the application are attached.

Based on the gas flow rate of 64,500 dscf/ton of acid, the stack gas flow rates on page 6 of the applications have also been changed.

5. The application states that the nitrogen oxide concentration in the tail gas stream of a typical sulfuric acid plant is in the range of 20 parts per million. What is the source of this information?

See response to Department question No. 3.

6. The uncontrolled emissions calculations for acid mist (shown on page 7c) need to be recalculated.

The referenced correction has been made in the attached amended application.

<u>General</u>

The change made by the Department on page 2 of the application is appreciated.

It is presumed from the statement regarding actual emissions that no further action is necessary as the emission increases are significant as calculated.

SECTION III: AIR POLLUTION SOURCES & CONTROL DEVICES (Other than Incinerators)

A. Raw Haterials and Chemicals Used in your Process, if applicable:

	Contami		Utilization	
Description	Тура	# Kt	Rate - lbs/hr	Rolato to Flow Diagram
Sulfur	Ash	0.005	71,000	1
•				
				•
				

8.	Process	Rato,	if	applicable: -	(See	Section	٧.	Item	L	Ì
----	---------	-------	----	---------------	------	---------	----	------	---	---

·1. Total Process Input Rate (16s/hr): 71,000 as sulf	·l. Iotal Process Inout Rate (1bs	a/hr}: /l.UUU as sullu
---	---------------------------------------	------------------------

2. Product Weight (lbs/hr): 232.975 as 93% Ho	2_	Product Weight (lbs/hr):		232.975 a	s 93% H2SC
---	----	--------------------------	--	-----------	------------

C. Airborno Contaminants Emitted: (Information in this table must be submitted for each emission point, use additional sheets as necessary)

Hzae of	Emisa	ion ^l	Allowed ² Emission Reto per	Allowable ³ Emission	Potent Emiss		Relate to Flow
.Conteminent	Haximum 16s/hr	Actual T/yr	Rule 17-2	lbs/hr	lbe/xx hr	T/yr	Diagram
S02	.433.3	1898	17-2.600(2)(1) 433.3	433.3	1898	2
N0x	4.2	18.4	17-2.630	4.2(1)	4.2	18.4	2
Acid Mist	16.2	71.2	17-2.600(2)(1) 16.2	270	1183	2
VE	10%		17-2.600(2)(1) 10%		-	2
(1) Expected	emissions;	no app	licable emiss	ion limiting	standard.		

¹ See Section V, Item 2.

²Reference applicable emission standards and units (e.g. Rulo 17-2.600(5)(b)2. Table II, E. (1) - 8.1 pounds per million BTU heat input)

³Calculated from operating rate and applicable standard.

⁴Emission, if source operated without control (See Section V, Item 3).

•	gire:	198.5			Stack D	Leact	or:	8.0	
as Flax	Rate: 140	,060 ACFH	116,460	OSCFH	Gas Exit	Coas	erature:	175	
•									
		·	·	"					
		SEC	TION IVE	INCINER	ATOR INFO	RHATI	ON		
		•	NOT	APPLICA	BLE				
Type of Wests					ge) (Path			Type (as (Solid By-	
					1				
Actual 16/hr ncinor- ated				·			·		
Incon- trolled lbs/hr)							· .		
		:			Desta	. Cana	osty (1he	·/hn)	
al Weigh	ht Incine: . Number o	ated (lbs/h	c) Operation	per dey				/hr)	
tal Weigh proximate nufacture	ht Incines Kumber o	etod (1be/h	r)	bec qsa		dey/w	rk	,	
roximato	ht Incines Kumber o	etod (1be/h	r)	per day	1 No	dey/s	rk	wks/yr	
tal Weigh proximate sufacture se Consti	ht Incines Number of	eted (lbe/h	Cperation	per day	1 Na	dey/s	rk	wks/yr	
imary Ch	nt Inciner	eted (lbe/h	Cperation	per day	1 Na	dey/s	rk	wks/yr	
imary Ch	Number of	Values of (ft)	Heat Re	per day Hode	1 Na	fuol	BTU/hr	wks/yr	· c
imary Ch	Number of	Yolumo (ft)	Heat Re (BTU/	per day Hode lease hr)	1 Na	fuol	BTU/hr Stack	Tomporatur (°F)	·
imary Ch condary ck Hoigh Flow Ra 50 or m	Number of the state of the stat	Yolumo (ft)3	Heat Re (BTU/	per day Hode lease hr)	I No	Fuol	Stack	Tomporatur (°F)	· ε

OER Form 17-1.202(1) Effective November 30, 1982 Proposed: 2600 tons per day 100% acid

S02 - 4.0 1b/ton

Mist - 0.15 1b/ton

Operating factor - 1.0

Emission Rates: (Each plant - as permitted and operated)

SO2: Hourly = $4.0 \text{ lb/ton } \times 2400/24 \text{ tons/hr}$ = 400 lb/hr.

> Annual = $400 \text{ lb/hr} \times 8760 \text{ hr/yr} \times 1/2000 \text{ lb/ton}$ = 1752 tpy

MIST: Hourly = 0.15 lb/ton x 2400/24 tons/hr = 15.0 lb/hr

Annual = 15.0 x 8760/2000 = 65.7 tpy

NOx: Based on 64500 dscf per ton of acid and 0.6×10^{-6} lb NOx per dscf (5 ppm, v/v)

Hourly = $64500 \text{ dscf/ton } \times 2400/24 \text{ ton/hr}$ $\times (0.6 \times 10^{-6}) \text{ lb/ft}^3$ = 3.9 lb/hr

Annual = $3.9 \text{ lb/hr} \times 8760/2000$ = 17.0 tpy

Emission Rates: (Each plant - as proposed)

S02: Hourly = $4.0 \text{ lb/ton } \times 2600/24 \text{ tons/hr}$ = 433.3 lb/hr.

> Annual = 433.3 lb/hr x 8760 hr/yr x 1/2000 lb/ton = 1898 tpy

MIST: Hourly = 0.15 lb/ton x 2600/24 tons/hr = 16.2 lb/hr

Annual = 16.2 x 8760/2000 = 71.2 tpy

NOx: Hourly = $64500 \text{ dscf/ton } \times 2600/24 \text{ ton/hr} \times (0.6 \times 10^{-6}) \text{ lb/ft}^3$ = 4.2 lb/hr

Annual = $4.2 \text{ lb/hr} \times 8760/2000$ = 18.4 tpy

NOTE: No other air pollutants are discharged from the C and D sulfuric acid plants.

3. Uncontrolled Emissions

- SO2 Controlled and uncontrolled emissions of SO2 are identical for a double absorption sulfuric acid plant.
- Mist The control efficiency of high efficiency mist eliminators is estimated to be 94 percent based on measurements at similar plants.

Hourly = 16.2 lb/hr controlled/(1-0.94)

 $= 270 \, lb/hr$

Annual = $270 \text{ lb/hr} \times 8760 \text{ hr/yr} \times 1/2000 \text{ lb/ton}$

= 1183 tons/yr

4. Control System

SO2 will be controlled by the existing two absorption towers and acid mist will be controlled with the existing high efficiency mist eliminators.

5. Control Efficiency

SO2 - Sulfur input to plant = 71000 lb/hr (as S)

Efficiency = (71000-216.7)x100/7100= 99.7%

- Mist High efficiency mist eliminators are estimated to be 94 percent efficient based on measurements made on similar double absorption plants.
- 6. Flow Diagram See attached.
- 7. Location Map See attached.
- 8. Site Map See attached.

ATTACHMENT 1B

ANNUAL AIR POLLUTANT EMISSION CHANGES RESULTING FROM THE PROPOSED SULFURIC ACID PLANT RATE INCREASES (1)

CF INDUSTRIES, INC. PLANT CITY PHOSPHATE COMPLEX HILLSBOROUGH COUNTY, FLORIDA

<u>POLLUTANT</u> Tons/Year)	<u>Sulfur</u>	Sulfuric Acid Plant C		
Tons/ rear)	<u> </u>			
02	1750	1750		
Present (actual) Proposed	1752 1898	1752 1898		
Annual Change	146	146		
Subtotal De minimis Increase	(2)	292 40		
ist Present (actual)	65.7	65.7		
Proposed	71.2	71.2		
Annual Change	5.5	5.5		
Subtotal De minimis Increase	(2)	11.0 7		
Ox Present (actual)	17.0	17.0		
Proposed`	18.4	18.4		
Annual Change	1.4	1.4		
Subtotal De minimis Increase	(2)	2.8 (3) 40		

- (1) Based on differences between present actual/permitted and proposed operating conditions.
- (2) Defined in 17-2.500(2)(e)2,FAC.
- (3) The emission rate increase of 2.8 tpy, when combined with NOx emission rate increases of 1.6 tpy and 8.3 tpy permitted in 1988 (and based on a stack gas concentration of 5 ppm, volume) is less than the de minimis emission rate increase of 40 tpy for NOx defined in 17-2.500(2)(e)2,FAC.