



**10 MW DEMONSTRATION
OF
GAS SUSPENSION ABSORPTION
FINAL PROJECT PERFORMANCE
AND
ECONOMICS REPORT**

**AirPol, Inc.
32 Henry Street
Teterboro, New Jersey 07608**

CLEAN COAL TECHNOLOGY III
10 MW DEMONSTRATION OF GAS SUSPENSION ABSORPTION
FINAL PROJECT PERFORMANCE AND ECONOMICS REPORT

PARTICIPANT

AirPol Inc.

Teterboro, New Jersey

Frank E. Hsu
Project Manager

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ABSTRACT

The 10 MW Demonstration of the Gas Suspension Absorption (GSA) program is a government and industry co-funded technology development. The objective of the project is to demonstrate the performance of the GSA system in treating a 10 MW slipstream of flue gas resulting from the combustion of a high sulfur coal. This project involves design, fabrication, construction and testing of the GSA system.

The Project Performance and Economics Report provides the nonproprietary information for the "10 MW Demonstration of the Gas Suspension Absorption (GSA) Project" installed at Tennessee Valley Authority's (TVA) Shawnee Power Station, Center for Emissions Research (CER) at Paducah, Kentucky.

The program demonstrated that the GSA flue-gas-desulfurization (FGD) technology is capable of achieving high SO₂ removal efficiencies (greater than 90%), while maintaining particulate emissions below the New Source Performance Standards (NSPS), without any negative environmental impact (section 6). A 28-day test demonstrated the reliability and operability of the GSA system during continuous operation. The test results and detailed discussions of the test data can be obtained from TVA's Final Report (Appendix A). The Air Toxics Report (Appendix B), prepared by Energy and Environmental Research Corporation (EERC) characterizes air toxic emissions of selected hazardous air pollutants (HAP) from the GSA process. The results of this testing show that the GSA system can substantially reduce the emission of these HAP. With its lower capital costs and maintenance costs (section 7), as compared to conventional semi-dry scrubbers, the GSA technology commands a high potential for further commercialization in the United States. For detailed information refer to The Economic Evaluation Report (Appendix C) prepared by Raytheon Engineers & Constructors.

POINT OF CONTACT

Frank E. Hsu
Project Manager
AirPol Inc.
32 Henry Street
Teterboro, New Jersey 07608
Tel. (201)-288-7070

Sharon K. Marchant
Project Manager
Pittsburgh Energy Technology Center
United States Department of Energy
Pittsburgh, Pennsylvania 15236
Tel. (412)-892-6008

Thomas A. Burnett
Project Specialist
Tennessee Valley Authority
Chattanooga, Tennessee 37402
Tel. (615)-751-3938

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LIST OF ABBREVIATIONS

ACFM	- actual cubic feet per minute
AST	- approach to saturation temperature
Btu	- British thermal unit
CCT	- Clean Coal Technology
CER	- Center for Emissions Research, Shawnee Fossil Plant
cf	- cubic feet
DOE	- Department of Energy
EERC	- Energy and Environmental Research Corporation
EPA	- Environmental Protection Agency
ESP	- electrostatic precipitator
FGD	- flue gas desulfurization
ft	- feet
GPM	- gallons per minute
gr	- grains
GSA	- Gas Suspension Absorption
HAP	- hazardous air pollutants
HCl	- hydrochloric acid
H ₂ O	- water
hr	- hour
in	- inch
kW	- kilowatts
lb	- pound
I.D.	- induced draft
MBtu	- million British thermal units
MW	- megawatt
NEPA	- National Environmental Policy Act
NSPS	- New Source Performance Standards
O ₂	- oxygen
PJBH	- pulsed jet baghouse
ppm	- parts per million by volume
PSIG	- pounds per square inch gauge
RCRA	- Resource Conservation and Recovery Act
SCF	- standard cubic feet
SCFM	- standard cubic feet per minute
SD	- spray dryer
SIP	- State Implementation Plan
SO ₂	- sulfur dioxide
TCLP	- toxic characteristic leaching procedure
TVA	- Tennessee Valley Authority
WG	- water gauge
WLFO	- wet limestone forced-oxidation

EXECUTIVE SUMMARY

The Clean Coal Technology Demonstration Program (CCT Program) is a government and industry co-funded technology development effort to demonstrate a new generation of innovative coal utilization processes in a series of full-scale, "showcase" facilities built across the country. These demonstrations are on a scale large enough to generate all the data, from design, construction, and operation, for technical/economic evaluation and future commercialization of the process.

The goal of the program is to furnish the U.S. energy marketplace with a number of advanced, more efficient, and environmentally responsive coal-using technologies. These technologies will reduce and/or eliminate the economic and environmental impediments that limit the full consideration of coal as a viable future energy resource.

To achieve this goal, a multi-phased effort consisting of five separate solicitations was administered by the Department of Energy (DOE). Projects selected through these solicitations have demonstrated technology options with the potential to meet the needs of energy markets and respond to relevant environmental considerations.

In response to these solicitations, AirPol Inc. with the assistance of the Tennessee Valley Authority (TVA), has demonstrated the Gas Suspension Absorption (GSA) technology in the Clean Coal Technology project entitled "10 MW Demonstration of Gas Suspension Absorption". AirPol performed this demonstration under a Cooperative Agreement awarded by DOE in October 1990. This project was selected in Round III of the Clean Coal Technology Program.

This low-cost retrofit project has demonstrated that the Gas Suspension Absorption technology can achieve more than 90% removal of the SO₂ from the flue gas on a coal-fired boiler application, while attaining a high utilization of lime reagent. The host site facility has been the Shawnee Fossil Plant, located at the TVA's Center for Emission Research (CER) in West Paducah, Kentucky.

The Gas Suspension Absorber was initially developed as a calciner for limestone in cement production. It has been used successfully to clean gases from commercial waste-to-energy plants in Denmark where it has also captured chloride emissions. Raw flue gas is provided to the GSA from Shawnee's unit 9 boiler which is configured to divert 7% of its total flue gas output to the GSA system. The diverted flue gas enters the bottom of the reactor and flows co-currently upwards with the recycled solids and fresh lime slurry. The lime slurry is fed into the reactor by means of a single spray nozzle. This spray nozzle is mounted in the throat of the reactor, such that the lime slurry is sprayed vertically upwards through the center of the reactor. The spray droplets, consisting of water and lime particles, coat the surface of the recycled solids, thus providing a medium with a large surface area for reaction. Acid gases such as SO₂ and HCl in the flue gas react with the lime particles on the surface of the recycled solids. At the same time, water evaporates from the surface of the solids, thereby simultaneously cooling the flue gas, and drying the solids. The dry solids, consisting of reaction products, namely, calcium sulfite,

calcium sulfate and calcium chloride, along with unreacted slaked lime and fly ash from the boiler are entrained in the flue gas and pass up through the reactor and into the cyclone. About 99% of the solids entering the cyclone are recycled back to the reactor via a feeder box, which provides temporary, in-process storage. This high concentration of solids being recycled through the reactor minimizes scaling due to its scouring effect on the reactor walls. Unused lime in the recycle solids can further react with acid gases in the flue gas, thus lowering the overall consumption of lime. The flue gas containing the remaining 1% of the solids leaving the cyclone enters an electrostatic precipitator (ESP) for final particulate collection. After passing through the ESP the cleaned flue gas is released to the atmosphere. The GSA system is designed to remove more than 90% SO₂ using high sulfur U.S. coal. Coal sulfur content during the demonstration ranged from 4 to 5 pounds of SO₂ per million Btu, or about 2.7 ~ 3.0 % sulfur by weight.

The GSA is distinguished from the average semi-dry scrubbing processes by its modest space requirement, simple means of introducing reagent to the reactor, direct means of recirculating unused lime, and low reagent consumption. The GSA system consists of the following major components:

- A gas suspension reactor with material recycle and lime slurry injection.
- A cyclone and feeder box for separating and recycling material back to the reactor.
- A dust collector which removes remaining fly ash and reaction products from the flue gas stream.
- A lime slurry preparation system.
- An ash storage and handling system.

In developing the general arrangement of the GSA system, design considerations were given to the following factors:

1. Minimizing material and construction cost by making the connecting duct system as compact as possible, while providing adequate gas flow pattern throughout the system.
2. Provide an enclosure for the most frequently serviced area of the GSA system. The enclosure will provide personnel protection in the injection lance area and the feeder box area, and shields the air sluice, slurry and water pipes from inclement weather.
3. The layout was designed to provide direct access to the lower operating area (injection nozzle level) and to save costs by utilizing the existing stair tower.

The GSA demonstration system was retrofitted to replace an existing spray dryer, which was used for testing prior to the GSA tests. The existing equipment that was suitable for the GSA system

was reused to minimize interface work and save equipment cost. The equipment reused includes the following:

- Air compressor
- Lime preparation system
- Electrostatic precipitator (ESP)
- Ash storage and handling system
- Motor control panel, which was modified to add additional circuit breakers.
- Foxboro computers and instrumentation

In view of the fact that the GSA outlet gas temperature is close to the saturation temperature of the flue gas, special design consideration was given in heating and insulation of the vessels and gas ducts to prevent condensation. Basically, all of the main equipment such as reactor, cyclone, feeder box and fabric filter as well as the ductwork were designed for external insulation with flat sheet aluminum lagging.

During normal operation, the GSA system is controlled by an automatic process control system which consists of three control loops: Recycled Solid Control Loop, Water Feed Rate Control Loop, and Lime Feed Rate Control Loop. The control system ensures that GSA works under maximum reagent utilization, proper reaction temperature, and minimum lime consumption.

The capital costs for the entire project were within the budget which was about \$7,717,200. As part of the DOE CCT Program, an economic evaluation of the GSA process was conducted using the same design and economic premises that were used to evaluate about 30 to 35 other FGD processes. The results show that the total capital requirements for the GSA process are substantially lower than those for the conventional wet limestone forced-oxidation (WLFO) scrubbing system (\$149/kw vs. \$216/kw). The substantially lower capital requirements are primarily due to the lower costs of the SO₂ absorption technology. Also, the levelized annual revenue requirements for the GSA process are 20 percent lower than those for the WLFO system.

All three of the major objectives of this demonstration were successfully achieved. Firstly, the GSA system demonstrated greater than 90 percent sulfur dioxide (SO₂) removal for a high-sulfur coal (i.e. greater than 4.5 lb SO₂/MBtu) application. Secondly, the emissions from the electrostatic precipitator (ESP) remained below the New Source Performance Standards for particulates (i.e. 0.03 lb/MBtu). Thirdly, the GSA plant demonstrated the reliability and operability of this technology by achieving 91 percent SO₂ removal during a 28-day period of continuous operation. It is obvious that this demonstration run truly fulfills the goal of the Clean Coal Technology Demonstration Program.

One of the objectives of this demonstration project is for AirPol to establish its capability in designing, fabricating, and constructing the GSA system so that the demonstrated technology can be effectively commercialized for the benefit of the U.S. electric utility and industrial markets. The progress of this demonstration project matches very well with the development of the utility FGD market. The GSA technology is now ready to be commercialized for the industry in order to meet the Phase II Clean Air Act Amendments compliance requirements.

1.0 INTRODUCTION

1.1 PURPOSE OF THE PROJECT PERFORMANCE AND ECONOMICS REPORT

The purpose of the Project Performance and Economics Report for "10 MW Demonstration of Gas Suspension Absorption" is to provide a technical account of the total work performed under the Cooperative Agreement awarded by U.S. Department of Energy (DOE). This report is based on recorded information from the demonstration run and various other tests with the incorporation of the modifications made during the test period. The report contains a comprehensive description of the results achieved, technical readiness, and participant's view and plan for commercialization of the technology demonstrated.

The scope of the report is limited to non-proprietary information. Its content is not sufficient to provide a complete tool in designing and constructing a replicate plant. This report, however, will serve as a reference for the design and application considerations involved in a commercial-scale facility. *The vital importance of this report is to present a practical application for this unique technology, Gas Suspension Absorption (GSA), to the public and provide a basic foundation for the upcoming commercialization usage.*

1.2 OVERVIEW OF THE PROJECT

1.2.1 Background and History of Project

This project is the first North American demonstration of the GSA system for flue gas desulfurization (FGD) for a coal-fired utility boiler. The purpose of the project was to demonstrate that the GSA system is able to remove more than 90% of the SO₂ from the flue gas, while achieving a high utilization of reagent lime.

The GSA process is a novel concept for FGD that was developed by AirPol's parent company, F.L. Smidth miljo a/s in Copenhagen, Denmark. The process was initially developed as a cyclone preheater system for cement kiln raw meal (limestone and clay). This innovative system provided both capital and energy savings by reducing the required length of the rotary kiln and lowering fuel consumption. The GSA system also showed superior heat and mass transfer characteristics and was subsequently used for the calcination of limestone, alumina, and dolomite. The GSA system for FGD applications was developed later by injecting lime slurry and the recycled solids into the bottom of the reactor to function as an acid gas absorber.

In 1985, a GSA pilot plant was built in Denmark to establish design parameters for SO₂ and hydrogen chloride (HCl) absorption for waste incineration applications. At this installation SO₂ and HCl removal efficiency were tested and found to be equal to or better than what could be obtained by competing processes. Shortly before that time, successful experimental results were obtained for SO₂ and HCl absorption in large laboratory scale experiments corresponding to a 2 MW unit. The engineering data gained from the bench and pilot testing were statistically treated by multiple linear regression analysis resulting in a formula, or mathematical model, predicting the performance of the GSA equipment as a function of the relevant process parameters. The first commercial GSA unit was installed at the KARA Waste-to-Energy Plant at Roskilde, Denmark, in 1988. Currently, there are more than a dozen GSA installations in Europe; most of them are municipal solid waste incinerator applications. The design criteria used for the 10 MW demonstration project were based upon the above mentioned engineering experience and data which were tailored and optimized to suit this particular application.

The three major types of fuels used in commercial and industrial boilers are natural gas, oil and coal, with the greatest quantity of pollutants being generated by high sulfur coal-fired boilers. Having reviewed and calculated emission levels from commercial/industrial boilers, EPA estimated that the uncontrolled emissions from boilers are projected to double the current amount by the year 2000. Presently all boilers with heat input capacity greater than 250 MW are subject to the New Source Performance Standards (NSPS). All smaller boilers are subject to State Implementation Plans (SIPs). The increased emphasis on SO₂ emission reduction by utility and industrial plants is required by the Clean Air Act Amendments of 1990. Consequently, a simple and economic FGD process, such as GSA, can be utilized as a viable alternative by the small to mid-size plants where a wet FGD system may not be feasible. Recognizing the vast potential market, AirPol was committed to additional research and development expenditures in the demonstration project in order to further develop the GSA technology for coal-fired boiler

application that will bring a significant share of this market in the 1990's.

1.2.2 Project Organization

This "10 MW Demonstration of Gas Suspension Absorption (GSA)" Program is a government and industry co-funded technology development project. The major participants in the execution of this project were AirPol Inc., DOE, TVA, and FLS miljo. The interrelationships among the team members are shown in Figure 1.2.2.

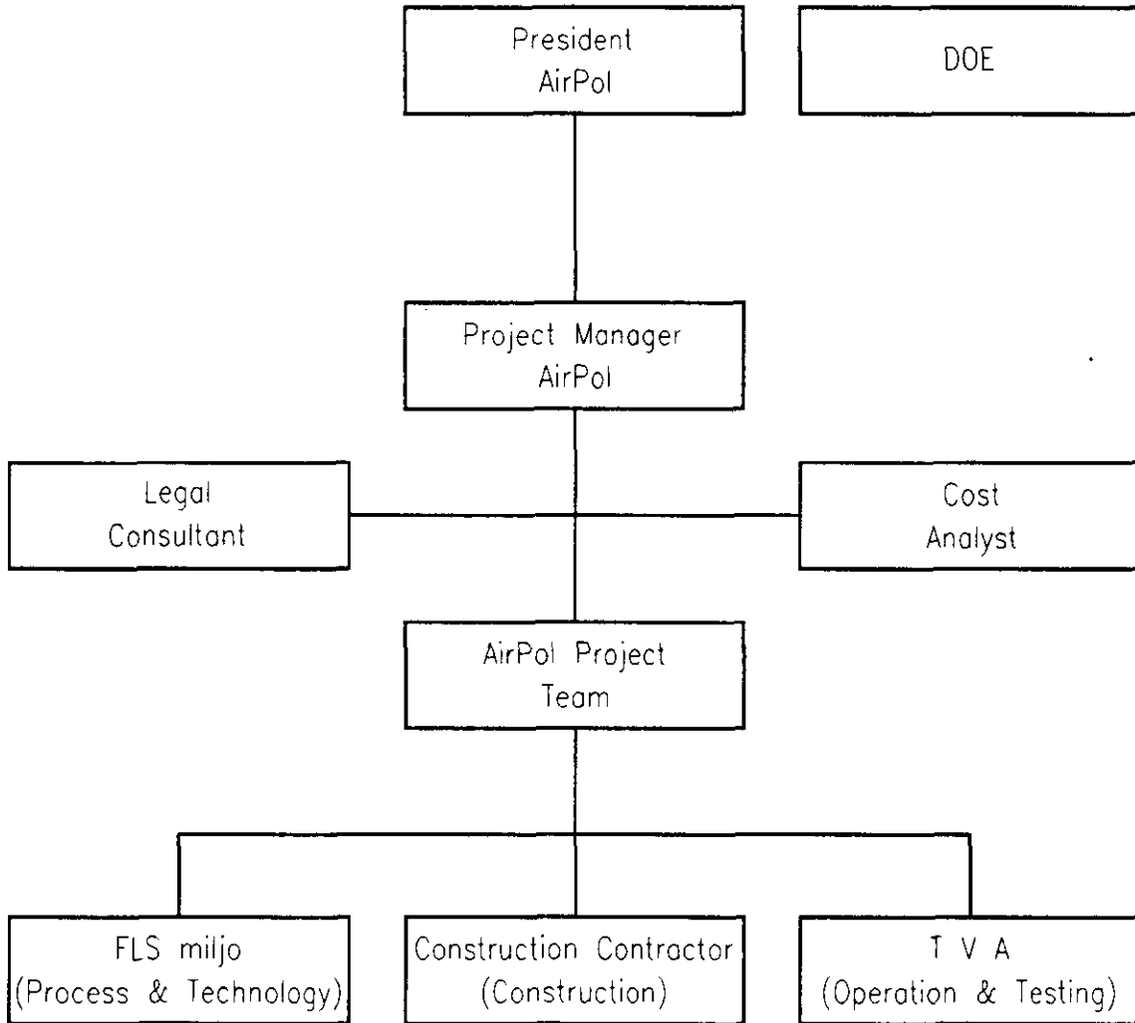
The DOE's Contracting Officer and Project Manager were responsible for all contract matters, technical liaison and monitoring of the project. AirPol took the lead in execution of this project, whereas TVA acted as a subcontractor to AirPol, provided the host site and all resources required for plant operation and testing services. FLS miljo, the GSA technology owner, provided technology transfer and technical assistance to AirPol on design, operation and testing of the demonstration system.

Throughout the course of this project, reports dealing with technical, cost and environmental aspects of the project were prepared by AirPol and provided to DOE. AirPol and TVA also prepared and published technical papers on the demonstrated technology, operating results, and its commercial advantages. The entire project funds were contributed by AirPol, DOE and TVA. The cost sharing contributions are listed as follows:

AirPol:	\$2,492,729
DOE:	\$2,315,259
TVA:	\$2,926,036
	<hr/>
	\$7,717,189

Figure 1.2.2

**PROJECT ORGANIZATION
10 MW DEMONSTRATION OF GSA**



1.2.3 Project Description

The innovative GSA process technology was developed and patented by F.L. Smidth miljo a/s in Copenhagen, Denmark in the 1980's for removing acid gases from the flue gases generated by many industrial processes. Presently, most of the GSA installations in Europe are municipal solid waste incinerator applications. The demonstration unit at TVA's CER is the first application of the GSA technology for flue gas desulfurization for a coal-fired boiler in U.S. The entire project involves design, fabrication, construction and testing of the GSA system. AirPol took the lead in execution of the project including in charging of engineering design, supervision of fabrication and construction, initial system start-up, and assistance in operation and testing activities.

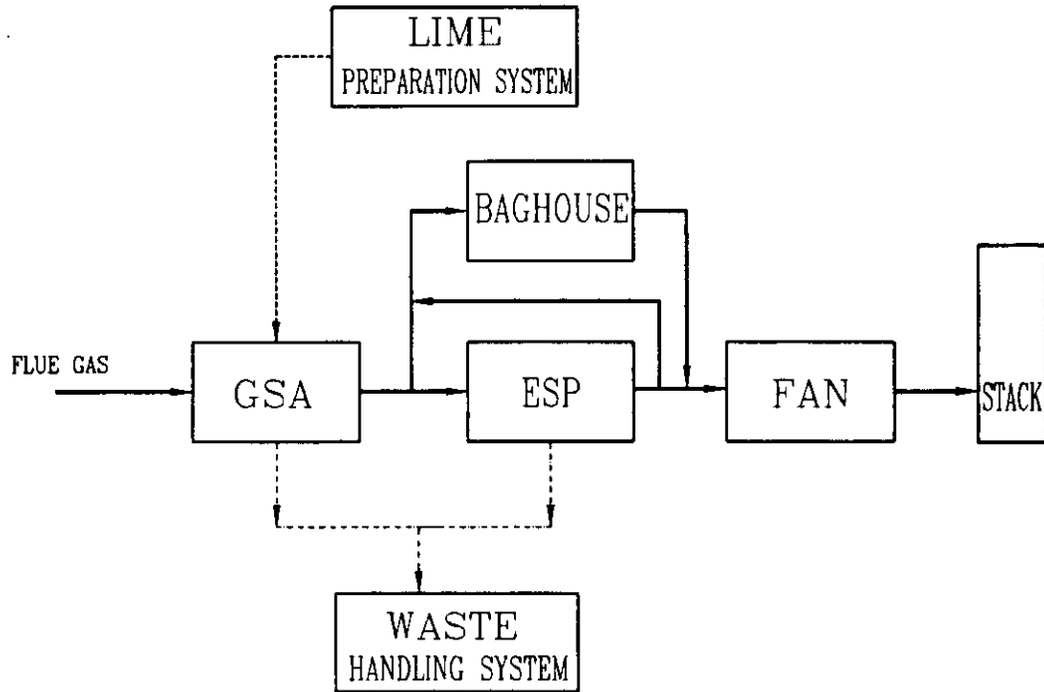
As AirPol's subcontractor, TVA took charge of the plant operation and testing services. At the beginning of the operation/testing phase a number of preliminary tests were conducted to determine the operating limits of the GSA demonstration system, and to define the relative importance of the various operating parameters. After the preliminary tests, a statistically-designed factorial test program was followed. The purpose of this factorial tests was to determine the effect of the process variables on the operation and SO₂ removal efficiency in the reactor/cyclone and the ESP/baghouse so as to optimize the GSA performance. The air toxics tests, which followed the factorial tests, were conducted to determine the capability of the GSA system in removing HCl, particulates and trace metals. A long-term, 28-day GSA demonstration run program was successfully completed after the air toxics tests. The objective of this program was to demonstrate that the GSA system (reactor/cyclone/ESP) could operate reliably and continuously, 24 hours/day, seven days/week, for a four-week period, while simultaneously achieving more than 90 percent SO₂ removal and maintaining the ESP emissions below the NSPS for particulates. The testing phase concluded with a 14-day pulsed jet baghouse (PJBH) continuous demonstration run of the GSA system. This test program was designed to confirm that the GSA system with an alternative arrangement, i.e. reactor/cyclone/baghouse, could still achieve the required 90+ percent removal efficiency for SO₂, and particulate emissions below NSPS.

The GSA system brings coal combustion gases into contact with a suspended mixture of solid, containing sulfur-absorbing lime and a spray of fresh lime slurry. After the lime absorbs the sulfur, the solids are separated from the gases in a cyclone and recycled back into the system where they capture additional sulfur. The cleaned flue gases are sent through a dust collector before being released into the atmosphere. The process block diagram for the demonstration project is shown in Figure 1.2.3.

The reagent required for the process of the GSA system can be either hydrated lime or quicklime slaked before feeding into the reactor. The major constituents of the by-product from the GSA process were inert material and calcium sulfite. Other components included calcium sulfate, calcium carbonate, excess calcium hydroxide, small quantities of calcium chloride and very little moisture, i.e., less than 1%. The average bulk density of the by-product was 66 lb/ft³. The texture of the material can be described as being similar to hour-glass-sand. The material is grey in color.

Figure 1.2.3

**PROCESS FLOW BLOCK DIAGRAM
GAS SUSPENSION ABSORPTION SYSTEM**



LEGEND:

- FLUE GAS
- - - - - LIME
- · - · - ASH/WASTE

1.2.4 Site

The project was conducted at the TVA's Center for Emissions Research (CER) located at Shawnee Fossil Plant, approximately 10 miles northwest of Paducah, in McCracken County, Kentucky. The plant is located on the south bank of the Ohio River on several hundred acres of river floodplain and a low upland terrace developed in thick deposits of unconsolidated clays, silts, and gravel. The active plant area is situated on this terrace, which lies above the 500-year floodplain. Over the past 15 years, the CER has served as a testground for FGD systems.

The Shawnee Fossil Plant currently operates 10 coal-fired boiler units with a total nameplate capacity of 1735 MW. Units 1-8 are fired with low-sulfur coal while units 9 and 10 are able to utilize a high-sulfur coal. Unit 9 supplies 7 % of its total flue gas to the GSA demonstration system. Units 1 through 9 are identical wall-fired Babcock and Wilcox boilers, each having a nameplate generating capacity of 175 MW, while unit 10 is a 160-MW Atmospheric Fluidized Bed Combustion boiler that was retrofitted in the 1980s.

Since the demonstration system is retrofitted into the existing test facility, the space requirement is only for the installation of the GSA reactor and cyclone. A space of 20 feet by 12 feet was used for the installation. Existing facilities such as the lime preparation system, stair tower, ESP, I.D. fan, and associated ductwork were reused for the GSA installation.

1.2.5 Project Schedule

The entire demonstration project was divided into three phases and tasks. AirPol began the design work on this project in November 1990, shortly after award of the Cooperative Agreement by DOE in October 1990. At the outset of the project, site access at the CER was delayed for one year by TVA to allow the completion of another project. That caused a one-year delay in this Clean Coal Technology project. The design phase of the GSA project was completed in December 1991. The fabrication and construction of the GSA unit was completed ahead of schedule in early September 1992. The planned operation and testing of the demonstration unit began in November 1992 and was completed in mid-March 1994. The final project schedule is presented in Table 1.2.5-1. As shown in Table 1.2.5-2 the GSA test program consists of the following five parts. The estimated total number of hours for each of these tests are also shown in the Table 1.2.5-2.

- The preliminary or start-up tests were conducted in November and December 1992.
- The factorial tests were performed between January and August 1993.
- The air toxics testing was finished between mid-September and mid-October 1993.
- The 28-day continuous GSA demonstration run was executed in late October and November 1993.

- The 14-day pulsed jet baghouse (PJBH) demonstration run was completed between late February and March 1994.

Table 1.2.5-1

PROJECT WORK BREAKDOWN STRUCTURE

PHASE	TASK	DESCRIPTIONS	TIME PERIOD
Phase I	ENGINEERING AND DESIGN		
	Task I	Project and Contract Management	11/01/90 - 12/31/91
	Task II	Process and Technology Design	11/01/90 - 09/30/91
	Task III	Environmental Analysis	11/01/90 - 09/30/91
	Task IV	Engineering Design	11/01/90 - 09/30/91
Phase II	PROCUREMENT AND CONSTRUCTION		
	Task I	Project and Contract Management	01/01/92 - 09/30/92
	Task II	Procurement and Supply of Material	01/01/92 - 04/30/92
	Task III	Construction and Commissioning	05/01/92 - 09/30/92
Phase III	OPERATION AND TESTING		
	Task I	Project Management	10/01/92 - 09/30/93
	Task II	Start-up and Training	10/01/92 - 10/14/92
	Task III	Experimental Testing and Reporting	10/15/92 - 10/31/94

Table 1.2.5-2

**TEST SCHEDULE
10 MW DEMONSTRATION OF GSA**

	1992			1993									1994					
	Oct	Nov	Dec	Jan	Feb	Mar	Apr	May	Jun	Jul	Aug	Sep	Oct	Nov	Dec	Jan	Feb	Mar
Preliminary Tests		<u>600 hr</u>																
Factorial Tests																		
-Basic Series				<u>1,000 hr</u>														
-Additional PJBH							<u>200 hr</u>											
-Replicate Series								<u>1,000 hr</u>										
Higher Sulfur Coal Tests											<u>500 hr</u>							
Air Toxics Tests												<u>400 hr</u>						
Demonstration Run													<u>672 hr</u>					
PJBH Demonstration Run																	<u>336 hr</u>	

1.3 OBJECTIVES OF THE PROJECT

The "10 MW GSA demonstration system" was installed and tested at the Tennessee Valley Authority the (TVA) Shawnee Fossil Plant at West Paducah, Kentucky. The new GSA system replaced the existing spray dryer that was installed previously as a test unit. The experience gained in designing, fabricating, and constructing the GSA equipment throughout the execution of this project will be used for future commercialization of the GSA system. Results of the operation and experimental testing will be used to further improve plant scale GSA design and operation.

Subsequent to optimization of the GSA system, air toxics testing was performed to determine the GSA's capability in removing hazardous air pollutants.

Along with the operation and testing of GSA, a 1 MW pulsed jet baghouse was tested to evaluate its long-term reliability and pollutant (both SO₂ and air toxics) removal performance. The baghouse was connected to the ESP to allow testing of the GSA system in one of three alternative arrangements: GSA with ESP only, GSA with baghouse only, and GSA with ESP followed by baghouse.

The specific technical objectives of the GSA demonstration project were to:

- Effectively demonstrate SO₂ removal performance in excess of 90% using a high-sulfur U.S. coal.
- Optimize recycle and design parameters to increase efficiencies of lime reagent utilization and SO₂ removal.
- Compare SO₂ removal efficiency and cost with existing Spray Dryer/ESP technology.
- To obtain data regarding the GSA's ability to remove air toxics from the waste gas stream (1) with ESP, (2) with baghouse, and (3) with baghouse following ESP.
- To compare the air toxics removal between a GSA with ESP and a GSA with baghouse.
- To compare the SO₂ removal between a GSA with ESP and a GSA with baghouse.
- To evaluate the merits of a baghouse following an ESP as a polishing unit in both SO₂ and air toxics removal.
- To evaluate the performance and long-term reliability of the baghouse in GSA applications.

1.4 SIGNIFICANCE OF THE PROJECT

The "10 MW Demonstration of Gas Suspension Absorption (GSA)" Project demonstrates the GSA's ability to effectively remove sulfur dioxide (SO₂) from coal-fired flue gas, while achieving a high utilization of reagent lime.

The GSA is distinguished from conventional semi-dry scrubbing processes by its modest space requirement, simple means of introducing reagent to the reactor, direct means of recirculating unused lime, and low reagent consumption.

The results from this demonstration project show that GSA is an effective, economic, and space efficient answer to the SO₂ removal performance needed by the U.S. utility industry. The importance and significance of this project is demonstrated by the following facts:

- The GSA system was retrofitted into an existing system with extremely tight available space which demonstrates the GSA's ability of being retrofitted into existing boiler systems.
- The GSA technology was successfully demonstrated at a coal-fired boiler plant with operating conditions that are typical of the average U.S. utility plant. Upon commercialization, this technology will have wide application to the utility industry.
- Commercialization of the GSA technology, as part of the objective of this project, will be carried out in time to meet the demand for FGD systems that meet the performance requirements required by the new Clean Air Act Amendments.
- The fact that a conventional spray dryer system has gone through similar tests on the very same boiler provides good comparison of the GSA with competing technologies.
- As part of the GSA demonstration program, a comparison between an electrostatic precipitator and baghouse was made. This comparison demonstrated GSA's flexible operation in conjunction with different types of dust collectors, and provided valuable information on the GSA performance with either type of dust collector.
- A special test program was conducted to determine GSA's ability in removing air toxics. This test program has indicated that the GSA system can address the air toxics problem faced by the U.S. utility industry.

1.5 DOE'S ROLE IN THE PROJECT

1.5.1 Innovative Clean Coal Technology Program

The Clean Coal Technology Demonstration Program (CCT Program) is a government and industry co-funded technology development effort to demonstrate a new generation of innovative coal utilization processes in a series of full-scale "showcase" facilities built across the country. These demonstrations will be on a scale large enough to generate all the data required for design, construction, and operation, and for technical/economic evaluation and future commercialization of the process.

The goal of the program is to furnish the U.S. energy marketplace with a number of advanced, more efficient, and environmentally responsive coal-using technologies. These technologies will reduce and/or eliminate the economic and environmental impediments that limit the full consideration of coal as a viable future energy resource.

To achieve this goal, a multi-phased effort consisting of five separate solicitations is administered by the Department of Energy. Projects selected through these solicitations will demonstrate technology options with the potential to meet the needs of energy markets and respond to relevant environmental considerations.

The third solicitation (CCT-III), issued in 1989, targeted those technologies capable of achieving significant reductions in the emission of SO₂ and/or NO_x from existing facilities to minimize environmental impacts, such as transboundary and interstate pollution, and/or provide for future energy needs in an environmentally acceptable manner.

In response to the third solicitation, AirPol Inc. submitted a proposal for the design, installation and testing of the GSA system at TVA's Center for Emission Research (CER). AirPol's proposal was selected as the result of DOE's evaluation in terms of technical advantage, cost effectiveness, technical readiness, and business and management performance potential. On July 25, 1990, a Cooperative Agreement was signed for the project entitled "10 MW Demonstration of Gas Suspension Absorption". The project was approved by Congress in October of 1990, and the Cooperative Agreement for the project was awarded by DOE on October 11, 1990.

1.5.2 Management Plan and Organization Chart

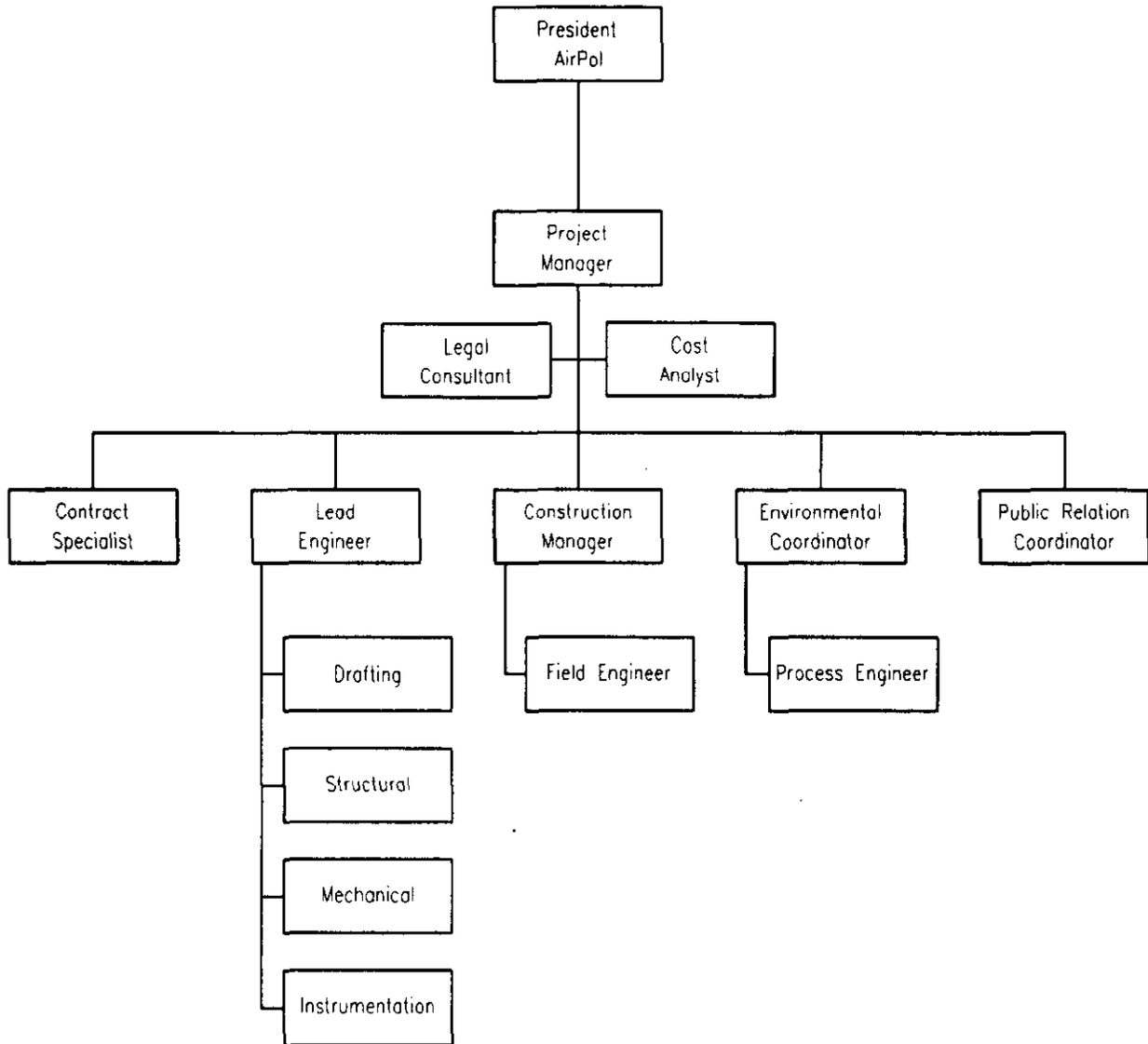
The DOE entered into the Cooperative Agreement with AirPol to conduct this project. The DOE was responsible for monitoring the project and all matters relating to the Cooperative Agreement through the DOE Contracting Officer.

AirPol Inc. was the leader, the center of communication and the major coordinator to all the parties participating in this project. AirPol's project execution team, as shown in Figure 1.5.2, consisted of the following members:

- **President - Manager in charge of the overall project, and the prime decision maker in all phases of the project.**
- **Project Manager - Responsible for the timely completion of all tasks required for the project. Acting as the focal point in steering the progress of the project, and in coordinating with DOE and TVA. Maintaining overall cost and schedule control of the project. Providing supervision and guidance to the project execution team. Coordinating with Purchasing Manager on all procurement tasks. Interfacing with the Process Specialist on all technological and environmental matters.**
- **Legal Consultant - Providing legal advice to the Project Manager.**
- **Contract Specialist - Responsible for all procurement tasks for the project. Compiling and disseminating project cost data to AirPol's President and the Project Manager for their review and analysis.**
- **Environmental Coordinator - Responsible for the preparation of all environmental information required by DOE to fulfill its obligation under National Environmental Policy Act (NEPA).**
- **Construction Manager - Responsible for the management and coordination of all construction and start-up related activities.**

Figure 1.5.2

**AIRPOL PROJECT ORGANIZATION
10 MW DEMONSTRATION OF GSA**



2.0 TECHNOLOGY DESCRIPTION

2.1 DESCRIPTION OF THE DEMONSTRATED TECHNOLOGY

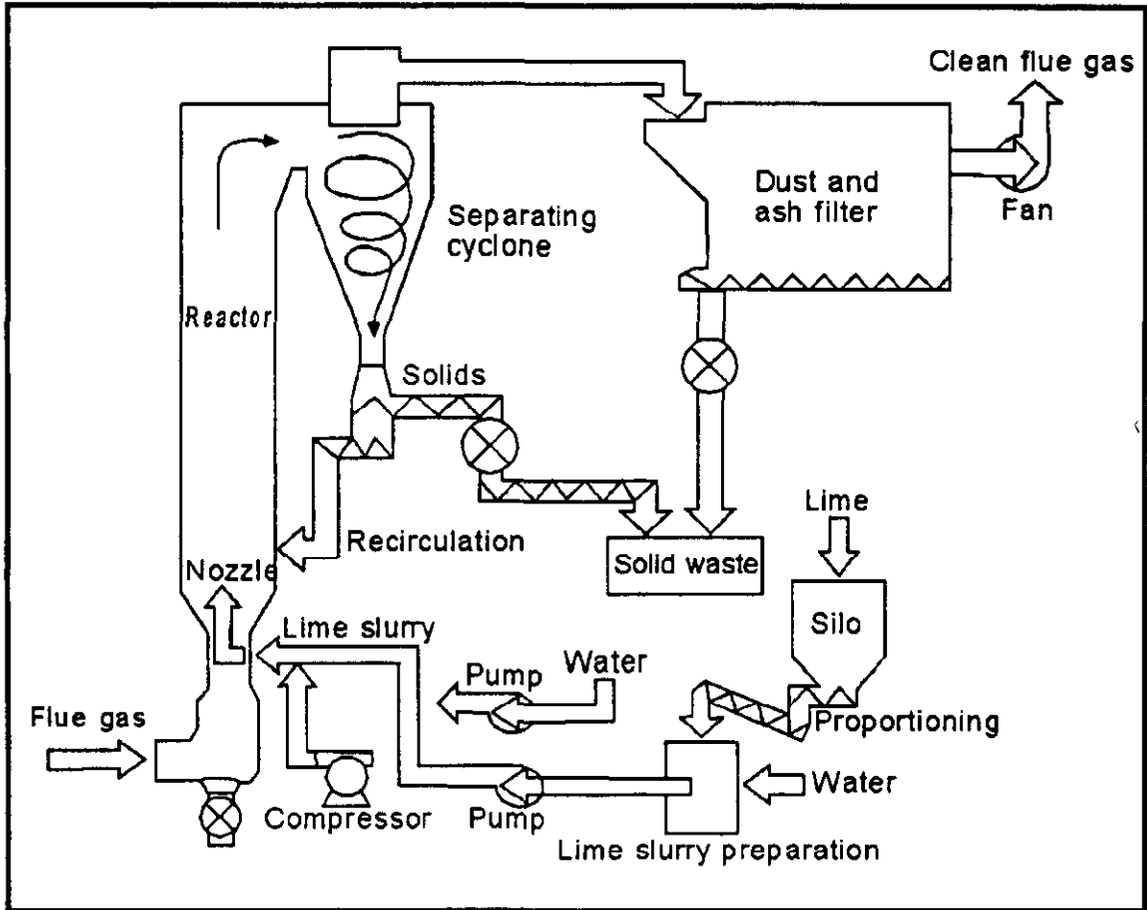
The primary objective for the installation of the GSA system at the TVA's CER was to demonstrate its ability to effectively remove sulfur dioxide (SO₂) from unconditioned flue gas. Raw flue gas was provided to the GSA from Shawnee's unit 9 which had been configured to divert 7% its total flue gas output to the GSA for the purpose of testing experimental scrubber technologies. A typical process flow diagram of Gas Suspension Absorption system is given in Figure 2.1. The key to the system's superior performance is the recirculation of solids. Typically, a solid particle will pass through the system about one hundred times before leaving the system. Another advantage of the GSA system is that a single spray nozzle is used to inject the fresh lime slurry. The only limitation for the GSA system is its 50 ~ 120% operating range. A minimum 50% of the designed gas capacity is required for keeping a desired gas flow velocity in the reactor and maintaining the solid particles suspended in the flue gas during chemical reactions. The major reason for setting the maximum flow limitation, 120% of the designed value, is that a sufficient gas retention time is needed in the reactor in order to allow a complete chemical reaction to take place between SO₂ and lime slurry, and to thoroughly dry the coating of lime slurry from the surface of the recycled solids to prevent clogging problems downstream. The GSA system is composed of four major process areas: SO₂ removal, dust collection, reagent preparation, and by-product handling.

2.1.1 SO₂ removal

The flue gas enters the bottom of the reactor where it is mixed with suspended solids and lime slurry which are being fed into the reactor by a single spray nozzle. The slurry is sprayed onto the recycled solids, which are suspended in the reactor by the flue gas. The lime in the slurry and the acid gas in the flue gas undergo a chemical reaction on the surface of the recycle solids. The partially cleaned flue gas leaves the reactor and enters a cyclone where the solids containing the calcium salts, ash, and unreacted lime are separated from the gas stream. About 99% of the solids collected in the cyclone are recycled back to the reactor. In this fashion any unused lime can further react with acid in the flue gas. This lowers the overall consumption of lime. The remaining 1% of the solids in the flue gas leaving the cyclone enters an ESP for final particulate collection before being discharged through the stack.

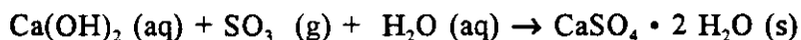
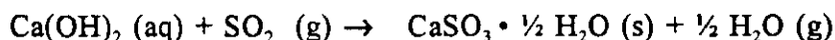
Figure 2.1

**PROCESS FLOW DIAGRAM
GAS SUSPENSION ABSORPTION SYSTEM**

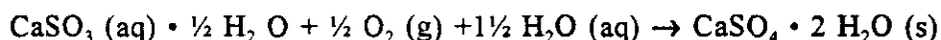
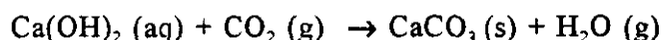


Process Chemistry

The primary overall reactions in the GSA system are:



The calcium hydroxide reacts with sulfur dioxide and sulfur trioxide to form calcium sulfite and calcium sulfate, respectively. In addition to the primary reactions the following secondary reactions also take place. Hydrogen chloride is removed and carbon dioxide is converted into calcium carbonate and water.



The major process design variables that affected the SO₂ removal are: stoichiometric ratio (Ca/S level), approach-to-saturation temperature, coal chloride level, flue gas flow rate, and recycle screw speed. Improvement of SO₂ removal performance in the GSA system can be achieved by increasing the stoichiometric ratio, increasing the coal chloride level, or lowering the approach-to-saturation temperature. The recycle feed rate and the flue gas flow rate individually have a minor effect on the SO₂ removal efficiency in the GSA system. These two variables have opposite effects on the SO₂ removal efficiency, i.e. increasing the recycle screw speed or decreasing the flue gas flow rate results in higher SO₂ removal efficiencies. The inlet fly ash loading also has a minor negative effect on SO₂ removal efficiency.

Reactor

The reactor plays a key role in the entire GSA system. The reactor vessel is of an up-flow fluidized design for handling of recirculated reaction products and fresh lime slurry. The reactor is constructed of carbon steel (ASTM A-36) and welded following AWS procedures. The unit is designed to incorporate 4 inches of insulation around the entire assembly. The unit also includes rod-out ports, access doors and a single view port.

The reactor contains a single dual-fluid nozzle designed to avoid plugging, erosion, and solids buildup. The nozzle is designed to permit the flow of lime slurry, water, and high pressure air for spraying the liquids. The design allows for removal and replacement of the complete nozzle assembly within 5 minutes without shutting down the system. One spare nozzle assembly is provided. The nozzle incorporates a replaceable orifice for replacement under routine maintenance.

Cyclone

The separating cyclone is sized to handle the cooled flue gas flow with high solids concentration from the reactor. Approximately 99% of the dry solids are collected by the cyclone and discharged to the recycle feeder box. Most of the collected solids will be recycled to the reactor. A small portion will leave the feeder box as by-product for disposal. The remaining 1% of the solids in the flue gas leaving the cyclone will be collected in the dust collector downstream as by-product. The cyclone is constructed of carbon steel (ASTM A-36) with an inspection door (outlet) and rod-out ports for servicing. The bottom cone of the cyclone discharges directly into the recycle feeder box.

The reactor/cyclone transition and cyclone outlet incorporates a fabric expansion joint located between flanges.

Recycle Feeder Box

Since a large portion of the solids collected in the cyclone are still reactive, a specially designed recirculating feeder box recycles approximately 99% of the solids back to the reactor via a multiple screw conveyor, while the remaining solids leave the system in the form of by-products, which consist of calcium sulfite, calcium sulfate, unreacted lime, and calcium chloride.

The feed system incorporates a triple screw conveyor with variable speed drive to discharge collected material back to the reactor and an overflow screw conveyor to discharge excess by-product for disposal.

The feeder box is designed for an external insulation with flat sheet aluminum lagging. All bearings are located outside of the insulation. Access doors are double wall, insulated type, with viewports. A ladder for access to the top of the unit is provided as part of the feeder box assembly. The lower portion of the feeder box is furnished with thermostatically-controlled electric heaters to prevent build-up.

2.1.2 Dust Collection

The flue gas leaves the separating cyclone and enters an electrostatic precipitator (ESP) for particulate collection. After passing through the ESP the cleaned flue gas is released to the atmosphere through a separate stack. The GSA system was designed to remove more than 90% SO₂ in flue gases from high sulfur coal. Coal sulfur content during the demonstration ranged from 4 to 5 pounds of SO₂ per million Btu, about 2.7 ~ 3.0% sulfur by weight.

Electrostatic Precipitator (ESP)

The process of particulate removal by the electrostatic precipitator (ESP) involves (1) the ionization of particle-laden gas flowing between electrodes, (2) the charging, migration, and collection of the solid particles on oppositely charged plates, and (3) the removal of the particles

from the plates.

The ESP installed at the CER is a relatively modern, four-field unit with 10 inch plate spacing and eight (8) parallel gas passages. The specific collection area of the ESP is about 440 ft²/kacfm under the cooled, humidified flue gas conditions. The aspect ratio is 1.6. Each field has a separate hopper and double-flap discharge valve for solids storage and removal. Both discharge and collector electrodes are rapped by tumbling hammers on a rotating shaft. A microprocessor-based system controls the voltage to the transformer-rectifier and sets the required rapping sequence.

Baghouse

A 1 MW pulsed jet type baghouse was incorporated to evaluate its long-term reliability and pollutant (SO₂, air toxic, and particulates) removal performance. The baghouse was connected to the ESP to allow testing of the GSA system in one of three alternative arrangements: GSA with ESP only, GSA with baghouse only, and GSA with ESP followed by baghouse.

Dust-laden gas enters the inlet manifold of the baghouse where a poppet valve system directs the flow to hopper areas under the zones or compartments. The dust-laden gas travels upward uniformly around the filter bags suspended above the hoppers in the zone. As the gas is drawn through the cage-supported filter bags, dust particles collect on the outside of the bags. Clean filtered gas continues upward inside the bags to the clean gas plenum section. The cleaned gas then passes through a poppet valve into the outlet exhaust manifold.

This baghouse was designed for an air to cloth ratio of 4.0 to 1. The bag material is Dralon-T used over a steel cage.

2.1.3 Reagent Preparation

The lime is delivered by trucks in the form of pebble lime (CaO), and pneumatically unloaded to a lime storage silo. The silo has a capacity of 107 tons of pebble lime, and is complete with the necessary level indication, bin vent filter, volumetric feeder, fill pipe, controls and instrumentation, caged access ladders, handrails and inspection doors for servicing. The lime is metered from the lime silo as needed to a detention type lime slaker. The lime and water rates to the slaker are proportioned to produce a smooth hydrated milk-of-lime slurry, of desired consistency. The lime slurry from the slaker is pumped to a storage tank. As needed in the process, the lime slurry is further pumped to the lime slurry feed tank. The slaked lime slurry, Ca(OH)₂, is then pumped to the dual-fluid nozzle at the base of the reactor.

2.1.4 By-Product Handling

By-products from the feeder box overflow and ESP/baghouse hoppers are discharged into the bucket elevator by screw conveyors. The bucket elevator conveys the material to a by-product storage silo. The solids in the silo are fed to the recycle mix tank with a capacity of 5,200 gallon

where the solids are mixed with water to form a 10% slurry. The resulting diluted slurry is pumped to the ash pond for ultimate disposal with the fly ash and bottom ash from the boilers.

2.2 DESCRIPTION OF THE DEMONSTRATED FACILITIES

Minimal ground space requirement is one of the salient features of the GSA system. As shown in Figure 2.1, the GSA system consists of:

- A fluidized reactor.
- A separating cyclone and feeder box for recycling material to the reactor.
- A dust collector removing particulates from the cleaned gas.
- A lime slurry preparation system.
- An ash storage and handling system.

Some existing equipment was reused to minimize interface work and save equipment cost. The equipment reused included the following:

- Air compressor.
- Lime slurry preparation system.
- Electrostatic precipitator (ESP).
- Ash handling system.
- Motor control panel modified to add additional circuit breakers.
- Instrumentation: Inlet and outlet gas flow indication; inlet and outlet flue gas SO₂ and O₂ monitors; temperature indication at the GSA inlet, cyclone outlet and ESP outlet; pressure indication at GSA inlet, ESP inlet and ESP outlet; lime slurry flow indication.

The existing Foxboro Control was used for the GSA system process control and data acquisition, but the start-up and shut-down sequence was manually performed. This is consistent with the previous operation of the spray dryer system and prefers by TVA operating/testing unit. The Foxboro Control was re-programmed to perform the GSA process control and monitoring as well as alarm annunciations. For more detailed information about system facilities please refer to AirPol's Public Design Report of section 3.5.

2.3 PROPRIETARY INFORMATION

The following is a list of the proprietary items:

- Dimensions of the fluidized reactor, cyclone, feeder box, ESP and baghouse.
- Detailed inner structure/configuration of the equipment.
- Critical process and equipment design parameters.
- Critical operation control logic and set points.
- Piping and instrument diagrams.
- Mathematical formula and correlations for SO₂ and particulate removal.

2.4 SIMPLIFIED PROCESS FLOW DIAGRAM

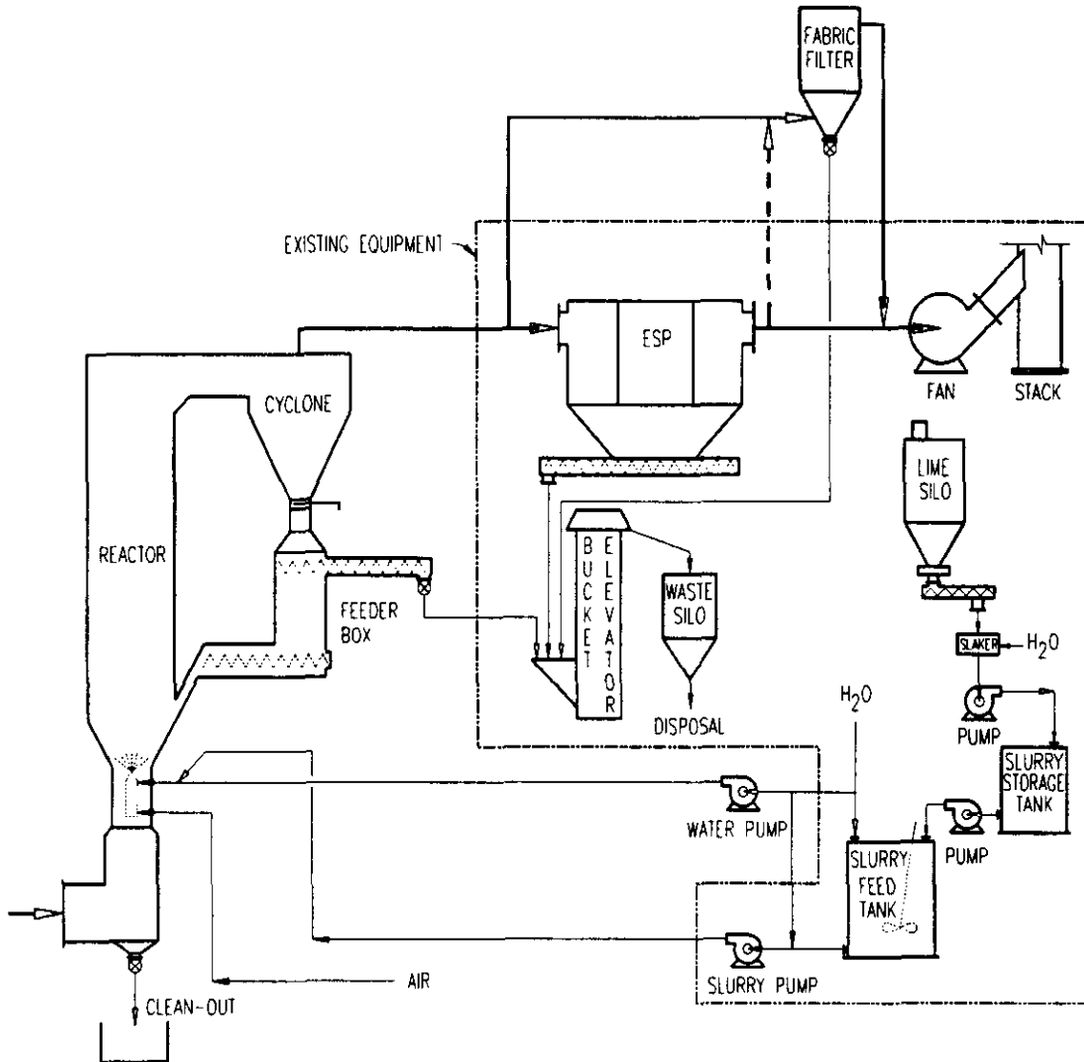
The Process Flow Diagram of "10 MW Demonstration of Gas Suspension Absorption (GSA) System" is shown in Figure 2.4. The figure shows how the GSA system was integrated with the equipment that could be utilized from the existing spray dryer (SD) test unit. Thus the electrostatic precipitator (ESP), the lime preparation system, and the ash storage and handling system were made part of the GSA test unit. The existing equipment is shown in Figure 2.4 within the limits of the phantom line.

The ductwork from the boiler to the spray dryer was re-routed to the inlet of the GSA reactor. The ductwork from the SD to the ESP was disconnected and a new duct connecting the GSA cyclone outlet with the ESP was installed.

No major problems were encountered during the installation of the GSA unit. Some minor field modifications of the equipment were made to accommodate the building steel structure, and the existing access platforms due to the very congested site conditions.

Figure 2.4

**PROCESS FLOW DIAGRAM
10 MW DEMONSTRATION OF GSA**



2.5 STREAM DATA

Table 2.5-1 shows data for three calculated design cases as well as the operating conditions for the 28-day continuous demonstration run. The GSA system was designed for an operating range from "Minimum Case" to "Maximum Case" with the actual operating condition expected to be as shown in the "Design Case" column. These operating parameters are close to the levels that can be expected for a GSA unit applied to a "normal" wall fired utility boiler burning pulverized high-sulfur coal and equipped with a Ljungstrom type air preheater.

The Operating Case proved to be very close to the Design Case with the exception of the Approach-to-Saturation-Temperature that was selected at 18 °F based upon preliminary test results. The removal efficiencies for both particulate and SO₂ exceeded the design data.

The lower portion of Table 2.5-1 shows the design and operating data for the condition where the baghouse was connected in series with the ESP. The design removal efficiencies for both particulate and SO₂ were also exceeded with this arrangement.

Figure 2.5 and Table 2.5-2 together presents the actual stream data for the "Design Case" including lime slurry, cooling water, compressed air for slurry atomization, and amount of by-product generated. Please refer to the AirPol's Public Design Report for detailed calculations of the design cases, and TVA's final technical reports for performance and operating results.

Table 2.5-1

SUMMARY TABLE FOR GSA PROCESS DATA

ITEM	MINIMUM CASE	DESIGN CASE	MAXIMUM CASE	OPERATING CASE
Gas Volume to System	12,600 SCFM	21,000 SCFM	23,100 SCFM	18,000~20,000 SCFM
Inlet Gas Temp.	320 °F	320 °F	320 °F	320 °F
AST Temp.	35 °F	35 °F	35 °F	18 °F
SO ₂ Removal Rate (%)	91.98	91.98	91.98	90.2~93.4
Particulate Removal Rate (%)	99.58	99.58	99.58	99.81~99.98
Baghouse in Series with ESP				
Gas Volume to System	12,600 SCFM	21,000 SCFM	23,100 SCFM	18,000~20,000 SCFM
Inlet Gas Temp.	320 °F	320 °F	320 °F	320 °F
AST Temp.	35 °F	35 °F	35 °F	18 °F
SO ₂ Removal Rate (%)	93.96	93.96	93.96	96.1~99.0
Particulate Removal Rate (%)	99.68	99.68	99.68	99.97~99.99

Table 2.5-2

PROCESS DATA FOR DESIGN CASE

Item	GSA Gas Inlet G1	ESP Gas Inlet G2	ESP Gas Outlet G3	Baghouse Gas Inlet G3a	Baghouse Gas Outlet G3b
Flow Rate (acfm)	35,300	30,770	30,716	5,000	5,282
Temperature (deg F)	320	161	158	158	152
Pressure (in WG)	-18	-26.5	-27.9	-27.9	-33.9
O ₂ (Vol %)	7.12	7.18	7.18	7.18	7.92
H ₂ O (Vol %)	7.66	13.69	13.7	13.7	13.7
SO ₂ (ppm)	1873	297	150	150	14
HCl (ppm)	19	0	0	0	0
Dust (gr/dscf)	2.806	2.035	0.012	0.012	0.000

Table 2.5-2 (Continued)

PROCESS DATA FOR DESIGN CASE

Item	Stack Gas Inlet G4	Stack Gas Outlet G5	GSA Atomiz. Air A1	PJBH In leakage Air A2
Flow Rate (acfm)	29,470	29,474	8.2	202.2
Temperature (deg F)	173	172	80	80
Pressure (in WG)	0.5	0.0	2767.8	0
O ₂ (Vol %)	7.3	7.3	-	-
H ₂ O (Vol %)	13.59	13.59	-	-
SO ₂ (ppm)	127	127	-	-
HCl (ppm)	0	0	-	-
Dust (gr/dscf)	0.010	0.010	-	-

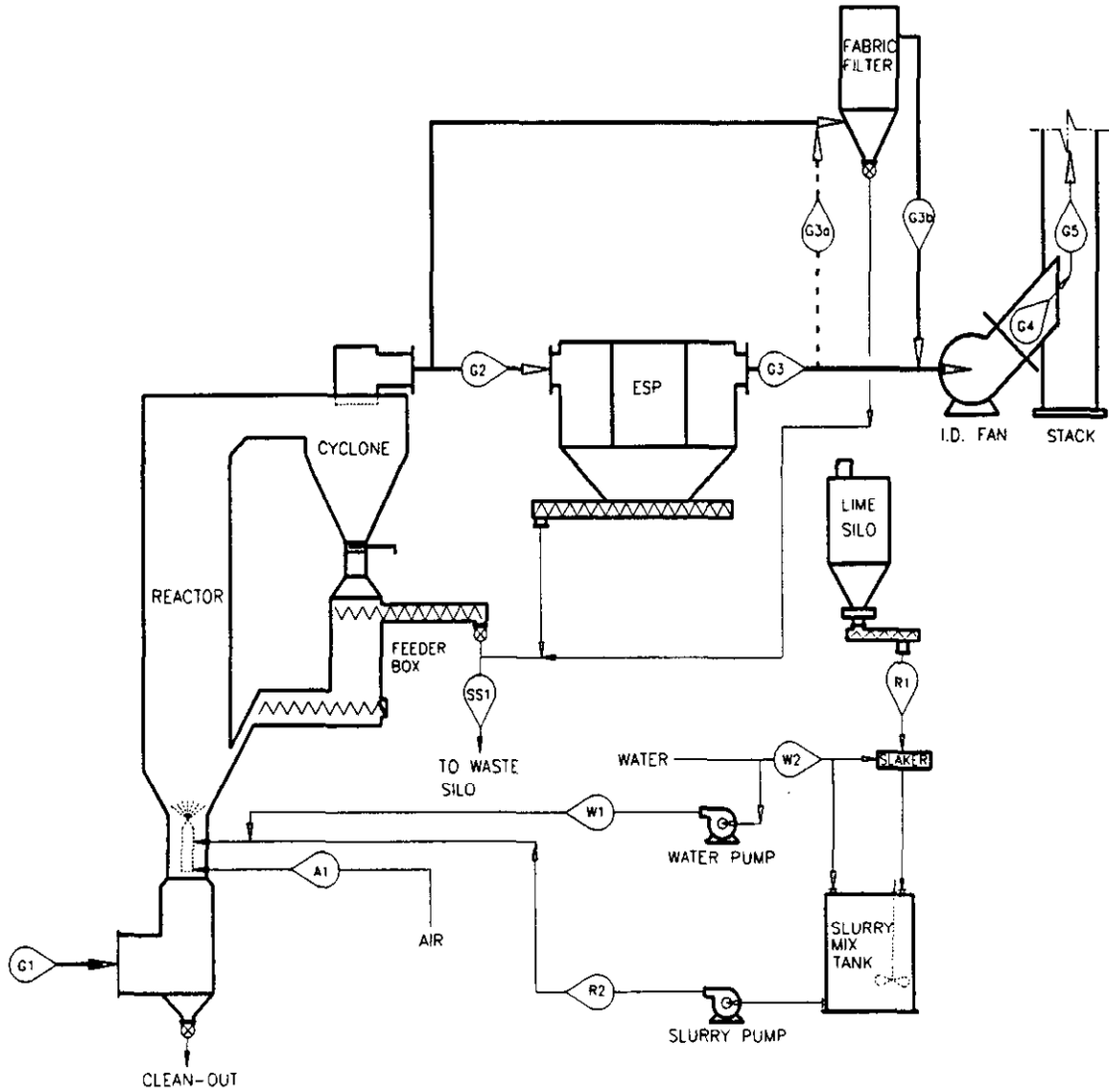
Table 2.5-2 (Continued)

PROCESS DATA FOR DESIGN CASE

Item	Pebble Lime Rate R1	Lime Slurry to GSA R2	GSA/PJBH Solid Waste SS1	Cooling Water to GSA W1	Lime Slurry Water W2
Flow Rate (lb/hr)	813.2	5241.4	1929.7	132.6	4193.2
Temperature (deg F)	-	78	-	68	68

Figure 2.5

**PROCESS CALCULATION DIAGRAM
10 MW DEMONSTRATION OF GSA**



2.6 PROCESS AND INSTRUMENTATION DIAGRAMS

There are three major control loops in this GSA control system as shown in the process control schematic diagram, Figure 2.6.

The first control loop, Recycled Solid Control, administers the flow of recycled solids to the reactor based on the amount of flue gas entering the system.

The second control loop, Feed Water Rate Control, adjusts the speed of the water pump and ensures that the flue gas is cooled sufficiently to optimize the chemical process. The amount of water added into the system is governed by the temperature of the flue gas exiting the cyclone.

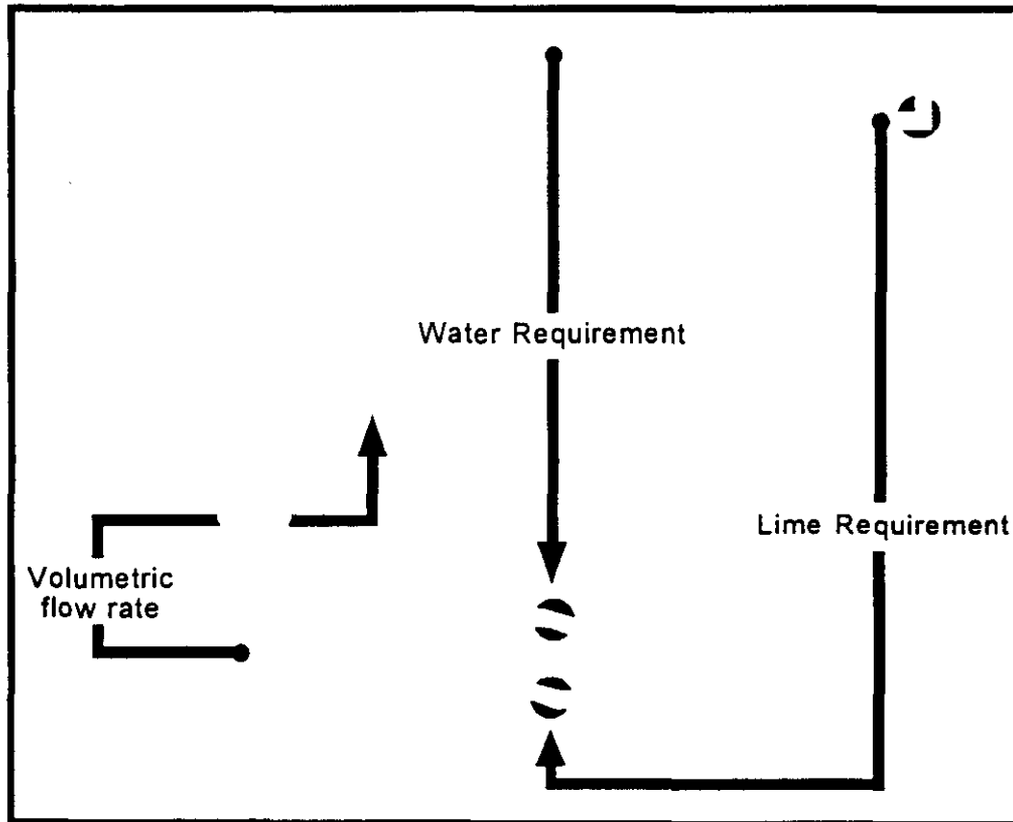
The third control loop, Lime Feed Rate Control, governs lime addition. This is accomplished by continuously monitoring the acid gas content in the outlet flue gas and comparing it with the required emission level. This control loop enables direct proportioning of the lime feed according to monitored results and further contributes to maintaining a low level of lime consumption.

The control parameters are established during initial start-up and can be adjusted during normal operation according to changes in operating conditions.

Any failure of mechanical or process equipment, such as pumps, motors, air compressors, etc. during operation, will be annunciated in the control room.

Figure 2.6

**PROCESS CONTROL SCHEMATIC DIAGRAM
GAS SUSPENSION ABSORPTION SYSTEM**



3.0 UPDATE OF THE PUBLIC DESIGN REPORT

Since no changes have been made to the design after the issuance of the Public Design Report, there is no update to the Public Design Report.

4.0 DEMONSTRATION PROGRAM

4.1 OPERATING PROCEDURES

During start-up of the GSA system the equipment in the system is started up in a sequential order as follows:

1. Plant ash conveying system
2. GSA ash conveying system
3. GSA heating system
4. ESP
5. I.D. fan
6. Solid recirculation system
7. Air compressor
8. Water pump
9. Lime slurry pump

During initial start-up, the system is "primed" by adding sand or fly ash to the feeder box. In addition, the boiler flue gas is allowed to flow through the GSA to enable some of the fly-ash to be collected by the cyclone to help fill the feeder box. Once the feeder box is 40% full, and a minimum flue gas flow rate has been achieved, the feeder box screw conveyor is started. At TVA, the solids recirculation rate was controlled by setting the speed on screw conveyor, although the option of using the first control loop also existed. With commercial applications the solids injection rate will be controlled by the first control loop as discussed in section 2.6. The air compressor, water pump and lime pump are started, in this sequence. The automatic control loops are now activated. The desired cyclone outlet temperature is maintained by controlling the speed on the water pump, and the SO₂ controls the lime slurry flow rate. At TVA, the third control loop, that controls the lime slurry flow to the GSA by adjusting the speed on the lime slurry pump, was set up with three options. The lime slurry flow could be adjusted according to the required:

1. SO₂ removal efficiency
2. Stoichiometric ratio
3. SO₂ emission at the stack. This option will be used on commercial GSAs.

When shutting down the GSA system, the start-up sequence is reversed. The lime slurry pump is stopped first and the slurry pipe is flushed with water. The water pump is then stopped. *By this time the nozzle will also be flushed out. The air compressor is stopped next. Recirculation of the solids continues for some time. This will ensure that the material is dry and warm. The slide gate under the cyclone is closed and the recirculated solids are allowed to collect in the cyclone hopper. The feeder box screw conveyor continues to run until the feeder box is empty. By collecting the solids in the cyclone during shut-down, the initial start-up procedure of accumulating the solids can be eliminated. The heating elements on the cyclone hopper will maintain a preset minimum temperature to prevent the material from caking. The heaters for the*

solid storage and handling system shall also remain on, until the system is completely emptied. A summary table of the simplified operating procedures are shown in Table 4.1-1.

Table 4.1-2 is an operator checklist for the major equipment and instrumentation on the demonstration unit. Most of the major instruments, including indicators, controllers, and alarms can be monitored and controlled from the control room.

Table 4.1-1

SIMPLIFIED OPERATING PROCEDURES

No.	Start-Up Procedure
1.	Plant Ash Conveying System
2.	GSA Ash Conveying System
3.	GSA Heating System
4.	ESP
5.	I.D. Fan
6.	Feeder Box Screw Conveyor
7.	Air Compressor
8.	Water Pump
9.	Lime Slurry Pump
No.	Shut-Down Procedure
1.	Lime Slurry Pump
2.	Water Pump
3.	Air Compressor
4.	Feeder Box Screw Conveyor
5.	I.D. Fan
6.	ESP
7.	GSA Ash Conveying System
8.	Plant Ash Conveying System
9.	GSA Heating System

Table 4.1-2

OPERATORS CHECKLIST

No.	Instrument	Instrument Location	Description
Reactor			
1.	Wet bulb thermometer	Reactor inlet duct	Measure gas saturation temp.
2.	Diff. press. transmitter	Reactor throat & vessel	Measure ΔP
Feeder Box			
1.	Screw speed control	F/B screw conveyor	Control solid reinjection rate
2.	Temp. indicator	Feeder box shell	Maintain feeder box temp.
3.	Speed sensor	F/B overflow screw	Indicate status of overflow screw
4.	Weight sensor	Feeder box bottom	Measure wt of material in F/B
Baghouse			
1.	Temp. indicator/controller	B/H inlet duct	Control water pump.
2.	Pulse sequence timer	Baghouse	Control air pulse rate & frequency
3.	Level indicator	Baghouse hopper	Control discharge valve
4.	Vibrator switch	Baghouse hopper	Monitor vibrator status
I. D. Fan			
1.	Speed sensor	ID fan	Indicate fan operation status
2.	Temp. indicator	ID fan motor	Monitor motor bearing temp.
3.	Vibration sensor	ID fan	Monitor fan vibration
4.	Damper position	ID fan	Monitor position of inlet damper

Table 4.1-2 (Continued)

OPERATORS CHECKLIST

No.	Instrument	Instrument Location	Description
Water Pump			
1.	Liquid flow indicator	Water pump discharge	Measure water rate to GSA
2.	Liquid press. indicator	Water pump discharge	Monitor water pressure
3.	Speed control	Motor on W/P	Controlled by B/H inlet temp.
Lime Preparation System			
1.	Liquid flow indicator	Lime slurry pump discharge	Measure lime slurry rate to GSA
2.	Liquid press. indicator	Lime slurry pump discharge	Monitor lime slurry pressure
3.	Speed control	Motor on L/P	Controlled by SO ₂ removal eff.
4.	Level indicator	Lime silo	Measure level of lime in silo
5.	Feed controller	Lime silo	Control silo discharge rate
6.	Liquid press. indicator	Slaker, water line	Monitor water press. to slaker
7.	Liquid flow indicator	Slaker, water line	Control lime/water ratio
8.	Temperature indicator	Slaker	Control emergency water to slaker
9.	Motion switch	Slaker agitator	Indicate agitator stopped
10.	Liquid level indicator	Lime slurry tank	Measure level of lime slurry
11.	Motion switch	Lime slurry tank agitator	Indicate agitator stopped

Table 4.1-2 (Continued)

OPERATORS CHECKLIST

No.	Instrument	Instrument Location	Description
Ash Storage and Handling System			
1.	Level indicator	Ash silo	Measure level of ash in silo
2.	Feed controller	Ash silo	Control silo discharge rate
3.	Motion switch	Screw conveyor	Indicate conveyor stopped
Others			
1.	Air press. indicator	Nozzle	Monitor air press. at nozzle
2.	Liquid press. indicator	Water line at nozzle	Maintain water press < air press.

4.1.1 Instrumentation and Data Acquisition

Instrumentation that is critical to the proper operation and control of the GSA process is basically composed of three (3) loops. The control of solid material recycle rate is the first control loop. A differential pressure indicator mounted in the reactor venturi throat reflects the inlet flue gas flow rate. The rate of solid recirculation back to the reactor is controlled in direct proportion to this prevailing inlet gas flow rate by adjusting the speed of the metering screws in the feeder box (via the variable frequency drive motor). A low flue gas flow rate alarm is provided to alert the operator. Should the gas flow rate drop to below 50% of the design value, a low-low flow alarm will be actuated, and the feeder box screw conveyor will trip. The feeder box screw conveyor is interlocked with the lime slurry and water pumps. Tripping of the feeder box screw conveyor will automatically stop the lime slurry and water pumps. At TVA, however, the flue gas flow rate entering the GSA is almost constant, hence it was decided to control the solids recirculation rate by setting the speed on the feeder box screw conveyor.

The second control loop is the flue gas temperature in the reactor. A temperature Indicator/Controller, located downstream of the cyclone, controls the flue gas temperature by adjusting the variable frequency drive on the motor of the pump supplying cooling water to the nozzle in the reactor. This controller also directly reflects and controls the chemical reactions taking place in the reactor. A high temperature alarm is furnished to notice a potential emergency condition. Also a low temperature alarm is provided to notify the possible condensation of moisture from the flue gas, and resulting acid gas corrosion, within the downstream equipment.

Acid gas emission control is the third critical loop among the entire GSA system control. This is accomplished by an acid gas emission Monitor/Controller in the exhaust stack and sending signal to control the speed of the variable frequency drive on the motor of the pump supplying lime slurry to the nozzle of the reactor. This simple control system will automatically maintain the required level of acid gas emissions while keeping the lime consumption at an absolute minimum. At TVA this third control loop was set-up with the option of being able to control the lime slurry flow rate by either the required SO₂ removal efficiency, the desired stoichiometric ratio or the SO₂ emission at the stack.

Table 4.1.1 is a list of the major instrumentation used on the demonstration plant, specifying both the manufacturer and model number. Block diagrams showing the interrelationships between the instrument and the equipment being controlled are presented in Figure 4.1.1-1 to Figure 4.1.1-3.

No major problems for instrumentation and data acquisition were encountered in the entire demonstration run. The only minor incident for the data acquisition was the lime slurry flowmeter drifting away from calibration during the 28-day demonstration run period. This problem was solved by recalibrating the lime slurry flowmeter.

Table 4.1.1

**INSTRUMENT SPECIFICATIONS
FOR CONTINUOUS MONITORS**

Equipment No.	Description	Manufacture	Model No.	Location
DPT-1	Reactor differential pressure	Rosemount	1151DR	Field
DPT-2	Reactor venturi differential pressure	Rosemount	1151DR	Field
PT-1	Inlet gas pressure	Rosemount	1151GP	Field
PT-5	Air pressure	Rosemount	1151GP	Field
PT-6	Water pressure	Rosemount	1151GP	Field
TT-1	Inlet gas temperature	Rosemount	244	Field
TT-2	Reactor temperature, Center	Rosemount	244	Field
WT-1	Feeder weight	Kistler-Morse	DISC II	Field
TIC-5	Cyclone heater control	Shimaden	SR27-2Y4000	Htr Cntl Encl
TIC-6	Feeder heater control	Shimaden	SR27-2Y4000	Htr Cntl Encl
AE-1	Photometric SO ₂ monitor	Du Pont	Series 460	Field
AE-2	Photometric O ₂ monitor	Du Pont	Series 460	Field

Figure 4.1.1-1

**BLOCK DIAGRAM FOR INTERRELATIONSHIPS
OF GSA INSTRUMENTATION AND CONTROL — I**

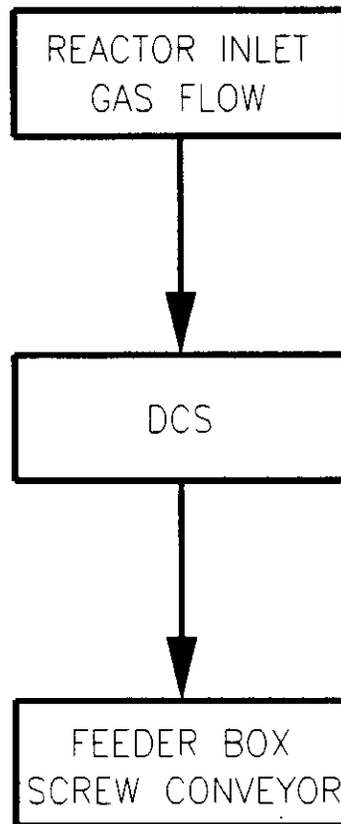


Figure 4.1.1-2

**BLOCK DIAGRAM FOR INTERRELATIONSHIPS
OF GSA INSTRUMENTATION AND CONTROL -- II**

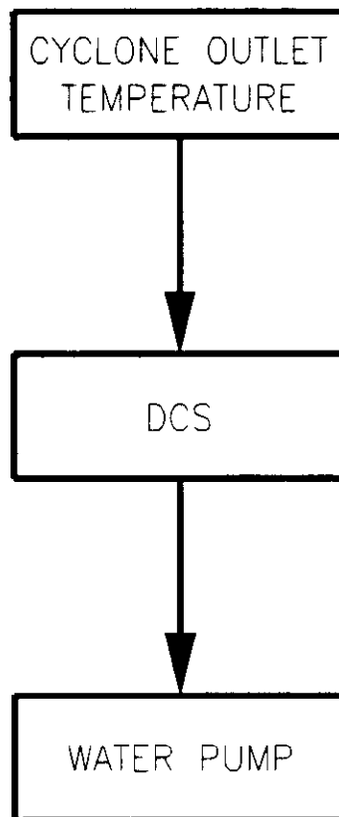
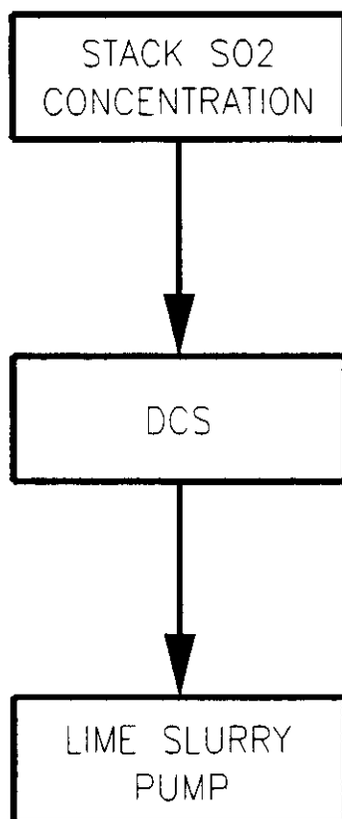


Figure 4.1.1-3

**BLOCK DIAGRAM FOR INTERRELATIONSHIPS
OF GSA INSTRUMENTATION AND CONTROL — III**



4.1.2 Test Methods

The overall test program for the GSA process consisted of five major phases: (1) the preliminary of start-up tests, (2) the factorial tests, (3) the air toxics tests, (4) the 28-day demonstration run and (5) the 14-day PJBH demonstration run. Among these test phases, gaseous streams, aqueous streams, and solid streams were monitored and analyzed by national standard methods. Table 4.1.2-1 shows gaseous stream monitoring summary including a detailed description of the test methods being used. A summary of the aqueous and solid stream monitoring is presented in Table 4.1.2-2 and 4.1.2-3, respectively.

For further information please refer to TVA's final report entitled "10 MW Demonstration of the Gas Suspension Absorption Process at TVA's Center for Emissions Research" in Appendix "B" of this report.

Table 4.1.2-1

GASEOUS STREAM MONITORING SUMMARY

Item	I	II	III	IV	V	Method
Gas flow	C	-	C	-	C	Using a Badger Meter mode PMT-U Lo-Loss venturi. Permanent press. loss 4.2% & diff press. 16" WG. venturi theory: Bernoulli's Theorem.
SO ₂ conc.	C	C	C	C	C	CEM. Using DuPont model 460 SO ₂ analyzers with a photometric analysis (ultraviolet absorption) technique. Instrument range: 0-400 ppm. The continuous sample of flue gas is extracted from the gas path through a 20 micron filter and transported to the sample cell through heat traced tubing (Dekoron model number 2250-24A10). A light source with a wavelength of 280 nanometer (nm) is used for the measuring channel and 578 nm for the reference channel.
O ₂ conc.	C	C	C	-	-	CEM. Using Thermox Instrument model WDGIII (Zirconium Oxide). When the sensing cell is red hot, a voltage is produced that is proportional to the ratio of the oxygen conc. of the gas reference side of the cell (ambient air) and the oxygen conc. of the sample. All (or a portion) of this voltage offset power supply is displayed on the panel meter as a logarithmic display of the oxygen concentration. Optionally, the display may be linearized. The sensing cell is a partial pressure device and responds directly to changes in the sample pressure.

Table 4.1.2-1 (Continued)

GASEOUS STREAM MONITORING SUMMARY

Item	I	II	III	IV	V	Method
Temp.	C	C	C	C	C	Using an Omega model PR13-2-100-1/4-24-E Resistance Temperature Detector (RTD). A platinum RTD inside a sheath is inserted into the flue-gas path. The resistance of the platinum is proportional to its temp. By applying a known voltage across the probe, measuring the current change, and using Ohm's law, a relationship between resistance and temperature is made.
Particulate loading	I	I	I	I	I	Using EPA Reference Method 5 or 17, depending on the particle loading. Method 5 uses a tared out-of-duct filter fitted to the end of a sampling nozzle/probe assembly. Method 17 uses a tared in-duct thimble filter, so the filtration occurs at actual duct gas temp. For both Methods 5 and 17, exhaust gas is drawn isokinetically through a nozzle into the sample system. Following filtration, both methods are identical. The sample gas flows through a series of impingers where moisture is removed. The impingers are followed by a dry-gas meter for volumetric flow measurement. Flue-gas velocity is measured by a pitot attached to the sample probe and isokinetic conditions are assured by selection of the proper nozzle size and appropriate sampling rates. The test methodology requires that a duct traverse of multiple sample points be made to obtain a representative particulate sample. The total particulate catch is determined using a post-test gravimetric analysis of the filter. Any material adhering to the inner walls of the nozzle, probe, filter housing, and connections upstream of the filter is removed through rinsing. This material is then collected and evaporated to dryness in a tared container and weighted.

Table 4.1.2-1 (Continued)

GASEOUS STREAM MONITORING SUMMARY

Item	I	II	III	IV	V	Method
Dewpoint temp.	I	-	-	-	-	Using an Omega model PR13-2-100-1/4-24-E RTD identical to that used for measuring temp except a wetted wick covers the end of the probe.
HCl conc.	TS	TS	TS	TS	TS	Using EPA Reference Method 26.
Pressure	-	-	-	C	C	Using Foxboro company model E13DLM Electronic Force-Balance gage operating on ranges from 1~205" WG. The gage consists of a single integrated dual-compartment unit. The high and low pressure connections are located on opposite sides of a twin-diaphragm capsule. The resulting differential pressure exerts a force on the capsule which is balanced by an opposing force from a feedback coil. Feedback coil output is amplified and transmitted giving pressure.
Air toxics	TS	TS	-	TS	TS	Using EPA Reference Method 5 or 17.

Notes: All items are supplemental parameters.

I = GSA system inlet

II = Reactor outlet/ESP inlet

III = ESP outlet

IV = Baghouse inlet

V = Baghouse outlet

C = Continuous measurements

CS = Continuous measurements for selected tests only (5 to 10% of the 100 to 120 tests)

I = Infrequent, two or three measurements per test

IS = Infrequent, two to three measurements per test for selected tests (5 to 10% of the tests)

TS = Four test conditions over approximately two-week period, triplicate samples

Table 4.1.2-2

AQUEOUS STREAM MONITORING SUMMARY

Item	Method
Lime slurry flow	Using a Foxboro 2800 series (ac Pulsed Coil Excitation) magnetic flow meter. The series 2800 magnetic flow meter (mag meter) consists of two units: (1) Mag flow tube and (2) Mag flow converter/transmitter. The operation of the mag meter is based on Faraday's law of electromagnetic induction; the voltage induced in a conductor of a known length moving through a magnetic field is proportional to the velocity of the conductor. In this application, the process fluid is the conductor. The fluid passes through a magnetic field induced by coils built around a section of the tube. Two metallic electrodes are mounted in the flow tube in contact with the fluid. A resultant voltage is developed across the electrodes which is directly proportional to the average velocity of the fluid.
Lime slurry concentration	Slurry samples are collected in accordance with NCER Method 2.1.1, "Slurry Samples." Slurries are collected from designated sampling points in clean, dry plastic bottles. The bottles are immediately capped and returned to the laboratory for analysis. The analysis follows one of two NCER laboratory procedures for determining this parameter: Method 2.3.1, "Percent Slurry Solids by LX-50 Analyzer," or Method 2.3.2, "Percent Slurry Solids by Gravimetric Determination." For Method 2.3.1, a homogeneous sample is drawn from a well-shaken container with a transfer pipette and delivered to the sample pan of an Arizona Instrument Corporation Model LX-50 percent solids analyzer. The LX-50 automatically weighs the sample, determines the solids content of the slurry, and presents the results on a digital display. For Method 2.3.2, a homogeneous sample is drawn from a well-shaken container with a transfer pipette and delivered to a pre-weighed sample dish. The sample is then dried to a constant weight in a microwave oven set on medium-low power or a convection oven set at 122°F.
Water flow	Using a Foxboro 2800 series (ac Pulsed Coil Excitation) magnetic flow meter. The description of the measurement system and the principle of operation are identical to that used for slurry flow.
Water analysis	The components to be determined are total alkalinity, total dissolved solids, pH and chlorine content. Analysis follows Shawnee Fossil Plant procedures and was performed by the plant's chemical laboratory.

Table 4.1.2-3

SOLID STREAM MONITORING SUMMARY

Item	Method
Coal analysis	Unit 9 coal sample collection and preparation activities followed ASTM D2234-86 and D2013-86. Proximate analyses followed ASTM procedure D3172-73 and ultimate analyses followed ASTM D271-64. The Shawnee Fossil Plant Chemical Laboratory determined heating value (wet and dry), and moisture, ash, sulfur, hydrogen, nitrogen, and chloride content on a daily basis. Volatile matter and fixed carbon content of a composite sample was determined by TVA's Power Service Center laboratory.
Lime analysis	An analysis was conducted of each shipment of lime to determine calcium oxide content following NCER procedure 2.2.9, "Available Lime Determination". The slaking rate, i.e., reactivity, was determined by NCER procedure 2.3.5, "Slaking Rate of Quicklime". These procedures are based on ASTM C110-76a and C25-81a, respectively.
By-product rate	Since the waste disposal system is operated on an intermittent basis to facilitate operations, most by-product rate determinations was made with a computerized material balance program.
By-product ash analysis	Regular laboratory analysis was performed to determine the specific constituents of the by-products. Samples were collected following NCER procedure 2.1.2", Collection and Preparation of Solids Samples". Analysis followed NCER procedure 2.2.1, "Dissolution of Samples and Determination of Acid Insolubles," for determination of calcium, magnesium, chloride, sodium and total sulfur.

5.0 TECHNICAL PERFORMANCE

The SO₂ removal and ESP particulate control results from the 10-MW GSA Clean Coal III demonstration project are discussed in the following sections. These discussions are organized according to the specific test series. The four test series that will be discussed are the two factorial test series, the basic and replicate tests, the 28-day demonstration run, and the 14-day pulse-jet baghouse (PJBH) demonstration run.

5.1 SO₂ REMOVAL PERFORMANCE

5.1.1 Factorial Tests

The tests from the statistically-designed factorial test plan were performed in two parts: the basic series of tests and the replicate series of tests. These factorial tests were designated as either the 2-AP or 3-AP series depending on the orientation and status of the 1-MW pulse-jet baghouse, which was tested concurrently with the GSA/ESP system. The test designation 2-AP was used to denote when either the PJBH was not operating or was operated in a series with the ESP (withdrawing a slipstream of flue gas from downstream of the ESP). All of the replicate series of tests were designed to be completed with the PJBH operating in series with the ESP. However, some of the basic tests were completed with the PJBH off-line and were designated as 2-AP series tests. The test designation 3-AP was used when the PJBH was operated in parallel with the ESP (withdrawing a slipstream of flue gas from upstream of the ESP). Most of the basic factorial tests were completed in this mode.

A total of 78 tests were performed during the factorial test phase. Not all of these tests, however, were part of the original factorial test plan. As an example, several tests were added during the factorial test phase to further evaluate the pulse-jet baghouse. Table 5.1.1-1 lists the 2-AP and 3-AP series tests that were conducted at operating conditions specified in the factorial test plan. These tests typically consisted of 12 to 24 hours of operation to reach steady state, followed by 24 to 48 hours of testing from which the test averages were developed. The data from 10 test segments will not be reported due to problems encountered during these test (tests 2-AP-05, 2-AP-10, 2-AP-14 (file 2), 2-AP-15 (files 1 & 2), 2-AP-16, 2-AP-93, 3-AP-15, 3-AP-60, and 3-AP-61). The problems include equipment operation which interfered with the GSA system achieving steady state conditions, calibration problems with process monitoring equipment, and/or an insufficient amount of data to develop representative test averages for specific operating conditions.

The SO₂ removal results for the tests conducted at baseline chloride levels (0.04 weight percent coal chloride) are presented in Table 5.1.1-2 for the 2-AP series tests and in Table 5.1.1-3 for the 3-AP series tests. Similarly, the SO₂ removal results are presented in Tables 5.1.1-4 and 5.1.1-5 for the chloride spiking tests (0.12 weight percent coal chloride equivalent) for the 2-AP and 3-AP series, respectively. As shown in these tables, the majority of the SO₂ removal occurs in the GSA reactor/cyclone. The ESP contribution to the total system SO₂ removal ranged from 1 to 7 percent.

Table 5.1.1-1

FINALIZED BASIC AND REPLICATE TESTS

<u>Basic Test Numbers</u>		<u>Replicate Test Numbers</u>	
<u>Planned</u>	<u>Actual</u>	<u>Planned</u>	<u>Actual</u>
2-AP-01	2-AP-01 3-AP-62	2-AP-71	2-AP-71
2-AP-04	2-AP-04	2-AP-74	2-AP-74
2-AP-05	3-AP-29	2-AP-75	2-AP-75
2-AP-08	3-AP-08	2-AP-78	2-AP-78
2-AP-03	2-AP-03 3-AP-03 3-AP-02	2-AP-73	2-AP-73
2-AP-07	2-AP-07	2-AP-77	2-AP-77
2-AP-06	2-AP-06	2-AP-76	2-AP-92
2-AP-09	2-AP-09 3-AP-12	2-AP-79	2-AP-79
2-AP-16	2-AP-16	2-AP-72	2-AP-72
2-AP-11	2-AP-11 3-AP-11	2-AP-81	2-AP-81
2-AP-10	2-AP-10	2-AP-80	2-AP-80
2-AP-17	2-AP-17	2-AP-82	2-AP-82
2-AP-18	3-AP-18	2-AP-88	2-AP-88
2-AP-19	2-AP-19 3-AP-19 2-AP-57	2-AP-89	2-AP-97
2-AP-20	3-AP-20 3-AP-13 3-AP-20	2-AP-86	2-AP-86
2-AP-21	3-AP-21	2-AP-87	2-AP-87
2-AP-22	2-AP-22 3-AP-22	2-AP-90	2-AP-90
2-AP-23	3-AP-23	2-AP-83	2-AP-83
2-AP-24	2-AP-24 3-AP-24	2-AP-84	2-AP-84
2-AP-25	2-AP-25	2-AP-85	2-AP-85

Table 5.1.1-2
 AIRPol GSA/ESP SO2 Removal Results Summary
 2-AP Series - Baseline Tests

Test No.	Cu/S Ratio	Reactor Inlet Temp (F)	Approach Temperature (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Injection Rate (lb/min)	Reactor Product Chloride Content (%)	Reactor/Cyclone SO2 Removal (%)	ESP SO2 Removal (%)	Total System SO2 Removal (%)
2-AP-09	1.0	320	8	14,000	30	0	0	70.6	5.6	76.2
2-AP-19	1.0	320	8	14,000	30	0	0	82.5	1.3	83.8
2-AP-72	1.0	320	8	20,000	45	7.3	0	66.9	4.1	71.0
2-AP-16	1.0	320	8	20,000	45	7.8	0.04	74.1	0.5	74.6
2-AP-11	1.3	320	8	14,000	30	5.0	0	88.4	3.7	92.1
2-AP-81	1.3	320	8	14,000	30	5.2	0	86.1	4.1	90.2
2-AP-10	1.3	320	8	20,000	45	0	0	88.0	3.4	91.4
2-AP-80	1.3	320	8	20,000	45	0	0.09	84.1	4.2	88.3
2-AP-01	1.0	319	18	14,000	30	0	0	67.1	3.5	70.6
2-AP-71	1.0	320	18	14,000	30	0	0.02	73.9	4.0	77.9
2-AP-78	1.0	320	18	20,000	30	7.2	0	63.6	4.1	67.7
2-AP-04	1.0	320	18	19,000	45	0	0	66.0	1.7	67.7
2-AP-74	1.0	320	18	20,000	45	0	0	69.1	2.1	71.2
2-AP-03	1.3	319	18	14,000	45	0	0	88.4	2.6	91.0
2-AP-73	1.3	320	18	14,000	45	5.1	0	82.2	2.3	84.5
2-AP-95	1.3	320	18	20,000	45	0	0.05	78.2	4.7	82.9
2-AP-96	1.3	322	18	20,000	45	0	0.21	83.2	3.4	86.6
2-AP-14	1.3	320	18	18,000	45	6.7	0.12	80.4	5.0	85.4
2-AP-63	1.3	320	18	20,000	45	7.2	0.04	76.1	5.9	82.0
2-AP-63	1.3	320	18	20,000	45	7.2	0.03	75.4	3.1	78.5
2-AP-88	1.0	320	28	14,000	30	0	0.04	63.9	2.2	66.1
2-AP-87	1.0	320	28	20,000	45	7.3	0	59.2	3.2	62.4
2-AP-86	1.3	320	28	14,000	45	0	0.03	75.5	2.0	77.5
2-AP-97	1.3	320	28	20,000	30	0	0	68.1	4.4	72.5
2-AP-19	1.3	320	28	20,000	30	7.1	0	61.9	5.2	67.1
2-AP-57	1.3	319	28	19,500	30	7.1	0	73.3	5.5	78.8

**Table 5.1.1-3
AirPol GSA/ESP SO2 Removal Results Summary
3-AP Series - Baseline Tests**

Test No.	Ca/S Ratio	Reactor Inlet Temperature (F)	Approach Temperature (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Injection Rate (lb/min)	Reactor Product Chloride Content (%)	Reactor/ Cyclone SO2 Removal (%)	ESP SO2 Removal (%)	Total System SO2 Removal (%)
3-AP-12	1.0	320	8	14,000	30	0	0	73.4	3.8	77.2
3-AP-58	1.0	320	8	14,000	30	0	0	87.8	3.7	91.5
3-AP-42	1.3	320	8	14,000	30	0	0	96.4	2.0	98.4
3-AP-11	1.3	319	8	14,000	30	5.2	0	98.6	0.7	99.3
3-AP-62	1.0	319	18	14,000	30	0	0.13	71.8	1.9	73.7
3-AP-08	1.0	320	18	20,000	30	7.0	0	57.2	6.9	61.1
3-AP-44	1.3	319	18	14,000	45	0	0.02	86.8	3.2	89.9
3-AP-03	1.3	319	18	14,000	45	5.0	0	78.3	4.0	82.3
3-AP-26	1.3	260	18	14,000	12	4.7	0	73.1	5.3	78.4
3-AP-27	1.3	260	18	14,000	30	4.7	0	81.1	3.8	84.9
3-AP-02	1.3	319	18	14,000	45	5.1	0	84.9	4.0	88.9
3-AP-18	1.0	319	28	14,000	30	0	0	63.3	3.4	66.7
3-AP-59	1.0	320	28	14,000	30	5.2	0	66.9	3.1	70.0
3-AP-21	1.0	319	28	20,000	45	7.3	0	54.7	6.0	60.7
3-AP-56	1.3	320	28	14,000	30	0	0	82.8	2.7	85.5
3-AP-20	1.3	320	28	14,000	45	0	0	83.2	3.3	86.5
3-AP-20	1.3	320	28	14,000	45	0	0.03	84.3	2.3	86.6
3-AP-13	1.3	319	28	14,000	45	0	0	79.8	5.0	84.8
3-AP-45	1.3	320	28	14,000	45	5.2	0	80.3	2.4	82.7
3-AP-19	1.3	320	28	20,000	30	7.4	0	69.9	5.3	74.2

Table 5.1.1-4
AirPol GSA/ESP SO2 Removal Results Summary
 2-AP Series - Chloride Spiking Tests

Test No.	Cx/S Ratio	Reactor Inlet Temperature (F)	Approach Temperature (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Inj. Rate (lb/min)	Reactor Product Chloride Content (%)	Reactor Cyclone SO2 Removal (%)	ESP SO2 Removal (%)	Total System SO2 Removal (%)
2-AP-28	1.0	320	18	14,000	40	5.0	0.39	77.6	5.1	82.7
2-AP-75	1.0	320	18	14,000	45	4.7	0.45	76.2	4.3	80.5
2-AP-17	1.0	320	18	20,000	30	0	0.75	76.8	2.6	79.4
2-AP-82	1.0	320	18	20,000	30	0	0.87	76.5	1.8	78.3
2-AP-07	1.3	320	18	14,000	30	5.4	0.29	91.0	3.9	94.9
2-AP-77	1.3	320	18	14,000	30	5.0	0.30	88.2	5.6	93.8
2-AP-98*	1.3	320	18	14,000	30	5.2	0.30	88.1	2.4	90.5
2-AP-06	1.3	320	18	20,000	45	0	0.47	88.8	4.9	93.9
2-AP-92	1.3	320	18	20,000	45	0	0.72	89.0	2.4	91.4
2-AP-91	1.3	320	18	20,000	45	7.2	0.32	87.6	2.1	89.7
2-AP-22	1.0	320	28	14,000	45	0	0.57	75.7	3.7	79.4
2-AP-90	1.0	320	28	14,000	45	0	0.86	68.9	1.0	69.9
2-AP-25	1.0	320	28	18,600	30	6.8	0.28	60.8	6.2	67.0
2-AP-94	1.0	320	28	20,000	30	0	0.81	65.4	0.7	66.1
2-AP-85	1.0	320	28	20,000	30	7.4	0.31	64.3	0.6	64.9
2-AP-84	1.3	320	28	14,000	30	0	0.60	79.4	3.0	82.4
2-AP-24	1.3	320	28	14,000	30	0	0.41	82.3	3.2	85.5
2-AP-83	1.3	320	28	20,000	30	7.3	0.26	70.2	9.7	79.9

* Test data has not been finalized.

**Table 5.1.1-5
AirPol GSA/ESP SO2 Removal Results Summary
3-AP Series - Chloride Spiking Tests**

Test No.	Ca/S Ratio	Reactor Inlet Temperature (F)	Approach Temperature (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Injection Rate (lb/min)	Reactor Product Chloride Content (%)	Reactor/Cyclone SO2 Removal (%)	ESP SO2 Removal (%)	Total System SO2 Removal (%)
3-AP-29	1.0	320	18	14,000	45	5.2	0.31	84.0	2.2	86.2
3-AP-22	1.0	320	28	14,000	45	0	0.57	76.7	3.4	80.1
3-AP-24	1.3	320	28	14,000	30	0	0.46	81.9	4.4	86.3
3-AP-23	1.3	320	28	19,200	45	7.2	0.26	79.3	4.1	83.4

5.1.2 Effect of Lime Stoichiometry and Approach Temperature

The SO₂ removal performance results from all of the 2-AP series tests conducted at baseline chloride levels (0.04 weight percent coal chloride) are presented in Figure 5.1.2-1. In the figure, the average total system (GSA + ESP) SO₂ removal is plotted for each test as a function of fresh lime stoichiometry with different symbols used for the three levels of approach temperature; 8, 18 and 28°F. Linear regression curves for each approach temperature are also plotted in the figure.

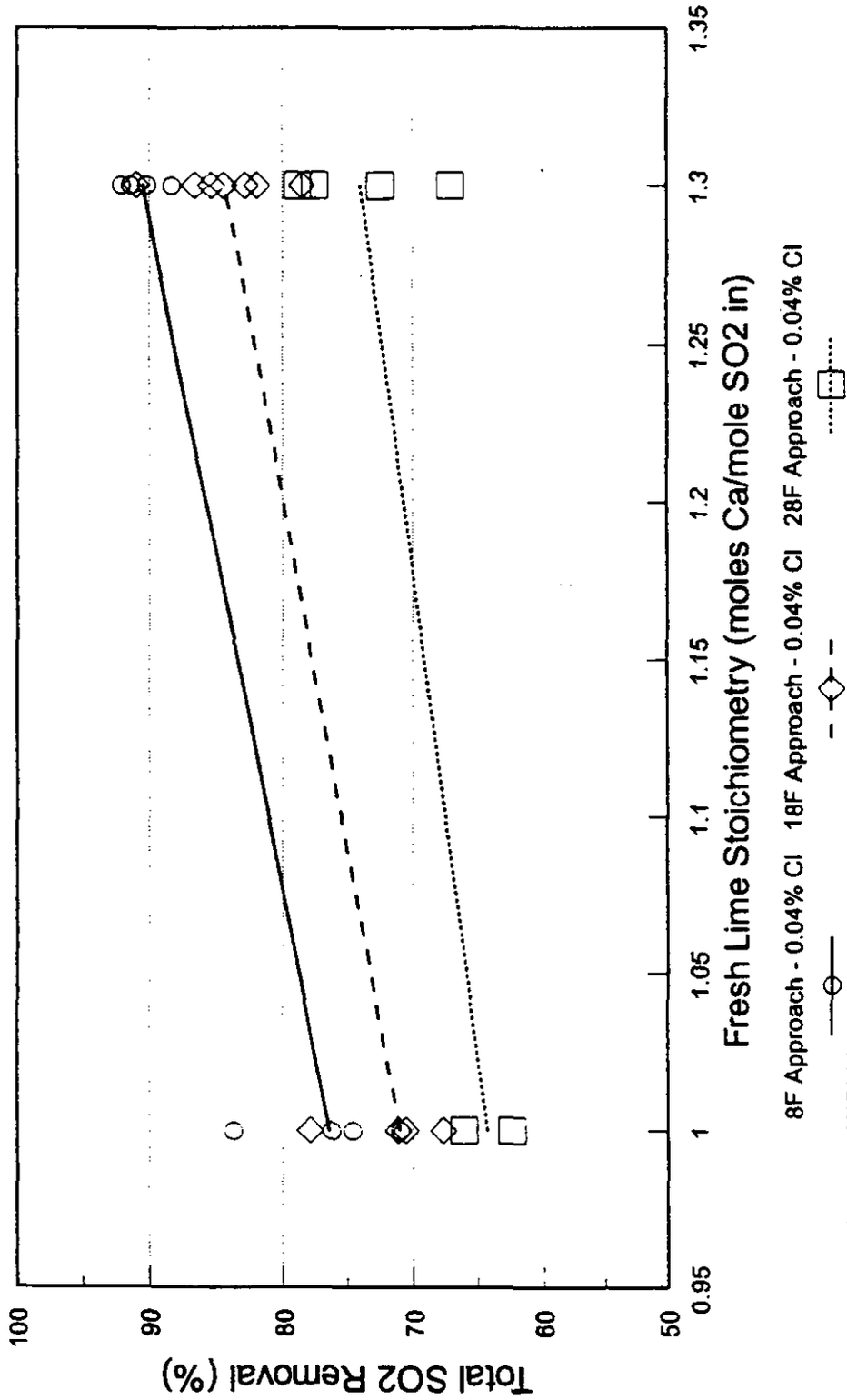
As shown in Figure 5.1.2-1, the total system SO₂ removal increases as the fresh lime stoichiometry is increased from 1.0 to 1.3 and the approach temperature is decreased from 28 to 8°F. The average total system SO₂ removal ranged from a low of approximately 62 percent at a 1.0 stoichiometry and a 28°F approach to a high of 92 percent at a 1.3 stoichiometry and an 8°F approach temperature. Based on the linear regression lines, the SO₂ removal increases approximately 9 to 13 percentage points as the stoichiometry is increased from 1.0 to 1.3. The increase in SO₂ removal as the approach temperature is reduced from 28 to 18°F is about 6 to 10 percentage points at the same fresh lime stoichiometry. A decrease in the approach temperature from 18 to 8°F results in a further increase in SO₂ removal of about 5 to 6 percentage points at the same fresh lime stoichiometry.

Figure 5.1.2-2 provides a similar plot of the data from the 3-AP series tests. In the figure, only the tests conducted at the lower flue gas flow rate of 14,000 scfm are plotted. Because only the 3-AP series tests at 14,000 scfm are plotted, the SO₂ removal performance is higher in these tests compared to the 2-AP series results presented in Figure 5.1.2-1. This higher SO₂ removal performance is presumably due to the increased flue gas residence time in the GSA reactor. Unlike the prior figure, the increase in SO₂ removal is greater when the approach temperature is decreased from 18 to 8°F (10 percentage points) compared to the increase when reducing the approach temperature from 28 to 18°F (2 to 5 percentage points).

Figure 5.1.2-1

AirPol GSA SO₂ Removal Results

2-AP Series Baseline Chloride Tests

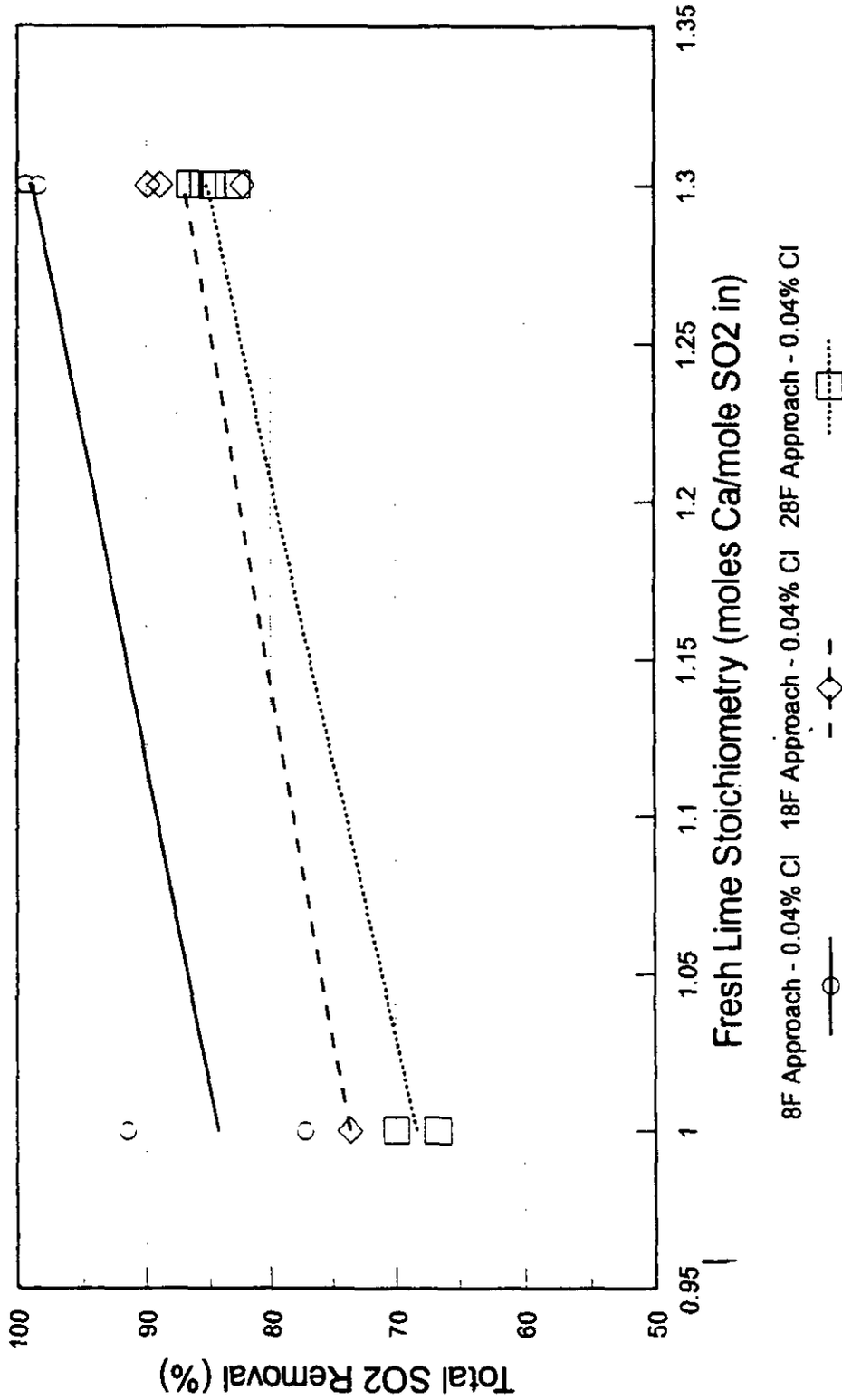


Notes: All tests conducted at a 320F inlet flue gas temperature and 25 wt % solids lime slurry.

Figure 5.1.2-2

AirPol GSA SO₂ Removal Results

3-AP Series Baseline Chloride Tests



Notes: All tests conducted at a 320F inlet flue gas temperature, 25 wt % solids lime slurry and 14,000 scfm.

5.1.3 Effect of Flue Gas Flow Rate

The flue gas flow rate through the GSA system was a significant variable in affecting the SO₂ removal performance. Figures 5.1.3-1, 5.1.3-2 and 5.1.3-3 present the results from the 2-AP series tests conducted at baseline chloride levels. In each figure, the average total SO₂ removal is plotted for each test as a function of fresh lime stoichiometry. The distinction is made in each figure for tests conducted at the two flue gas flow rate levels, 14,000 and 20,000 scfm. Linear regression lines are plotted for each flue gas flow rate. Figure 5.1.3-1 plots data for tests conducted at an 8°F approach temperature, while Figures 5.1.3-2 and 5.1.3-3 plot data for tests conducted at an 18 and 28°F approach temperatures, respectively.

In all three figures, the SO₂ removal performance is lower at the higher flue gas flow rate, 20,000 scfm. The decrease in performance ranges from approximately 2 to 8 percentage points based on the linear regression lines. The lower SO₂ removal performance at 20,000 versus 14,000 scfm was also observed in the 2-AP series tests conducted with calcium chloride spiking. Figures 5.1.3-4 and 5.1.3-5 provide similar plots of the average total SO₂ removal as a function of fresh lime stoichiometry for tests conducted at an 18 and 28°F approach temperature, respectively. Similar to the baseline chloride tests, the SO₂ removal decreased from approximately 2 to 9 percentage points as the flue gas flow rate increased from 14,000 to 20,000 scfm.

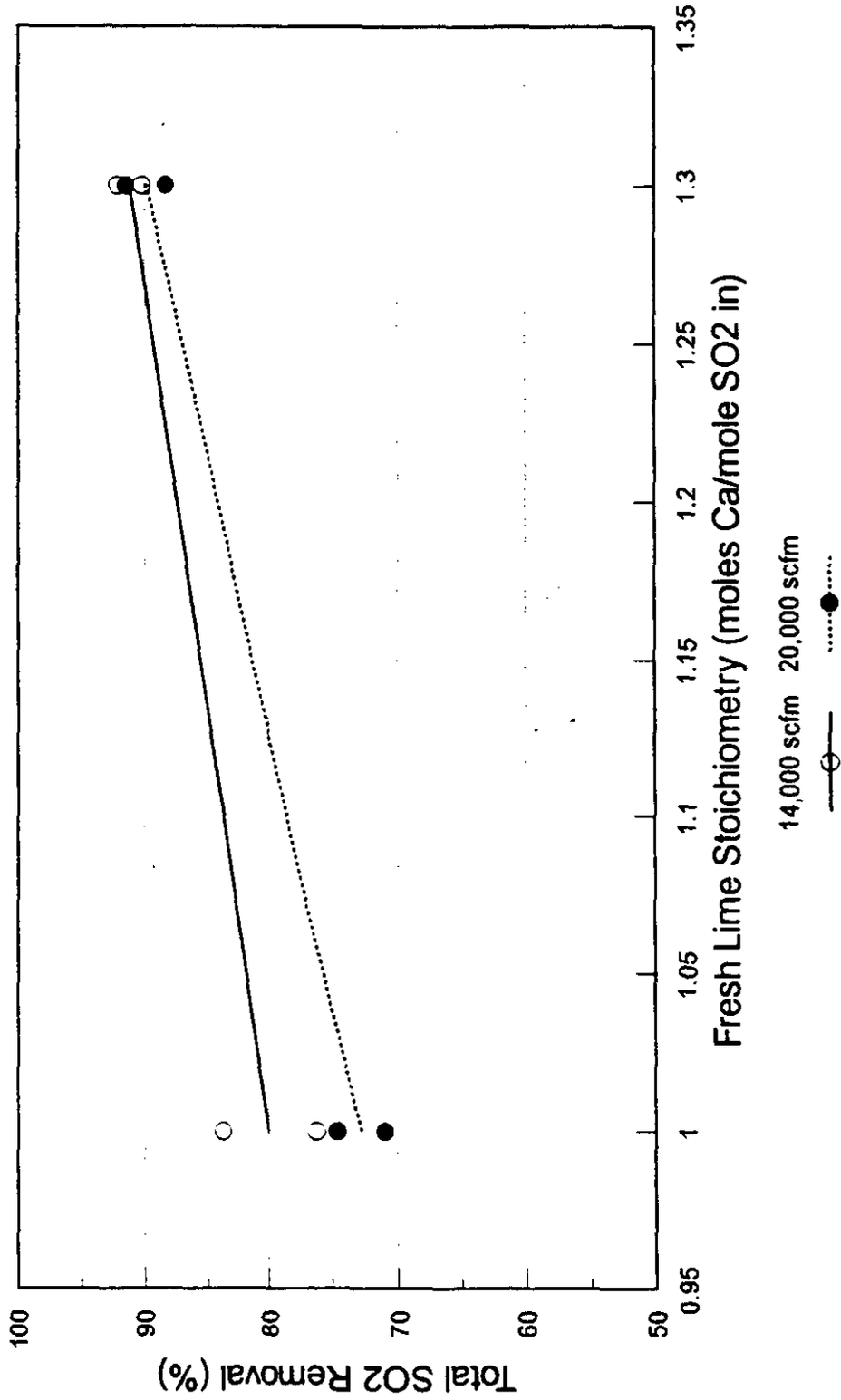
This same effect was also observed in the 3-AP series tests. Figure 5.1.3-6 plots the average total SO₂ removal as a function of fresh lime stoichiometry for the 3-AP series tests conducted at baseline chloride levels. In the figure, the tests conducted at an 18 and 28°F approach temperature and at flue gas flow rates of 14,000 and 20,000 scfm are plotted. Only the regression lines are plotted for the test data at a flue gas flow rate of 14,000 scfm in order to more readily distinguish the data points from the tests conducted at 20,000 scfm. All three tests conducted at the higher flue gas flow rate resulted in decreased SO₂ removal performance. Based on the linear regression lines, the decrease in SO₂ removal was approximately 10 percentage points.

The increase in SO₂ removal at the lower flue gas flow rate is presumably due to the increased residence time in the GSA absorber. The residence time increases from approximately 3.9 seconds at a flue gas flow rate of 20,000 scfm to 5.5 seconds at 14,000 scfm. Although this is only a 1.6 second differential, it represents a 41 percent increase in residence time which is significant. The effect of residence time in the GSA absorber, especially at these low residence times, may be more significant compared to other dry scrubbing technologies such as spray drying because the cyclone downstream of the GSA absorber removes over 90 percent of the solids/sorbent from the flue gas stream, thus eliminating any further reaction of the sorbent with the flue gas SO₂.

Figure 5.1.3-1

AirPol GSA SO₂ Removal Results

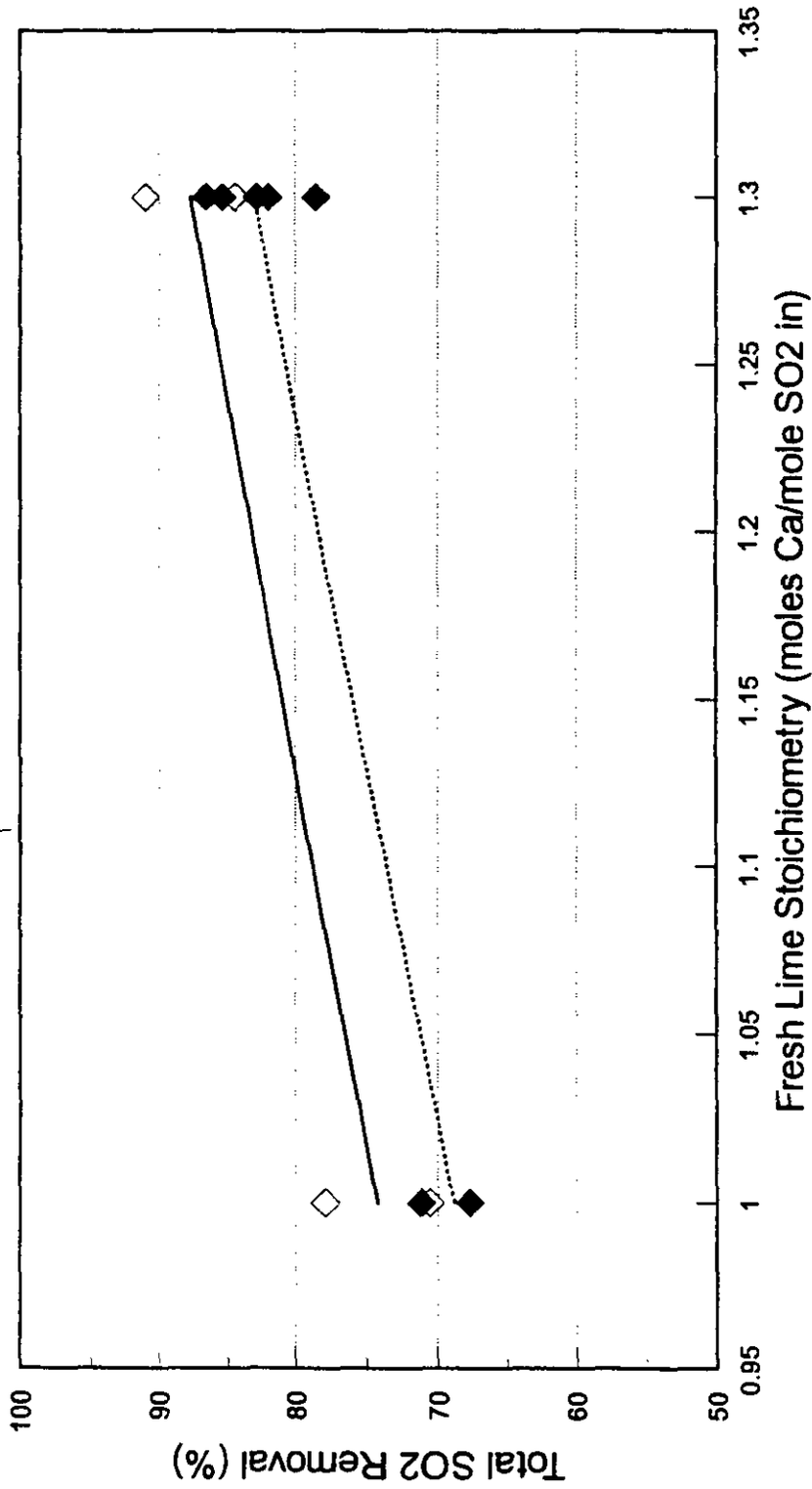
2-AP Series 8F Approach Tests



Notes: All tests conducted at a 320F Inlet
flue gas temperature, 8F approach temperature,
and at a 0.04% coal Cl level.

Figure 5.1.3-2

AirPol GSA SO₂ Removal Results 2-AP Series 18F Approach Tests

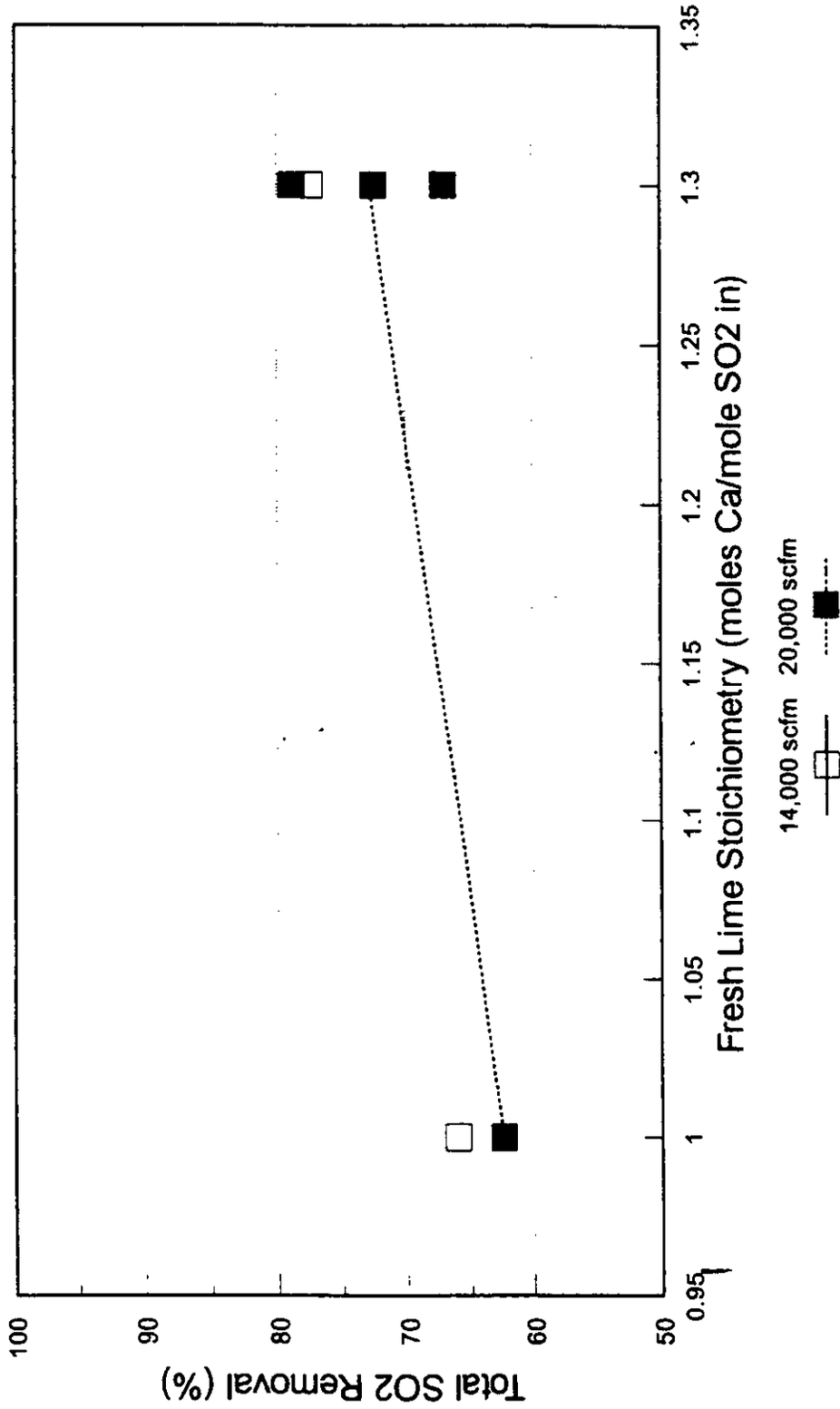


Notes: All tests conducted at a 320F inlet flue gas temperature, 18F approach temperature, and at a 0.04% coal Cl level.

Figure 5.1.3-3

AirPol GSA SO₂ Removal Results

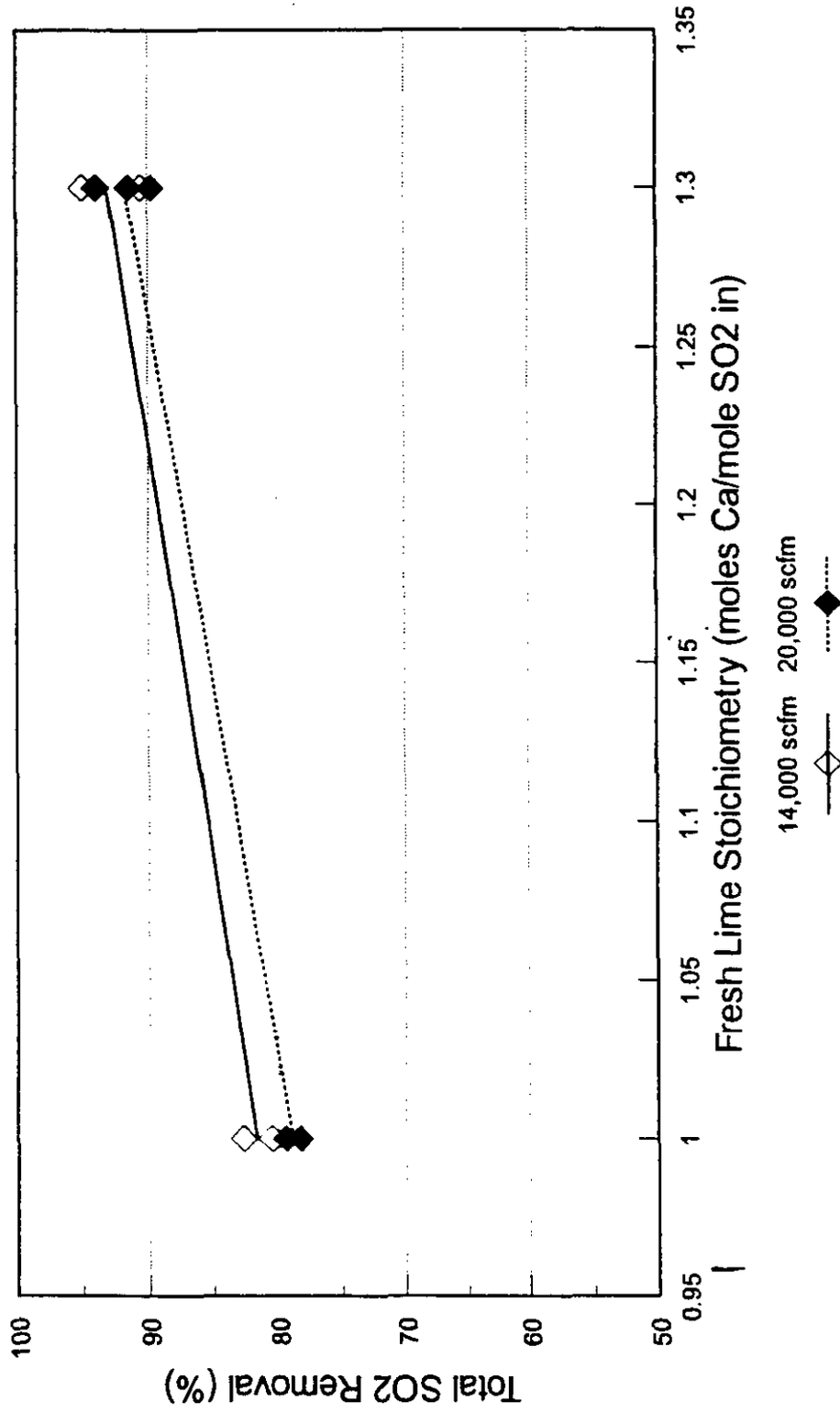
2-AP Series 28F Approach Tests



Notes: All tests conducted at a 320F inlet flue gas temperature, 28F approach temperature, and at a 0.04% coal Cl level.

Figure 5.1.3-4

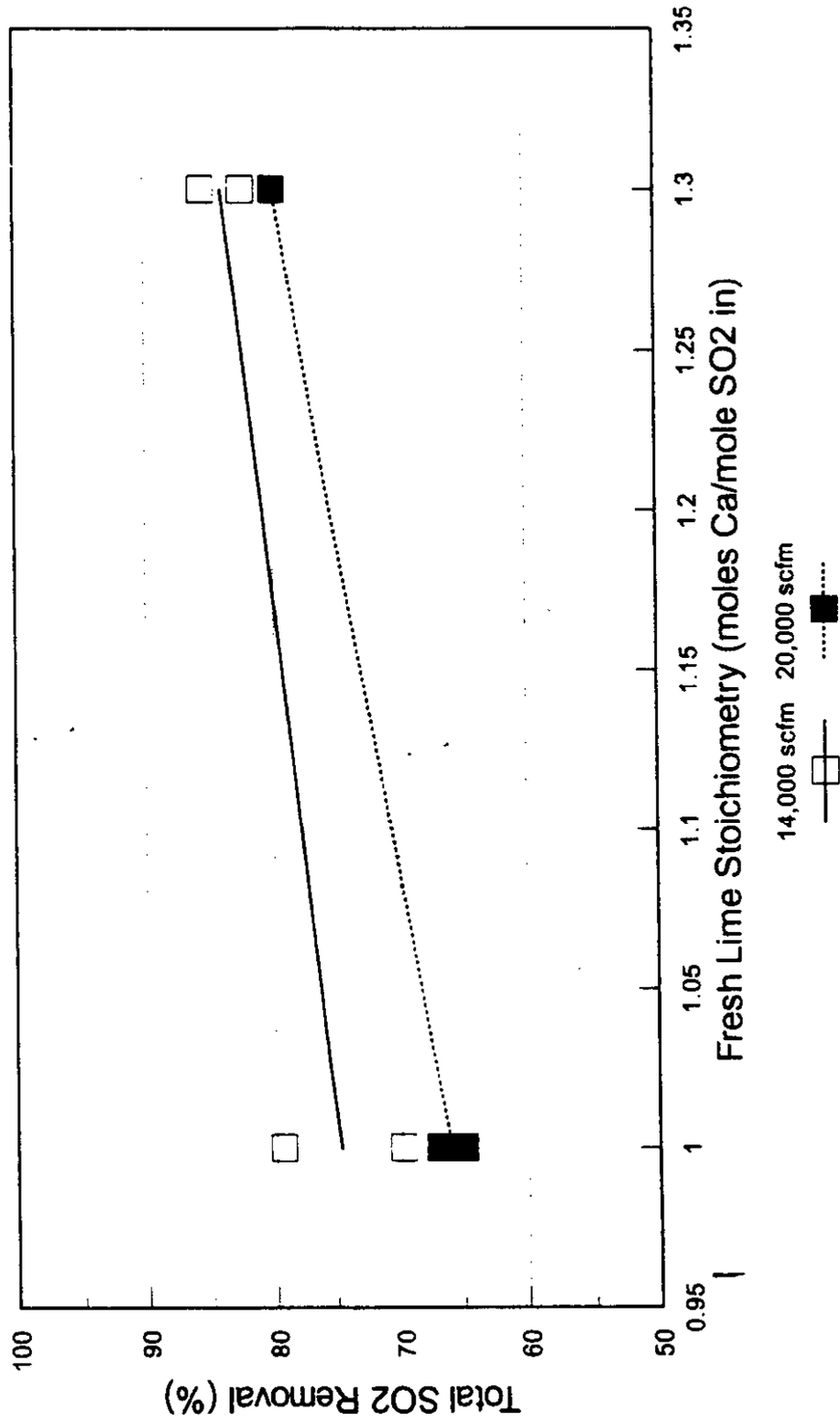
AirPol GSA SO₂ Removal Results 2-AP Series 18F Approach/Chloride Spiking Tests



Notes: All tests conducted at a 320F inlet flue gas temperature, 18F approach temperature, and at a 0.12% coal Cl level.

Figure 5.1.3-5

AirPol GSA SO₂ Removal Results 2-AP Series 28F Approach/Chloride Spiking Tests

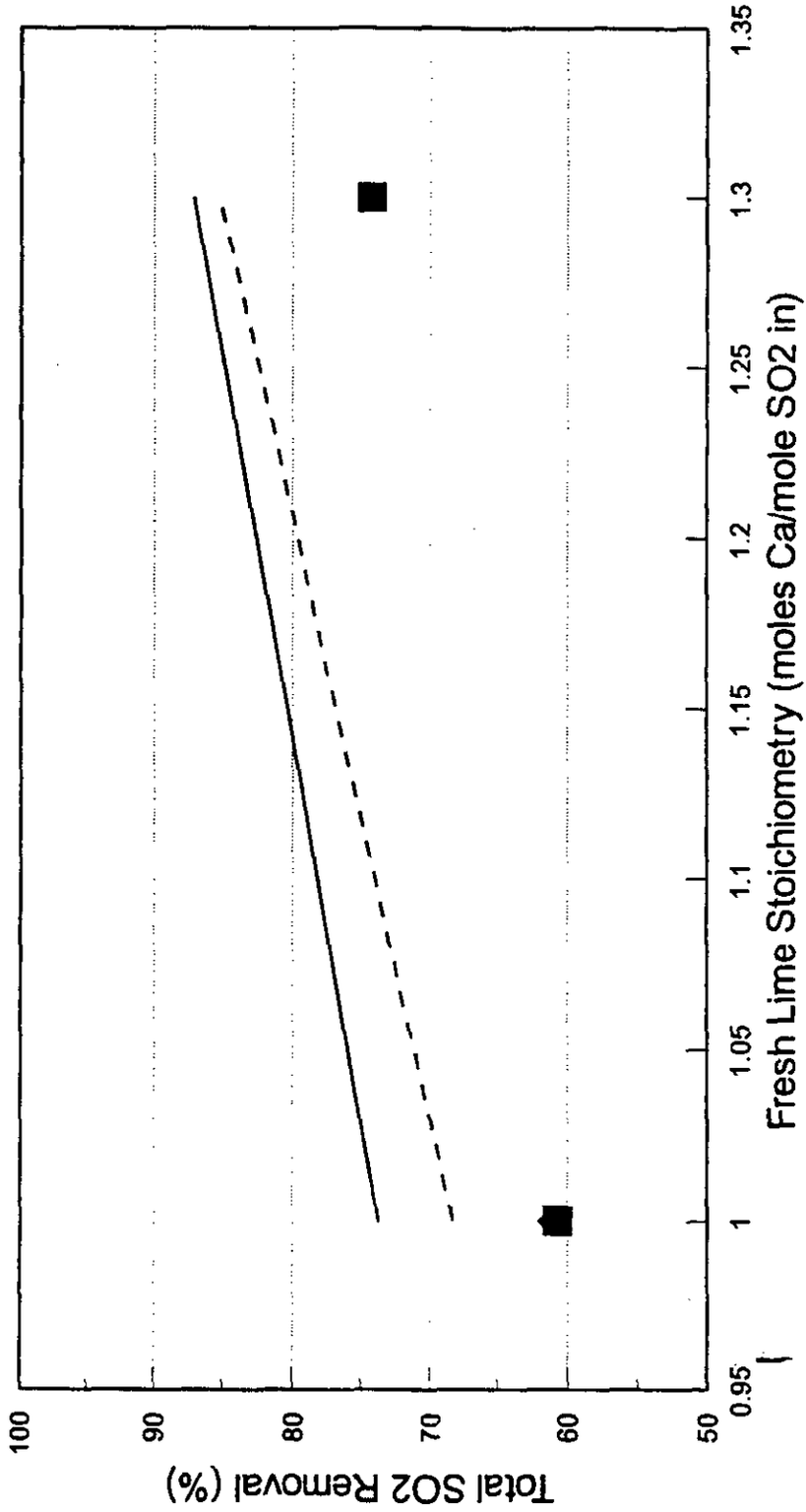


Notes: All tests conducted at a 320F inlet
flue gas temperature, 28F approach temperature,
and at a 0.12% coal Cl level.

Figure 5.1.3-6

AirPol GSA SO2 Removal Results

3-AP Series Baseline Chloride Tests



18F Approach - 14,000 scfm 28F Approach - 14,000 scfm 18F Approach - 20,000 scfm 28F Approach - 20,000 scfm

Notes: All tests conducted at a 320F Inlet flue gas temperature and 25 wt % solids lime slurry.

5.1.4 Effect of Chloride Spiking

Similar to prior dry scrubbing studies, calcium chloride spiking was found to have a beneficial effect on SO₂ removal in the GSA system. Figure 5.1.4-1 presents the data from the 2-AP series tests conducted with calcium chloride spiking to simulate scrubbing flue gas resulting from the combustion of a 0.12 weight percent chlorine coal. In the figure, the average total SO₂ removal is plotted as a function of fresh lime stoichiometry for tests conducted at an 18 and 28°F approach temperature. The average total system SO₂ removal ranged from a low of approximately 65 percent at a 1.0 stoichiometry and a 28°F approach to a high of 94 percent at a 1.3 stoichiometry and an 18°F approach temperature. Based on the linear regression lines, the SO₂ removal increases approximately 12 percentage points as the stoichiometry is increased from 1.0 to 1.3. The increase in SO₂ removal as the approach temperature is reduced from 28 to 18°F is about 10 percentage points. No chloride spiking tests were completed below an 18°F approach temperature because of the potential for solids build-up/pluggage problems.

The baseline chloride results for the 2-AP series tests are compared with the chloride spiking test results in Figures 5.1.4-2 and 5.1.4-3. Figure 5.1.4-2 presents the data at an 18°F approach temperature and Figure 5.1.4-3 presents the 28°F approach test results. The distinction is made in the figures for tests conducted at 14,000 and 20,000 scfm. Compared to the baseline chloride results, the higher chloride level improves SO₂ removal by about 4 to 10 percentage points at a stoichiometric ratio of 1.0. At a stoichiometric ratio of 1.3, the increase in SO₂ removal is comparable, ranging from about 4 to 9 percentage points.

An increase in SO₂ removal with calcium chloride addition was also observed in the 3-AP series tests. Figure 5.1.4-4 presents the data for tests conducted at an 18 and 28°F approach temperature. Only the regression lines are plotted for the baseline chloride test data in order to more readily distinguish the data points from the chloride spiking tests. The tests conducted at a 1.0 stoichiometry exhibited approximately a 12 to 13 percentage point increase in total average SO₂ removal. The one test conducted at a 1.3 stoichiometry, however, did not show any improvement. This latter result is somewhat unexpected.

5.1.5 Effect of Other Operational Variables

The other operational variables, such as recycle screw speed and inlet fly ash loading, also had an effect on SO₂ removal. The influence of these variables, however, was less than the effect of stoichiometry, approach temperature, and flue gas flow rate (residence time). In addition, two tests were conducted at a lower inlet flue gas temperature of 260°F.

5.1.6 Lime Utilization

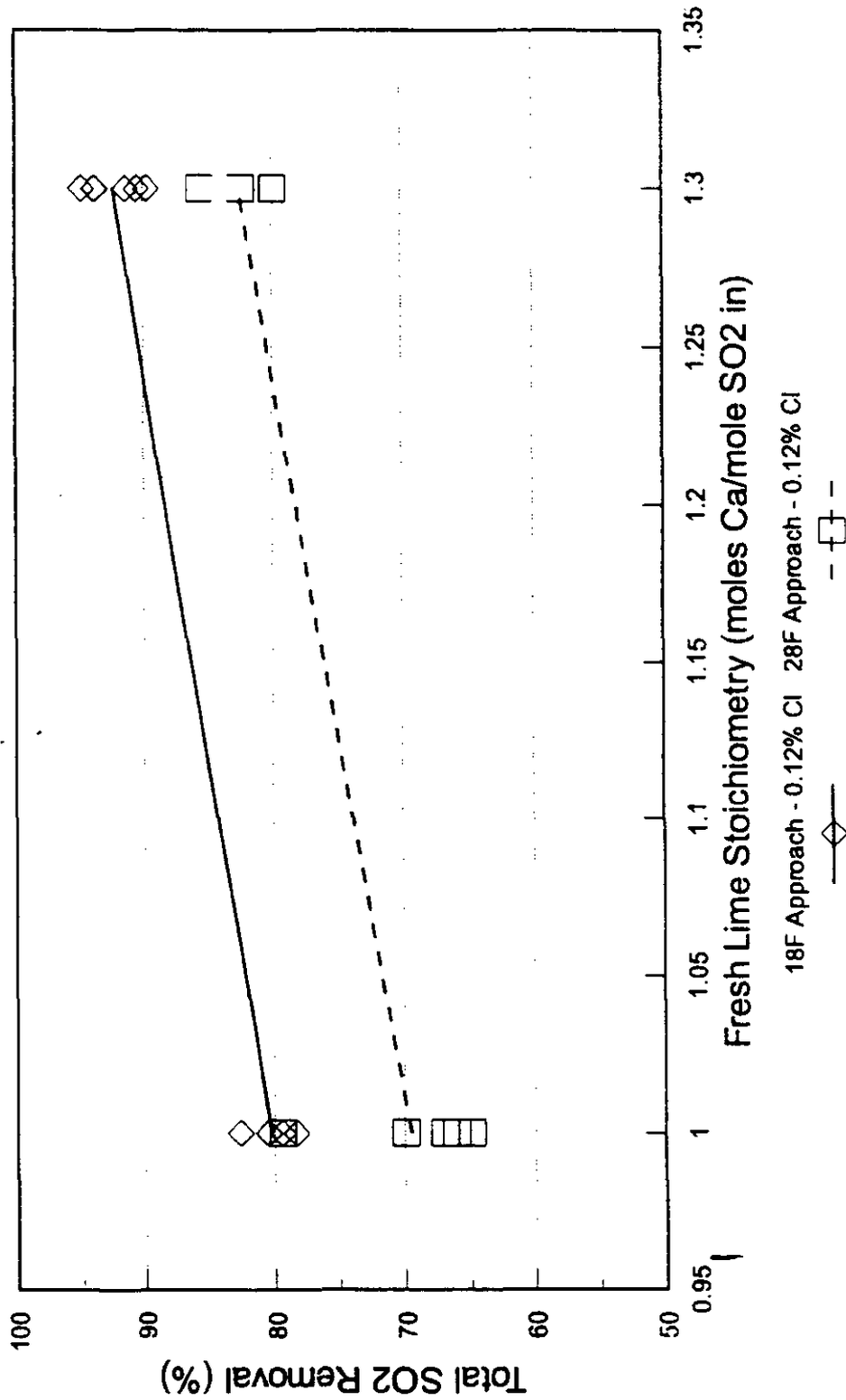
The total system (GSA + ESP) calculated lime utilizations based on the process data ranged from 50 to 84 percent during the factorial tests. The lime utilization is calculated by dividing the total system SO₂ removal by the fresh lime stoichiometry. The lowest lime utilization rates were for tests conducted at the higher approach temperature (28°F) and higher fresh lime stoichiometry

(1.30 moles $\text{Ca}(\text{OH})_2$ /mole inlet SO_2). Decreasing the approach temperature and/or the fresh lime stoichiometry improved the GSA system lime utilization. Calcium chloride spiking also improved the lime utilization compared to tests conducted at the same operating conditions at baseline (0.04 weight percent) coal chloride levels.

The lime or calcium utilization was also determined analytically for three sample locations; the recycle feeder box solid samples, and the solids from the first field ESP hopper and a composite from ESP fields 2 through 4. Typically, the measured calcium utilization for the reactor recycle solids would either be lower or fall in between the measured calcium utilization values for the ESP solids. The highest calcium utilization values were measured for the solids from ESP hoppers 2 through 4. This is to be expected due to the additional SO_2 removal that occurs in the ESP.

Figure 5.1.4-1

AirPol GSA SO₂ Removal Results 2-AP Series Chloride Spiking Tests

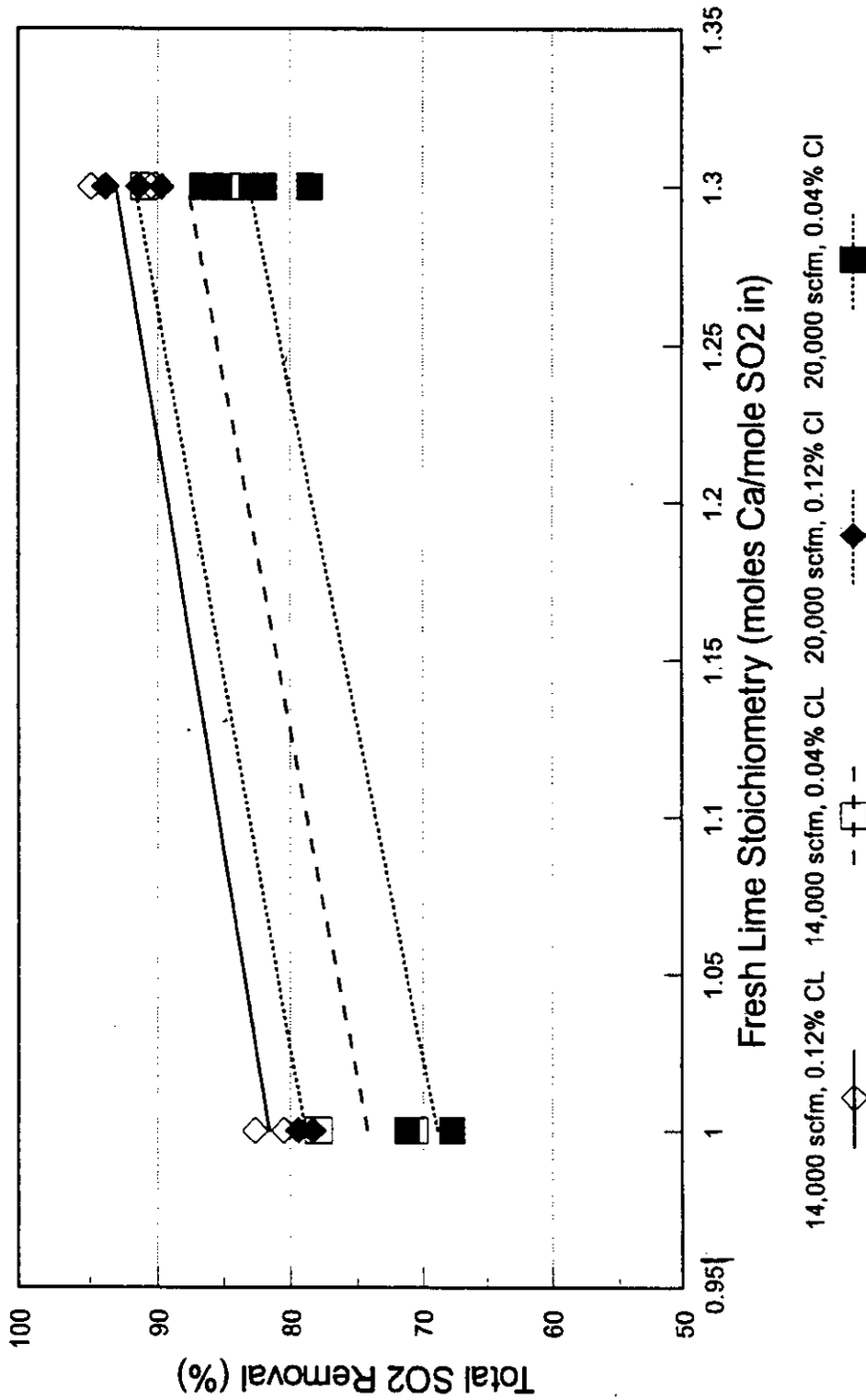


Notes: All tests conducted at a 320F inlet flue gas temperature and 25 wt % solids lime slurry.

Figure 5.1.4-2

AirPol GSA SO₂ Removal Results

2-AP Series 18F Approach Tests

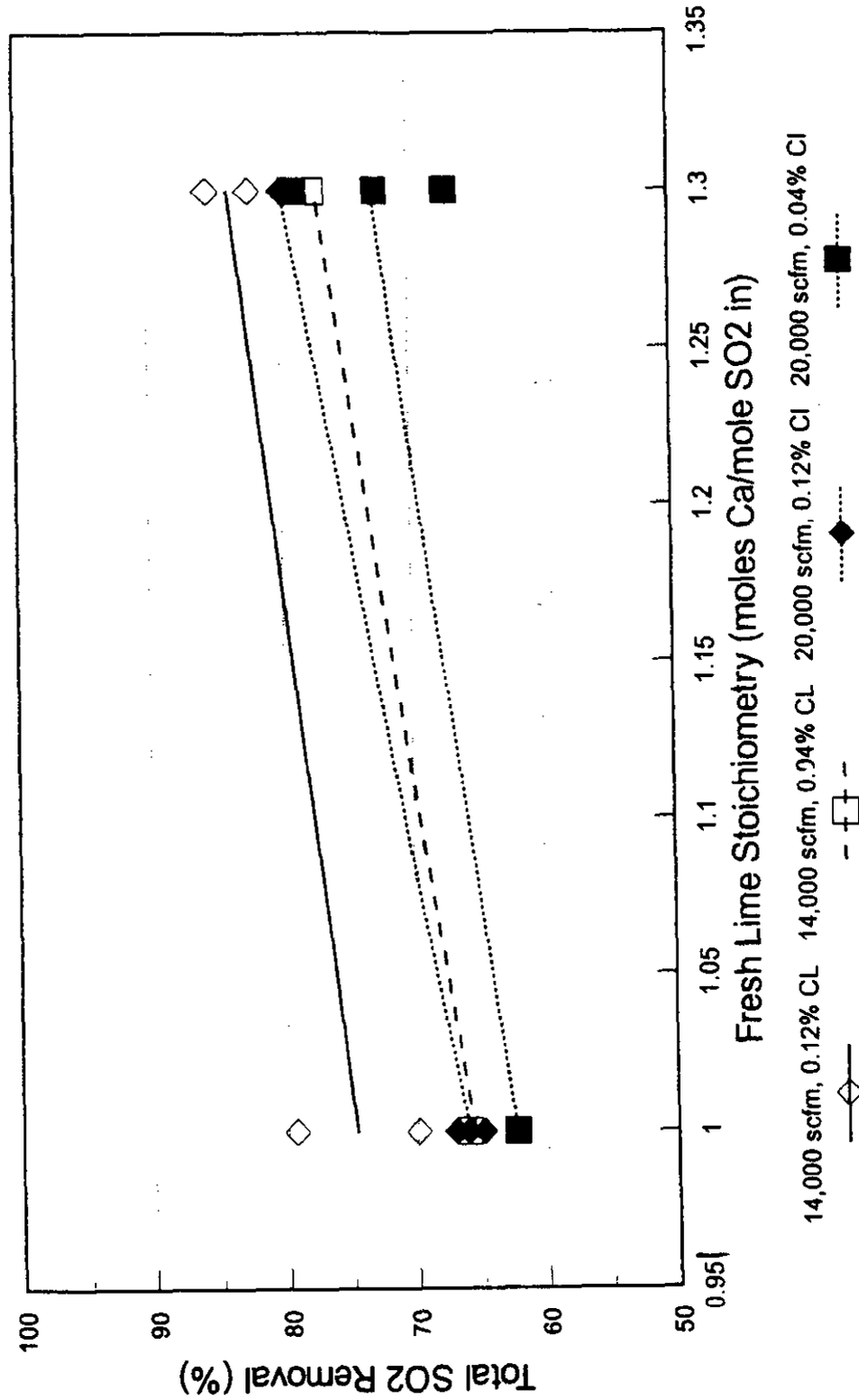


Notes: All tests conducted at a 320°F inlet flue gas temperature, 18F approach temperature.

Figure 5.1.4-3

AirPol GSA SO2 Removal Results

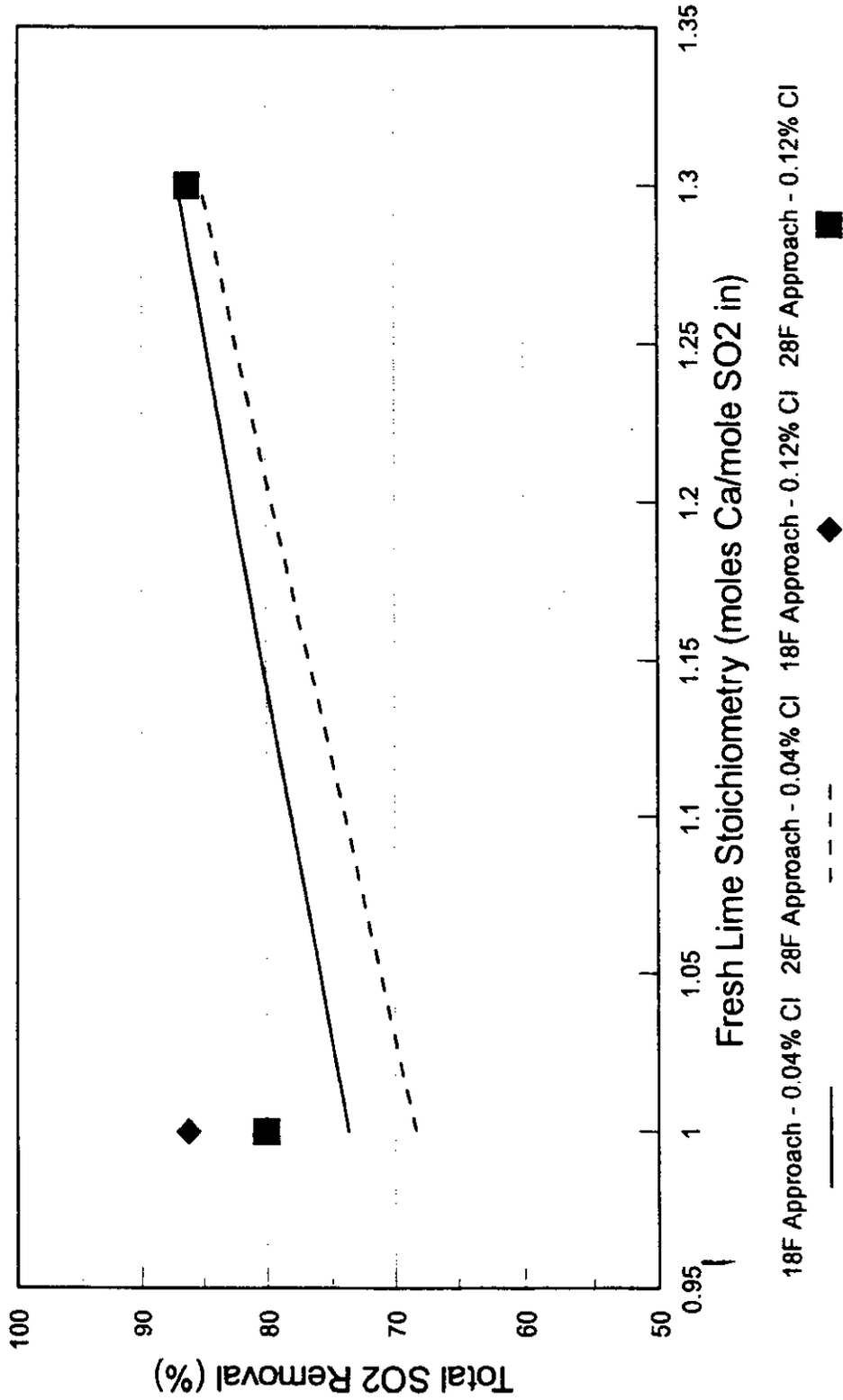
2-AP Series 28F Approach Tests



Notes: All tests conducted at a 320F inlet flue gas temperature, 28F approach temperature.

Figure 5.1.4-4

AirPol GSA SO₂ Removal Results 3-AP Series Baseline/Chloride Spiking Tests



Notes: All tests conducted at a 320F inlet flue gas temperature, 25 wt % solids lime slurry and 14,000 acfm.

5.1.7 Comparison with 10-MW Spray Dryer

Prior to conducting the AirPol GSA demonstration, approximately five years of research and development were conducted at the CER evaluating a 10 MW spray dryer/ESP system. A comparison of the spray dryer/ESP SO₂ removal results and the AirPol SO₂ removal results is presented in Figure 5.1.7. In the figure, the total system SO₂ removal is plotted as a function of fresh lime stoichiometry, which is defined the same for both systems. The results plotted in the figure are for tests that were conducted at a 320°F inlet flue gas temperature, an 18°F approach temperature, a flue gas flow rate of approximately 20,000 scfm at the inlet venturi, and at baseline (low) chloride levels. The spray dryer results plotted in the figure are from tests 5-F-03, -50, -53, -65, -68, -69, -70, and 5-F-71. Also plotted in the figure is a regression line based on the spray dryer model developed by TVA. This model was reported in an April 18, 1991 memorandum entitled, "Preliminary Results of the Remodeling of the Chloride Evaluation Data".

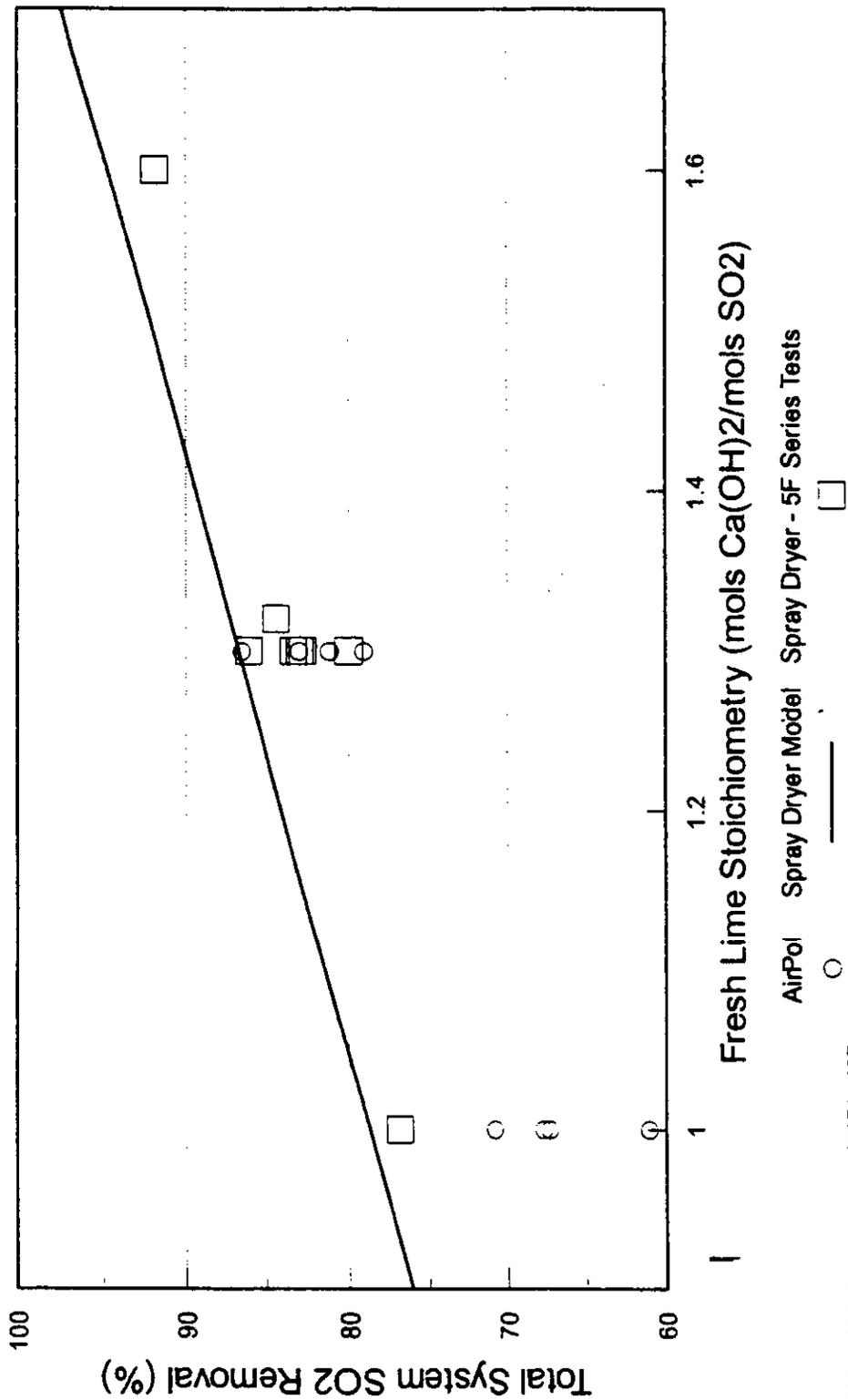
The individual spray dryer test results plotted in Figure 5.1.7 are slightly lower than the regression model because other data at different test conditions were included when developing the model. Based on the data in the figure, the AirPol SO₂ removal performance appears to be lower than the spray dryer/ESP results at a fresh lime stoichiometry of 1.0. At a stoichiometry of 1.3, the AirPol and individual spray dryer test results are virtually identical. The limited test results available at the lower lime stoichiometry do not provide an explanation for the "poorer" performance in the GSA system.

The comparable SO₂ removal in the GSA system means that the GSA technology has another advantage over the conventional spray dryer technology. The flue gas residence time in the GSA reactor is much lower than in the spray dryer (3 versus 10-12 seconds). This means that one can achieve comparable SO₂ removal performance in the GSA system, but with a much smaller (and hence cheaper) reactor. This is a significant advantage for the GSA technology, particularly if additional testing at the lower lime stoichiometry indicates that the apparent differences in performance are due to variability in the data and the limited number of tests.

The fact that the GSA circulating bed provides a very effective reactor for heat and mass transfer is an important factor. The dry recycle solids making up this circulating bed also means that only a small, single two-fluid nozzle is needed to inject the fresh lime slurry. The spray dryer technology, in contrast, requires a single, larger rotary atomizer or multiple two-fluid or rotary atomizers to introduce the combined lime/recycle slurry. Thus, essentially the same SO₂ removal performance can be achieved with a single atomizer.

Figure 5.1.7

Comparison of Spray Dryer and AirPol SO₂ Removal Performance



Notes: All tests conducted at 320F in, 18F approach temperature and 3% S/O.02 to 0.04% Cl coal, and approximately 20,000 scfm in.

5.1.8 28-Day Demonstration Run

As part of the Clean Coal III GSA test plan, a 28-day (approximately 690 hours) demonstration run was performed at one set of operating conditions. This run began on October 25 and was completed on November 24 with one short outage due to a boiler tube leak. The operating conditions selected for this demonstration run were an overall system SO₂ removal set point of 91 percent, an 18°F approach temperature, 20,000 scfm flue gas flow rate at the inlet venturi, 320°F inlet flue gas temperature, 30 rpm recycle screw speed, a fly ash injection rate equivalent to 1.5 gr/acf, and calcium chloride spiking to simulate scrubbing flue gas from a boiler firing a 0.12 weight percent chlorine coal. This demonstration run was divided into 9 test segments to keep the data files manageable. The length of these test segments varied from 1 to 7 days. With the exception of the last two test segments, 1-DR-07 and 1-DR-08, the fresh lime stoichiometry was allowed to fluctuate to meet the target SO₂ removal. The fresh lime stoichiometry was fixed at 1.40 and 1.45 moles Ca(OH)₂/mole inlet SO₂ for test segments 1-DR-07 and 1-DR-08, respectively.

The 1-MW pulse-jet baghouse was down during all of these test segments with the exception of test 3-DR-04. The PJBH was operated for approximately 30 hours during this test segment before being shut down due to failure of approximately 1/3 of the bags. The original plan had been to operate the PJBH during the last two weeks of the demonstration run, but this plan was abandoned.

A summary of the average operating conditions and SO₂ removal performance for the DR series test segments is presented in Table 5.1.8. With the exception of test segments 1-DR-06 and 1-DR-07, the average total system SO₂ removal for all of the test segments was greater than 90 percent. The average fresh lime stoichiometry required to achieve this SO₂ removal varied from 1.32 to 1.58 moles Ca(OH)₂/mole inlet SO₂ and the lime utilization rates ranged from 58 to 69 percent during the demonstration run.

The fluctuation in fresh lime stoichiometry is illustrated more clearly in Figure 5.1.8, which plots the average daily stoichiometry during the demonstration run. As shown in the figure, the average daily stoichiometry ranged from 1.4 to 1.6 moles Ca(OH)₂/mole inlet SO₂. During the last three days of the demonstration run, the stoichiometry was fixed at values of 1.40 and 1.45 moles Ca(OH)₂/mole inlet SO₂.

The demonstration run test conditions were selected based on the results from the prior factorial tests i.e., greater than 90 percent overall SO₂ removal at a reasonable, 1.3 moles Ca(OH)₂/mole inlet SO₂ stoichiometry. These results were obtained in test 2-AP-06, which was conducted in March, and tests 2-AP-91 and 2-AP-92, which were conducted in June. However, during the demonstration run, fresh lime stoichiometries greater than 1.4 moles Ca(OH)₂/mole inlet SO₂ were required to achieve over 90 percent overall SO₂ removal.

There are several possible explanations for this discrepancy in required stoichiometry to achieve greater than 90 percent SO₂ removal. Part of this discrepancy is probably due to unit 9 firing a

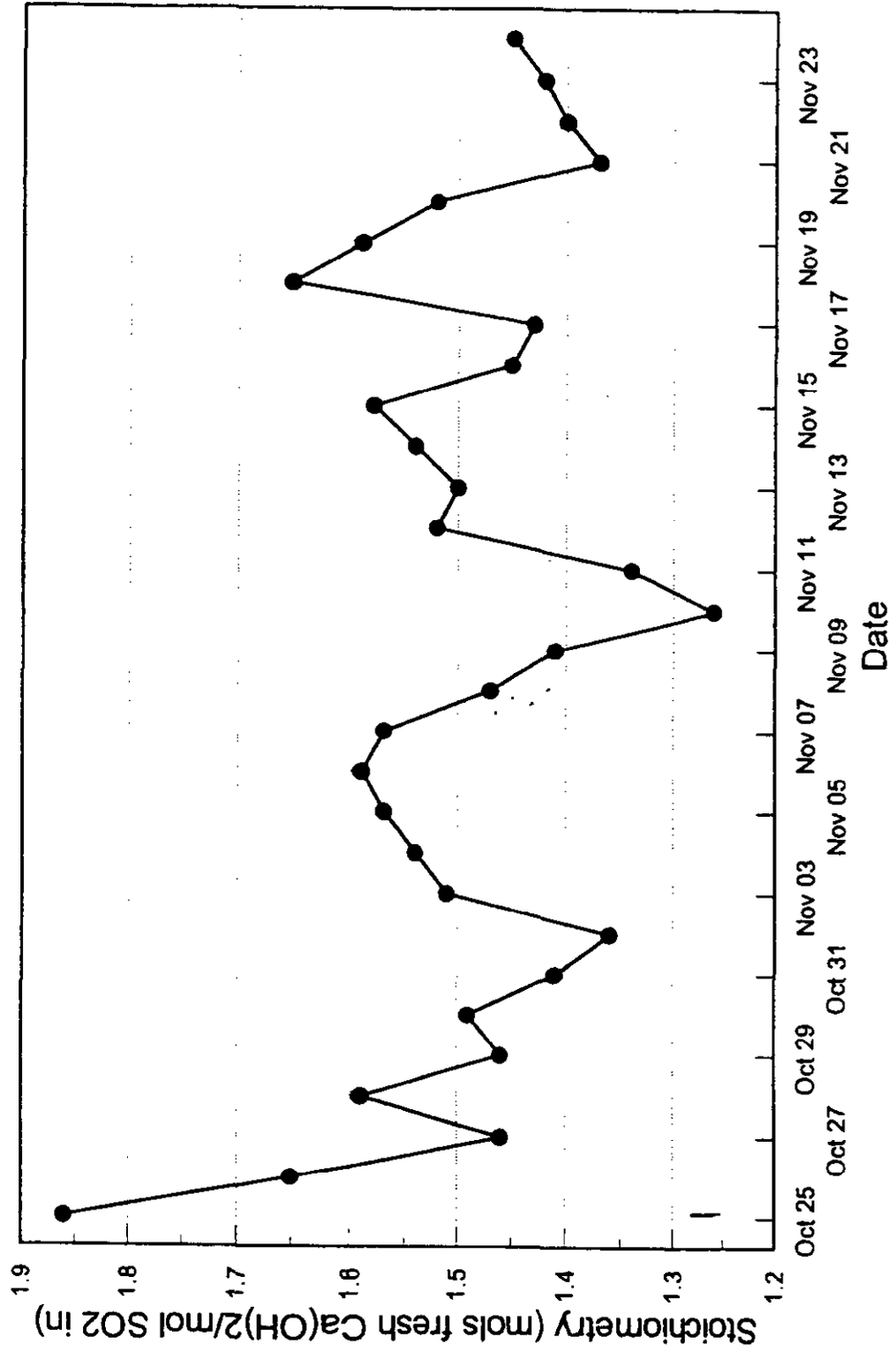
higher sulfur coal during some of the test segments. Approximately one week into the demonstration run on October 31, the supply of Andalex coal was exhausted and the unit was switched to a higher sulfur Warrior coal. The unit continued to burn this higher sulfur coal until November 9. The unit also briefly burned this higher sulfur coal again on November 11, 18, and 22. Based on data from prior tests, an increase in inlet SO₂ concentration resulted in decreased SO₂ removal performance. Thus, the higher lime stoichiometries during these periods, i.e., 1.5-1.6 moles Ca(OH)₂/mole inlet SO₂, were not completely unexpected and these high lime stoichiometries are not a major concern. Also, some of the demonstration test segments were conducted at lower solids chloride levels compared to the factorial tests. Late in the demonstration run, it was discovered that the fly ash loading to the GSA reactor may have been higher than originally planned. The source of this excess ash may have been the higher ash levels in the flue gas entering the GSA system. This higher ash level would dilute both the chloride and the alkalinity levels in the system.

Table 5.1.8
AirPol GSA/ESP SO2 Removal Results Summary
 DR Series - 28 Day Demonstration Run

Test No.	Ca/S Ratio	Reactor Inlet Temperature (F)	Approach Temperature (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Injection Rate (lb/min)	Reactor Product Chloride Content (%)	Reactor/Cyclone SO2 Removal (%)	ESP SO2 Removal (%)	Total System SO2 Removal (%)
1-DR-01	1.46	320	18	20,000	30	7.4	0.27	89.2	1.7	90.9
1-DR-02	1.47	319	18	20,000	30	7.0	0.34	89.0	1.3	90.3
1-DR-03	1.58	319	18	20,000	30	7.3	0.21	91.1	0.5	91.6
1-DR-04	1.48	319	18	20,000	30	7.1	0.19	88.0	3.7	91.7
3-DR-04	1.32	320	18	20,000	30	7.3	0.32	85.7	4.8	90.5
1-DR-05	1.52	319	18	20,000	30	7.2	0.33	87.2	3.0	90.2
1-DR-06	1.43	320	18	20,000	30	7.3	0.30	87.2	2.7	89.9
1-DR-07	1.40	320	18	20,000	30	7.3	0.30	77.0	6.9	83.9
1-DR-08	1.45	320	18	20,000	30	7.4	0.41	88.3	5.2	93.5

Figure 5.1.8

28-Day Demonstration Run Average Daily Stoichiometry



5.1.9 14-Day Pulse-jet Baghouse Demonstration Run

As mentioned in the previous section, the pulse-jet fabric filter was to be operated for two weeks in parallel with the ESP during the 28-day demonstration run. However, due to the failure of the PJBH bag fabric, the PJBH was not operated during this time period. Therefore, the demonstration run conditions were repeated, beginning in February, with the PJBH in operation to evaluate PJBH performance over a longer period of time at one set of operating conditions.

The operating conditions for the PJBH demonstration run were an overall system SO₂ removal set point of 91 percent, an 18°F approach temperature, 20,000 scfm flue gas flow rate at the inlet venturi, 320°F inlet flue gas temperature, 30 rpm recycle screw speed, and calcium chloride spiking to simulate scrubbing flue gas from a boiler firing a 0.12 weight percent chlorine coal. One difference in the operating conditions for the PJBH demonstration run compared to the prior, 28-day demonstration run, was a lower fly ash injection rate. During the 28-day demonstration run, the fly ash injection rate was set to achieve an inlet particulate concentration of 1.5 gr/acf to the GSA reactor. This concentration does not include the fly ash already present in the flue gas. The desired total particulate concentration is 2.0 gr/acf. Since the particulate concentration from Unit 9 is higher while firing the Andalex coal (approximately 1.0 gr/acf versus 0.5 gr/acf with previously fired coals), the fly ash injection rate set point was reduced to 1.0 gr/acf for the PJBH demonstration run.

All of the PJBH demonstration run test segments were conducted while Unit 9 fired medium-sulfur, low-chloride Andalex coal (2.8% S/0.04% Cl). Mississippi pebble lime was used for all the tests and the lime slurry solids concentration set point was 25 percent. The ESP was operated with all four fields in service and the baffle was in place in the fourth field ESP hopper during all test segments.

The PJBH demonstration run was divided into 4 test segments to keep the data files manageable. The length of these test segments varied from 4 to 5 days. One segment of the 1-PJ series demonstration run was completed in February and three test segments were completed during the month of March. A summary of the average operating conditions and SO₂ removal performance is presented in Table 5.1.9-1 for all of the PJBH demonstration test segments.

A plot of the average daily fresh lime stoichiometry during the demonstration run is presented in Figure 5.1.9-1. As shown in the figure, there were two periods when the stoichiometry was significantly higher than the overall demonstration run average of 1.40 moles Ca/mole of inlet SO₂. The first period was from February 28 through March 1. The high stoichiometry during this period was due to the lime flow meter calibration. Based on a flow meter calibration on March 1, the lime flow meter was indicating 4 percent higher than the actual lime flow rate. Therefore, the reported stoichiometry was 4 percent higher than the actual stoichiometry for some period prior to the March 1 calibration. Based on the data, the reported stoichiometry for February 28 may have also been influenced by the lime flow meter calibration.

The second period in which the fresh lime stoichiometry was significantly higher than the overall

test average was after the pilot unit outage from March 8 to March 10. When the PJBH demonstration run resumed on March 12, the fresh lime stoichiometry was very high averaging over 1.7 moles Ca/mole of inlet SO₂. The high stoichiometry required to achieve the 91 percent SO₂ removal set point is probably due to the low chloride concentration in the recycle and ESP solids during the first part of this test segment. Several of the recycle and ESP solid samples on March 12, had either none or very low levels of chloride. The reactor and ESP solids chloride levels did not reach steady state until approximately mid-day on March 13. Therefore, the data from March 12 was not used in developing the test segment averages.

Also influencing the fresh lime stoichiometry was the wet bulb temperature used during the demonstration run. There were several periods during the demonstration run when the approach temperature control was switched between the manual wet bulb measurements and the continuous wet bulb monitor (CWBM). This switching was due to problems with the CWBM in which the two measurements deviated by more than 3°F. Test data in which an inaccurate wet bulb temperature was used for approach temperature control was removed prior to developing the reported test results.

Figure 5.1.9-2 provides a plot of the average total system SO₂ removal for each PJBH demonstration run test segment. The total system SO₂ removal for both the GSA/ESP and the GSA/PJBH configurations are plotted in the figure. The GSA/ESP total SO₂ removal averaged 91.2 percent during the demonstration run. The GSA/PJBH total SO₂ removal, presented in Table 5.1.9-2, was significantly higher and averaged 97.7 percent. Since the GSA/PJBH configuration provides much higher SO₂ removal performance compared to the GSA/ESP, the stoichiometry to achieve 91 percent overall SO₂ removal would be lower than the average of 1.40 moles Ca/mole of inlet SO₂ for the GSA/ESP configuration.

The average total system lime utilization for both the GSA/ESP and GSA/PJBH configurations is presented in Figure 5.1.9-3. The GSA/ESP total system lime utilization averaged 66.1 percent during the demonstration run. The GSA/PJBH total system lime utilization was 4.4 percentage points higher, due to the higher SO₂ removal across the PJBH, and averaged 70.5 percent. Therefore, the GSA/PJBH configuration would be more cost effective in terms of reagent utilization in comparison to the GSA/ESP configuration.

Figure 5.1.9-4 presents the calculated reactor lime utilization and the measured reactor recycle solids calcium utilization for each PJBH demonstration run test segment. As shown in the figure, the calculated and measured utilizations are almost identical for all four test segments. This very good agreement helps to validate the reported SO₂ removal results.

Table 5.1.9-1
AirPol GSA/ESP SO2 Removal Results Summary
 PJ Series - 14 Day Demonstration Run

Test No.	Ca/S Ratio	Reactor Inlet Temperature (F)	Approach Temperature (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Injection Rate (lb/min)	Reactor Product Chloride Content (%)	Reactor/Cyclone SO2 Removal (%)	ESP SO2 Removal (%)	Total System SO2 Removal (%)
1-PJ-01	1.37	319	18	20,000	30	4.9	0.32	88.9	2.5	91.4
1-PJ-02	1.41	319	18	20,000	30	4.9	0.32	88.2	2.7	90.9
1-PJ-03	1.34	320	18	20,000	30	4.9	0.29	88.2	3.1	91.3
1-PJ-04	1.43	319	18	20,000	30	4.9	0.30	90.8	0.5	91.3

Figure 5.1.9-1

14-Day PJBH Demonstration Run

Average Daily Stoichiometry

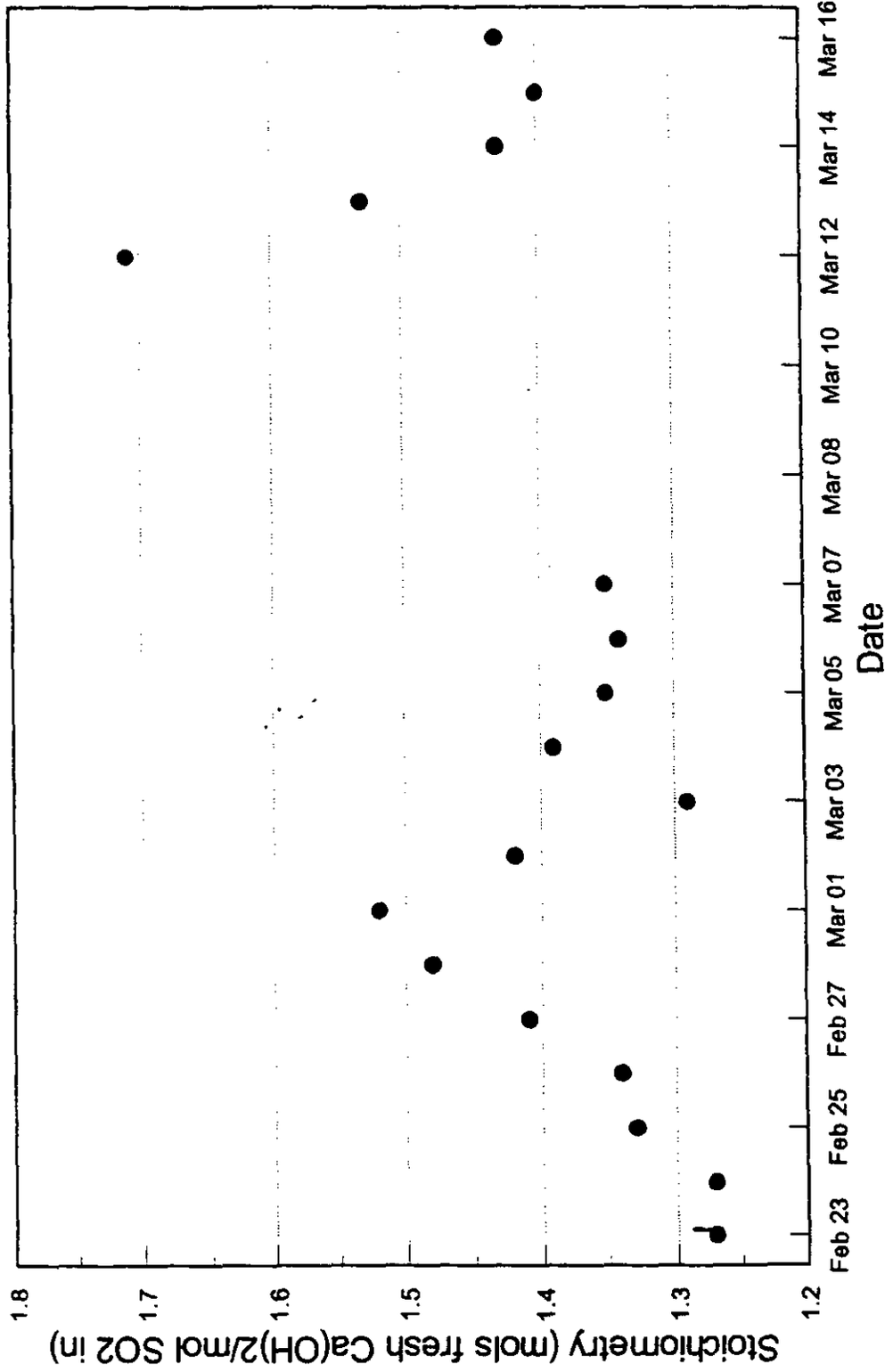


Figure 5.1.9-2

14-Day PJBH Demonstration Run Total System SO2 Removal

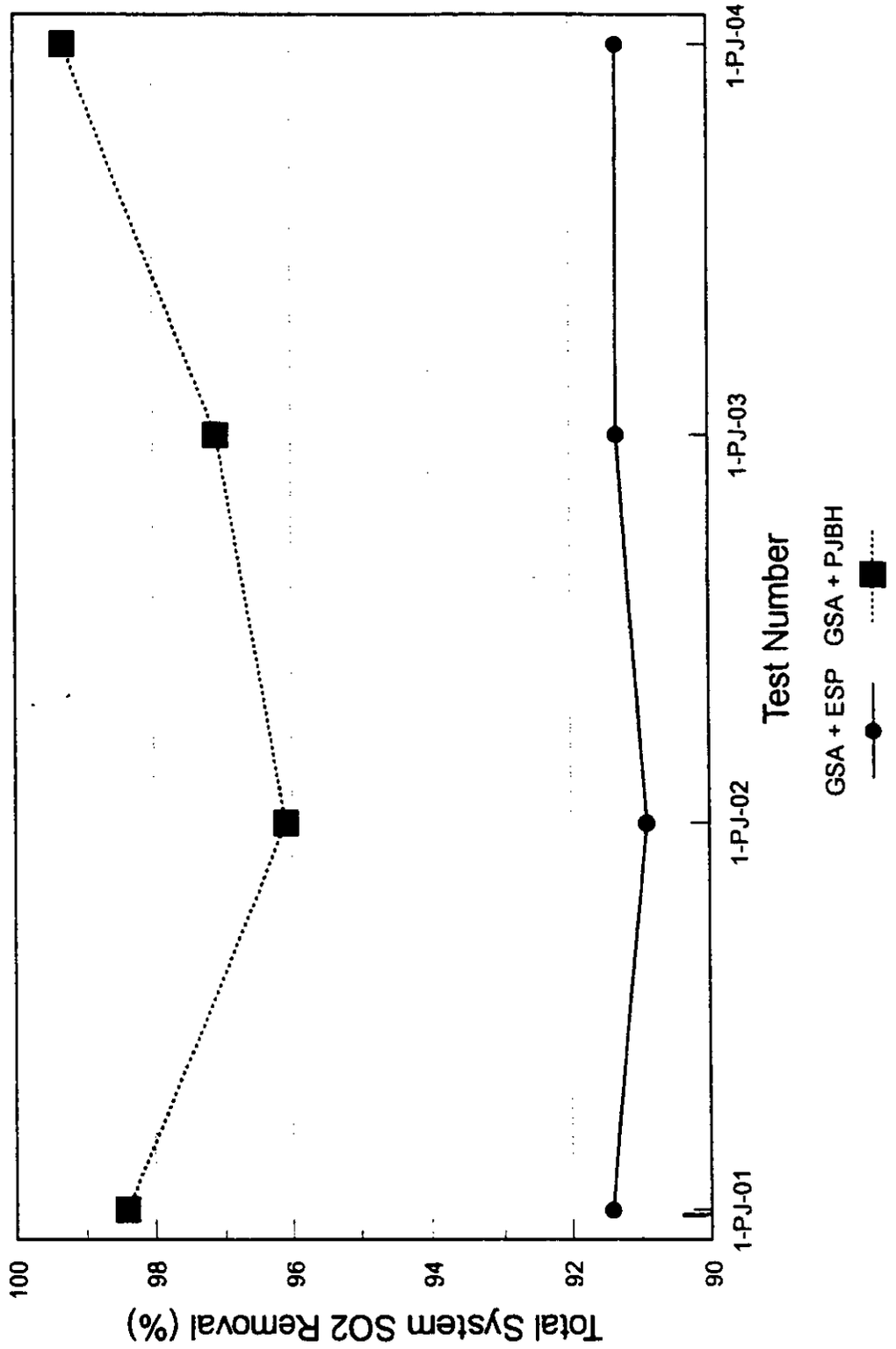


Table 5.1.9-2
AirPol GSA/PJBH SO2 Removal Results Summary
 PJ Series - 14 Day Demonstration Run

Test No.	Ca/S Ratio	Reactor Inlet Temperature (F)	Approach Temperature (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Injection Rate (lb/min)	Reactor Product Chloride Content (%)	Reactor Cyclone SO2 Removal (%)	PJBH SO2 Removal (%)	Total System SO2 Removal (%)
1-PJ-01	1.37	319	18	20,000	30	4.9	0.32	88.9	9.5	98.4
1-PJ-02	1.41	319	18	20,000	30	4.9	0.32	88.2	7.9	96.1
1-PJ-03	1.34	320	18	20,000	30	4.9	0.29	88.2	8.9	97.1
1-PJ-04	1.43	319	18	20,000	30	4.9	0.30	90.8	8.5	99.3

Figure 5.1.9-3

14-Day PJBH Demonstration Run

Total System Lime Utilization

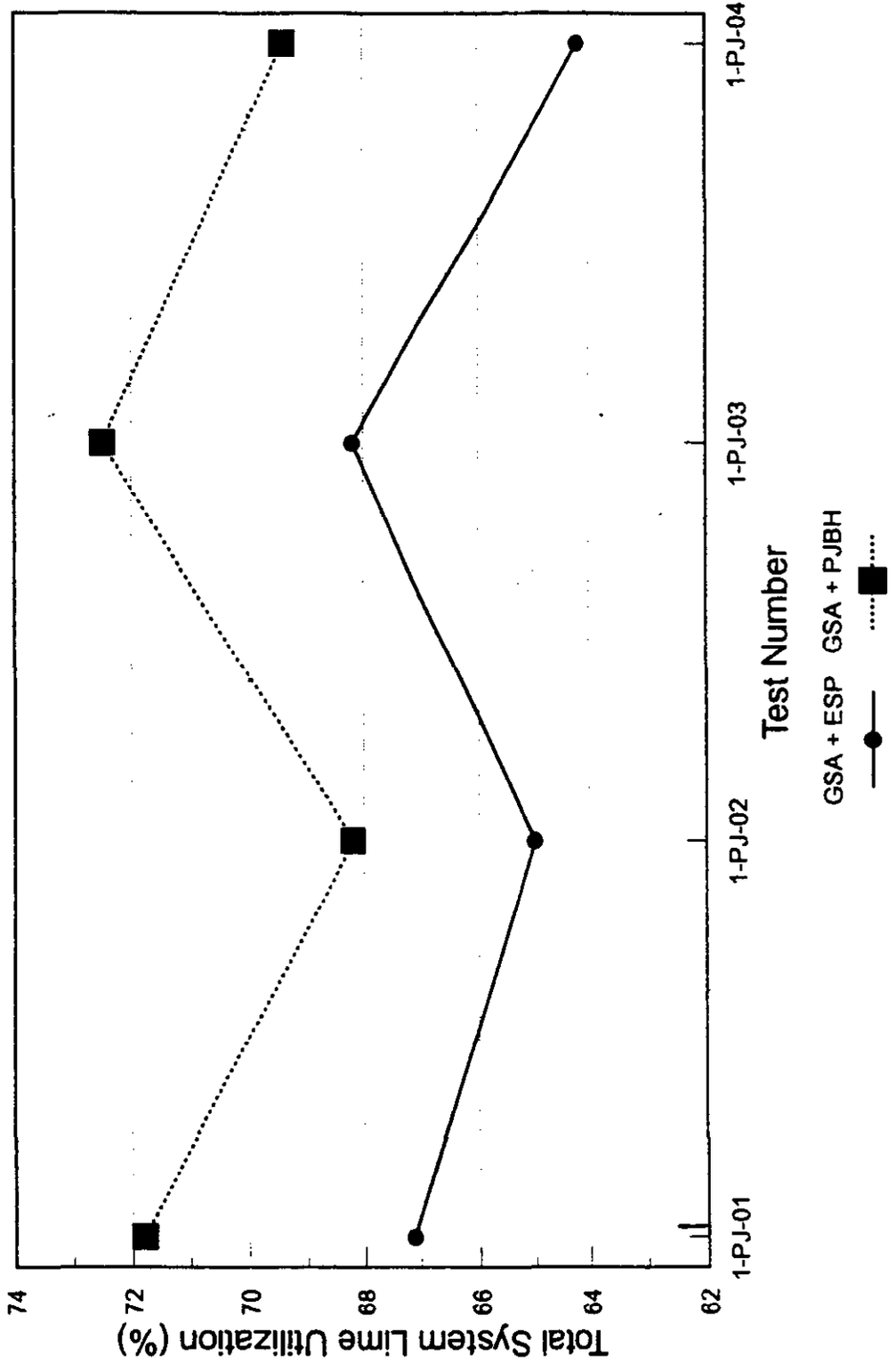
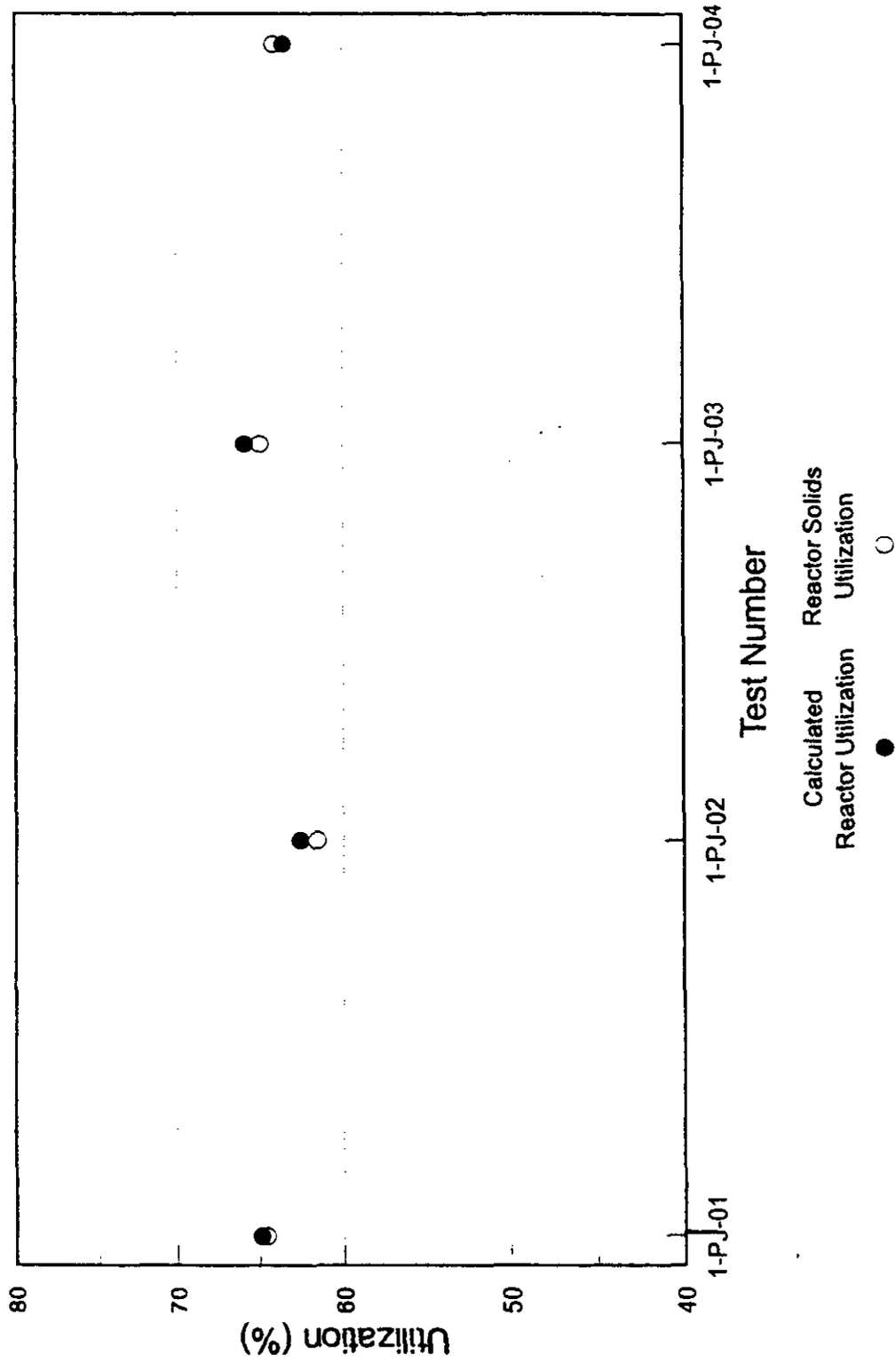


Figure 5.1.9-4

Comparison of Calculated and Measured Reactor Lime Utilizations - 1-PJ Series Tests



5.2 ESP PARTICULATE CONTROL PERFORMANCE

5.2.1 Factorial Tests

The ESP particulate control results for the tests conducted at baseline chloride levels are presented in Table 5.2.1-1 for the 2-AP series tests and in Table 5.2.1-2 for the 3-AP series tests. Similarly, the particulate control results are presented in Tables 5.2.1-3 and 5.2.1-4 for the chloride spiking tests for the 2-AP and 3-AP series, respectively.

The ESP particulate removal results for all of the 2-AP and 3-AP series tests are plotted in Figure 5.2.1-1. In the figure, the ESP emissions in pounds per million Btu (lb/MMBtu) are plotted as a function of ESP specific collection area (SCA). The baseline chloride and calcium chloride spiking test data are separated in the figure. In addition, linear regression lines for each data set are plotted in the figure. In the figure, the outlet emissions typically range from 0.005 to 0.015 lb/MMBtu and they do not appear to decrease with increasing SCA for the baseline tests, as would normally be expected. This could be explained if the emissions were dominated by non-ideal effects, such as sneakage, rapping reentrainment, low resistivity reentrainment, etc., limiting ESP performance. However, for the chloride spiking tests, there does appear to be a decrease in emissions with increasing SCA. If the emissions from baseline test conditions are limited by non-ideal effects, chloride spiking would help to overcome the limitation by making the collected solids more cohesive and improving their ability to stick to the collection plates.

Similar to Figure 5.2.1-1, Figure 5.2.1-2 plots the ESP particulate collection efficiency as a function of ESP specific collection area for all of the 2-AP and 3-AP series tests. The particulate collection efficiency is typically above 99.9 percent. Similar to the particulate emissions, the ESP efficiency does not improve with increasing SCA at baseline conditions but does appear to improve slightly with calcium chloride spiking.

Table 5.2.1-1 AirPol GSA/ESP Particulate Control Performance Results Summary
2-AP Series - Baseline Tests

Test No.	Cw/S Ratio	Reactor Inlet Temp (F)	Approach Temp (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Injection Rate (lb/min)	Reactor Product Chloride Content (%)	ESP Fields	ESP SCA (ft ² /acfm)	ESP Inlet Mass Loading (gr/acf)	ESP Outlet Mass Loading (gr/acf)	ESP Eff Mass Basis (%)	ESP Eff Conc Basis (%)	ESP Emissions (lb/MMBtu)
2-AP-09	1.0	320	8	14,000	30	0	0	4	673	2,598	0.0059	99.77	99.75	0.021
2-AP-79	1.0	320	8	14,000	30	0	0	4	651	2,380	0.0004	99.99	99.98	0.001
2-AP-72	1.0	320	8	20,000	45	7.3	0	4	453	3,938	0.0018	99.96	99.95	0.006
2-AP-16	1.0	320	8	20,000	45	7.8	0.04	4	463	4,455	0.0012	99.97	99.97	0.004
2-AP-11	1.3	320	8	14,000	30	5.0	0	4	664	4,823	0.0019	99.96	99.96	0.007
2-AP-81	1.3	320	8	14,000	30	5.2	0	4	643	3,584	0.0012	99.97	99.96	0.004
2-AP-10	1.3	320	8	20,000	45	0	0	4	460	3,404	0.0027	99.92	99.92	0.009
2-AP-80	1.3	320	8	20,000	45	0	0.09	4	453	3,242	0.0006	99.99	99.98	0.001
2-AP-01	1.0	319	18	14,000	30	0	0	4	652	2,785	0.0017	99.94	99.94	0.006
2-AP-71	1.0	320	18	14,000	30	0	0.02	4	641	2,806	0.0024	99.93	99.91	0.008
2-AP-78	1.0	320	18	20,000	30	7.2	0	4	458	3,266	0.0015	99.96	99.95	0.005
2-AP-04	1.0	320	18	19,000	45	0	0	4	486	2,870	0.0028	99.91	99.90	0.009
2-AP-74	1.0	320	18	20,000	45	0	0	4	445	2,553	0.0022	99.93	99.91	0.007
2-AP-03	1.3	319	18	14,000	45	0	0	4	638	3,810	0.0041	99.90	99.88	0.014
2-AP-73	1.3	320	18	14,000	45	5.1	0	4	641	4,566	0.0046	99.91	99.89	0.017
2-AP-95	1.3	320	18	20,000	45	0	0.05	4	448	2,447	0.0014	99.95	99.94	0.005
2-AP-96	1.3	322	18	20,000	45	0	0.21	4	448	2,477	0.0026	99.91	99.89	0.009
2-AP-14	1.3	320	18	18,000	45	6.7	0.12	4	491	5,644	0.0030	99.95	99.94	0.010
2-AP-63	1.3	320	18	20,000	45	7.2	0.04	4	456	4,681	0.0026	99.95	99.94	0.009
2-AP-63	1.3	320	18	20,000	45	7.2	0.03	4	447	5,069	0.0039	99.93	99.92	0.013
2-AP-88	1.0	320	28	14,000	30	0	0.04	4	642	2,283	0.0047	99.80	99.78	0.015
2-AP-87	1.0	320	28	20,000	45	7.3	0	4	446	2,995	0.0017	99.95	99.94	0.006
2-AP-86	1.3	320	28	14,000	45	0	0.03	4	629	3,065	0.0022	99.94	99.93	0.007
2-AP-97	1.3	320	28	20,000	30	0	0	4	442	2,042	0.0089	99.45	99.37	0.029
2-AP-19	1.3	320	28	20,000	30	7.1	0	4	463	4,443	0.0028	99.95	99.93	0.010
2-AP-57	1.3	319	28	19,500	30	7.1	0	4	463	4,665	0.0030	99.94	99.93	0.010

**Table 5.2.1-2
AirPol GSA/ESP Particulate Control Performance Results Summary
3-AP Series - Baseline Tests**

Test No.	Ca/S Ratio	Reactor Inlet Temp (F)	Approach Temp (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Injection Rate (lb/min)	Reactor Product Chloride Content (%)	ESP Fields	ESP SCA (ft ² /kacfm)	ESP Inlet Mass Loading (gr/acf)	ESP Outlet Mass Loading (gr/acf)	ESP Eff Mass Basis (%)	ESP Eff Conc Basis (%)	ESP Emissions (lb/MMBtu)
3-AP-12	1.0	320	8	14,000	30	0	0	4	857	3.352	0.0091	99.78	99.70	0.030
3-AP-58	1.0	320	8	14,000	30	0	0	4	835	3.022	0.0016	99.96	99.94	0.005
3-AP-42	1.3	320	8	14,000	30	0	0	4	864	3.645	0.0038	99.92	99.89	0.012
3-AP-11	1.3	319	8	14,000	30	5.2	0	4	840	-	-	-	-	-
3-AP-62	1.0	319	18	14,000	30	0	0.13	4	850	-	-	-	-	-
3-AP-08	1.0	320	18	20,000	30	7.0	0	4	543	3.797	0.0024	99.95	99.93	0.009
3-AP-44	1.3	319	18	14,000	45	0	0.02	4	829	0.606	0.0022	99.71	99.59	0.008
3-AP-03	1.3	319	18	14,000	45	5.0	0	4	861	4.896	0.0026	99.96	99.95	0.010
3-AP-26	1.3	260	18	14,000	12	4.7	0	4	911	3.561	0.0045	99.91	99.87	0.016
3-AP-27	1.3	260	18	14,000	30	4.7	0	4	893	-	-	-	-	-
3-AP-02	1.3	319	18	14,000	45	5.1	0	4	823	5.303	0.0024	99.97	99.95	0.008
3-AP-18	1.0	319	28	14,000	30	0	0	4	844	2.932	0.0020	99.95	99.94	0.007
3-AP-59	1.0	320	28	14,000	30	5.2	0	4	844	-	-	-	-	-
3-AP-21	1.0	319	28	20,000	45	7.3	0	4	457	4.412	0.0026	99.95	99.94	0.010
3-AP-56	1.3	320	28	14,000	30	0	0	4	855	-	-	-	-	-
3-AP-20	1.3	320	28	14,000	45	0	0	4	815	4.765	0.0040	99.92	99.90	0.014
3-AP-20	1.3	320	28	14,000	45	0	0.03	4	826	3.535	0.0042	99.91	99.88	0.014
3-AP-13	1.3	319	28	14,000	45	0	0	4	839	3.323	0.0025	99.94	99.92	0.009
3-AP-45	1.3	320	28	14,000	45	5.2	0	4	805	5.600	0.0018	99.97	99.97	0.006
3-AP-19	1.3	320	28	20,000	30	7.4	0	4	525	3.024	0.0027	99.93	99.91	0.009

**Table 5.2.1-3
AirPol GSA/ESP Particulate Control Performance Results Summary
2-AP Series - Chloride Spiking Tests**

Test No.	Ca/S Ratio	Reactor Inlet Temp (F)	Approach Temp (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Injection Rate (lb/min)	Reactor Product Chloride Content (%)	ESP Fields	ESP SCA (ft ² /kacfm)	ESP Inlet Mass Loading (gr/acf)	ESP Outlet Mass Loading (gr/acf)	ESP Eff Mass Basis (%)	ESP Eff Conc Basis (%)	ESP Emissions (lb/MMBtu)
2-AP-28	1.0	320	18	14,000	40	5.0	0.39	4	646	4.972	0.0027	99.95	99.94	0.009
2-AP-75	1.0	320	18	14,000	45	4.7	0.45	4	635	4.516	0.0028	99.95	99.94	0.010
2-AP-17	1.0	320	18	20,000	30	0	0.75	4	452	2.461	0.0033	99.89	99.86	0.011
2-AP-82	1.0	320	18	20,000	30	0	0.87	4	448	2.585	0.0015	99.95	99.94	0.004
2-AP-07	1.3	320	18	14,000	30	5.4	0.29	4	642	5.089	0.0025	99.95	99.95	0.008
2-AP-77	1.3	320	18	14,000	30	5.0	0.30	4	626	3.471	0.0118	99.96	99.95	0.005
2-AP-98	1.3	320	18	14,000	30	5.2	0.30	4	641	4.544	0.0029	99.95	99.95	0.007
2-AP-06	1.3	320	18	20,000	45	0	0.47	4	444	3.853	0.0037	99.91	99.90	0.012
2-AP-92	1.3	320	18	20,000	45	0	0.72	4	445	3.313	0.0008	99.98	99.98	0.003
2-AP-91	1.3	320	18	20,000	45	7.2	0.32	4	452	4.532	0.0018	99.97	99.96	0.006
2-AP-22	1.0	320	28	14,000	45	0	0.57	4	626	-	-	-	-	-
2-AP-90	1.0	320	28	14,000	45	0	0.86	4	628	2.431	0.0020	99.93	99.91	0.007
2-AP-25	1.0	320	28	18,600	30	6.8	0.28	4	472	4.763	0.0041	99.92	99.91	0.014
2-AP-94	1.0	320	28	20,000	30	0	0.81	4	443	2.400	0.0039	99.86	99.83	0.013
2-AP-85	1.0	320	28	20,000	30	7.4	0.31	4	441	3.147	0.0025	99.93	99.92	0.008
2-AP-84	1.3	320	28	14,000	30	0	0.60	4	623	2.630	0.0012	99.96	99.95	0.004
2-AP-24	1.3	320	28	14,000	30	0	0.41	4	631	3.471	0.0033	99.91	99.90	0.011
2-AP-83	1.3	320	28	20,000	30	7.3	0.26	4	431	3.359	0.0026	99.93	99.92	0.009

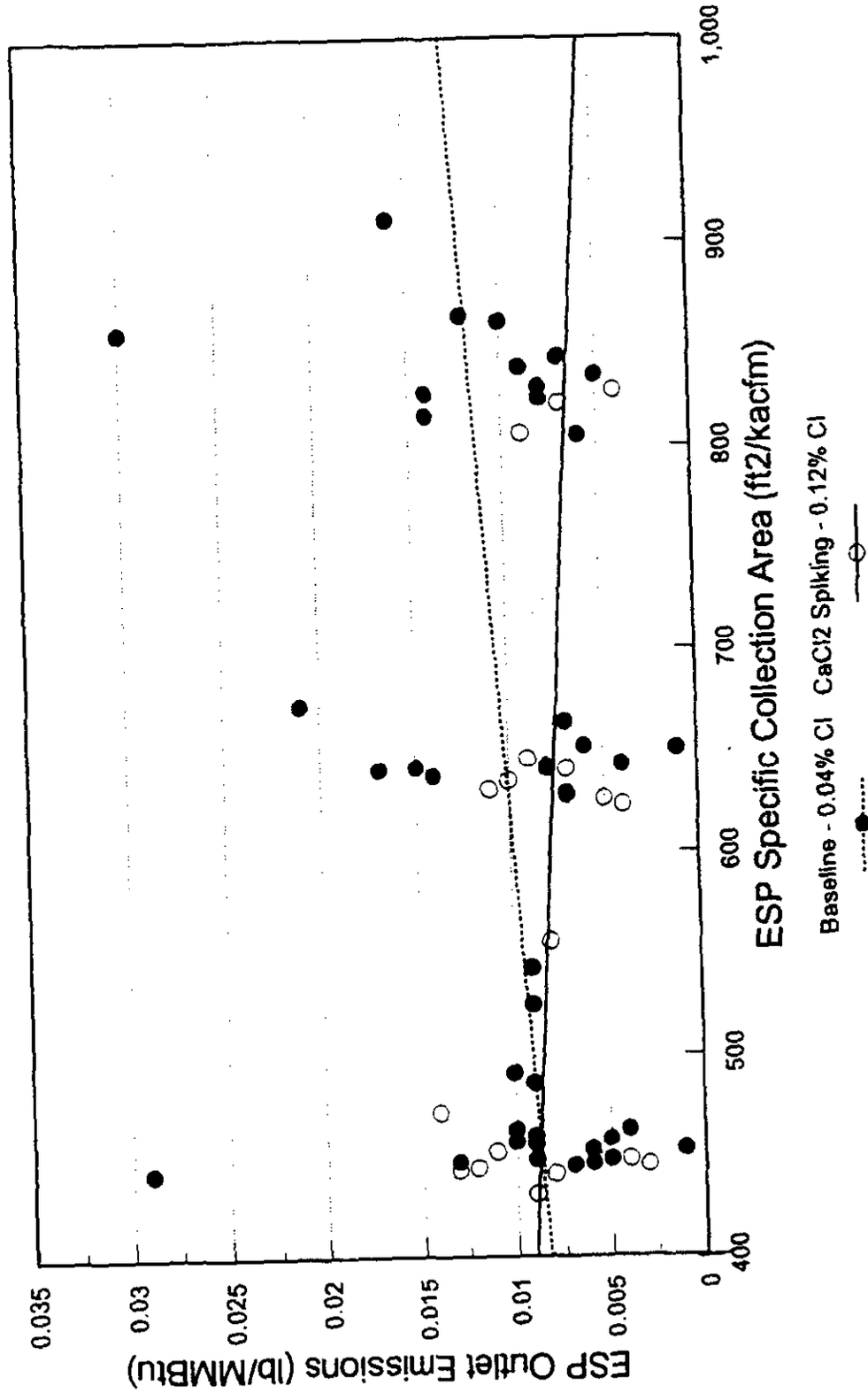
**Table 5.2.1-4
AirPol GSA/ESP Particulate Control Performance Results Summary
3-AP Series - Chloride Spiking Tests**

Test No.	Ca/S Ratio	Reactor Inlet Temp. (F)	Approach Temp. (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Injection Rate (lb/min)	Reactor Product Chloride Content (%)	ESP Fields	ESP SCA (ft ² /kacfm)	ESP Inlet Mass Loading (gr/acf)	ESP Outlet Mass Loading (gr/acf)	ESP Eff. Mass Basis (%)	ESP Eff. Conc. Basis (%)	ESP Emissions (lb/MMBtu)
3-AP-29	1.0	320	18	14,000	45	5.2	0.31	4	806	5.357	0.0024	99.96	99.95	0.009
3-AP-22	1.0	320	28	14,000	45	0	0.57	4	827	3.201	0.0013	99.97	99.96	0.004
3-AP-24	1.3	320	28	14,000	30	0	0.46	4	821	3.321	0.0020	99.95	99.94	0.007
3-AP-23	1.3	320	28	19,200	45	7.2	0.26	4	556	5.091	0.0025	99.96	99.95	0.008

Figure 5.2.1.1-1

AirPol GSA ESP Performance Results

ESP Particulate Emissions versus SCA

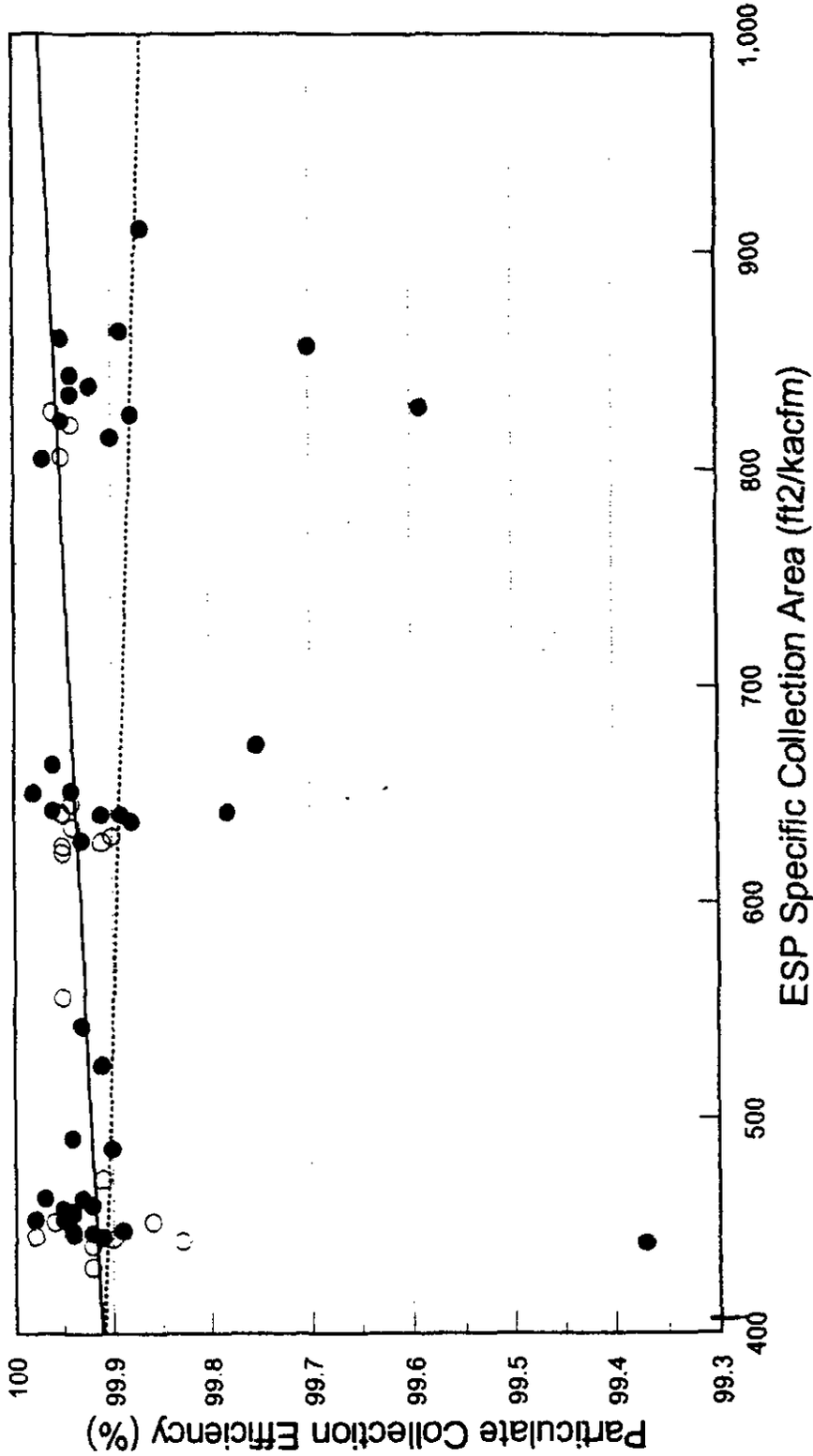


Notes: Data from 2-AP and 3-AP series tests.

Figure 5.2.1-2

AirPol GSA ESP Performance Results

Particulate Collection Efficiency versus SCA



Notes: Data from 2-AP and 3-AP series tests.

5.2.2 Comparison with 10-MW Spray Dryer

Figures 5.2.2-1 and 5.2.2-2 compare the ESP particulate control performance of the AirPol GSA system and the prior spray dryer system. Figure 5.2.2-1 plots the ESP particulate emissions as a function of SCA for both baseline and chloride spiking tests. Similar particulate emissions are observed for both systems at SCAs ranging from 400 to 500 ft²/kacfm. Figure 5.2.2-2, which plots the ESP particulate removal as a function of SCA, also shows that the ESP removal for both systems is approximately the same at SCAs of 400 to 500 ft²/kacfm. These figures also point out the deterioration in particulate control performance at SCAs below 400 ft²/kacfm for the spray dryer system. It is important to determine whether a similar deterioration will be observed with the AirPol system, since most FGD retrofit applications involving ESPs would be in the 200 to 400 ft²/kacfm SCA size range. There were indications during the demonstration run, which is discussed in the following sections, that the ESP performance will deteriorate at lower SCA levels.

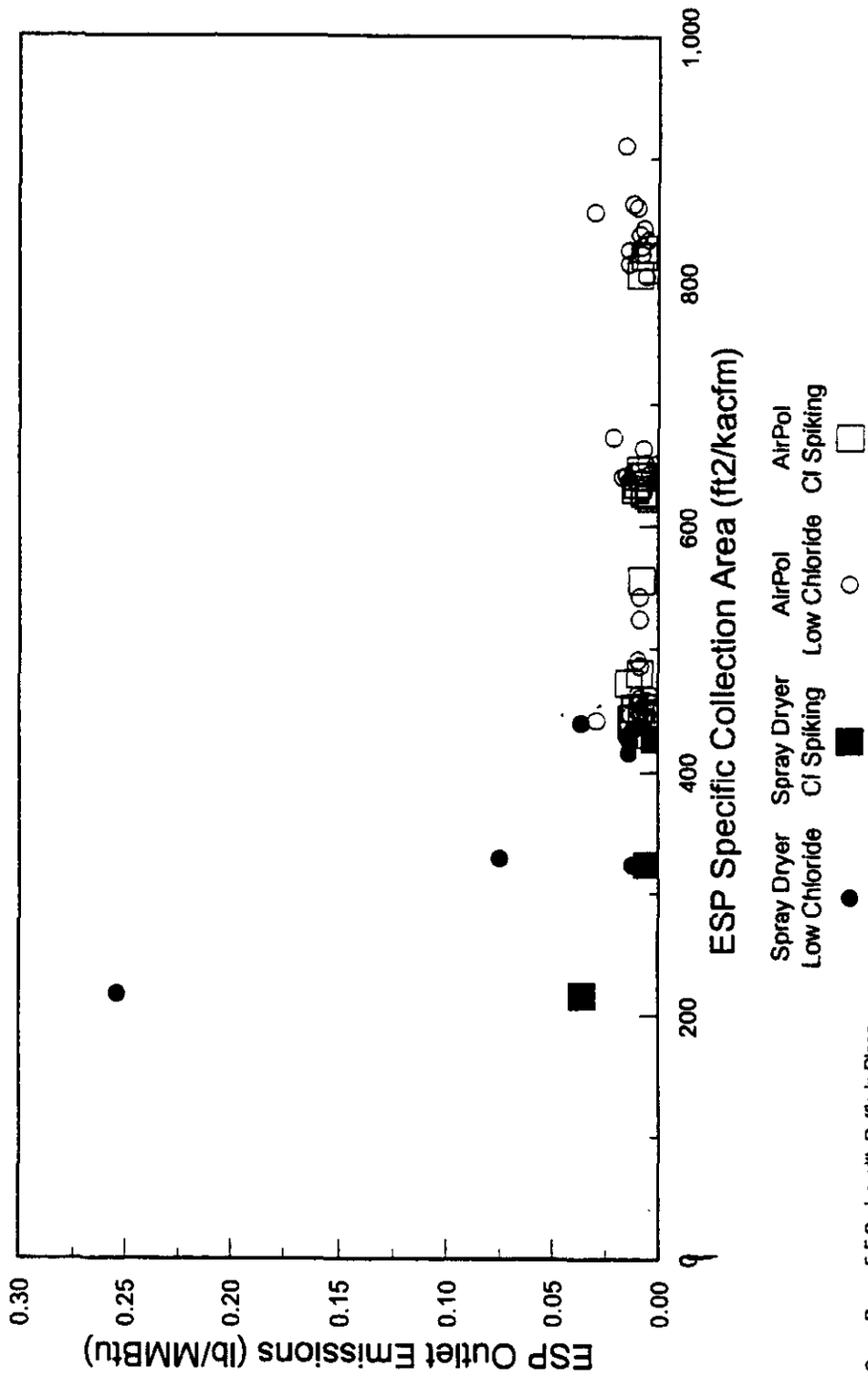
The major difference between these technologies is that the GSA system has a cyclone installed immediately downstream of the GSA reactor to reduce the inlet grain loading entering the ESP. The inlet grain loadings entering the ESP during the GSA testing ranged from 3-5 gr/acf versus 6-10 gr/acf during the spray dryer testing. These lower inlet grain loadings mean that the ESP can achieve the required emission regulations with a lower particulate removal efficiency than would be required with the spray dryer system, which is another advantage for the GSA technology.

However, the cyclone removes the larger particles and a higher proportion of the particles entering the ESP are the smaller, more difficult to remove particles. This larger proportion of smaller particles may contribute to the lower current levels in the first field of the ESP (i.e., current suppression) that were noted in the GSA testing. There were lower currents in the first field of the ESP during the spray dryer testing, but not to the low levels seen in the GSA testing. The higher proportion of smaller particles may also have contributed to the apparent increase in the total ESP emissions that seem to be due to the non-ideal effects as shown by the insensitivity of the emissions to significant changes in the ESP SCA above 400 ft²/kacfm.

Figure 5.2.2-1

Comparison of Spray Dryer and AirPol

ESP Particulate Control Performance

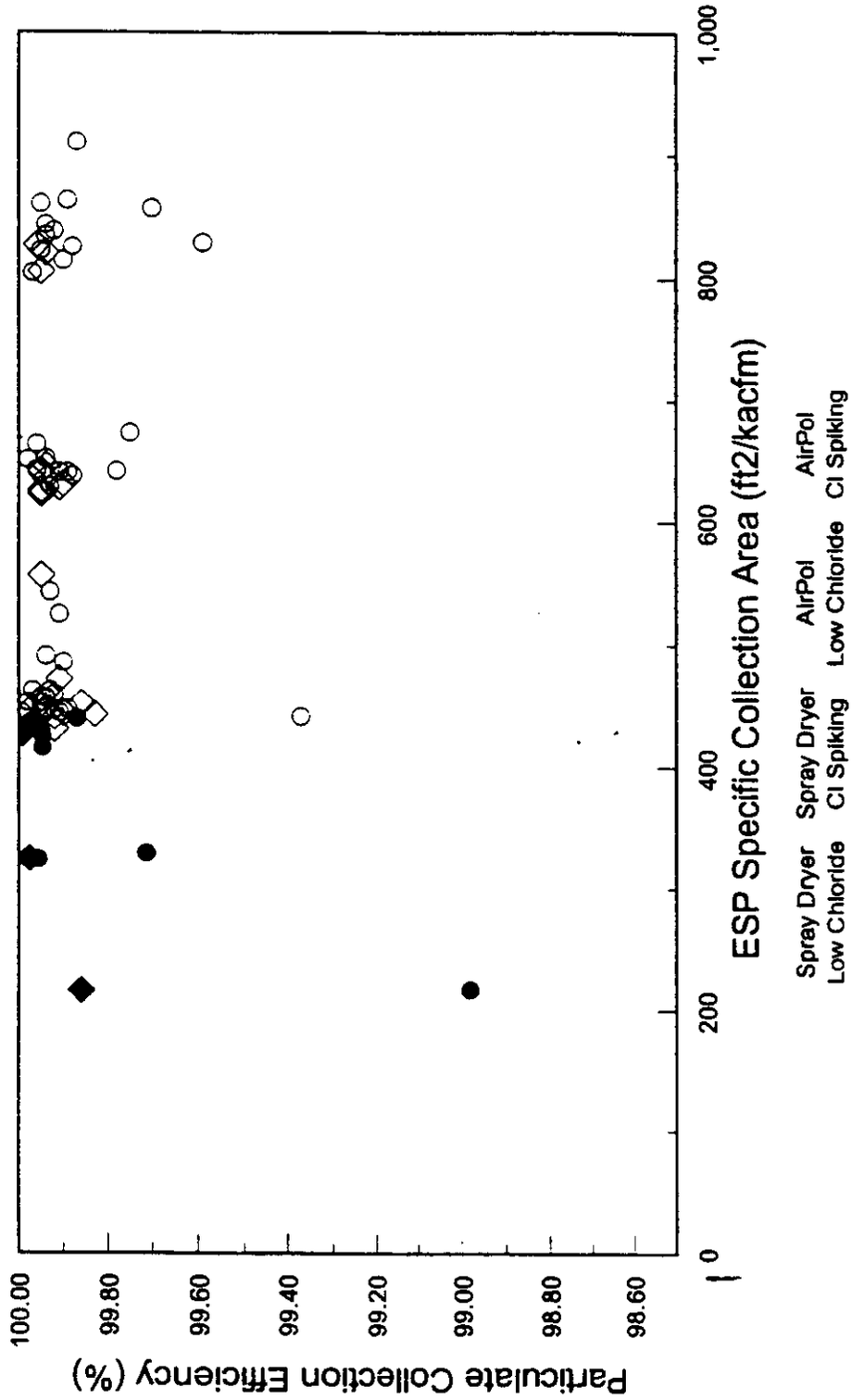


Spray Dryer: 5-F Series with Baffle in Place

Figure 5.2.2-2

Comparison of Spray Dryer and AirPol

ESP Particulate Collection Efficiency



Spray Dryer: 5-F Series with Baffle in Place

5.2.3 28-Day Demonstration Run

A summary of the ESP particulate control results for the 28-day demonstration run is presented in Table 5.2.3. Based on these results, there was a significant decrease in ESP particulate control performance during the 28-day demonstration run. This decrease in performance is illustrated in Figures 5.2.3-1 and 5.2.3-2. In Figure 5.2.3-1, the ESP particulate collection efficiency is plotted for each test segment. Included in the figure is both the test average and the individual mass loading test results. The average ESP particulate collection efficiency was greater than 99.95 percent through the first five test segments (1-DR-01 to 3-DR-04). The last three test segments, 1-DR-05 through 1-DR-07, exhibited poorer performance with the particulate collection efficiency averaging approximately 99.90 percent. Effectively, the particulate penetration doubled (0.05 versus 0.10 percent penetration) during the last three test segments.

Figure 5.2.3-2 presents the ESP particulate emissions for each test segment. Similar to Figure 5.2.3-1, both the test average and individual test data are presented. Concurrent with the poorer ESP particulate removal efficiency, an increase in ESP particulate emissions was also observed for the last three test segments as the ESP emissions increased from approximately 0.006 to 0.015 lb/MMBtu.

The poorer ESP particulate control performance is due to a decrease in power levels in all fields. The reduction in power levels in the first two fields was due to solids build-up on the hopper ridges between fields 1 and 2 and between fields 2 and 3. The reduction in power levels in fields 3 and 4 were due to the increased particulate loading to these fields. A more complete discussion of the ESP operation during the demonstration run is presented in the next section.

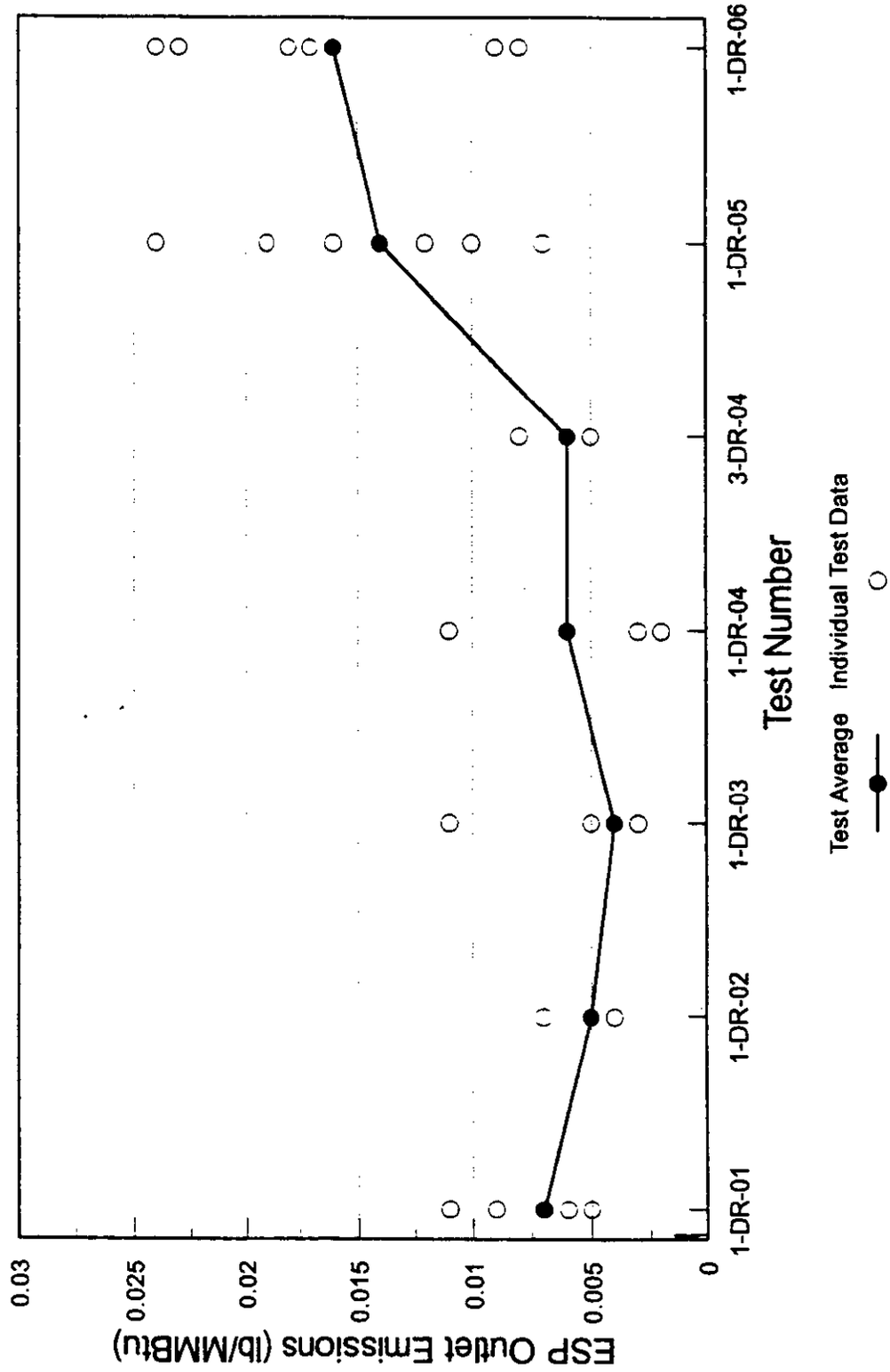
**Table 5.2.3
AirPol GSA/ESP Particulate Control Performance Results Summary
DR Series - 28 Day Demonstration Run**

Test No.	Ca/S Ratio	Reactor Inlet Temp (F)	Approach Temp (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Injection Rate (lb/min)	Reactor Product Chloride Content (%)	ESP Fields	ESP SCA (tq/kacfm)	ESP Inlet Mass Loading (gr/acf)	ESP Outlet Mass Loading (gr/acf)	ESP Eff Mass Basis (%)	ESP Eff Conc Basis (%)	ESP Emissions (lb/MMBtu)
1-DR-01	1.46	320	18	20,000	30	7.4	0.27	4	440	4.608	0.0021	99.96	99.95	0.007
1-DR-02	1.47	319	18	20,000	30	7.0	0.34	4	449	4.612	0.0015	99.97	99.97	0.005
1-DR-03	1.58	319	18	20,000	30	7.3	0.21	4	439	4.563	0.0013	99.97	99.97	0.004
1-DR-04	1.48	319	18	20,000	30	7.1	0.19	4	450	4.253	0.0017	99.96	99.96	0.006
3-DR-04	1.32	320	18	20,000	30	7.3	0.32	4	465	4.226	0.0019	99.96	99.95	0.006
1-DR-05	1.52	319	18	20,000	30	7.2	0.33	4	434	4.858	0.0042	99.91	99.91	0.014
1-DR-06	1.43	320	18	20,000	30	7.3	0.30	4	436	4.387	0.0050	99.89	99.89	0.016
1-DR-07	1.40	320	18	20,000	30	7.3	0.30	4	425	-	-	-	-	-
1-DR-08	1.45	320	18	20,000	30	7.4	0.41	4	441	-	-	-	-	-

Figure 5.2.3.3-2

28-Day Demonstration Run ESP Performance Results

ESP Particulate Emissions



5.2.4 14-Day Pulse-jet Baghouse Demonstration Run

Similar to the 28-day demonstration run, the ESP particulate collection efficiency deteriorated during the 14-day PJBH demonstration run. A summary of the ESP particulate control results for the 14-day PJBH demonstration run is presented in Table 5.2.4-1. This deterioration in performance is shown in Figures 5.2.4-1 and 5.2.4-2. In Figure 5.2.4-1, the ESP particulate removal is plotted for each test segment. Both the test segment average and the individual mass load removal efficiencies are plotted in the figure. As shown in the figure, the average ESP particulate removal efficiency decreased from 99.96 percent in test segment 1-PJ-01 to 99.89 percent in test segment 1-PJ-04. As would be expected with a decrease in the particulate removal efficiency, the ESP particulate emissions increased during the demonstration run. The increase in particulate emissions is presented in Figure 5.2.4-2. As shown in the figure, the average particulate emissions increased from 0.006 to 0.017 lb/MMBtu. Both figures indicate that the ESP particulate control performance was still deteriorating at the conclusion of the demonstration run.

The reason for the deterioration in ESP particulate control performance is not clear. In the prior 28-day demonstration run, the deterioration in particulate control performance was attributed to solids build-up on the hopper ridges between fields 1 and 2 and fields 2 and 3 electrically shorting out fields 1 and 2. However, this did not occur during the PJBH demonstration run. The ESP was inspected on March 9, between test segments 1-PJ-03 and 1-PJ-04, and on March 25. Although some solids build-up was observed on the hopper ridges during these inspections, it did not extend up into the plates and wires.

During the March 9 inspection, however, the wires in the first field were heavily coated with solids. Some of the wires had solids build-up to 3/4 inch thick. The cause of the build-up was that the wires were not being rapped in the first field due to failure of the coupling between the rapper drive motor and the rappers. Apparently the first field wires had not been rapped since February 1, when the rapper drive motor failed and was subsequently replaced. However, even after the first field rappers were repaired, the ESP particulate control performance continued to deteriorate. Therefore, it does not appear that the build-up on the first field wires was influencing the ESP particulate control performance.

Contrary to the ESP particulate control performance, the PJBH did not exhibit a decrease in performance during the demonstration run. Based on the data in Table 5.2.4-2, the particulate removal efficiency and outlet emissions averaged 99.99 percent and 0.0017 lb/MMBtu, respectively. Figures 5.2.4-3 and 5.2.4-4 compare the ESP and PJBH particulate control performance during the demonstration run. In Figure 5.2.4-3, the average ESP and PJBH particulate removal efficiency for each test segment is plotted. Figure 5.2.4-4 plots the average ESP and PJBH outlet particulate emissions for each test segment. As shown in each figure, the PJBH particulate control performance was superior to the ESP. In addition, the PJBH particulate control performance did not deteriorate during the demonstration run.

**Table 5.2.4-1
AirPol GSA/ESP Particulate Control Performance Results Summary
PJ Series - 14 Day Demonstration Run**

Test No.	Ca/S Ratio	Reactor Inlet Temp (F)	Approach Temp (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Injection Rate (lb/min)	Reactor Product Chloride Content (%)	ESP Fields	ESP SCA (ft ² /kacfm)	ESP Inlet Mass Loading (gr/acf)	ESP Outlet Mass Loading (gr/acf)	ESP Eff Mass Basis (%)	ESP Eff Conc Basis (%)	ESP Emissions (lb/MMBtu)
1-PJ-01	1.37	319	18	20,000	30	4.9	0.32	4	531	4.533	0.0017	99.97	99.96	0.006
1-PJ-02	1.41	319	18	20,000	30	4.9	0.32	4	534	4.188	0.0027	99.94	99.93	0.010
1-PJ-03	1.34	320	18	20,000	30	4.9	0.29	4	536	4.543	0.0042	99.92	99.90	0.014
1-PJ-04	1.43	319	18	20,000	30	4.9	0.30	4	520	4.439	0.0049	99.91	99.89	0.017

Figure 5.2.4-1

14-Day PJBH Demonstration Run ESP Performance

ESP Particulate Removal

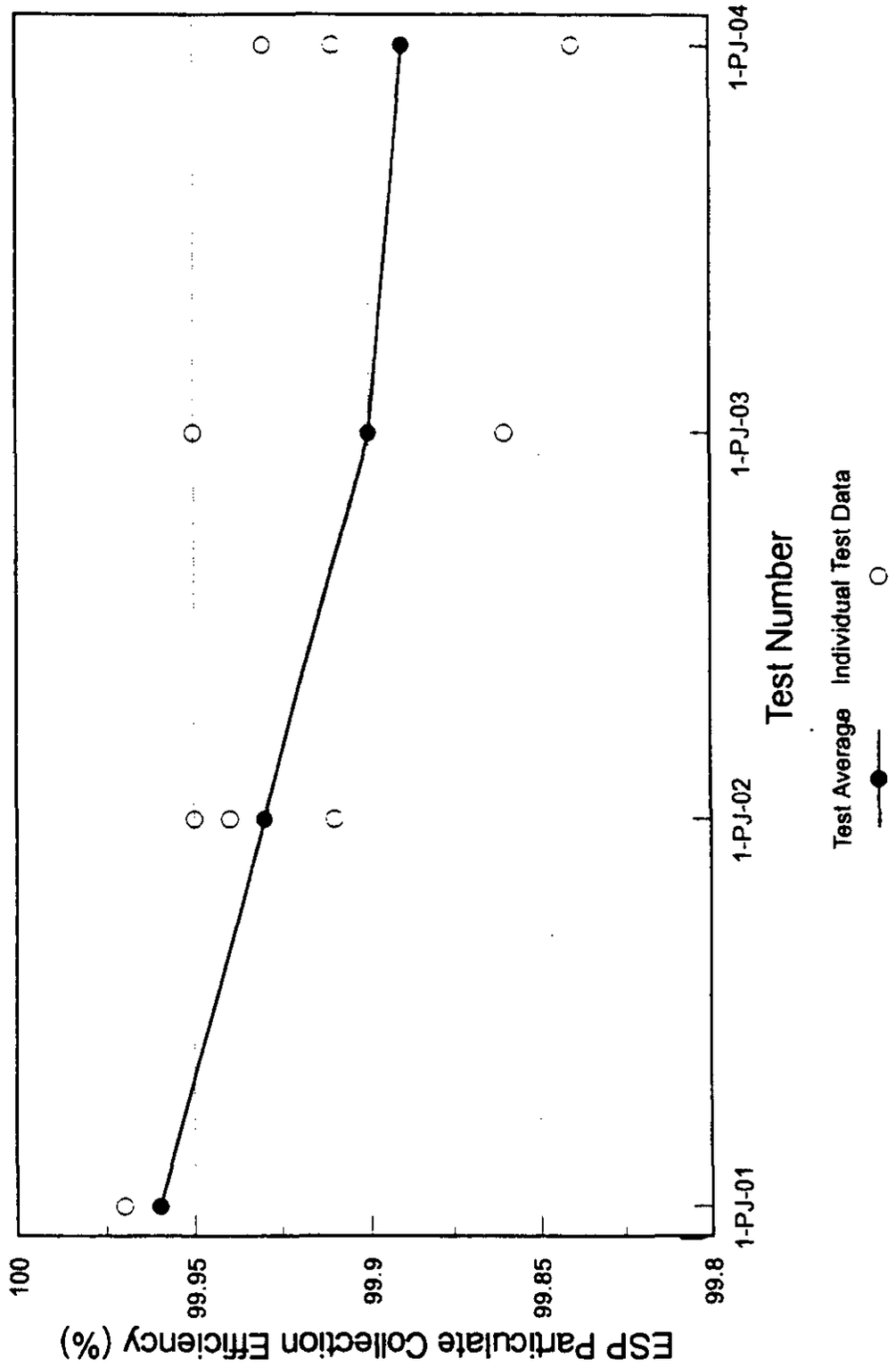


Figure 5.2.4-2

14-Day PJBH Demonstration Run ESP Performance ESP Particulate Emissions

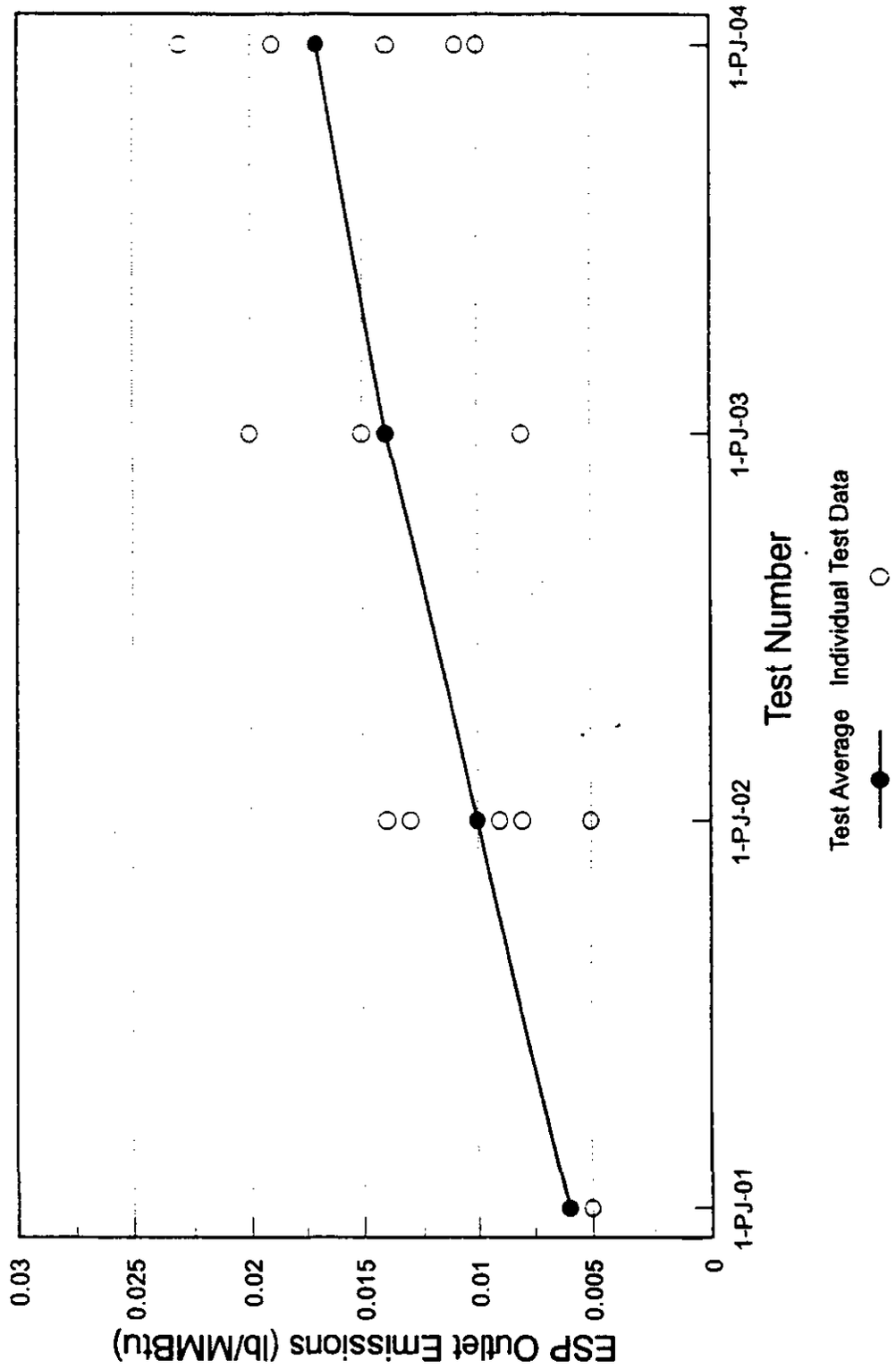


Table 5.2.4-2
AirPol GSA/PJBH Particulate Control Performance Results Summary
 PJ Series - 14 Day Demonstration Run

Test No.	Ca/S Ratio	Reactor Inlet Temp (F)	Approach Temp (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Injection Rate (lb/min)	Reactor Product Chloride Content (%)	PJBH Inlet Mass Loading (gr/scf)	PJBH Outlet Mass Loading (gr/scf)	PJBH Eff. Basis (%)	PJBH Emissions (lb/MMBtu)
1-PJ-01	1.37	319	18	20,000	30	4.9	0.32	3.009	0.0003	99.99	0.001
1-PJ-02	1.41	319	18	20,000	30	4.9	0.32	3.369	0.0007	99.98	0.002
1-PJ-03	1.34	320	18	20,000	30	4.9	0.29	3.383	0.0008	99.98	0.002
1-PJ-04	1.43	319	18	20,000	30	4.9	0.30	3.442	0.0003	99.99	0.001

Figure 5.2.4-3

14-Day PJBH Demonstration Run ESP and PJBH Particulate Removal

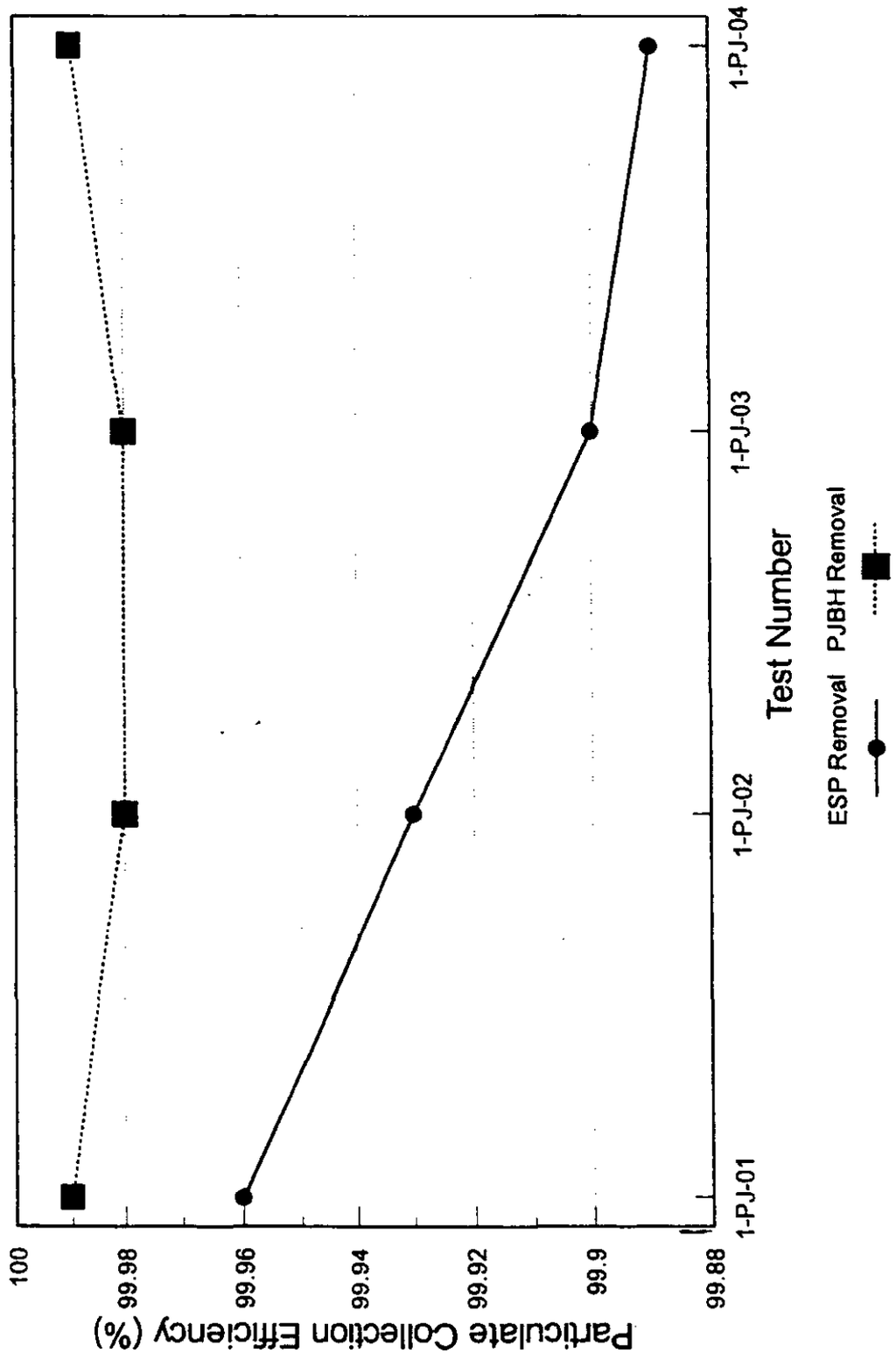
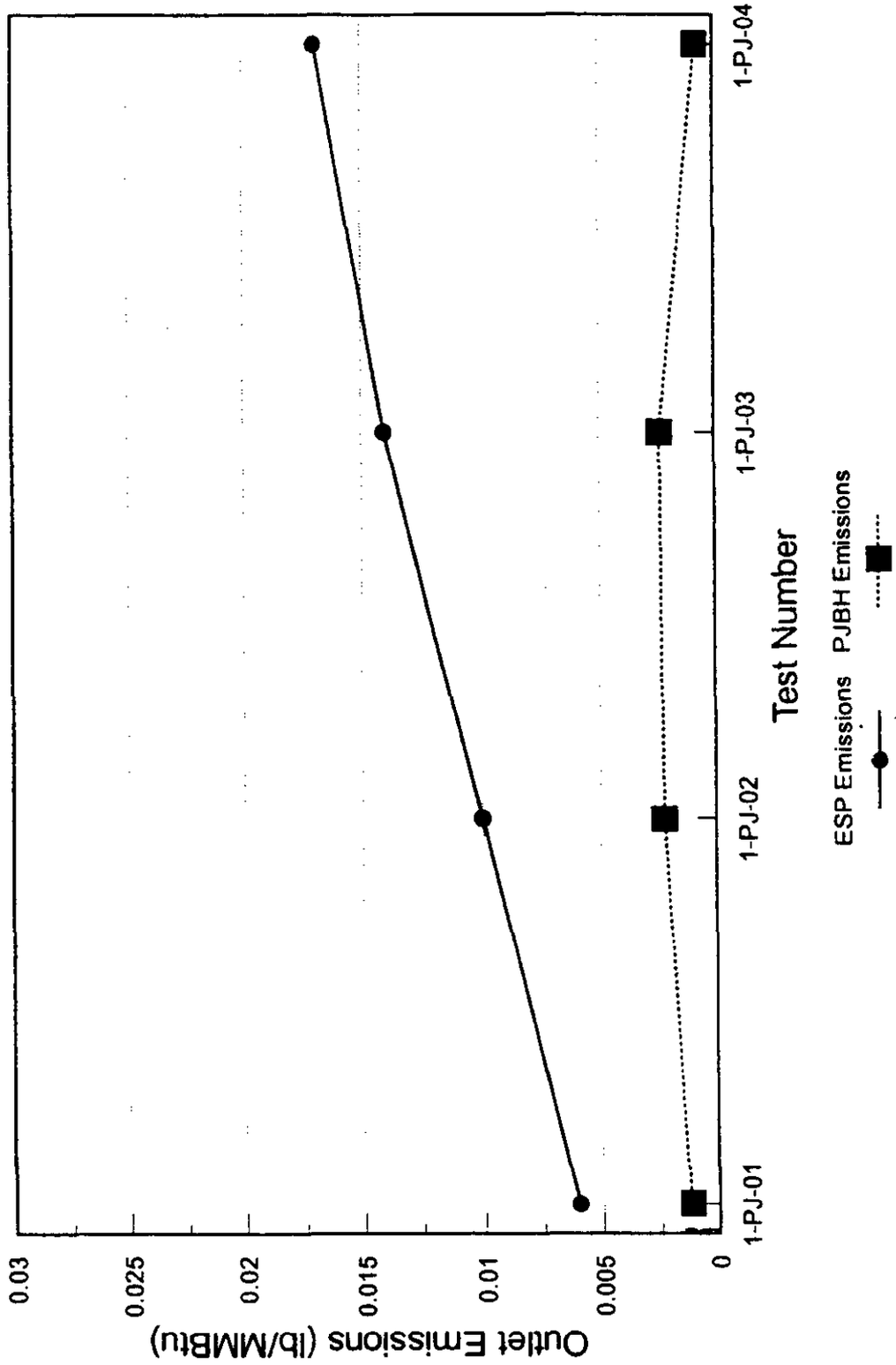


Figure 5.2.4-4

14-Day PJBH Demonstration Run

ESP and PJBH Particulate Emissions



5.3 ESP OPERATION

5.3.1 Factorial Tests

The average secondary voltage and current for the tests conducted at baseline chloride levels (0.04 weight percent coal chloride) are presented in Table 5.3.1-1 for the 2-AP series tests and in Table 5.3.1-2 for the 3-AP series tests. Similarly, the average secondary voltage and current are presented in Tables 5.3.1-3 and 5.3.1-4 for the chloride spiking tests (0.12 weight percent coal chloride equivalent) for the 2-AP and 3-AP series, respectively.

The changes in variable levels during the factorial tests resulted in changes in ESP operation. Specifically, the secondary current in field 1 and sometimes in fields 2 and 3 would be suppressed depending on the test conditions. The current suppression was greater during tests conducted at 20,000 scfm flue gas flow rate and at approach temperatures of 28°F compared to similar tests conducted at 14,000 scfm and lower approach temperatures. The current suppression is thought to be partially due to changes in particle size distribution due to the change in gas flow rate/velocity and changes in particle resistivity due to the change in approach temperature. Changes in other variable levels did not have as significant effect on the secondary current suppression. In addition, the secondary current suppression in the first field did not influence the ESP particulate control performance.

Table 5.3.1.1 Average AirPol ESP Secondary Currents and Voltages
2-AP Series - Baseline Tests

Test No.	Cu/S Ratio	Reactor Inlet Temp (F)	App. Temp (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Inj. Rate (lb/min)	Coal Chloride Content (%)	First Field Current (mA)	Second Field Current (mA)	Third Field Current (mA)	Fourth Field Current (mA)	First Field Voltage (kV)	Second Field Voltage (kV)	Third Field Voltage (kV)	Fourth Field Voltage (kV)
2-AP-09	1.0	320	8	14,000	30	0	0.03	175	195	196	193	52	48	48	43
2-AP-79	1.0	320	8	14,000	30	0	0.04	43	196	197	194	41	45	46	40
2-AP-72	1.0	320	8	20,000	45	7.3	0.04	97	196	195	194	49	46	47	41
2-AP-16	1.0	320	8	20,000	45	7.8	0.04	143	195	196	196	52	46	47	43
2-AP-11	1.3	320	8	14,000	30	5.0	0.03	185	197	197	194	51	45	45	42
2-AP-81	1.3	320	8	14,000	30	5.2	0.04	182	196	197	194	51	43	45	40
2-AP-10	1.3	320	8	20,000	45	0	0.03	55	192	196	194	46	49	46	41
2-AP-80	1.3	320	8	20,000	45	0	0.04	68	195	197	195	49	46	45	40
2-AP-01	1.0	319	18	14,000	30	0	0.03	59	196	196	*	46	48	48	44
2-AP-71	1.0	320	18	14,000	30	0	0.04	179	196	179	196	54	45	46	43
2-AP-78	1.0	320	18	20,000	30	7.2	0.04	155	196	196	196	54	45	48	42
2-AP-04	1.0	320	18	19,000	45	0	0.03	12	196	196	195	42	52	47	44
2-AP-74	1.0	320	18	20,000	45	0	0.04	95	195	196	195	51	48	48	42
2-AP-03	1.3	319	18	14,000	45	0	0.03	79	196	198	*	50	46	46	43
2-AP-73	1.3	320	18	14,000	45	5.1	0.04	190	196	197	195	53	43	46	40
2-AP-95	1.3	320	18	20,000	45	0	0.04	122	196	197	195	54	45	45	39
2-AP-96	1.3	322	18	20,000	45	0	0.04	83	196	198	194	51	45	45	39
2-AP-14	1.3	320	18	18,000	45	6.7	0.04	67	197	197	194	51	47	45	40
2-AP-63	1.3	320	18	20,000	45	7.2	0.04	56	193	196	196	49	49	47	42
2-AP-63	1.3	320	18	20,000	45	7.2	0.04	136	196	196	196	56	47	47	43
2-AP-88	1.0	320	28	14,000	30	0	0.04	166	196	196	196	54	47	47	42
2-AP-87	1.0	320	28	20,000	45	7.3	0.04	42	195	196	196	51	49	48	41
2-AP-86	1.3	320	28	14,000	45	0	0.04	40	196	196	195	48	45	46	40
2-AP-97	1.3	320	28	20,000	30	0	0.04	9	149	196	194	45	50	48	40
2-AP-19	1.3	320	28	20,000	30	7.1	0.03	11	187	196	193	46	54	48	42
2-AP-57	1.3	319	28	19,500	30	7.1	0.04	32	190	196	196	51	51	48	43

* Faulty current meter.

Table 5.3.1-2
Average AirPol ESP Secondary Currents and Voltages
3-AP Series - Baseline Tests

Test No.	Cu/S Ratio	Reactor Inlet Temp (F)	App. Temp (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Inj. Rate (lb/min)	Coal Chloride Content (%)	First Field Current (mA)	Second Field Current (mA)	Third Field Current (mA)	Fourth Field Current (mA)	First Field Voltage (kV)	Second Field Voltage (kV)	Third Field Voltage (kV)	Fourth Field Voltage (kV)
3-AP-12	1.0	320	8	14,000	30	0	0.04	133	196	195	198	48	47	47	42
3-AP-58	1.0	320	8	14,000	30	0	0.04	125	191	195	195	46	38	45	40
3-AP-42	1.3	320	8	14,000	30	0	0.04	129	191	196	196	47	42	43	41
3-AP-11	1.3	319	8	14,000	30	5.2	0.04	158	195	197	195	48	43	44	41
3-AP-62	1.0	319	18	14,000	30	0	0.04	191	196	197	195	51	45	46	42
3-AP-08	1.0	320	18	20,000	30	7.0	0.03	174	196	196	196	55	48	49	43
3-AP-44	1.3	319	18	14,000	45	0	0.04	144	182	196	193	49	43	45	42
3-AP-03	1.3	319	18	14,000	45	5.0	0.03	195	197	196	196	51	45	47	42
3-AP-26	1.3	260	18	14,000	12	4.7	0.03	199	196	196	195	51	46	47	43
3-AP-27	1.3	260	18	14,000	30	4.7	0.03	198	197	196	195	51	45	47	42
3-AP-02	1.3	319	18	14,000	45	5.1	0.03	140	196	197	197	51	45	46	41
3-AP-18	1.0	319	28	14,000	30	0	0.03	73	196	197	198	49	48	48	43
3-AP-59	1.0	320	28	14,000	30	5.2	0.04	104	197	196	195	49	46	47	42
3-AP-21	1.0	319	28	20,000	45	7.3	0.03	21	193	196	196	46	52	48	42
3-AP-56	1.3	320	28	14,000	30	0	0.04	99	197	198	195	49	44	45	40
3-AP-20	1.3	320	28	14,000	45	0	0.03	27	194	198	197	45	47	45	41
3-AP-20	1.3	320	28	14,000	45	0	0.04	51	196	197	194	47	45	45	42
3-AP-13	1.3	319	28	14,000	45	0	0.03	30	193	196	197	47	48	47	42
3-AP-45	1.3	320	28	14,000	45	5.2	0.04	121	196	198	196	50	45	45	42
3-AP-19	1.3	320	28	20,000	30	7.4	0.03	9	179	196	197	45	52	48	42

Table 5.3.1-3
Average AirPol ESP Secondary Currents and Voltages
2-AP Series - Chloride Spiking Tests

Test No.	Ca/S Ratio	Reactor Inlet Temp (F)	App. Temp (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Inj. Rate (lb/min)	Coal Chloride Content (%)	First Field Current (mA)	Second Field Current (mA)	Third Field Current (mA)	Fourth Field Current (mA)	First Field Voltage (kV)	Second Field Voltage (kV)	Third Field Voltage (kV)	Fourth Field Voltage (kV)
2-AP-28	1.0	320	18	14,000	40	5.0	0.12	33	193	196	195	37	46	47	43
2-AP-75	1.0	320	18	14,000	45	4.7	0.12	44	195	195	194	40	46	47	41
2-AP-17	1.0	320	18	20,000	30	0	0.12	25	143	189	193	37	46	47	42
2-AP-82	1.0	320	18	20,000	30	0	0.12	20	144	187	195	38	46	47	43
2-AP-07	1.3	320	18	14,000	30	5.4	0.12	20	195	197	195	36	46	46	42
2-AP-77	1.3	320	18	14,000	30	5.0	0.12	33	195	197	196	37	44	45	40
2-AP-98	1.3	320	18	14,000	30	5.2	0.12	158	196	196	195	50	43	44	41
2-AP-06	1.3	320	18	20,000	45	0	0.12	17	110	186	195	34	48	47	43
2-AP-92	1.3	320	18	20,000	45	0	0.12	50	160	194	195	46	46	45	43
2-AP-91	1.3	320	18	20,000	45	7.2	0.12	12	158	192	195	35	48	45	43
2-AP-22	1.0	320	28	14,000	45	0	0.12	15	177	197	194	37	49	45	41
2-AP-90	1.0	320	28	14,000	45	0	0.12	160	197	196	196	52	46	47	43
2-AP-25	1.0	320	28	18,600	30	6.8	0.12	18	195	196	194	46	52	49	44
2-AP-94	1.0	320	28	20,000	30	0	0.12	86	195	196	195	55	48	48	43
2-AP-85	1.0	320	28	20,000	30	7.4	0.12	103	195	196	195	53	48	49	43
2-AP-84	1.3	320	28	14,000	30	0	0.12	183	196	196	195	53	44	46	42
2-AP-24	1.3	320	28	14,000	30	0	0.12	33	196	197	194	47	47	46	42
2-AP-83	1.3	320	28	20,000	30	7.3	0.12	49	195	196	196	52	48	47	40

Table 5.3.1-4
Average AirPol ESP Secondary Currents and Voltages
3-AP Series - Chloride Spiking Tests

Test No.	Ca/S Ratio	Reactor Inlet Temp (F)	App. Temp (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Inj. Rate (lb/min)	Coal Chloride Content (%)	First Field Current (mA)	Second Field Current (mA)	Third Field Current (mA)	Fourth Field Current (mA)	First Field Voltage (kV)	Second Field Voltage (kV)	Third Field Voltage (kV)	Fourth Field Voltage (kV)
3-AP-29	1.0	320	18	14,000	45	5.2	0.12	124	196	197	193	43	44	46	42
3-AP-22	1.0	320	28	14,000	45	0	0.12	14	194	197	195	38	48	45	41
3-AP-24	1.3	320	28	14,000	30	0	0.12	95	197	197	195	49	45	46	42
3-AP-23	1.3	320	28	19,200	45	7.2	0.12	9	189	196	196	41	53	47	43

5.3.2 28-Day Demonstration Run

As mentioned previously, the ESP particulate removal performance deteriorated during the 28-day demonstration run. The reason for the poorer performance is due to a decrease in power levels in all four ESP fields. The most significant decrease was in the first two fields. Table 5.3.2 summarizes the average secondary current and voltage for each demonstration run test segment. Figures 5.3.2-1 and 5.3.2-2 illustrate the decrease in power levels. In Figure 5.3.2-1, the average secondary current is plotted for each test segment. Similarly, the average secondary voltage for each test segment is presented in Figure 5.3.2-2. In both figures, a significant drop in power levels was observed in field 1 during test 1-DR-04 and in field 2 during test 3-DR-04. A hypothesis for the increase in power levels in the first field after test 1-DR-02 is that the increase was due to cleaning the field during the November 1 through 3 outage.

Although the power levels decreased during test 3-DR-04, the reported ESP particulate control performance for this test segment was equivalent to the prior four test segments. This is because the mass loading tests, which were conducted on November 9, were conducted while the secondary current and voltage were decreasing in the second field. This is illustrated in Figure 5.3.2-3, which shows a daily plot of the secondary current for each field. As shown in the figure, the secondary current in field 2 was dropping during the day on November 9. There was also a slight drop in current level observed in field 3 on November 9. However, a more significant drop in secondary current levels was observed in field 3 on November 10. Also shown in the figure is a slight drop in average secondary current for field 4 on November 10. The secondary current in fields 3 and 4 were lower after November 10 and remained at the lower levels for the remainder of the demonstration run.

The reason for the drop in power levels in the ESP fields appears to be due to solids build-up shorting out the first two fields. Solids build-up was observed on the hopper ridge beams between the first and second field and the second and third field hoppers. The build-up, which was observed during an ESP inspection on November 29, extended up into the plates and wires approximately 6 to 8 inches. The reduction in power levels in fields 3 and 4 was probably due to the increase in particulate loading resulting from the poorer particulate collection performance in fields 1 and 2.

Table 5.3.2
Average AirPol ESP Secondary Currents and Voltages
1-DR Series

Test No.	Ca/S Ratio	Reactor Inlet Temp (F)	App. Temp (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Inj. Rate (lb/min)	Coal Chloride Content (%)	First Field Current (mA)	Second Field Current (mA)	Third Field Current (mA)	Fourth Field Current (mA)	First Field Voltage (kV)	Second Field Voltage (kV)	Third Field Voltage (kV)	Fourth Field Voltage (kV)
1-DR-01	1.46	320	18	20,000	30	7.4	0.27	71	193	196	197	52	47	46	41
1-DR-02	1.47	319	18	20,000	30	7.0	0.34	46	180	195	198	45	49	47	41
1-DR-03	1.58	319	18	20,000	30	7.3	0.21	75	170	196	198	52	46	46	41
1-DR-04	1.48	319	18	20,000	30	7.1	0.19	8	171	196	197	38	53	47	42
3-DR-04	1.32	320	18	20,000	30	7.3	0.32	9	38	179	195	35	40	49	41
1-DR-05	1.52	319	18	20,000	30	7.2	0.33	13	24	178	195	33	40	50	41
1-DR-06	1.43	320	18	20,000	30	7.3	0.30	13	20	177	194	34	37	49	40
1-DR-07	1.40	320	18	20,000	30	7.3	0.30	11	27	189	196	36	41	51	41
1-DR-08	1.45	320	18	20,000	30	7.4	0.41	12	15	169	192	34	38	49	40

Figure 5.3.2-1

28-Day Demonstration Run ESP Performance

Average Secondary Currents

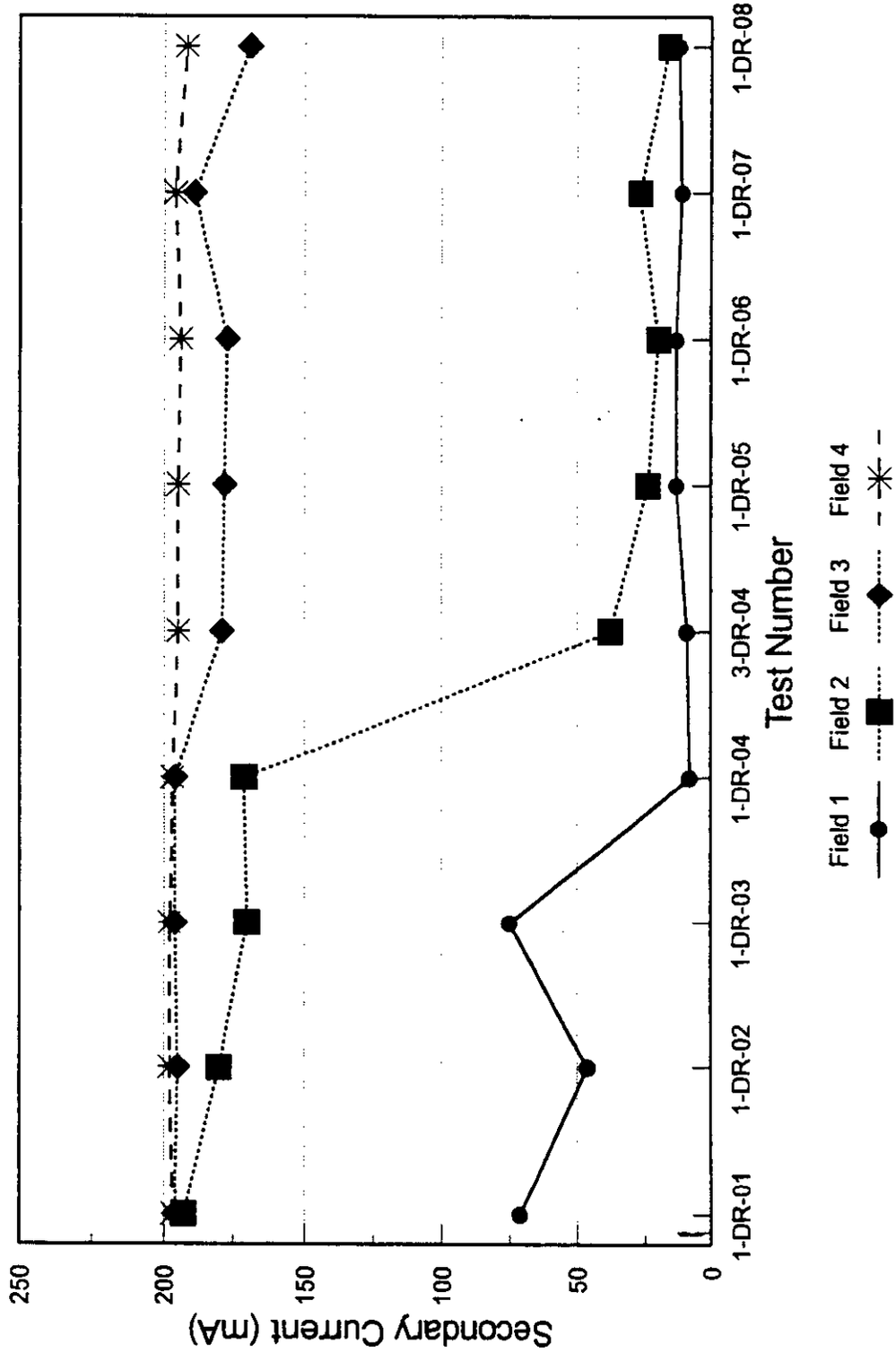


Figure 5.3.2.2-2

28-Day Demonstration Run ESP Performance

Average Secondary Voltages

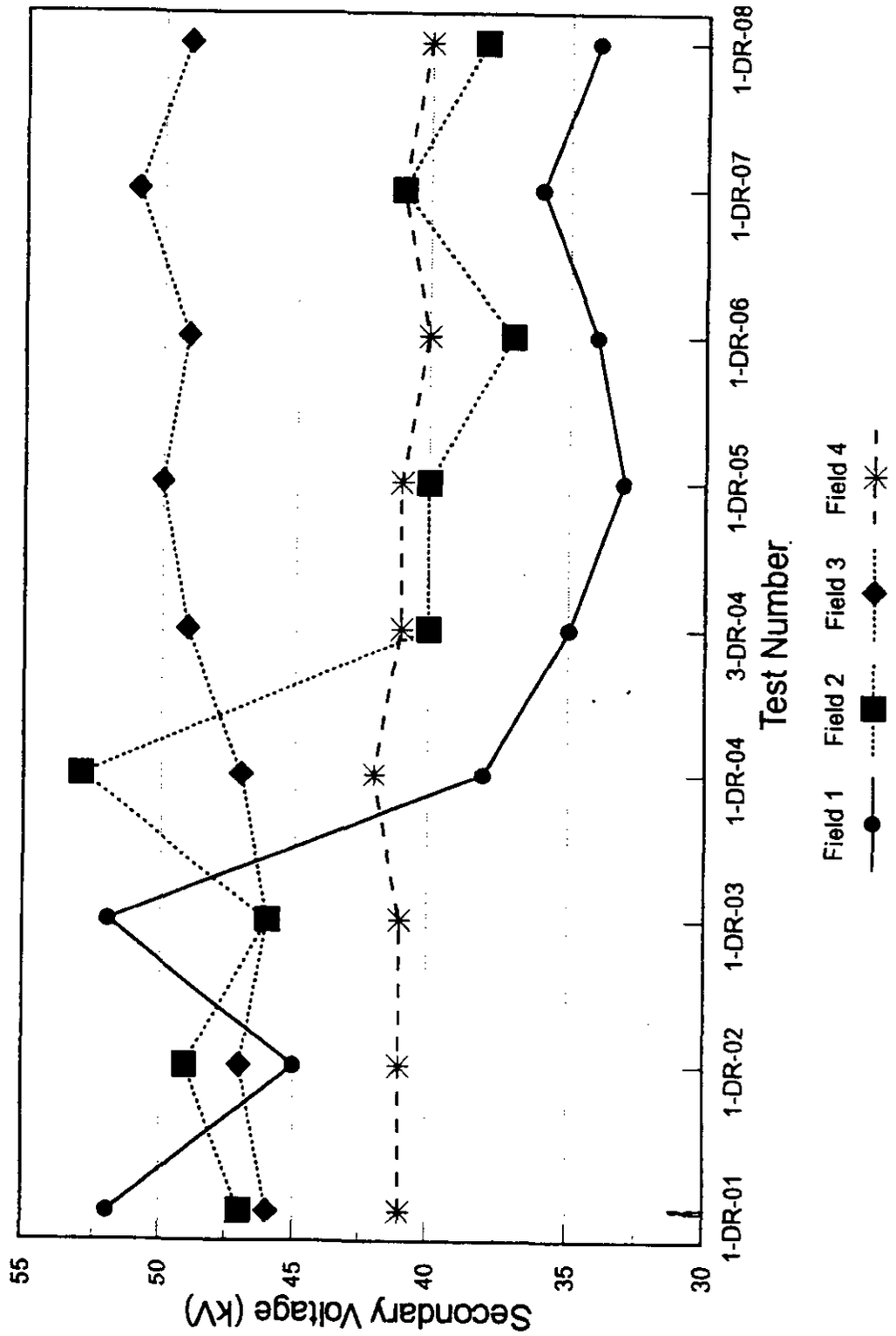
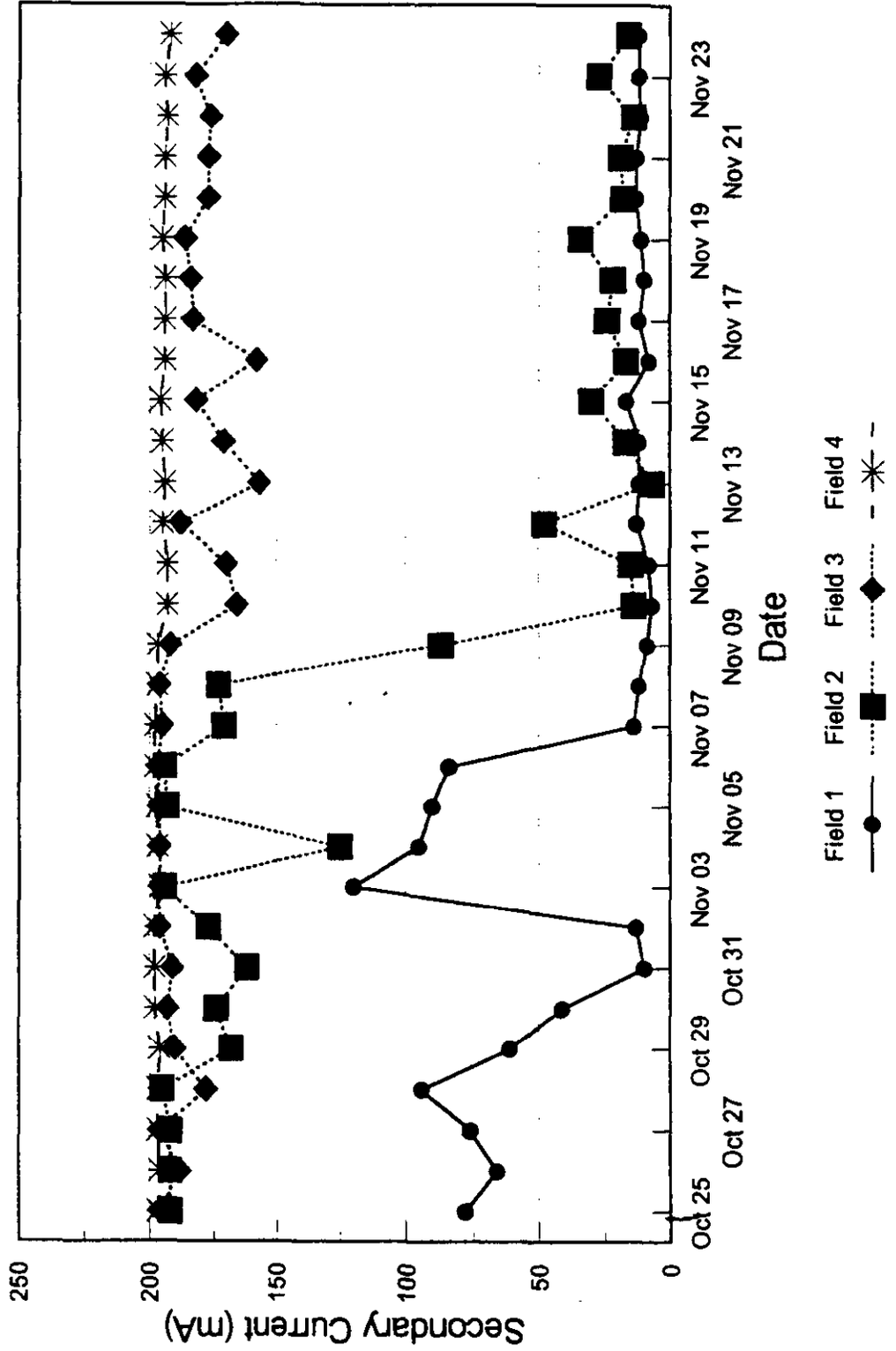


Figure 5.3.2-3

28-Day Demonstration Run ESP Performance

Average Daily Secondary Currents



5.3.3 14-Day Pulse-jet Baghouse Demonstration Run

The average secondary currents and voltages for each test segment during the PJBH demonstration run are presented in Table 5.3.3. As noted previously, the solids build-up on the first field wires due to failure of the rapper drive may have affected the first field secondary current during the first three test segments. Figures 5.3.3-1 and 5.3.3-2 present the secondary current levels for each field during the demonstration run. In Figure 5.3.3-1, the average secondary current levels are plotted for each test segment. The figure shows a significant increase in the average first field secondary current during test 1-PJ-04 compared to the prior test segments when the first field wires were not being rapped. Figure 5.3.3-2, which plots the average daily secondary currents during the PJBH demonstration run, also shows a significant increase in the first field current for the last test segment, 1-PJ-04. However, there was a sharp decrease in the first field secondary current on the last day of the demonstration run. Also observed during the demonstration run was a decrease in the second field secondary current from February 27 to March 3. No explanation was found for this decrease in secondary current in the second field.

Table S.3.3
Average AirPol ESP Secondary Currents and Voltages
1-PJ Series

Test No.	Ca/S Ratio	Reactor Inlet Temp (F)	App Temp (F)	Flue Gas Flow Rate (scfm)	Recycle Screw Speed (rpm)	Reactor Inlet Fly Ash Inj. Rate (lb/min)	Coal Chloride Content (%)	First Field Current (mA)	Second Field Current (mA)	Third Field Current (mA)	Fourth Field Current (mA)	First Field Voltage (kV)	Second Field Voltage (kV)	Third Field Voltage (kV)	Fourth Field Voltage (kV)
1-PJ-01	1.37	319	18	20,000	30	4.9	0.12	21	155	197	198	45	46	45	41
1-PJ-02	1.41	319	18	20,000	30	4.9	0.12	25	112	198	196	45	43	45	41
1-PJ-03	1.34	320	18	20,000	30	4.9	0.12	31	183	197	196	45	47	44	42
1-PJ-04	1.43	319	18	20,000	30	4.9	0.12	130	199	198	195	50	43	44	41

Figure 5.3.3-1

14-Day PJBH Demonstration Run ESP Performance

Average Secondary Currents

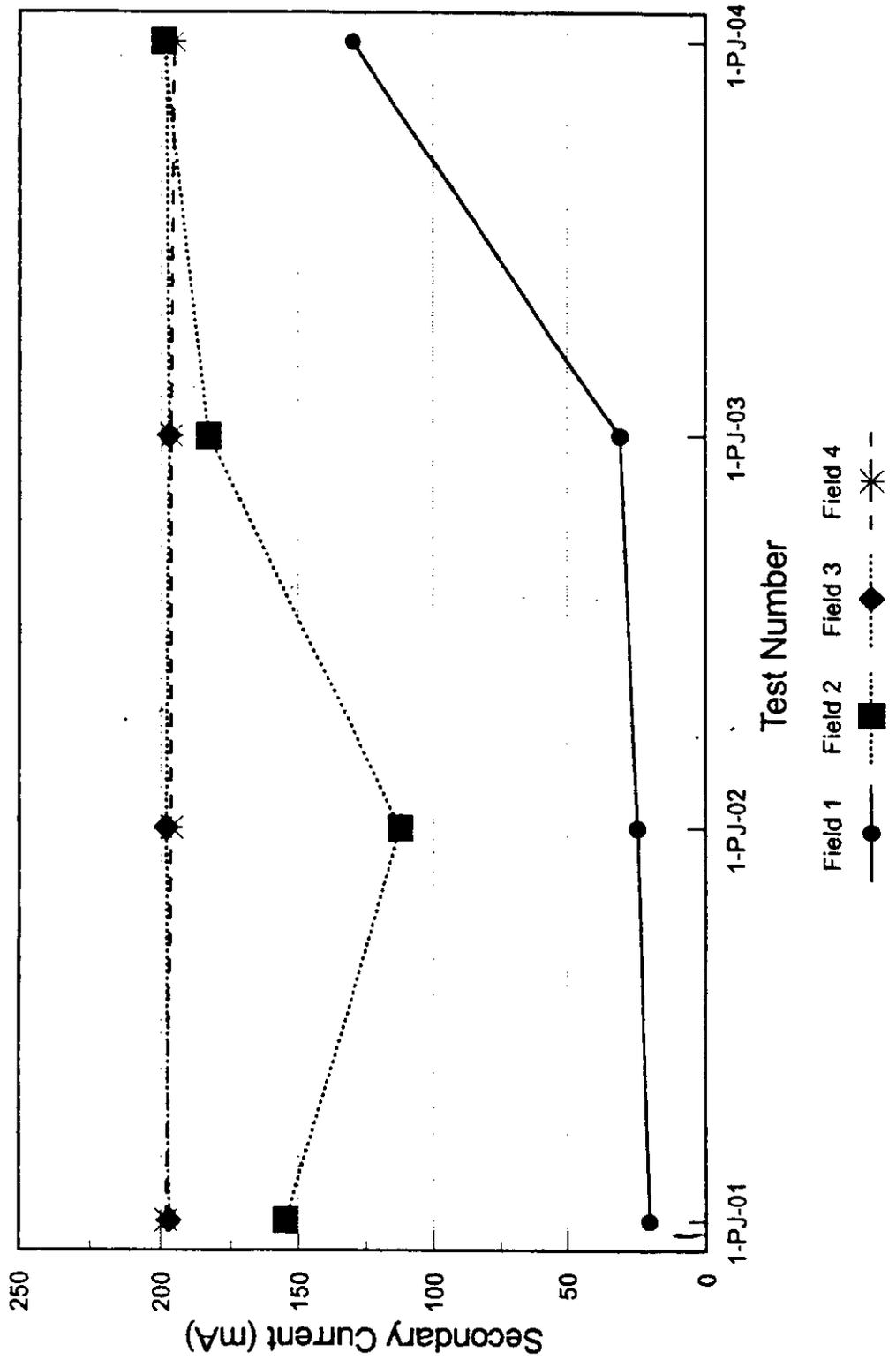
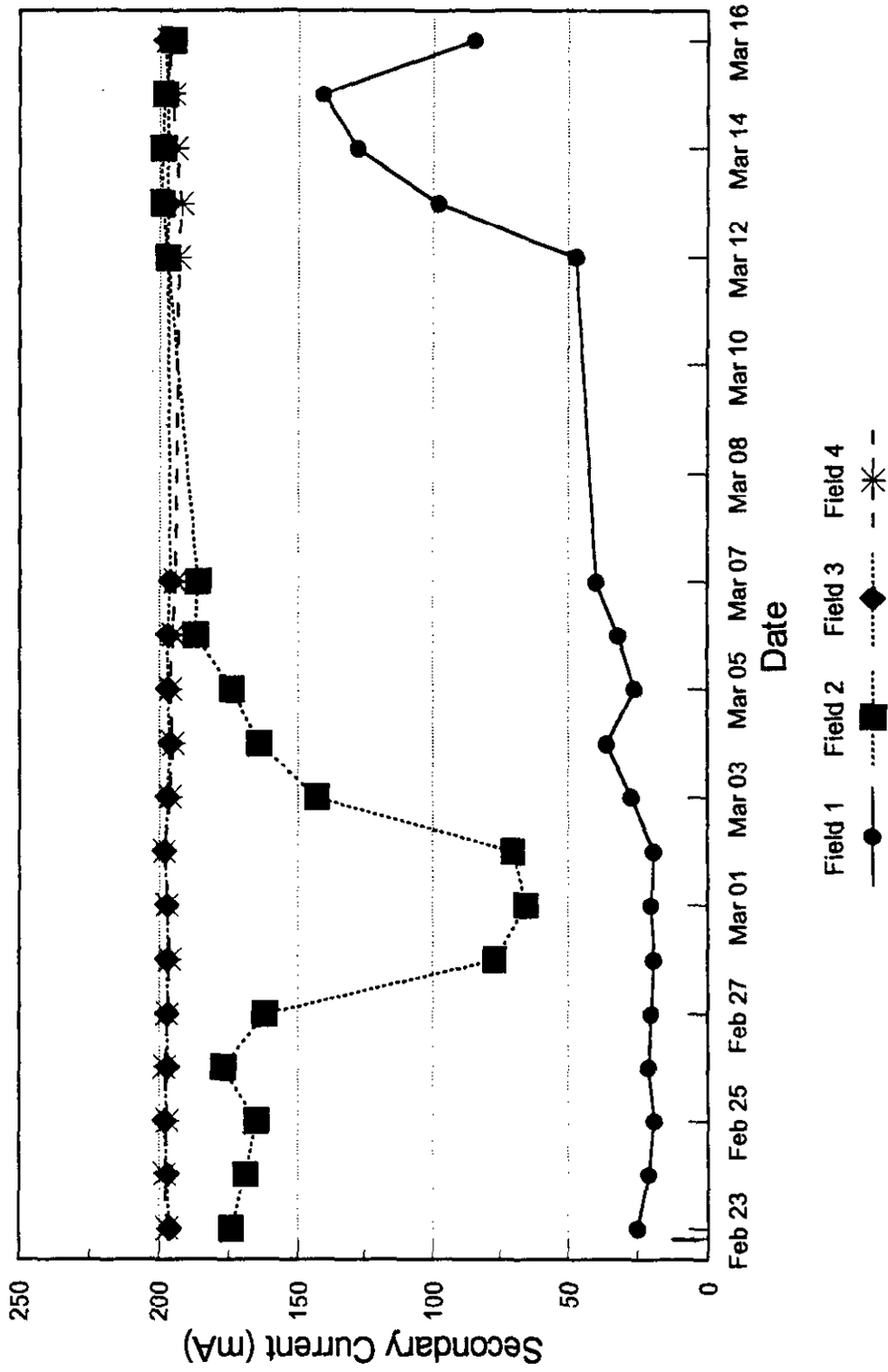


Figure 5.3.3-2

14-Day PJBH Demonstration Run ESP Performance

Average Daily Secondary Currents



5.4 CORRELATION OF RESULTS

The following discussions are related to the correlations determined in the factorial test. The purpose of this factorial testing was to determine the effect of the process variables on the operation and SO₂ removal efficiency in the reactor/cyclone, the ESP, the PJBH, and the overall system so as to optimize the GSA performance.

Given the large number of major process design variables and the limited amount of test time available, only two levels for most of the variables were selected to cover the range of primary interest for a utility FGD application. To further reduce the length of the factorial test plan, but still retain the quality control on the test results, a half-factorial design with a full set of replicate tests was used for the GSA testing.

The major variables, as determined from the preliminary tests, were approach-to-saturation temperature, lime stoichiometry, fly ash loading, coal chloride level, flue gas flow rate, and recycle screw speed. Two levels were determined for nearly all of the variables. The one exception was the approach-to-saturation temperature where three levels were defined, but the third level was only run for those tests at the lower coal chloride level. The variables and their selected levels are shown in the Table 5.4.

A total of 78 factorial tests were performed. During 51 of these tests the baghouse was either off-line or pulling flue gas from the ESP outlet (in series arrangement), and during 27 of these tests the baghouse was pulling flue gas from the ESP inlet (in parallel arrangement).

The lime stoichiometry level, which was tested at 1.00 and 1.30 moles Ca(OH)₂/mole inlet SO₂, seems to have the most significant effect on the SO₂ removal efficiency in the GSA system. The approach-to-saturation temperature, evaluated at three levels 8, 18, and 28 °F for the low coal chloride conditions and at two levels 18 and 28 °F for the higher coal chloride conditions, appear to be the second most important variable in the GSA system in terms of the overall system SO₂ removal efficiency.

The third most important variable seems to be the chloride level in the system. Two coal chloride levels were tested, the baseline coal chloride level of 0.02 to 0.04% and the equivalent of a 0.12% coal chloride level. The higher chloride level was achieved by spiking the feed slurry with a calcium chloride solution.

For further detailed information please refer to TVA's final report entitled "10 MW Demonstration of the Gas Suspension Absorption Process at TVA's Center for Emissions Research" in Appendix "B" of this report.

Table 5.4

**MAJOR VARIABLES AND LEVELS
FOR FACTORIAL TEST**

Variable		Level
Approach-to-saturation temperature	°F	8*, 18, and 28
Ca/S	moles Ca(OH) ₂ /mole inlet SO ₂	1.00 and 1.30
Fly ash loading	gr/acf	0.5 and 2.0
Coal chloride level	%	0.02 and 0.12
Flue gas flow rate	kscfm	14 and 20
Recycle screw speed	rpm	30 and 45
* --- 8 °F level run only at the low-chloride level		

6.0 ENVIRONMENTAL PERFORMANCE

6.1 IMPACT ON THE ENVIRONMENT

Environmental impacts associated with the GSA system are minimal. The consequence of both construction and operation of the project on various environmental conditions are discussed as follows:

6.1.1 Land Impacts

Land impacts are insignificant since no additional land outside the TVA Shawnee Steam Plant boundaries was required for the GSA unit and the unit was constructed on previously impacted land between an existing spray dryer building and an electrostatic precipitator.

6.1.2 Air, Water Quality and Solid Waste Impacts

According to the environmental monitoring report, the GSA system averaged greater than 90 percent SO₂ removal efficiency over the course of the demonstration run, even when the boiler switched to a higher sulfur coal. Meanwhile, the particulate emission rate for the ESP remained well below the NSPS for particulates (0.03 lb/MBtu) throughout the run.

In accordance with the compliance monitoring results, the GSA demonstration system does not generate additional aqueous waste over the amount discharged from the plant during normal operations.

The solid waste by-product resulting from the operation of the GSA unit has the same composition as the spray dryer by-product. In keeping with the existing practices, these solids are diluted with water to generate a slurry containing approximately 10% solids before being pumped to an existing ash pond for ultimate disposal. Changes in ash pond effluent as a result of the operation of the GSA is negligible. Table 6.1.2 presents the results of a lechate test performed on GSA waste product by GAI Consultant Inc. The results show that the leaching characteristics of GSA waste is well within the Resource Conservation and Recovery Act (RCRA) limits.

6.1.3 Ecological Impacts

No adverse ecological impacts to either terrestrial or aquatic environments are expected for the GSA project. The GSA unit is constructed on previously disturbed land located beyond the 500 year flood plain of the Ohio River. Effluent from its operation constitutes less than 0.001% of the total Shawnee Steam Plant waste water balance discharged to the Ohio River. Although wetlands are present within the broader confines of the Shawnee Steam Plant, the GSA unit does not discharge any material to any wetlands or lakes.

Table 6.1.2

**RESULTS OF LEACHATE TEST OF GSA DISPOSAL MATERIAL
10 MW DEMONSTRATION OF GSA**

Contaminants	RCRA Regulatory Limits mg/l	TCLP Test Results mg/l
Arsenic	5.0	0.039
Cadmium	1.0	0.156
Chromium	5.0	0.15
Lead	5.0	0.81
Mercury	0.2	<0.0002
Selenium	1.0	0.033
Silver	5.0	0.05
Barium	100.0	0.22

Note:

RCRA = Resource Conservation and Recovery Act

TCLP = Toxic Characteristic Leaching Procedure

6.1.4 Health and Safety Impacts

The health and safety requirements applicable to the construction and operation of the GSA demonstration project include “construction” and “general industry” standards of the Federal Occupational Safety and Health Act in 29 CFR Parts 1910 and 1926, respectively. These standards include requirements related to walking and working surfaces, means of ingress and egress, operation of powered equipment, adequate ventilation, noise exposure controls, fire protection, and electrical equipment safeguards. Shawnee Fossil Plant employees are already trained in work protection and safety procedures under existing TVA guidelines. The current procedures are adequate to ensure that applicable standards are not exceeded. Contractors have complied with all site rules and regulations concerning health and safety.

The demonstration project has not required the storage and/or use of any “extremely hazardous substances” as defined under the Superfund Amendments and Reauthorization Act (SARA) Title III, or Emergency Planning and Community Right-to-Know program. Thus, no SARA Title III emergency planning notification is applicable to this project.

6.1.5 Impact Summary

The majority of the potential additional environmental consequences resulting from the installation, operation and testing of the GSA can be categorized as insignificant because TVA's existing SD/ESP and AirPol's GSA process are essentially identical and the GSA replaced the SD/ESP. However, two potential positive environmental impacts are identifiable: (1) the GSA consumes less lime than the existing system; (2) and will, therefore, generate less solid waste by-product. As a whole, the environmental benefits resulting from the commercial implementation of GSA systems are extraordinary.

6.2 WASTE STREAMS AND THEIR DISPOSAL

The by-products from the demonstration system are cleaned gas exiting the stack and waste material discharged from the reactor and cyclone. The cleaned gas is discharged into the atmosphere. The solid by-product is mixed with the ESP ash, diluted with water to generate a slurry containing approximately 10% solids and pumped to an existing ash pond for dewatering and ultimate disposal with other ash. Changes in ash pond effluent quality or quantity as a result of the operation of the GSA are infinitesimal.

The GSA system is designed to treat a slip stream from Unit 9 Boiler of 20,000 SCFM of boiler flue gas. The average composition of the by-product from the GSA system is as follows:

Ca(OH) ₂	1.4 %
CaCO ₃	9.0 %
CaSO ₃	44.0 %
CaSO ₄	19.4 %
Acid Insoluble	26.2 %

The quantity of waste is approximately 1,060 lb/hr, being sluiced with 17,000 gallons of water per day. The once-through cooling water used by the entire plant and returned to the Ohio River is 1.5 billion gallons of water per day. Since the waste from the GSA unit was the same as that currently discharged to the ash ponds and had only 0.001 % of the total water discharge to the Ohio River, the demonstration project did not show any environmental impact or impact on current operational practices.

The SO₂ loading was 389 lbs/hr in the inlet gas stream and 31 lbs/hr in the outlet. The dust loading was 501 lbs/hr in the inlet gas stream and 2.18 lbs/hr in the outlet gas stream. This level of particulate discharge into the atmosphere is below the NSPS maximum limit of 0.03 lb/MBtu.

6.3 POTENTIAL ENVIRONMENTAL CONCERNS

Successful removal of hazardous air pollutants was expected in the design of the GSA demonstration system. Air toxics tests were conducted by Energy and Environmental Research Corporation (EERC). The results of the air toxics testing show that the GSA process is capable of removing HF, particulate and trace metals. For detailed analytical data of air toxics testing please refer to a separate monitoring report prepared by EERC.

7.0 ECONOMICS

The economic evaluation was performed by Raytheon Engineers & Constructors. The economic evaluation basis was a pulverized coal-fired plant producing a nominal 300 MW (net). Operating conditions for the boiler were assumed to be typical of modern units. Table 7.0 is a brief summary of performance and economics of the GSA technology based upon Raytheon's report presented for a brief review of this project. The detailed information of this section is available in the *Economic Evaluation Report* presented by Raytheon Engineers & Constructors in Appendix "A" of this report. Appendix "A" consists of two sections describing the results of the analysis presented in the Raytheon's report.

The Section 3 of the appendix "A" is the general criteria of the evaluation, which incorporates both process and economic design criteria.

The specific economic evaluation for AirPol's Gas Suspension Absorber (GSA) is presented in Section 20 of the Raytheon's report.

Table 7.0

SUMMARY OF PERFORMANCE AND COST DATA

Power Plant Attributes	Units		Value		
Plant capacity, net	MW		300		
Power produced, net	10 ⁹ kWh/yr		2.628		
Capacity factor	%		100		
Plant life	year		30		
Coal feed	10 ⁶ tons/yr		0.975		
Sulfur in coal	wt %		2.6		

Emissions Control Data	Units	SO₂	NO_x	TSP	PM₁₀
Removal efficiency	%	90	N/A	98.9	N/A
Emissions standard	lb/10 ⁶ Btu	1.2	N/A	0.03	N/A
Emissions without controls	lb/10 ⁶ Btu	3.94	N/A	2.88	N/A
Emissions with control	lb/10 ⁶ Btu	0.394	N/A	0.029	N/A
Amount removed	tons/yr	22,338	N/A	36,410	N/A

	Current Dollars		Constant Dollars	
Levelized Cost of Power	Factor	Mills/kWh	Factor	Mills/kWh
Capital Charge	0.16	5.0	0.124	3.7
Fixed O&M Cost	1.314	2.3	1.000	1.6
Variable Operating Cost	1.314	3.1	1.000	2.3
Total Cost	-	10.4	-	7.6
Levelized Cost--SO₂ Basis	Factor	\$/ton removed	Factor	\$/ton removed
Capital Charge	0.16	291	0.124	213
Fixed O&M Cost	1.314	129	1.000	94
Variable Operating Cost	1.314	182	1.000	132
Total Cost	-	602	-	439

8.0 COMMERCIALIZATION POTENTIAL AND PLANS

8.1 MARKET ANALYSIS

8.1.1 Applicability of the Technology

The equipment has been demonstrated to operate at the same design requirements as previously experienced in the incineration industry where the GSA has established a position as a lower cost semi-dry type of scrubber. The technology offers solutions for many of the problems experienced in the use of conventional spray dryers for SO₂ control for utility and industrial boiler installations. The GSA demonstration unit operated with very little buildup, without corrosion and reached removal levels above 90% at reasonable consumption of the lime reagent.

The GSA has some superior features that make it attractive to the utility industry. The unit has a very small footprint when compared to either wet systems or conventional dry scrubbers. The collected byproduct is easier to handle due to a very low moisture level (less than 1%) and low leachability. The spray is accomplished with a single large orifice nozzle that can be serviced on line. The power consumption is less than competing technologies due to minimized air consumption, limited material handling systems and the design does not require fine atomization of the lime slurry spray.

The unit may be used for any size of boiler as it is best installed as a modular system. It will be more attractive for systems less than 300 MW especially for retrofit installations. The unit has both low capital and low operating costs when compared with wet lime stone scrubbing and should be well received for industrial boilers.

The unit has been tested for air toxics removal and achieved some superior results. This could help in the development of the market especially for new and smaller retrofit applications.

8.1.2 Market Size

The domestic market for this technology appears to be limited at the present time. The current unknown status of clean air act regulations as to level of cleaning requirements and the timing for meeting the requirement offers a serious problem in entering the market. Will the requirements be limited to only larger units? How much improvement in gas cleaning is needed by these units to reach compliance? These uncertainties makes the market extremely difficult to quantify at this time. Currently we project the major market for GSA to develop between the year 2000 to 2010.

AirPol anticipates a market size of one (1) domestic order for a smaller boiler over the next five (5) years. The market should then grow at a 50 percent rate per year over the following ten (10) years. The order for the GSA technology are expected to grow from \$3 ~ 5 million to \$15 ~ 30 million.

8.1.3 Market Barriers

The utility industry has been slow to accept new technologies until it is demonstrated for application with scaled up capacity. AirPol has secured its first installation on a 50 MW municipal boiler where extensive testing will be performed during the first six (6) months of operation. The unit will begin operation in 1996 and should support acceptance by the utility industry for smaller installations by 1997.

The technology will compete with fuel switching and the cost of purchasing SO₂ allowances. We expect this to be the major barrier to the introduction of this or other new technologies.

8.1.4 Economic Comparison with Competing Technologies

The economics of the GSA system is projected as a general comparison to a conventional spray dryer and a wet system.

As part of the DOE CCT Program, an economic evaluation of the GSA process was conducted using the same design and economic premises that were used to evaluate about 30 to 35 other FGD processes. The relative process economics for the GSA system were evaluated for a moderately difficult retrofit to a 300 MW boiler burning a 2.6 percent sulfur coal. The design SO₂ removal efficiency was 90 percent.

The resulting capital cost estimates (in 1990 dollars) are shown in Table 8.1.3-1 along with those for the conventional wet limestone, forced-oxidation (WLFO) scrubbing system. The total capital requirements for the GSA process are substantially lower than those for the WLFO (\$149/kw vs. 216/kw). Since the presumed accuracy of these estimates is +/- 10 percent, this lower capital requirement estimate for the GSA process is significant. The substantially lower capital requirements are primarily due to the lower costs in the SO₂ absorption technology.

The levelized annual revenue requirements for the two processes (in 15-year levelizing) are shown in Table 8.1.3-2. The GSA system operating cost differs most in the reagent cost. The reagent cost for GSA process is 2.8 times that of WLFO due to the higher lime cost. However, this higher reagent costs are offset by lower power cost and lack of steam requirement for re-heating. Overall, the levelized costs for GSA are 20 percent lower than those for the WLFO process.

A comparison of the space to be occupied by the GSA to the spray dryer unit shows that the space requirement of a GSA is much lower than that of a spray dryer. Owing to its comparatively simple design and fewer components, the GSA capital cost is lower than that of the spray dryer. Meanwhile, based on the fact that the GSA requires less power and has better lime utilization than a spray dryer, the GSA will also have lower operating cost.

The details of the comparison can be found in Appendix "A", Economic Evaluation Report.

Table 8.1.3-1

TOTAL CAPITAL INVESTMENT COMPARISON

1990 \$, 300 MW, 2.6% sulfur coal

Area	Description	GSA (\$/kW)	WLFO (\$/kW)
10	Reagent Feed	25.1	36.7
20	SO ₂ Removal	38.2	71.1
30	Flue Gas Handling	18.0	24.0
60	Solids Handling	4.6	6.7
70	General Support	1.2	1.9
80	Additional Equipment	4.1	4.0
Total Process Capital		91.2	144.4
General Facilities		9.1	14.4
Engineering and Home Office Fees		9.1	14.4
Project Contingency		19.3	25.1
Process Contingency		8.3	3.5
Total Plant Cost		137.0	201.8
Total Cash Expended		133.7	197.0
Allowance for Funds During Construction		7.5	11.1
Total Plant Investment		141.2	208.1
Royalty Allowance		0.5	0.7
Preproduction Costs		5.2	6.7
Inventory Capital		1.9	0.7
Initial Catalysts and Chemicals		0.0	0.0
Total Capital Investment		148.8	216.2

Table 8.1.3-2

LEVELIZED COSTS

15-year levelizing, 300 MW, 2.6% sulfur coal

Description	GSA (Mills/kWh)	WLFO (Mills/kWh)
Fixed Operating Costs		
Operating Labor	0.53	0.66
Maintenance	1.37	1.74
Administrative and Support Labor	0.32	0.41
Total	2.22	2.81
Variable Operating Costs		
Reagent	1.82	0.65
Solids Disposal	0.85	0.57
Steam	-	0.55
Electricity	0.45	1.16
Total	3.13	2.93
Fixed Charge		
Capital	5.0	7.30
Total	10.35	13.04

8.2 COMMERCIALIZATION PLANS

A set of commercial steps were established by AirPol as part of this program and are in the process of being carried out. They consists of the following elements:

1. Analyze the test data from the test program to determine the operating capabilities for SO₂ removal of the GSA versus conventional spray dryer scrubbers.
2. Evaluate the pilot plant operating results to determine the mechanical design and maintenance requirements to be incorporated in the design of larger systems. In addition, the operating and maintenance cost will be evaluated and compared with competitive designs.
3. Develop a presentation based on the test program results, equipment experience, previous incineration experience (1988 ~ Present) and requirements of the utility marketplace. The presentation will present past experience, test results, capital costs, operating costs and disposal considerations.
4. Develop an estimating capability for bidding and building complete GSA - FGD facilities for smaller industrial boilers, transfer of the technology and for bidding larger USA utility plants. The units will be modular and include computerization of the process calculations, equipment descriptions, cost estimates and general arrangement drawings.
5. Make presentations to people, organizations and conferences within the DOE CCT program to find potential demonstration plants for installation of the GSA technology. The goal is to secure at least one (1) other demonstration installation.
6. Explore export possibilities to developing markets where low capital cost and new technology will be accepted and evaluated along side of developed proven technologies. The goal is to secure at least two (2) installations (hard coal and brown coal).
7. Use the TVA pilot installation and the first full scale demonstration plant(s) and overseas installations to qualify for large USA FGD utility applications.
8. Develop capabilities to service the full FGD utility market by bidding and building FGD process plants based on the prior commercialization work.

AirPol has completed the steps (1) to (5) and is currently working on steps (6) and (7). Some comments as to the results are appropriate prior to discussing the market potential as projected in this point of the commercialization.

In step (1) the initial test pointed out positive aspects of the GSA in removing high SO₂ loading from a coal fired plant. The loading was about 1800 PPM which is in the high sulfur range (1500 ~ 3000 PPM). The GSA was able to reach higher levels of removal than conventional spray dryers while operating at lower approaches to saturation, where the removal rates are enhanced.

When chloride is added the GSA was able to reach even higher levels of removal. Overall the GSA can be expected to equal or exceed the removal efficiency of a conventional semi-dry wheel type spray dryers.

In step (2) the more interesting parts of the test program began to emerge. Significant problems were not experienced in the operation of the unit compared to the operating experience of conventional semi-dry scrubbers. Start-up and commissioning prior to testing was accomplished in less than one month while previous experience at the test facility suggested an anticipate three (3) to six (6) month time frame. Solid build-up did not occur and no mechanical operating difficulties were encountered.

Steps (3) and (4) are ongoing and have reached an acceptable level for securing orders. The presentation and cost estimation will continue to develop and change through the life of the GSA.

Step (5) has been accomplished in the securing of a 50 MW municipal order. The unit will be operational in the second half of 1996.

In step (6) AirPol is active in several export markets and expects to receive the first orders during 1995. The same market barriers exist but the market is more active and compliance dates during 1997 through 1999 are a driving force.

We project full market development in steps (7) and (8) for the GSA will occur after the year 2000 when compliance with phase two of the Clean Air Act Amendment of 1990 must be achieved.

9.0 CONCLUSIONS AND RECOMMENDATIONS

The 10 MW Demonstration of Gas Suspension Absorption project was completed successfully and on time. Following conclusions are drawn from the entire demonstration run.

1. The GSA/ESP process can achieve more than 90 percent SO₂ removal efficiencies at modest stoichiometric ratio (1.30 moles Ca(OH)₂/mole inlet SO₂) and a close approach-to-saturation temperature (8 °F) when treating flue gas resulting from the combustion of a 2.6 percent sulfur, low-chloride (0.02 ~ 0.04 percent) coal.
2. The GSA/ESP process can achieve 90+ percent SO₂ removal efficiencies at a modest stoichiometric ratio and a higher approach-to-saturation temperature (18 °F) with slightly higher levels of chlorine in the coal (0.12 percent).
3. Most of the SO₂ removal efficiency occurs in the reactor/cyclone with relatively low SO₂ removals (2 ~ 5 percentage points) in the ESP.
4. The enhanced mass and heat transfer characteristics of the GSA reactor allows high SO₂ removal efficiencies to be achieved at a very low flue gas residence time. The GSA reactor/cyclone also operates at a high flue gas velocity (20 ~ 25 ft/sec). Thus, the GSA reactor is only one-third to one-fourth the size of the conventional spray dryer vessel, which reduces the capital investment for the GSA system.
5. The expected enhancing effect of chlorine on the SO₂ removal efficiency in the GSA/ESP process was documented. Even modest coal chloride levels (0.12 percent), which are typical of many coals, can provide this effect.
6. The GSA/ESP process has very low particulate emission rates, i.e. well below the NSPS for particulates, when a four-field ESP with a specific collection area (SCA) of 440 ft²/kacfm is used as the particulate control device.
7. The SO₂ removal efficiency in the GSA/PJBH system was typically about 3 ~ 5 percentage points higher than that achieved in the GSA/ESP system at the same test conditions.
8. The GSA system produces a by-product material containing very low moisture levels. This material contains both fly ash and unreacted lime and thus, with the addition of water, undergoes a pozzolanic reaction and can be disposed of in a landfill.
9. The GSA system has no wet/dry interface and the entire system is fabricated from carbon steel rather than high-cost alloy material.
10. The GSA system has lower capital costs and comparable revenue requirements to those for a conventional wet limestone forced-oxidation (WLFO) system for a 2.6 percent sulfur coal application. At lower coal sulfur levels, the GSA system will have lower revenue

requirements than the comparable WLFO system.

As presented in the this report, the GSA process has been designed with proper considerations for existing site conditions, cost economization, environmental impacts and operation concerns. The demonstration unit has achieved all the projected performance and will be commercialized in time for the intended market.

It is concluded that the results of the 10 MW Demonstration of Gas Suspension Absorption have met the goal of the Clean Coal Technology Program.

10.0 REFERENCES

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10. Environmental Monitoring Report; 10 MW Demonstration of Gas Suspension Absorption, Frank E. Hsu, June 1994.
11. The Clean Coal Technology Program 10 MW Demonstration of Gas Suspension Absorption for Flue Gas Desulfurization, Third Annual CCT Conference, Frank E. Hsu, September 1994.

11.0 APPENDIXES

Appendix "A": "Economic Evaluation Report" by Raytheon Engineers & Constructors

Section 3: General Criteria

Section 20: AirPol Gas Suspension Absorber (GSA)

Appendix "B": "10 MW Demonstration of the Gas Suspension Absorption Process at TVA's Center for Emissions Research" by Tennessee Valley Authority

APPENDIX "A"

ECONOMIC EVALUATION REPORT

Section 3: General Criteria

Section 20: AirPol Gas Suspension Absorber (GSA)

By Raytheon Engineers & Constructors

**Prepared for the United States Department of Energy
Under Cooperative Agreement No. DE-F22-90PC90542**

**Cleared by Office of Patent Counsel
Chicago Operations Office
U.S. Department of Energy**

SECTION 3

GENERAL CRITERIA

SYSTEM DESIGN CRITERIA

General design criteria were established for this study to maintain consistency from case to case analyses, as well as with other EPRI-sponsored projects and criteria. Design criteria and sparing requirements were established jointly by UE&C and EPRI. Both process and economic criteria are applicable to all processes evaluated. Criteria specific to each process are presented in the individual FGD system sections. It must be emphasized that the use of premises or assumptions different from EPRI's criteria could alter the comparative ranking of process costs. The dry injection processes involve much less equipment relative to wet FGD. The application of maintenance, as well as process, and project contingency factors based on percentages of capital cost will tend to favor lower capital processes unless their percentages are adjusted upward dramatically to reflect the inherent risk of these lesser developed processes. There is no data on the dry injection processes to validate lower maintenance and operating cost. The items discussed in this section are as follows:

- Changes From Previous EPRI Estimates
- Process Design Criteria
 - Generating Plant
 - Coal and Flue Gas
 - Sulfur Dioxide Removal Requirements
 - Boiler Capacity
 - General Equipment
 - FGD System Battery Limits
 - Raw Materials
 - Solids Disposal
- Economic Design Criteria
 - Capital Estimate Summary
 - Operating Cost Summary

CHANGES FROM PREVIOUS EPRI ESTIMATES

The design and economic criteria for the current cost estimates have changed from those used in previous EPRI cost estimates. The principal changes include the following:

Coal Sulfur - reduced from 4.0% to 2.6%, which matches the Department of Energy coal used in the Clean Coal Technology program. The future computer model will permit sensitivity calculations over the range of 0.5% to 6.0% sulfur.

Engineering - reduced from 12.5% to 10% of total process capital to reflect the experience and knowledge gained from the first generation of FGD systems installed at utility power generating stations. This factor may have to be adjusted upward if legislation leads to a large demand for domestic architect/engineering services that are currently in short supply.

Maturing Technology - Many of the problems with the first generation of scrubbers have been solved, and overseas designs have incorporated numerous advancements. As the industry benefits from the learning curve, design improvements are being incorporated into newer designs. The resulting cost reduction reflects reduced contingency, less component sparing, in-situ oxidation, and reduced reheat.

Financial/Accounting Practice - with the reduced unit size (from 500 MW to 300 MW), the construction period was reduced from three to two years, thereby resulting in a lower adjustment for Allowance for Funds During Construction. In addition, when incorporating the charge for FGD power consumption, the plant net kw are no longer reduced. FGD power is now charged strictly as an operating cost at 50 mills/kwh.

PROCESS DESIGN CRITERIA

Generating Plant

The generating plant chosen for study is a new one-unit, pulverized coal-fired plant producing a nominal 300 MW (net). Operating conditions for the boiler are assumed to be typical of modern units.

The plant is assumed to be located near Kenosha, Wisconsin, at an elevation of 600 ft. above sea level. Atmospheric pressure used for flue gas composition and volume determination is 29.4 in. Hg (99.2 kPa). An ambient temperature range of -25° to 105°F (aver. 60°F) was assumed. Structural criteria are those for Seismic Zone 1 as defined by the Uniform Building Code. Plant design life was assumed to be 30 years for new plant installations (wet and LSD systems) and 15 years for retrofits (dry injection processes).

A fifteen year plant life was selected for those older units which would typically require lower SO₂ removal efficiency in a system-wide compliance strategy. The wet LSF0 and LSD processes were evaluated on both new 30-year and retrofit 15-year plant lives for comparison to the dry FGD economic evaluations.

The reference plant is equipped with a separate particulate removal device (cold, rigid frame electrostatic precipitator) upstream of all wet FGD systems. Dry scrubbing systems are also assumed to use the existing ESP for simultaneous removal of fly ash and SO₂ absorption reaction products. The particulate collection device is assumed to be designed to meet current federal New Source Performance Standards (NSPS) (0.03 lb/10⁶ Btu).

Coal and Flue Gas

The coal used for the base case analysis in the study is a medium sulfur, deep mined, Pennsylvania bituminous coal. The coal analysis is presented in Table 3-1. The coal selected is consistent with the base coal specified in the 1989 EPRI Technical Assessment Guide, EPRI P-6587-L (1).

Boiler performance parameters used for calculating flue gas flow and composition are tabulated in Table 3-2.

Combustion calculations using the coal analysis from Table 3-1 and combustion parameters from Table 3-2 result in the flue gas composition and flow rates shown in Table 3-3. Raw material and utility consumption rates calculated in this study are based on full load flue gas conditions at 100 percent load.

TABLE 3-1
DESIGN COAL ANALYSIS

Proximate Analysis, %	<u>Pittsburgh Seam Freeport Medium Sulfur</u>
Moisture	6.0
Ash	9.1
Fixed Carbon	48.7
Volatile Matter	36.2
Ultimate Analysis, %	
Chlorine	0.12
Moisture	6.0
Carbon	71.3
Hydrogen	4.8
Nitrogen	1.4
Sulfur	2.6
Ash	9.1
Oxygen	4.8
Btu/lb	13,100
Grindability (Hargrove)	59
Sulfur Content lbs/10 ⁶ Btu	1.98

TABLE 3-2
BOILER PERFORMANCE PARAMETERS

Excess Air	
Furnace Outlet	20%
FGD System Inlet	37%
Ambient Air Humidity (80°F, 60% Relative Humidity)	0.013 lb H ₂ O/lb Dry Air
FGD System Inlet Flue Gas Pressure	1 in. w.g. greater than FGD System Pressure Drop
Total Coal Sulfur Converted to SO ₂	98.4%
SO ₂ Removed with Ash	0% assumed
Boiler Efficiency	88.5% for Design Coal
ID Fan Outlet Temperature	282°F

TABLE 3-3
 FLUE GAS CHARACTERISTICS
 FOR ONE 300 MW UNIT

	Freeport, Pennsylvania Medium Sulfur	
	%-Volume	Lb/Hr
O ₂	5.37	186.0 x 10 ³
CO ₂	12.14	578.7 x 10 ³
SO ₂	0.17	11.45 x 10 ³
SO ₃	0.002	140
HCl	0.007	270
H ₂ O	7.55	147.2 x 10 ³
N ₂ /NO _x	74.761	2,267.8 x 10 ³
 Total	 100	 3,191.6 x 10 ³
 Fly Ash Produced		 8.4 tph
 SCFM*		 0.685 x 10 ⁶
ACFM		1.030 x 10 ⁶

*Standard conditions are defined as 60°F and 29.92 in. Hg

Sulfur Dioxide Removal Requirements

The latest federal New Source Performance Standards (NSPS) require a 90 percent overall reduction in potential SO₂ emissions for the medium sulfur coal (2.6 percent sulfur in the design coal) used in this study. The removal efficiency is to be based on a 30-day rolling average. This study criteria assumes a nominal FGD design efficiency of 90 percent, on a continuous 100 percent load basis. For those processes which cannot achieve 90% removal efficiency, the reported maximum SO₂ removal capability is assumed. These nominal efficiency values are used to develop material balances which provide the basis for calculation of the annual operating cost and equipment sizing.

The reduction efficiency of an FGD system may, in actual practice, temporarily fall below the NSPS limits due to plant upsets and operational problems. The study makes several conservative assumptions which should offset these periods of low performance:

1. No credit is given in this study to sulfur entrained in the fly ash or bottom ash. Consequently the actual removal rate is higher than the design values.
2. No allowance is given for the fact that utilities can compensate for low removal efficiencies by:
 - increasing the SO₂ removal efficiency above the NSPS limit during partial load and non-peak periods.
 - increasing the absorptive capacity of the scrubbing solution. This involves increasing the pH of a lime, limestone or sodium system or increasing the concentration of other active species.
 - increasing the L/G ratio.
 - operating the spare absorber module.

Boiler Capacity

The base case boiler capacity data shown in Table 3-4 were compiled from an update to the EPRI TAG report(2). The base case net plant heat rate includes a four percent allowance for plant auxiliaries, but does not reduce the net plant output due to FGD steam or power consumption. The steam and power usage developed for each FGD process are treated strictly as operating costs using the economic criteria presented later in this section. These Economic Design Criteria describe in more detail the method used to account for the steam and

TABLE 3-4
- BOILER OPERATING PARAMETERS

	<u>Medium Sulfur</u>
Average Steam Cycle Heat Rate, Btu/kwh	8,246
Average Boiler Efficiency, %	88.5
Average Gross Heat Rate, Btu/kwh	9,318
Average Net Heat Rate, Btu/kwh	9,722
Average Heat Input to Boiler, 10 ⁶ Btu/hr	2916.5
Average Coal Heat Value, Btu/lb	13,100
Average Coal Burn Rate, tph	111.3
Net Output, MW (excluding FGD system energy consumption)	300

power requirements of each FGD process. The combustion conditions described in Table 3-4 result in a coal flow rate and flue gas flow rate equal to that used in EPRI report CS-1428, "FGD Economic and Design Factors" (3).

General Equipment

Certain equipment requirements and criteria are common to all medium sulfur coal, wet FGD processes. It has been assumed that scrubbed gas reheat of 25°F over the saturated gas temperature is required for the wet FGD system base case analysis. Reheat is provided using steam-heated ambient air. Reheat steam coils use both latent heat and sensible heat by subcooling the condensate to 200°F, prior to its return to the boiler feedwater cycle. All wet systems include 100 percent flue gas bypass duct work. Ducts are sized for 3600 fpm gas velocity for normal operation. Dry and semi-dry FGD systems are assumed to require no flue gas reheat, with the exception of the LIFAC process.

Combined induced draft (ID)/booster fans have been included for all systems. The fans provide the required differential pressure to overcome the FGD system pressure drop requirements. Two fans are assumed to operate simultaneously in parallel. The total flange-to-flange pressure drop of the FGD system determines the total fan pressure requirement.

Equipment redundancy has been incorporated into all base case FGD systems to increase reliability. Equipment typically is spared if its loss would require immediate system shutdown. Absorber vessels include 3-50 percent capacity modules. Pumps are 100 percent spared where the loss of a single pump would require shutdown. No sparing is used where a bank of recycle pumps supply several spray levels in an absorber. An uninstalled warehouse spare pump has been included in these cases. Items such as tanks, bins, silos, agitators, and heat exchangers are not spared. By-product production and additive feed systems are assumed to contain spare critical components for continuous operation and are not spared as a system. It is also assumed that, should shutdown be necessary, surge capacity is included in the by-product and additive systems to allow repair without shutdown of the FGD system.

Equipment sparing and sizing criteria is given in Table 3-5.

TABLE 3-5
EQUIPMENT SPARING AND SIZING CRITERIA

<u>Equipment Item</u>	<u>Sparing/Capacity</u>	<u>Sizing Criteria</u>
Absorbers	3 @ 50%	Spray Tower - 10 ft/s gas velocity, 10 sec gas residence time*
Recycle Tanks	3 @ 50%	6 minute retention time (except CT121 & Pure Air)
Mills and Hydration Systems	2 @ 100%	Ball Mill (Lime) - 90%-200 mesh 7 hp/tph Ball Mill (Limestone) - 90%-200 mesh 18 hp/tph Roller Mill - 95%-325 mesh
Silos	2 @ 50% (Bulk Storage) 1 @ 100% (All others)	Raw reagent bulk storage - 60 days Prepared reagent and product solids - 3 days
Bins	2 @ 67% 1 @ 100%	Raw reagent and fixative - 30 hrs Fly ash and recycle solids - 8 hrs
Tanks	1 @ 100%	Slurry tanks - 8 hrs, 90% full Water tanks - 30 min, 90% full Rxn mix tanks - 6 min, 90% full
Pumps	2 @ 100%**	70% efficiency
Fans	2 @ 50% (ID Booster) 1 @ 100% (Reheat Air)	80% efficiency, 120% design ACFM, 140% design pressure drop
Compressors	2 @ 100%	75% efficiency
Ductwork	100% capacity for all ductwork including flue gas bypass	3600 fpm gas velocity
Vacuum Filter	2 @ 100%	LSFO, LS/DBA - 150 lb/hr/ft ² LS/INHIBIT - 125 lb/hr/ft ² LDA, MagLime - 100 lb/hr/ft ²
Thickener	2 @ 67%	LSFO, LS/WB, LS/DBA - 2 ft ² /TPD dry LS/INHIBIT - 5 ft ² /TPD dry LDA - 18 ft ² /TPD dry MagLime - 20 ft ² /TPD dry
Pneumatic Conveyor System	1 @ 100% (1 spare blower)	-----
Heat Exchangers	1 @ 100%	-----

*Pure Air Process cocurrent spray tower absorber is sized at 18 ft/s gas velocity and 1.7 seconds gas residence time.

**No installed spares are assumed for recycle spray pumps. Only one warehouse spare is included for each absorber.

System Battery Limits

General. Battery limits are required to determine the scope of equipment and subsequent cost impact on each of the FGD processes. It is sometimes difficult to differentiate between the cost of FGD and particulate removal systems. This difficulty arises from the following interrelationships:

- Spray dryer and sorbent injection processes use a particulate collector to perform the dual function of removing fly ash and the reaction products. The particulate collector cost is not included.
- The solids collected in the ESP (or FF) downstream of a dry FGD process is a mixture of fly ash and absorbent product. This mixture is pneumatically conveyed from the particulate device to storage silos and transported to the landfill.
- Processes producing a sludge typically combine fly ash with the sludge as a stabilizing agent. This mixture of fly ash and sludge is transported to the landfill.
- Portions of flue gas duct are shared by both particulate removal and FGD equipment.

Because these areas are shared by both systems, the capital and operating cost of particulate removal and flue gas desulfurization cannot be easily separated. A systematic methodology is needed to obtain the FGD cost exclusive of the particulate removal cost. This study will use the following approach:

- The existing particulate system is assumed to be adequate for any increased grain loading caused by the FGD retrofit.
- A differential cost is developed for any additional cost associated with removal of solids from the existing particulate device. This could include installation of extra insulation and/or the replacement of a wet ash handling system with a dry pneumatic conveying system.
- This differential cost is added to the cost of the FGD system to derive the total FGD system cost.

FGD System Battery Limits. The cost for the FGD system is developed within the following battery limits.

- Duct work - From the plant air preheater outlet to the chimney including all dampers, support steel, manifolds, etc.
- ID fan - That portion of the capital and operating cost of ID/booster fans necessary to provide the pressure to convey the flue gas through the FGD system and associated ductwork is considered.

- Raw water - Generating plant raw water system at the boundary of the FGD system and the generating plant. Supply conditions are 50 psig and 60°F. The assumed chemical analysis is given in Table 3-6.
- Steam - Process steam is assumed to be available from the generating plant turbine cycle at main steam, cold reheat, and crossover extraction points. Minimum available pressures and corresponding temperatures at each extraction point are assumed to exceed:

Main Steam	2400 psig, 1000°F
Intermediate	418 psig, 572°F
Crossover	84 psig, 665°F
- Electrical power - Power supply to the FGD system is assumed to be available from the generating plant transformer yard to the FGD plant boundary as 34.5 kV feeders. All necessary electrical equipment for voltage reduction and distribution is included in the FGD system.
- Emergency power - Sufficient emergency power is available from the generating plant emergency power system, including an emergency power bus for the FGD system. The FGD system includes switching and distribution for critical service items.
- Solids disposal - Solids disposal includes transportation and ultimate placement in a landfill for the base case.

Reagents and Raw Materials

Raw material compositions assumed for the FGD process evaluations are listed in Table 3-7.

Solids Disposal

Depending on the process an FGD system can have many product streams:

- Prescrubber purge - A prescrubber purge stream is required in some FGD processes to remove fly ash, SO₂, chloride, and/or to presaturate the gas prior to entering the absorber. Each prescrubber has an acidic blowdown stream which is disposed of with the fly ash or pumped to the cooling tower blowdown pond at no cost to the FGD process.
- Fly ash - The fly ash and reaction products collected by the ESP are pneumatically conveyed to fly ash disposal silos for storage. These solids are unloaded from the silo by rotary unloaders and transported one mile by truck to the landfill for disposal. The landfill average placement depth is 30 feet. Cost for fly ash disposal is not included in the operating cost.

TABLE 3-6

RAW WATER CHEMICAL ANALYSIS

<u>Constituent</u>	<u>ppm</u>
SiO ₂	2.1
Fe	0.04
Mn	0.0
Ca	35
Mg	10
Na	4.1
K	1.0
HCO ₃	134
CO ₃	0
SO ₄	19
Cl	6.5
F	0.1
NO ₃	0.4
Dissolved Solids	159
Hardness (as CaCO ₃)	129
Non-carbonate Hardness	18
pH	8.0

TABLE 3-7

REAGENT AND RAW MATERIALS COMPOSITIONS

Limestone	-	CaCO ₃ , 94% by weight (dry basis)
Lime	-	CaO, 90% by weight (dry basis)
Soda Ash	-	Na ₂ CO ₃ , 99.8% by weight (dry basis)
Formic Acid	-	90% by weight
Dibasic Acid	-	90% by weight
Sulfur Emulsion	-	70% by weight sulfur in water
MagLime	-	5-8% MgO, 90% CaO by weight (dry basis)

- Fly ash/absorber product mixture - Fly ash plus the absorber product solids are collected in the ESP in dry or semi-dry processes. This mixture is pneumatically conveyed to silos for storage. The solids are unloaded from the silo by rotary unloaders and transported one mile by truck to the landfill. A lined landfill is used for sodium based systems and an unlined landfill is used for lime based systems. The landfill depth is 30 feet.
- Gypsum - Slurry product from wet systems containing a high ratio of sulfate to sulfite (99%+) is dewatered and trucked to an unlined landfill located one mile from the plant and impounded to a height of 30 feet.
- Sludge from wet systems containing a high ratio of sulfite to sulfate is neutralized/stabilized with lime and fly ash and trucked one mile to a landfill site. An unlined landfill is used for all sludge disposal sites, except for the lime dual alkali and MagLime processes where the site is lined with an impermeable membrane. The average landfill depth is 30 feet.

Additional criteria and guidance for solids disposal are available from EPRI (4).

ECONOMIC DESIGN CRITERIA

The economic criteria for all evaluated FGD processes were standardized to provide consistent economic comparisons. The cost development and breakdown follow EPRI Economic Premises (included in Section 4 as drawn from the 1989 TAG), and are based on January 1990 dollars with a January 1, 1990 plant startup.

The base case plant net output shown in Table 3-4 does not account for the varying quantities of steam and power consumed by each FGD process. The steam and power requirements draw energy from the boiler, which would otherwise be included in the net output of the plant. Consequently, adding the FGD steam and power usage to the base case heat requirements reduces the net MW output of the plant. However, accounting for this energy consumption by only reducing the net plant MW will not result in a change in the levelized cost in terms of \$/ton of SO₂ removed. Therefore, steam and power usage is reflected only as a direct cost to the plant (50 mills/kwh and \$2.85 - \$5.30/1000 lbs steam depending on its pressure). No reduction in net plant power output is assumed for the cost calculations.

None of the capital and operating costs for the particulate removal process have been included in the FGD system economic analysis. Only costs for required ESP or FF modifications are included. A cold-side ESP (560 SCA) is assumed to operate upstream of the wet FGD systems (and downstream of the dry FGD systems) to ensure that NSPS for particulate are met and that the ash loading of the flue gas stream entering the scrubber system is held to an acceptable level. A reverse gas fabric filter has been assumed to operate downstream of the absorber for the lime spray dryer (LSD) FGD system only. No cost differential for this FF has been assumed in the LSD cost estimate.

Capital Estimate Summary

The bases and items included for each component of the capital estimate are summarized in the following sections.

Process Capital. Process Capital is the total constructed cost of on-site FGD and related facilities, including direct and indirect construction costs.

Table 3-8 is a listing of those items included in the estimate of process capital. FGD system capital cost components are listed in Table 3-9.

The equipment size is determined from mass balance data developed for each FGD process. Quotes were obtained from three or four bidders for all major pieces of equipment. Prices were then cross-checked with a national computerized material cost tracking network. Bulk quantities and installation labor charges were developed for each equipment item over a wide range of sizes to arrive at installation factors for each piece of equipment. Labor manhours were adjusted for local productivity rates (Kenosha, WI for the base case).

The process capital estimate is broken down on an area basis for each process. All processing areas are not applicable for each process, but some are common to all processes. The cost areas are listed in Table 3-10.

General Facilities or Off-Site Capital. The general facilities include roads, office buildings, shops, laboratories, etc. EPRI guidelines for this cost item are a 5 to 20% share of the process capital cost. A value of 10% of the total process capital was chosen for this study.

Engineering and Home Office Fees. An estimate of engineering, home office overhead and fees is included for costs representative of this type of plant. EPRI guidelines for this cost item are 10 to 15% of the process capital for this cost. A value of 10% was chosen for this study.

Project Contingency. Project contingency covers additional equipment or other costs which would result from a more detailed design. Project contingency factors will be based on the EPRI Class III cost estimate guidelines (10 to 20%) and applied on an area-by-area basis. Higher contingency factors will be applied to those equipment items of special design, and lower factors to those capital purchases that are standard, off-the-shelf items. In general, a value of 15% will be used for all standard equipment.

Process Contingency. A process contingency is applied to a new technology in an attempt to quantify the design uncertainty and cost of a commercial-scale system. The contingency is related to the level of process development. The process contingencies are expressed as percentages of the area process capital

TABLE 3-8
ITEMS INCLUDED IN PROCESS CAPITAL COST ESTIMATE

Earthwork
 Concrete
 Buildings and Structures
 Process Equipment
 Piping
 Electrical
 Painting
 Instruments and Controls
 Insulation
 Direct Field Labor
 Indirect Field Costs
 Payroll Taxes
 Insurance, Bonds
 Construction Supplies
 Temporary Facilities
 Construction Equipment
 Vendor Fees

TABLE 3-9
FGD SYSTEM CAPITAL COST ESTIMATE COMPONENTS

<u>Capital Investment</u>	<u>\$/kw</u>
Process Capital (includes sales tax)	A
General Facilities	B
Engineering and Home Office Fees	C
Project Contingency	D
Process Contingency	E
Total Plant Cost, TPC	= A+B+C+D+E
Total Cash Expended, TCE	TPC x Adjust factor*
AFDC (Allowance for Funds During Construction)	F
Total Plant Investment, TPI	= TCE + F
Royalty Allowance	G
Preproduction Costs	H
Inventory Capital	I
Initial Catalyst and Chemicals	J
Land	K
Total Capital Requirement, TCR (Capital Investment Jan., 1990, Startup Jan., 1990)	<u>TPI+G+H+I+J+K</u>

*Adjustment Factor for TCE per table 3-4 of EPRI TAG (P6587-L, Reprinted in Section 4 of this report)

TABLE 3-10

COST AREAS FOR PROCESS CAPITAL BREAKDOWN

<u>Area</u>	<u>Description</u>
10	Reagent Feed System
20	SO ₂ Removal System
30	Flue Gas System
40	Regeneration System
50	By-product System
60	Solids Handling System
70	General Support Area
80	Miscellaneous Equipment

- 10: Reagent Feed System - all equipment required for storage, handling and preparation of raw materials, reagents, and additives used in each process.
- 20: SO₂ Removal System - equipment required for SO₂ scrubbing, such as the absorption tower, recirculation pumps and other associated equipment.
- 30: Flue Gas System - duct work and fans required for flue gas distribution to the SO₂ scrubbing system, plus gas reheat as required.
- 40: Regeneration System - specific to regenerable reagent systems, equipment used to regenerate spent absorbent for return to the process, plus any preconditioning system for SO₂ or H₂S off gas.
- 50: By-product System - production equipment for salable process by-products and storage facilities for the final products.
- 60: Solids Handling System - equipment required for fixation, treatment, and transportation of all sludge/dry solids materials produced by each scrubbing process.
- 70: General Support Area - additional equipment required to support FGD system operation such as makeup water and instrument air.
- 80: Miscellaneous Equipment - This area will include plant modifications necessitated by the addition of the FGD system. Also included are costs for electrical equipment tie-ins and other associated systems.

costs. Based on EPRI guidelines, 2 to 50% factors will be applied to the total capital investment requirement for each cost area. Since some FGD systems utilize common equipment in some areas and new, untested designs in others, the process contingency is applied by cost area. Generic reagent handling systems would receive a low process contingency, while a new regeneration system tested only at the pilot plant level would receive a high process contingency within the same process cost estimate.

Sales Tax. Sales tax is included in the process capital estimate.

Total Plant Cost (TPC). The total plant cost is the sum of the process capital, general facilities, engineering and home office fees, and contingencies.

Royalty Allowance. The royalty allowance was established by EPRI at 0.5 percent of the process capital.

Preproduction Costs. Preproduction costs are intended to cover operator training, equipment checkout, major changes in plant equipment, extra maintenance, and inefficient use of materials during plant startup.

Preproduction costs are estimated as follows:

- One-month fixed operating costs (operating and maintenance labor, administrative and support labor, and maintenance materials).
- One-month of variable operating costs at full capacity (chemicals, water, and other consumables, and solids disposal charges). Full capacity estimates of the variable operating costs will assume operation at 100% load.
- Two percent of the Total Plant Investment, TPI (to cover expected changes and modifications to equipment required to bring the FGD system up to full capacity).

Inventory Capital. The inventory capital includes the value of raw materials and other consumables on a capitalized basis. The inventory capital is estimated using the raw material supply based on 100% capacity operation during 60 days.

Initial Catalyst and Chemicals. The initial cost of any catalyst or chemicals contained in the process equipment is included. Those materials in storage are included in inventory capital.

Allowance for Funds During Construction (AFDC). The schedule for engineering, procurement, and construction of a 300 MW Wet FGD system is assumed to be two years for installation of the entire system. The Total Plant Cost (TPC) is given in January, 1990 dollars. For an escalation rate of 5% per year, the TPC is multiplied by 0.0548 to calculate the allowance for interest expenses during a two year construction period.

Total Cash Expended (TCE). The total cash expended is an estimate of the total cash expended over the construction duration of the FGD system. The TCE includes only the escalation up to various dates of expenditures and does not include the AFDC.

Total Plant Investment (TPI). The total plant investment is the sum of the total cash expended and the allowance for interest during construction.

Total Capital Requirement (TCR). All capital necessary to complete the project including AFDC, raw material inventory, royalties and preproduction costs.

Land. FGD system land requirements include the plant site area and disposal area. Land has not been included as a separate line item. Land requirements are similar for wet processes as a group and for dry processes as a group. Land costs will vary with site location.

Operating Cost

Operating costs for the FGD systems are separated into fixed and variable operating costs. Fixed operating costs include operating and maintenance labor, maintenance materials, and administrative/support labor. Factors for these costs are based on EPRI Economic Premises. Variable operating costs include consumables such as fuel, water, power, chemicals, and solids disposal. Table 3-11 summarizes the criteria used for these operating costs.

TABLE 3-11
OPERATING COST CRITERIA

Fixed Operating Costs	Unit	Rate
Operating Labor	Man-hrs	\$20.00 (Jan. 1990)
Maintenance Labor - Slurry Handling	\$/yr	3.2% of Process Capital
- Liquid Handling	\$/yr	1.2% of Process Capital
Maintenance Material - Slurry Handling	\$/yr	4.8% of Process Capital
- Liquid Handling	\$/yr	1.8% of Process Capital
Administrative & Support Labor	\$/yr	30% of Operating and Maintenance Labor

Variable Operating Costs	Unit	Jan. 1990 Cost/Unit	Additional Freight \$/t	30-Yr Level Factor	30-Yr Levelized Cost/Unit
Fuel Oil (No. 6)	gal	\$ 0.41	Incl	1.613	\$ 0.66
Raw water	1000 gal	\$ 0.60	Incl	1.613	\$ 0.97
Cooling water	1000 gal	\$ 0.16	Incl	1.613	\$ 0.26
Power	kwh	50 mills	Incl	1.668	83 mills
Methane	1000 ft ³	\$ 3.00	Incl	1.668	\$ 5.00
Lime	ton	\$ 55.00	Incl	1.613	\$ 88.72
Limestone	ton	\$ 15.00	Incl	1.613	\$ 24.20
Soda Ash	ton	\$ 93.00	\$ 43.00	1.613	\$ 219.37
Magnesia	ton	\$ 232.00	\$135.00	1.613	\$ 591.97
Ammonia	ton	\$ 145.00	\$ 5.50	1.613	\$ 242.76
Sulfur Emulsion	ton	\$ 220.00	Incl	1.613	\$ 354.86
Dibasic Acid	ton	\$ 360.00	Incl	1.613	\$ 580.68
Formic Acid	ton	\$ 800.00	\$20 (est.)	1.613	\$1322.66
Allied Catalyst	ton	\$2500.00	\$45.00	1.613	\$4105.09
Claus Catalyst	ton	\$1000.00	Incl	1.613	\$1613.00
Disposal Charges,					
Dry Solids (lined)	ton (dry)	\$ 9.29	Incl	1.613	\$ 14.98
Fly ash (unlined)	ton (dry)	\$ 8.00	Incl	1.613	\$ 12.90
Fly ash with Nahcolite	ton (dry)	\$ 11.14	Incl	1.613	\$ 17.97
Sludge (trucked to lined landfill)	ton (dry)	\$ 9.25	Incl	1.613	\$ 14.92
Sludge (trucked to unlined landfill)	ton (dry)	\$ 8.15	Incl	1.613	\$ 13.15
Sludge (ponded)	ton (dry)	\$ 6.00	Incl	1.613	\$ 9.68
Gypsum (pumped & stacked)	ton (dry)	\$ 4.75	Incl	1.613	\$ 7.66
Condensate	1000 lb.	\$ 0.77	Incl	1.613	\$ 1.24
Steam					
0-70 psia	1000 lb.	\$ 2.85	Incl	1.668	\$ 4.75
70-250 psia	1000 lb.	\$ 3.50	Incl	1.668	\$ 5.84
250-400 psia	1000 lb.	\$ 5.30	Incl	1.668	\$ 8.84
By-product Credit					
Sulfur	Long ton	\$ 90.00	Incl	1.613	\$ 145.17
Sulfuric Acid	ton	\$ 50.00	Incl	1.613	\$ 80.65
Liquid Sulfur Dioxide	ton	\$ 230.00	Incl	1.613	\$ 371.00
Gypsum	ton	\$ 2.00	Incl	1.613	\$ 3.23

Note: 1000 gal = 3.785 m³, t (short) = 0.9072 t (SI),
1000 ft³ = 28.32 m³, 1000 lb = 453.6 kg

Assumed 100 mile shipping distance for all chemical reagents except where noted.

Levelized Cost

The capital and operating costs determined for each FGD system are converted to a levelized cost using a fixed charge rate for capital costs, and appropriate levelization factors for operating costs. Levelization factors for processes evaluated over a 30-year service life are listed in Table 3-11. Levelization factors for 15-year service lives are 1.371 for labor and other variable operating costs with the exception of power and steam (L.F. = 1.4). A fixed charge rate specific to the plant life and discount rate assumed is multiplied times the total capital requirement to determine the capital charges for inclusion in the levelized cost estimate. Both constant and current dollar totals are presented in each case. The total levelized cost of each FGD system is presented as mills/kwh, \$/kw-yr and in terms of \$/ton SO₂ removed.

References:

1. "Technical Assessment Guide: Volume 1: Electricity Supply 1989," EPRI Report No. P-6587-L, Revision 6 (September 1989).
2. "FGD Economic and Design Factors," EPRI Report No. CS-1428 (April 1980).
3. "FGD Sludge Disposal Manual," EPRI Report No. CS-1515 (September 1980).

Section 20

AirPol GAS SUSPENSION ABSORBER (GSA)

SYSTEM DESCRIPTION

The AirPol Gas Suspension Absorber (GSA) FGD system is a dry, throwaway process. The evaluation presented in this section is done on the basis of a retrofit plant with an expected 15-year service life. The AirPol GSA is intended for retrofit as well as new plants.

In this system, hot flue gas enters a venturi-shaped bottom of a vertical reactor where it contacts a mixture of suspended solid particles, made up of hydrated lime reagent, fly ash and recycled reaction products. Fresh lime slurry is sprayed directly into the suspended bed of particles. The water in this slurry humidifies the flue gas to improve SO₂ removal efficiency. Humidification would also improve downstream ESP performance due to conditioning of the flue gas. A relatively high gas velocity (24 fps compared to wet FGD towers operating at 9-10 fps) is required to fluidize and convey the solids in the reactor, further promoting SO₂ reaction with the lime solids. The spent solids and fly ash are carried out of the top of the absorber and 99% of the solids exiting the reactor are collected in a cyclone. Most of the solids collected in the cyclone are recirculated back to the system to achieve greater sorbent utilization. The remaining solids in the flue gas are captured downstream in the ESP. These waste solids are transported to a landfill for disposal.

Process Description

A moving fluid bed is created in the GSA reactor by flue gas flowing upward through a suspended bed of fine-grained solids. Unlike a classical fluidized bed where the particles retain a top horizontal surface within the reactor, the gas velocity is high enough to carry the solids out of the reactor where they are then largely captured and reintroduced to the reactor. By this mechanism, the solids may be rewetted providing a large reactive surface.

The classical fluidized bed was first used in the 1920's for coal gasification in the Winkler generator. Industrial applications of the circulating fluid bed (CFB) began in the 1960's, where it was initially used for alumina calcination in the aluminum industry. "Gas suspension absorption" or GSA was first developed in Europe as a method to calcine limestone for use in cement production. When used in power plant applications, the GSA process shows the potential to combine some of the economic benefits of lime spray dryers with sulfur dioxide removal efficiencies close to those of wet scrubbing processes (i.e., greater than 95% removal capability).

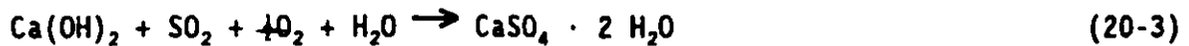
In the AirPol GSA system, there are several process parameters which promote high desulfurization efficiencies. One parameter is the very large concentration of fly ash, reaction products, and unreacted lime that build up inside the reactor. This concentration will normally be as high as 500-1500 g/Nm³ (200-800 grains/acf) which is 50-100 times as much as would be present in a conventional spray dryer. The advantages of this high concentration of solids in the reactor is two fold; (1) it enables rapid evaporation of the water from the lime slurry coating the surface of each particle; (2) it prevents solid build-up on the reactor walls, avoiding a common problem for spray dryers. Another parameter is the relatively long time the reagent solids are in contact with the flue gas. This is achieved by recycling the lime particles through the absorber up to 100 times.

Chemistry and Criteria

Raw lime (in pebble form) is slaked with an excess of water to form a calcium hydroxide slurry, by the following reaction:



For the base case evaluation, the flue gas enters the gas suspension absorber at 277°F. The slaked lime slurry is injected into the reactor through a dual-fluid nozzle in the venturi throat. As the flue gas flows up through the reactor, it mixes intimately with the lime reagent to form calcium salt reaction products according to the following equations:



The predominant sulfation reaction in the absorber is Equation 20-2, producing hydrated calcium sulfite. As the sorbent particles pass through the system, some of the unreacted lime reacts with the carbon dioxide and hydrogen chloride in the flue gas to form calcium carbonate and calcium chloride:



The spent sorbent, unreacted reagent, and fly ash exiting the GSA reactor are collected downstream by a cyclone-type collector and ESP or Fabric Filter.

Table 20-1 summarizes the specific process design criteria developed for the economic evaluation of the AirPol GSA process.

Process Design

The AirPol GSA system will be discussed according to Figure 20-1 in the following order:

- Reagent Feed System, Area 10
- SO₂ Removal System, Area 20
- Flue Gas System, Area 30
- Solids Handling System, Area 60
- General Support Equipment, Area 70
- Additional Equipment, Area 80

Equipment listings for each area are given at the end of this section.

Table 20-1
AirPol GAS SUSPENSION ABSORBER (GSA)
PROCESS DESIGN CRITERIA*

Flue Gas Handling Area Criteria

Flue gas flow rate**	1,020,000 ACFM @ 277°F
GSA pressure drop	5.6 in. H ₂ O
Mechanical collector pressure drop	3.4 in. H ₂ O
Duct and chimney pressure drop	1 in H ₂ O
Total system pressure drop (including mechanical collector)	10.0 in. H ₂ O
Flue gas temp. (air heater outlet)**	277°F
Approach to adiabatic saturation temperature	18°F

SO₂ Removal Area Criteria

SO ₂ removal (2.6% S Coal)**	90%
Absorber - type	Gas suspension absorber
- capacity**	3 @ 50% capacity (2 op, 1 spare)
Gas residence time in absorber	3.1 sec
Gas velocity in absorber	24 ft/sec
Slurry water injection - type	Dual-fluid nozzle
- air pressure	100 psi
Mechanical collector - type	Separating Cyclone
- capacity	6 @ 25% capacity (4 op, 2 spare)
Total solids recycle	<u>0.067 lb solids</u> Ft ³ Flue Gas
Interstitial water retained by solids	Less than 1%

Reagent Feed Area Criteria

Reagent - purchased	Pebble lime (10% inerts)
- absorber feed	Lime slurry
Lime storage silo capacity**	60 days
Lime feed bin capacity**	30 hours
Slaking system - type	Detention Slaker
- capacity**	3 @ 50% capacity (2 op, 1 spare)
Reagent feed rate	<u>1.30 moles Ca in feed</u> mole inlet SO ₂

Solids Handling Area Criteria

Solids silo capacity (total)**	3 days
Sulfite/sulfate mole ratio	4/1

General Support Area Criteria

Makeup water tank capacity**	30 min
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*For one 300 MW unit

**Identical for all processes evaluated

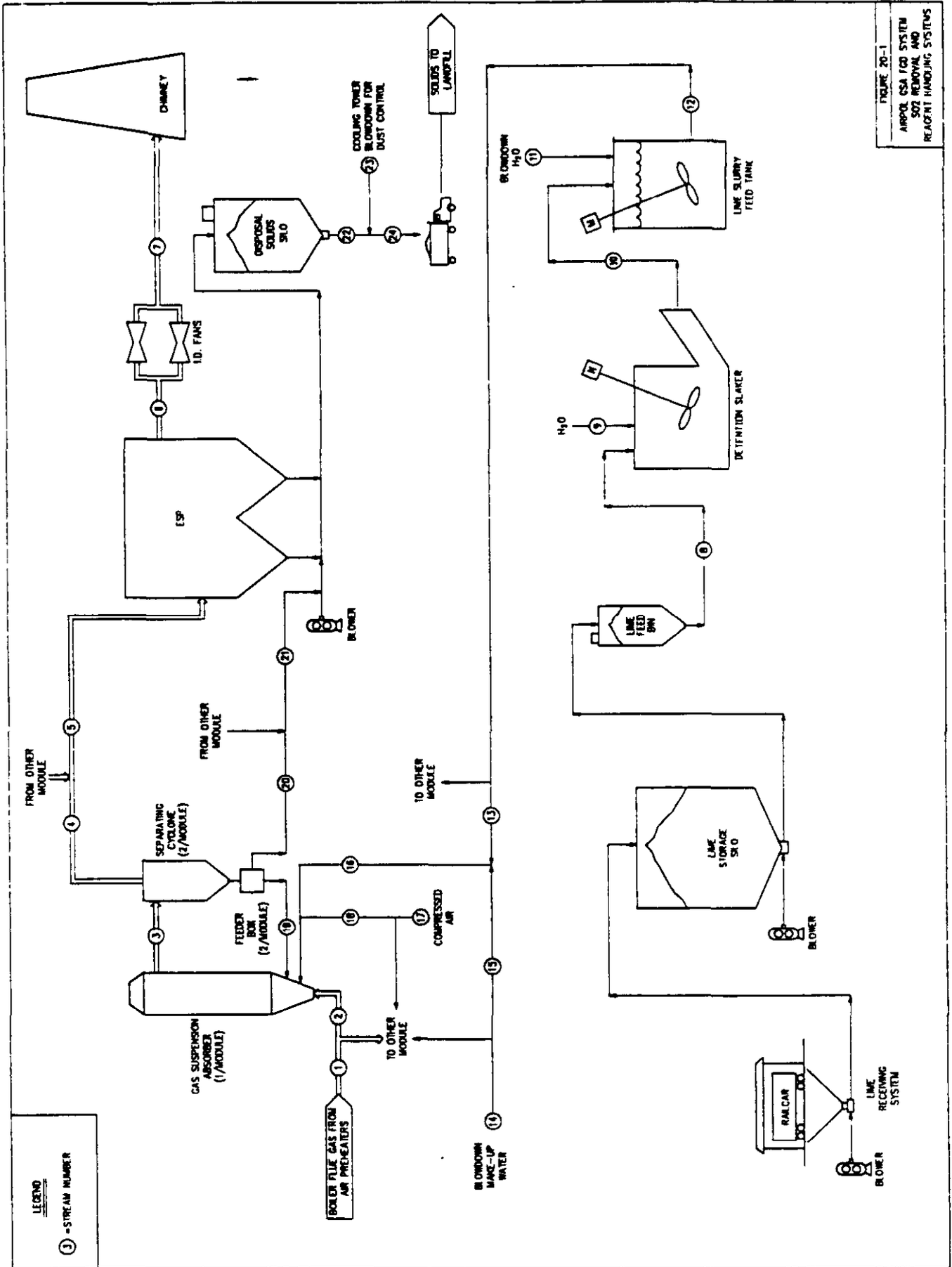


FIGURE 20-1
AIRPOL CSA/CO SYSTEM
SO2 REMOVAL AND
REAGENT HANDLING SYSTEMS

Table 20-2

**MATERIAL BALANCE
AirPol GSA PROCESS
(FOR FIGURE 20-1)**

Stream Number ----->	1	2	3	4	5	6	7		
Stream Name ----->	Flue Gas Downstrm of Air Heater	Flue Gas to GSA Reactor	Flue Gas after GSA Reactor	Flue Gas after Cyclone	Total Flue Gas Upstrm of ESP	Flue Gas Downstrm of ESP	Flue Gas to Chimney		
Temperature, °F	277	277	145	145	145	145	145		
Pressure, " H2O	-12.0	-12.0	-17.6	-21.0	-21.0	-24.0	1.0		
K SCFH	685	343	359	355	711	711	711		
K ACFM	1,019	510	445	444	868	895	839		
CO2, K Lb/Hr	578.7	289.4	288.7	288.7	577.5	577.5	577.5		
N2, K Lb/Hr	2,267.6	1,133.8	1,136.1	1,136.1	2,272.3	2,272.3	2,272.3		
SO2, K Lb/Hr	11.5	5.7	0.6	0.6	1.1	1.1	1.1		
O2, K Lb/Hr	186.1	93.1	93.4	93.4	186.9	186.9	186.9		
HCl, K Lb/Hr	0.3	0.1	0.0	0.0	0.0	0.0	0.0		
Other Gases, K Lb/Hr	1.4	0.7	0.6	0.6	1.3	1.3	1.3		
H2O, K Lb/Hr	147.2	73.6	120.0	109.6	219.3	219.1	219.1		
Fly Ash, (solids) K Lb/Hr	16.8	8.4	2,087.0	20.9	41.7	0.087	0.087		
Total, K Lb/Hr (Gas Only)	3,192.8	1,596.4	1,639.5	1,629.1	3,258.3	3,258.1	3,258.1		
Stream Number ----->	8	9	10	11	12	13	14	15	
Stream Name ----->	Lime to Slaker	Water to Slaker	Slaked Lime from Ball Mill	Dilute H2O to Slurry Feed Tank	Slurry from Slurry Feed Tank	Slurry to GSA Reactor	Blowdown Makeup Water	Blowdown Water to Each GSA	
Temperature, °F	60	60	60	60	80	80	60	60	
Flow, GPM	--	64	67	56	120	60	29	15	
Wt % Solids	100%	0%	40%	0%	25.0%	25.0%	0.0%	0%	
Inerts, K Lb/Hr	1.5	--	1.45	--	1.5	0.7	--	--	
Ca(OH)2, K Lb/Hr	--	--	17.25	--	17.3	8.6	--	--	
CaO, K Lb/Hr	13.1	--	--	--	--	--	--	--	
H2O, K Lb/Hr	--	32.3	28.1	28.1	56.1	28.1	14.55	7.3	
Total, K Lb/Hr	14.5	32.3	46.8	28.1	74.8	37.4	14.6	7.3	
Stream Number ----->	16	17	18	19	20	21	22	23	24
Stream Name ----->	Slurry to Spray Nozzle	Atomizing Air from Compressor	Atomizing Air to each GSA	Recycle Solids to each GSA	Disposal Solids from Cyclone	Total Solids from Cyclones	Disposal Solids to Pugmill	Blowdown Water for Dust Cntrl	Disposal Solids to Landfill
Temperature, °F	80	60	60	100	100	100	100	60	100
Flow, GPM	74	--	--	--	--	--	--	23	--
Wt % Solids	20.9%	--	--	99.5%	99.5%	99.5%	99.5%	0.0%	80.0%
CaSO3*1/2H2O, K Lb/Hr	--	--	--	734.1	0.9	1.8	16.6	--	16.6
CaSO4*2H2O, K Lb/Hr	--	--	--	255.3	0.3	0.6	5.8	--	5.8
Flyash + Inerts, K Lb/Hr	0.7	--	--	806.3	1.0	2.0	18.3	--	18.3
Ca(OH)2, K Lb/Hr	8.6	--	--	124.7	0.2	0.3	2.8	--	2.8
CaCO3, K Lb/Hr	--	--	--	124.7	0.2	0.3	2.8	--	2.8
CaCl2, K Lb/Hr	--	--	--	18.4	0.0	0.05	0.4	--	0.4
H2O, K Lb/Hr (Steam)	35.3	--	--	10.4	0.0	0.03	0.2	11.4	11.7
Conveying Air, K Lb/Hr	--	6.0	3.0	--	--	--	--	--	--
Total Solids, K Lb/Hr	44.7	6.0	3.0	2074.0	2.6	5.1	47.0	11.4	58.4

NOTES : 1. K = Thousands (1000's) 2. GSA = Gas Suspension Absorber

The capital and operating costs for the ESP have not been included in the FGD system economic analysis. However, the cost for the mechanical cyclone collector installed at the absorber outlet is included. An existing cold-side ESP is assumed to operate downstream of the FGD system and that it is adequately sized to ensure that the preretrofit particulate emission rate is maintained. All of the fly ash and reaction product solids collected in the ESP are conveyed pneumatically to the solids disposal silo for storage, then transported by truck to the landfill disposal area.

Reagent feed system, area 10.

- Reagent storage and handling - The lime is delivered by rail car in the form of pebble lime. The pebble lime is discharged into two below-grade hoppers. The hoppers are pneumatically unloaded, and the contents stored in two bulk storage silos (total capacity of 60 days). The storage silos are fabricated of concrete and are water- and air-tight. The silos are equipped with baghouse dust collectors capable of handling all of the transport air flow from the pneumatic systems. The lime is pneumatically conveyed from the bulk storage silos to two day bins (total capacity of 30 hrs.). The bins are equipped with bag filters designed to handle the exhaust conveying air.
- Feed preparation and storage - From the day bins, the pebble lime is gravimetrically conveyed to one of three 50% capacity detention slakers. The pebble lime entering the slaker contains 10% inert material. Blowdown water is introduced which hydrates the pebble lime; the unreacted materials are removed from the bottom of the slaking tank by a traveling grate and discarded with the FGD solids materials. The slaked lime, Ca(OH)_2 , exiting the slaker flows to an agitated lime slurry feed tank (8-hr capacity) where it is diluted with blowdown water to 25 percent solids and allowed to stabilize. The slaked lime slurry is then pumped to the dual fluid nozzles at the base of each reactor.

SO₂ removal system, area 20.

- The SO₂ removal system for this evaluation includes three venturi-entry GSA reactors, each capable of handling 50% of the total flue gas flow. This allows two reactors to be on line at 100% boiler load while the third is on standby or undergoing maintenance. Each reactor is sized to have a gas retention time of 3.1 seconds.
- The slaked lime slurry is mixed with additional water and is then injected into the base of each operating reactor via an injection lance assembly. For this design, the reagent ratio is assumed to be 1.30 moles CaO required per mole of inlet SO₂ to achieve 90% SO₂ removal efficiency. A single dual fluid nozzle located in the center of each reactor disperses the slurry as a spray of fine droplets. The injection lance can be withdrawn and serviced during normal operation without loss in SO₂ absorption efficiency due to the large excess of reagent solids present in the fluid bed. This normal maintenance service is done once per week and takes 5-10 minutes.

Compressed air (at 100 psig) is used to shatter the slurry droplets exiting the nozzles, producing fine droplets which will evaporate quickly in the reactor. The compressed air is supplied by one of two centrifugal air compressors (one operating and one spare). The feed rate of lime slurry is automatically controlled by the outlet SO_2 concentration and as a function of percent SO_2 removal required. The flow of additional water to the nozzles is regulated to maintain the outlet flue gas temperature at approximately 18°F above the flue gas adiabatic saturation temperature. The humidification enhances SO_2 removal efficiency and conditions the flue gas for better ESP performance.

- Flue gas enters the bottom of the reactor at 277°F . The solid reactant particles flow upward, concurrently with the flue gas. In the reactor, the lime hydrate particles react with the SO_2 and SO_3 in the flue gas to form the product salts. The product salts, fly ash and unreacted lime discharge from the top of the reactor along with the flue gas.
- The solids are separated from the flue gas by two cyclone mechanical collectors located at the exit of each reactor. The cyclones remove approximately 99% of the solids from the gas. Approximately 99% of these solids are recirculated back to the reactor. The amount of solids returned to the reactor is controlled via a mechanical screw feeder located in the bottom of a feeder box. A second conveyor located at the top of the feeder box removes excess solids from the system (approximately 1%) which are then sent to the disposal silo.
- The recirculated solids enter the reactor just above the point of lime slurry injection. The particles are coated with lime and provide a large surface area for SO_2 absorption. In addition, unreacted lime is returned to the fluid bed from the solids collected by the cyclone for optimum utilization of the reagent.
- The gas leaving the cyclone continues to the ESP (or FF) where most of the remaining solids are removed. For a new plant, the selection of a new particulate removal device (baghouse or ESP) can be made independently of the required SO_2 removal efficiency since SO_2 removal in the particulate collection device is relatively low.

Flue gas system, area 30.

- The flue gas exits the air preheater at 277°F and flows to the GSA reactors. In the absorbers the gas is humidified and reacts with the reagent particles. The particle-laden flue gas exits the absorber at 145°F . The absorber causes an increase in pressure drop of about 5.6 inches of water. An additional pressure drop of 3.4 inches of water also occurs across the cyclone separators installed at the reactor outlet.
- A bypass system around both the GSA reactor and cyclone particulate collection system is included for plant startup or emergency shutdown. ID/booster fans located between the ESP and stack operate in parallel to maintain sufficient pressure to overcome the pressure loss from the FGD system and additional duct.

- The cost to install six inches of additional insulation have been included as ESP modifications to prevent the temperature of the gas in the ESP from dropping below the saturation temperature, which could result in condensation due to the increased water vapor downstream of the GSA reactors.

Solids handling system, area 60.

- Approximately 1% of the solids collected in the cyclones is combined with solids removed from the ESP hoppers and pneumatically conveyed to the disposal silos. It was assumed that the existing ash handling system would be demolished due to insufficient capacity and/or wet sluicing operation. A new positive pressure pneumatic conveying system is used to transfer the solids from the particulate collector to the storage silos. A new silo was included in the estimate to handle the incremental solids capacity. The product solids are mixed with water in two 67% pugmills for dust control (to about 20% moisture) and loaded into off-highway trucks by either a shuttle loader (belt conveyor) or a front-end loader. The trucks transport the solids to a landfill disposal area located one mile from the plant site. A bulldozer at the disposal area spreads and compacts the material to an average depth of 30 feet. The loading, transfer, and landfill area equipment operate one shift/day, 5 days/week. Disposal rates are presented in Table 20-3.

General support equipment, area 70.

- The AirPol GSA system requires the following support equipment: instrument air compressor, makeup water system, and control room.

Additional equipment, area 80.

- Additional equipment - Onsite electric power equipment including transformers and grounding that is required to supply electricity to the FGD system.

TECHNICAL EVALUATION

The AirPol GSA process is a relatively new SO₂ removal technology for a utility application. However, the successful industrial operation of the GSA in Denmark has demonstrated its potential to become a commercially viable process. As part of the Department of Energy (DOE) Clean Coal Technology Program, the AirPol GSA system has been installed as a 10-MW demonstration unit at the Tennessee Valley Authority's (TVA) Shawnee Power Station.

With AirPol GSA, considerable improvement in the SO₂ removal efficiency is obtained compared to most other dry processes. The major differences between the GSA and a typical Circulating Fluid Bed (CFB) design are as follows:

Table 20-3
**AirPol GSA PROCESS
DISPOSAL RATES AND COMPOSITION
FOR ONE 300 MW UNIT
(Operating at 100 Percent Load)**

<u>Component</u>	<u>tph</u>
CaCO ₃	1.4
Ca(OH) ₂	1.4
CaCl ₂	0.2
Fly Ash and Inerts	9.2
CaSO ₃ · 1/2H ₂ O	8.3
CaSO ₄ · 2H ₂ O	<u>2.9</u>
Total Dry Solids to Disposal	23.4
Daily Maximum Volume*	14,020 ft ³
Annual Maximum Volume (100% load)	118 Acre-ft
Plant Life Volume (15 yrs)**	1,150 Acre-ft

*Based on a solids density of 80 lb/ft³ compacted (20 wt. percent moisture).
**Calculated with a 0.65 load factor over the plant life.

- 1) The GSA introduces the wet lime slurry and recycled particles so that the surface of the particles becomes coated with a thin film of slurry that then absorbs the gaseous contaminants and dries quickly and thoroughly. The CFB design introduces the reagent as a dry material and maintains a circulating fluid bed for absorption of contaminants.
- 2) The GSA recirculates material from a specially designed cyclone(s) with feeder box(s). The CFB design recirculates material from a precipitator.
- 3) GSA can be followed by either a fabric filter or precipitator.

Results to date show removal efficiencies approaching those of wet scrubbing processes.

The advantages and disadvantages of the AirPol GSA process as compared to the Limestone Forced Oxidation process are summarized in Table 20-4. The major advantages claimed are the following:

- Maintenance costs are significantly lower due to reduced scaling and corrosion when compared to wet FGD systems. The GSA systems require only a small stream of scrubbing slurry to be pumped into the absorber as compared to the large volume of scrubbing slurry recycled in wet systems. This small, alkaline stream contacts the gas entering the absorber rather than the walls of the system. In wet systems, the walls of the absorbers, tanks, and pipes are subject to corrosion because of the continuous contact with low pH slurries. The high pH slurry and dry solids product inherent in the GSA allow the use of mild steel materials of construction for the GSA vessel and slurry tanks. In contrast, wet FGD systems frequently require rubber liners or alloy liners at various locations in the system.
- The smaller volume of liquid scrubbing medium required in dry systems results in reduced pumping requirements, significantly lowering power consumption.
- Power requirements are significantly reduced for the AirPol system over a wet FGD system due to the elimination of large recycle pumps and the entire dewatering system.
- Dry solids are produced that can be handled by conventional fly ash systems, resulting in the elimination of the dewatering equipment and a reduction in the associated operating and maintenance requirements.
- Reheat is not required in the AirPol system because the gas is not saturated when it exits the absorber. This reduces capital cost and eliminates reheat steam consumption.

Table 20-4

TECHNICAL EVALUATION
ADVANTAGES/DISADVANTAGES
OF THE AirPol GSA PROCESS
(Compared to the Limestone Forced Oxidation Process)

<u>Item</u>	Dis- advantage	Same	Advantage
1. Process			
a. Complexity of operation			X
b. Oxidation of sulfite to sulfate		N/A	
c. Load following capability			X
d. Adaptability to flue gas temperature changes		X	
e. Capability of using cooling tower blowdown process water		X	
f. Stability of process		X	
g. Extreme vessel pressure		X	
h. High equipment operating temperature		X	
i. Use of liquid fuel or natural gas		X	
j. Material handling characteristics		X	
k. Reagent availability		X	
l. Space requirements for FGD equipment			X
2. Operation and Maintenance Requirements			
a. Labor requirements			X
b. Solids deposition		X	
c. Equipment corrosion, scaling			X
d. Equipment erosion		X	
e. Reagent feed rate			
3. Effect on Net Plant Heat Rate			
a. Power consumption			X
b. Steam usage			X
4. Disposal			
a. Land requirements	X		
b. Reactivity of solids	X		
c. Solids handling characteristics			X
5. Use of Exotic Materials of Construction			X
6. Operational Hazard			
a. High temperature		X	
b. High pressure		X	
c. Use of hazardous chemicals		X	

Note: N/A = Not Applicable

- High chloride concentrations in the slurry will typically reduce the SO₂ removal efficiency in wet systems. In the GSA process, chloride has been found to enhance SO₂ removal.
- Cooling tower blowdown can be used for all slurry dilutions after completing the slaking of the lime reagent, with virtually no adverse effects on system performance. In some wet scrubber applications, cooling tower blowdown makeup water can have adverse effects on system SO₂ removal performance, particularly those with high-chloride content.
- The AirPol system is potentially well suited for retrofit installations due to its reduced space requirements and ease of installation. The GSA reactor is smaller in diameter than conventional absorbers due to the increased flue gas velocity through the reactor.

AirPol GSA System disadvantages compared to the Forced Oxidation FGD System are as follows:

- The potential exists for adverse impact on ESP performance due to the increased particulate loading and change in ash resistivity. Additional specific collecting plate area (SCA) may be needed to maintain particulate emission levels at or below required limits.
- The AirPol process requires a higher reagent feed ratio (to achieve the desired removal efficiency) compared to a conventional wet alkali system. In addition, lime is more expensive than limestone. However, relatively low levels of chloride in the coal (0.1%) can reduce reagent consumption. Calcium chloride spiking, can also significantly reduce reagent consumption. The savings in reagent consumption is partially offset by the costs of installing a calcium chloride injection system as well as the cost of the calcium chloride itself. Lowering gas outlet temperature can also reduce reagent consumption, but this presents greater potential for duct deposition and plugging.
- The potential for duct deposition and plugging problems may exist due to flue gas and solids humidification. Pluggage of the AirPol absorbers could cause negative pressure transients due to the downstream location of the ID fans. However, GSA systems operating since 1988 have experienced no plugging problems. Additionally, during the 13 month testing period at TVA's Center for Emissions Research, not a single plugging problem related to high-moisture levels in the solids has been experienced, even at approach-to-saturation temperatures as low as 8°F in the reactor/cyclone.
- No full-scale operating system exists in the United States.
- The AirPol process may not be able to use cooling tower blowdown water in the lime slaking process, but instead may require fresh water for this step. High chloride levels in the cooling tower water can adversely effect the slaking process. However, cooling tower blowdown can be used for slurry dilutions after completing the slaking of the reagent. For high-sulfur applications the fresh water used for lime slaking can represent as much as half of the system water makeup requirement. In contrast, wet

limestone scrubbers (not producing a marketable gypsum byproduct) can generally use cooling tower water for limestone grinding circuits, and for most other makeup water applications, although, water with high chloride content can also have adverse effect on wet scrubber performance.

- A higher solids recycle rate is required relative to other dry technologies.
- The largest GSA reactor installed to date treats 114,000 ACFM of flue gas, equivalent to approximately 35 MW.

COMMERCIAL STATUS

The GSA technology was developed by FLS Miljo A/S of Denmark for removing acid gases from the flue gas generated by many industrial processes. The first commercial GSA system was installed on a municipal solid waste incinerator in Denmark in 1988. This technology is currently being used at several waste incinerator applications in Europe to remove both HCl and SO₂ from flue gas. The largest unit sold to-date is a recent order at an iron pelletizing installation at LKAB¹ in Sweden. The GSAs for one pelletizing line will handle a total of 228,000 SCFM. Two reactors will be installed per line, each reactor more than 12 feet in diameter. An installation list is shown in Table 20-5.

As part of the DOE Clean Coal III Technology program, AirPol, Inc., a U.S. subsidiary of FLS Miljo A/S, has built a GSA demonstration plant at TVA's Center for Emissions Research (CER). The demonstration is being conducted on a 10 MWe slip stream from a 150 MWe coal-fired boiler at the Shawnee plant near Paducah, Kentucky. The unit burns a high-sulfur (2.7 percent) eastern bituminous coal. Testing began in November 1992 and ended in late November 1993. The results of the testing includes the following:

¹LKAB is the Swedish iron ore processing company: Luossavaara Kirunavaare AB. Luossavaara Kirunavaare in northern Sweden is the world's largest underground iron ore mine.

Table 20-5
INSTALLATION LIST FOR AirPol GSA PROCESS

<u>UNIT AND LOCATION</u>	<u>UNIT SIZE</u>	<u>FUEL TYPE</u>	<u>FILTER TYPE</u>
Kara #4 Roskilde, Denmark	177 TPD	Waste Incinerator	ESP
Slagelse #1 Slagelse, Denmark	144 TPD	Waste Incinerator	Baghouse
Seas Stignaes, Denmark	15 MW (Pilot Plant)	Coal-Fired Boiler	ESP
Reno Nord #1 & 2 Aalborg, Denmark	212 TPD	Waste Incinerator	ESP
Reno Nord #3 Aalborg, Denmark	320 TPD	Waste Incinerator	Baghouse
Aarhus Nord #3 Aarhus, Denmark	212 TPD	Waste Incinerator	Baghouse
Slagelse #2 Slagelse, Denmark	106 TPD	Waste Incinerator	Baghouse
Kara #1 Roskilde, Denmark	200 TPD	Waste Incinerator	ESP
Kara #3 Roskilde, Denmark	160 TPD	Waste Incinerator	Baghouse plus Wet Scrubber
Shawnee Unit #9 DOE/TVA Paducah, Kentucky	10 MW Slipstream	Coal-Fired Boiler	ESP
San Ying Enterprises, CO Taipei, Taiwan, R.O.C.	100 TPD	Waste Incinerator	Baghouse plus Wet Scrubber
LKAB Kiruna, Sweden	(114 K ACFM per GSA)	Iron Ore Sintering	ESP
Hamilton Hamilton, OH	50 MW	Coal-Fired Boiler	Baghouse

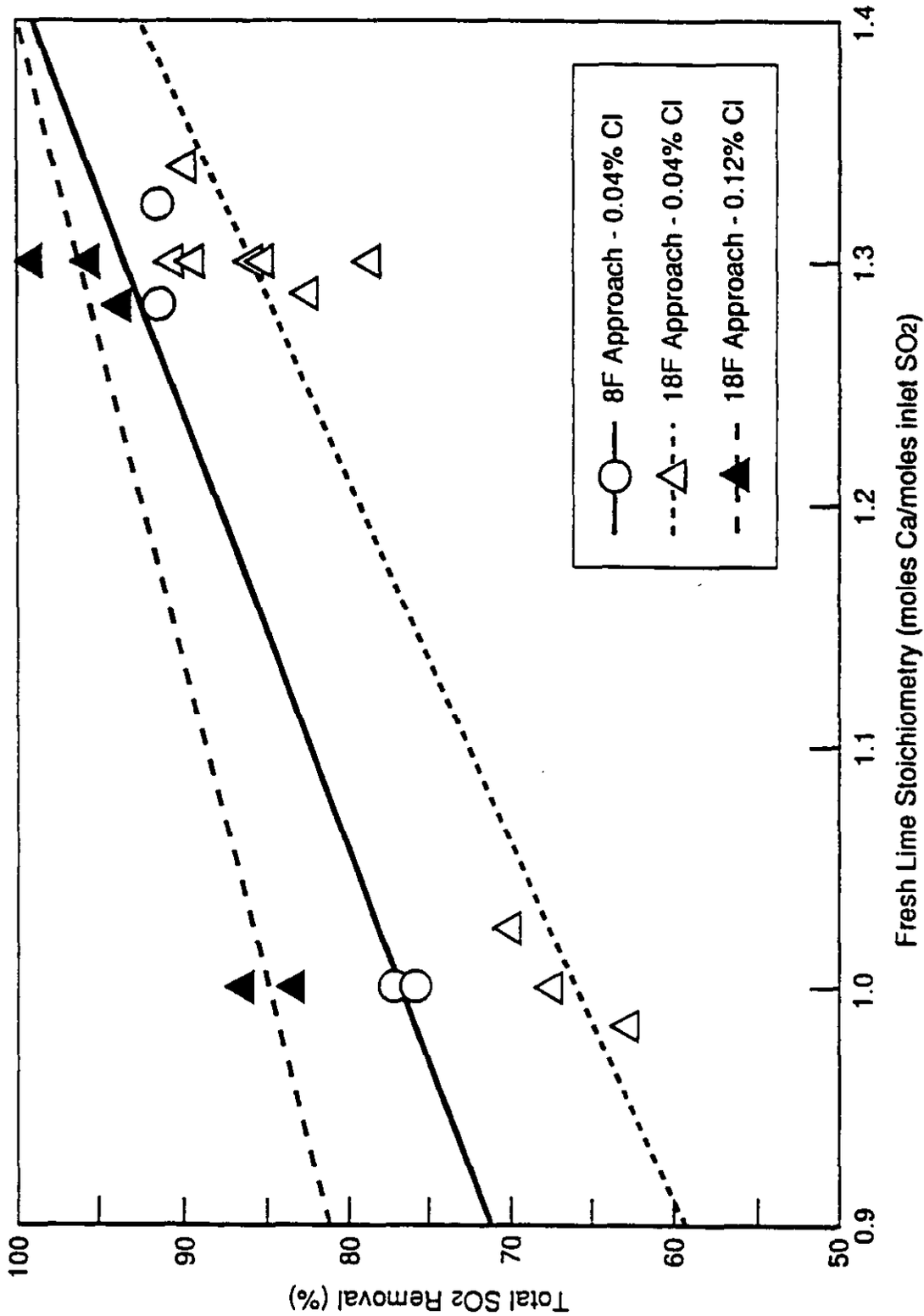
- Efficiencies approaching 100% at reagent ratio of 1.4 and 5°F approach to saturation temperature. (Additional test results are shown in Figure 20-2.)
- Able to operate at 8°F approach to saturation without build-up.
- No build-up encountered during operation.
- Improved particulate collection efficiency by the installed precipitator compared to other competitive systems.
- High GSA availability.

The project was co-funded by TVA, DOE, and AirPol. This testing marked the first application of this technology in the U.S.

The AirPol GSA system will also be installed on the 50 MW boiler at the Hamilton station, owned by the City of Hamilton, OH. The GSA process was chosen to allow the Hamilton boiler to burn high sulfur Ohio-mined coals while meeting state and federal pollution control regulations. The projected start-up date for this facility is during the third quarter of 1995.

ECONOMIC EVALUATION

For this evaluation, the AirPol GSA FGD process is a dry throwaway system retrofit to a pulverized coal-fired boiler. This process is compared in this section to the commercially developed Limestone Forced Oxidation (LSFO) FGD process retrofit to a similar coal-fired boiler. The base case economic evaluation of the AirPol process is presented in Table 20-7, 20-8, and 20-9. As stated in the General Design Criteria, the gross plant output is 313 MW, resulting in a nominal 300 MW unit after subtracting out plant auxiliary power, exclusive of FGD energy consumption. Raw material and utility consumptions were calculated for the AirPol process operating at 100 percent load and are presented in Table 20-6. The FGD system power consumption is addressed only as an operating cost with no reduction in net plant output. Cost sensitivities to various parameters are also presented. These curves were developed by adjusting capital and operating costs with changes in a single input, and re-evaluating the capital and levelized costs. The sensitivity curves are plots of the results for four additional parameter values, combined with the base case value.



Notes: All tests conducted at a 320°F inlet flue gas temperature

Figure 20-2 Overall System SO₂ Removal Results from the AirPol GSA Demonstration

Source: "10-MW Demonstration of the AirPol Gas Suspension Absorption Flue Gas Desulfurization Process," Presented at the 1993 SO₂ Control Symposium, Boston, MA, August 1993.

Table 20-6

**AirPol GSA SYSTEM
RAW MATERIAL AND UTILITY CONSUMPTION
FOR ONE 300 MW UNIT
(Operating at 100 Percent Load)**

<u>Item</u>	<u>Quantity</u>	
Pebble Lime @ 1.30 Stoichiometric Feed Ratio	7.3 tph	
Cooling Tower Blowdown Water	170 gpm	
<u>Power (Operating Hp and Equivalent kW Consumed)</u>	<u>Hp</u>	<u>kW</u>
Area 10 - Reagent Feed System	230 Hp	(170 kW)
Area 20 - SO ₂ Removal System	330 Hp	(250 kW)
Area 30 - Flue Gas System	1,760 Hp	(1,310 kW)
Area 60 - Solids Handling System	230 Hp	(170 kW)
Area 70 - General Support Equipment	20 Hp	(10 kW)
Total	2,570 Hp	(1,910 kW)

Table 20-7

**AirPol GSA SYSTEM
TOTAL CAPITAL REQUIREMENT
(1990 \$, 300 MW, 2.6% S Coal)**

<u>Area</u>	<u>Description</u>	<u>\$/kW</u>
10	Reagent Feed System	25.1
20	SO ₂ Removal System	38.2
30	Flue Gas Handling System	18.0
60	Solids Handling System	4.6
70	General Support Equipment	1.2
80	Additional Equipment	<u>4.1</u>
	Total Process Capital	91.2
	General Facilities	9.1
	Engineering and Home Office Fees	9.1
	Project Contingency	19.3
	Process Contingency	<u>8.3</u>
	Total Plant Cost (TPC)	<u>137.0</u>

	Total Cash Expended (TCE)	133.7
	Allowance for Funds During Construction (AFDC)	<u>7.5</u>
	Total Plant Investment (TPI)	141.2
	Royalty Allowance	0.5
	Preproduction Costs	5.2
	Inventory Capital	1.9
	Initial Catalyst and Chemicals	<u>0.0</u>
	Total Capital Requirements (TCR - Moderate Retrofit)	<u>148.8</u>

Note: 2 + 1 spare modules (50% capacity each)
Accuracy = ± 20 percent

Table 20-7 shows the capital costs for the AirPol system. The overall project contingency for the AirPol process was calculated as 14 percent, while the overall process contingency was calculated as approximately 6.1 percent of the total plant cost. These values are higher than LSFO due to the lack of large scale U.S. installations. The weighted average maintenance factor (annual cost of maintenance material and labor) was estimated to be 4.1 percent of the total plant cost in this evaluation. The total capital requirement represents the differential cost to the utility for the FGD system, resulting in an estimated capital cost of 149 \$/kW for a moderately difficult retrofit situation.

The fixed and variable operating costs are shown in Table 20-8. First-year costs and levelized costs in terms of both constant dollars and current dollars are presented. Constant dollar analysis assumes no inflation and 0.3 percent real escalation on power. Current dollar analysis assumes a 5 percent annual inflation rate and 0.3 percent real escalation on power.

The levelized cost for the AirPol process (Table 20-9) is calculated by the "present worth" method (as defined in EPRI Economic Premises, Section 4), assuming a 15-year plant operating life. The FGD system levelized cost is 7.6 mills/kwh (constant dollars). This levelized cost is the sum of the fixed and variable operating costs (Table 20-8), plus the cost of capital over the 15-year plant life.

When compared with the limestone forced oxidation process, the AirPol process operating cost differs most in the reagent cost (see Table 20-8). The reagent cost is 2.8 times that of limestone forced oxidation due to the higher lime cost and reagent ratio. However, the higher reagent costs are offset by lower power costs and lack of steam requirement for reheat. Overall, the levelized cost is 20 percent lower than the limestone forced oxidation process.

A comparison of the total process capital for this system with that of Limestone Forced Oxidation also indicates the major capital investment requirements for both processes are for the SO₂ absorption equipment. Comparison of the total capital requirement for both processes indicates that the AirPol process will require approximately 31 percent less initial investment than a limestone forced oxidation system retrofitted on a 2.6% sulfur coal-fired boiler.

Table 20-8
AirPol GSA SYSTEM
OPERATING COSTS
(300 MW, 2.6% S Coal)

	1st Year mills/kWh*	Constant Dollars** 15-year Levelized mills/kWh	Current Dollars*** 15-year Levelized mills/kWh
<u>Fixed Operating Costs</u>			
Operating Labor	0.39	0.39	0.53
Maintenance Labor and Material	1.00	1.00	1.37
Administration and Support Labor	<u>0.23</u>	<u>0.23</u>	<u>0.32</u>
Total Fixed Operating Costs	1.62	1.62	2.22
<u>Variable Operating Costs</u>			
Pebble Lime	1.33	1.33	1.82
Solids Disposal	0.62	0.62	0.85
Additional Power	<u>0.32</u>	<u>0.33</u>	<u>0.45</u>
Total Variable Operating Costs	2.27	2.28	3.12

* January 1990 dollars.

** No inflation, 0.3% escalation for power.

*** 5% annual inflation, 0.3% escalation for power.

Note: Accuracy = ± 20 percent

Table 20-9

**AirPol GSA SYSTEM
OPERATING COST (1990-2005)
(300 MW, 2.6% S Coal)**

	<u>Fixed O&M</u>	<u>Variable Operation</u>	<u>Fixed Charge</u>	<u>Total</u>
First year: 1990 \$	2,762,100	3,882,100	10,938,900	17,583,100
\$/kW-yr	9.2	12.9	36.5	58.6
mills/kWh	1.6	2.3	6.4	10.3
\$/ton SO ₂	94	132	372	598
Levelized Costs - Constant Dollars:				
\$/kW-yr	9.2	13.0	20.8	43.0
mills/kWh	1.6	2.3	3.7	7.6
\$/ton SO ₂	94	132	213	439
Levelized Cost - Current Dollars (Inflation = 5%):				
\$/kW-yr	12.6	17.8	28.6	59.0
mills/kWh	2.3	3.1	5.0	10.4
\$/ton SO ₂	129	182	291	602

Note: Accuracy = ± 20 percent

Coal sensitivity of the AirPol GSA process to various parameters are shown in Figures 20-2 through 20-21. Parameters analyzed include the following:

- Coal sulfur content
- Lime stoichiometric feed rate
- Flue gas flow rate, acfm
- Solids disposal cost
- Lime cost
- Power cost
- Maintenance factor variations
- Inflation rate
- Unit size
- SO₂ removal efficiency
- Retrofit difficulty
- Plant life

The levelized costs for the AirPol process are very sensitive to coal sulfur content as shown in Figure 20-2. Over the 0.5 to 6 percent sulfur range investigated, the levelized cost (mills/kWh) increases approximately 13 percent with a 1 percent increase in coal sulfur content. This high sensitivity can be attributed to the increased cost of capital investment in the reagent handling areas and corresponding increase in fixed costs and variable operating costs including lime, waste disposal, and power.

Reagent feed rate has a significant effect on operating costs. A 25 percent increase in stoichiometry corresponds to an 8.4 percent increase in levelized cost. Reagent cost also is important, with a 25 percent change from the base case cost resulting in a 5 percent difference in levelized cost. Maintenance cost differences affect the levelized cost by 6 percent for a 50 percent deviation from the base case value.

Other parameters analyzed for this process include flue gas flow rate, solids disposal cost, power cost, and inflation rate. None of these variables were found to significantly affect the overall levelized cost.

The capital cost for AirPol GSA system with no spare capacity is 126 \$/kW for a moderately difficult retrofit situation. This represents a 15 percent savings compared to the 50 percent spare absorber philosophy adopted in this evaluation. The 15-year FGD system levelized cost with no spare capacity is 6.8 mills/kWh (constant dollars).

In conclusion, the economic evaluation of the AirPol GSA process indicates that the lower capital requirement of the GSA absorber and lower labor and maintenance costs results in a 20% lower overall levelized cost compared to an LSFO system. From the sensitivity analyses, the levelized cost for the AirPol process is a major function of the coal sulfur content, reagent cost/feed rate, and maintenance cost and is less significantly affected by flue gas flow rate, solids disposal cost, power cost, and inflation rate.

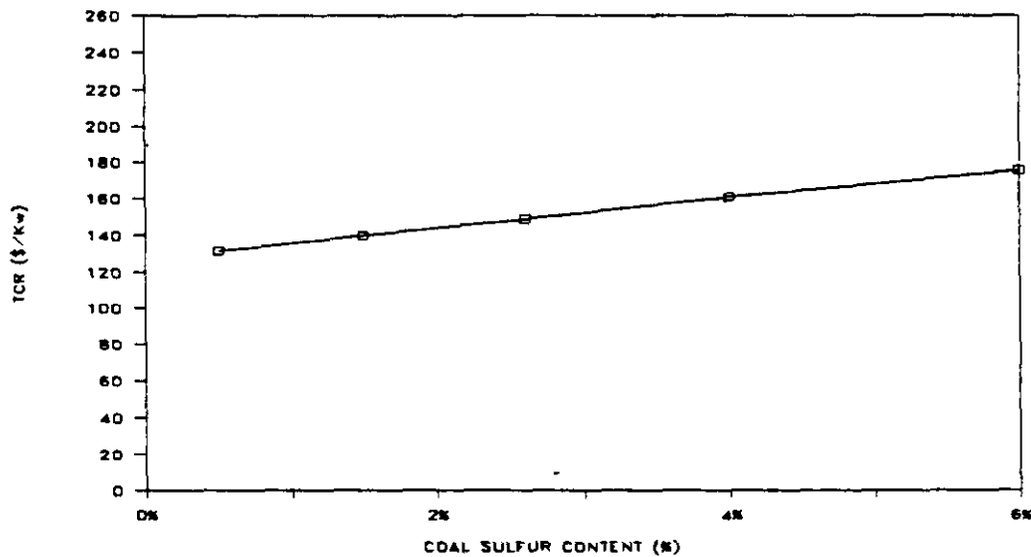
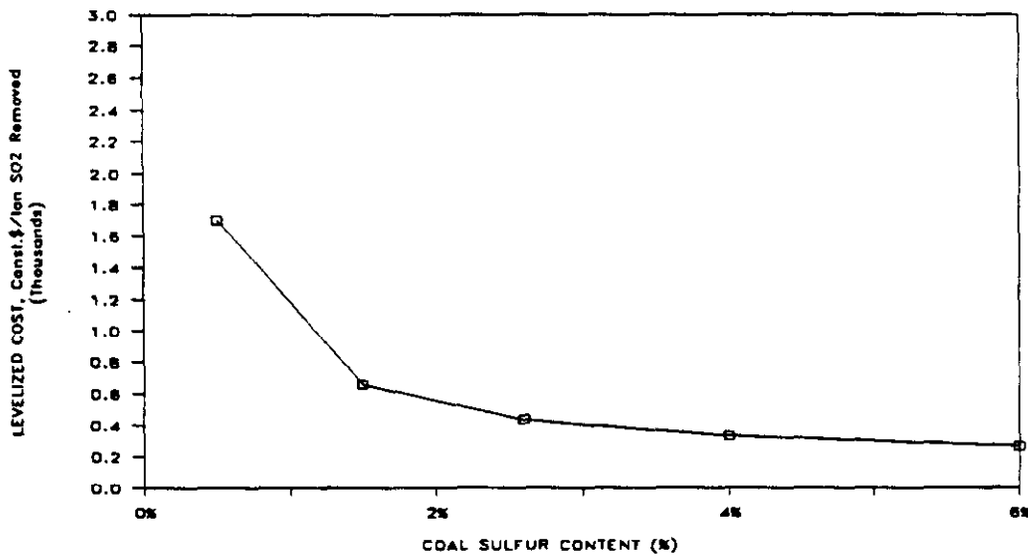
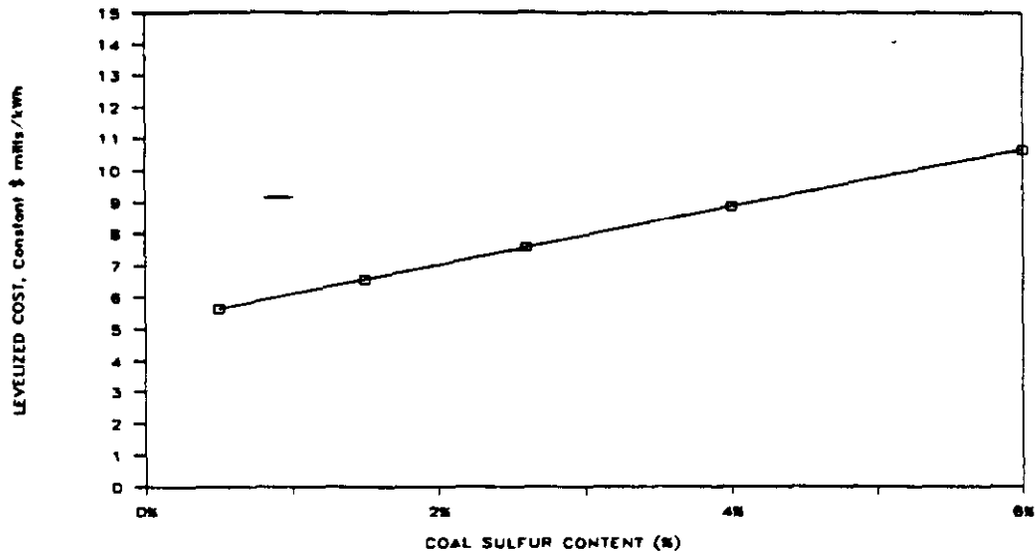


Figure 20-3. AirPol GSA Sensitivity Curves

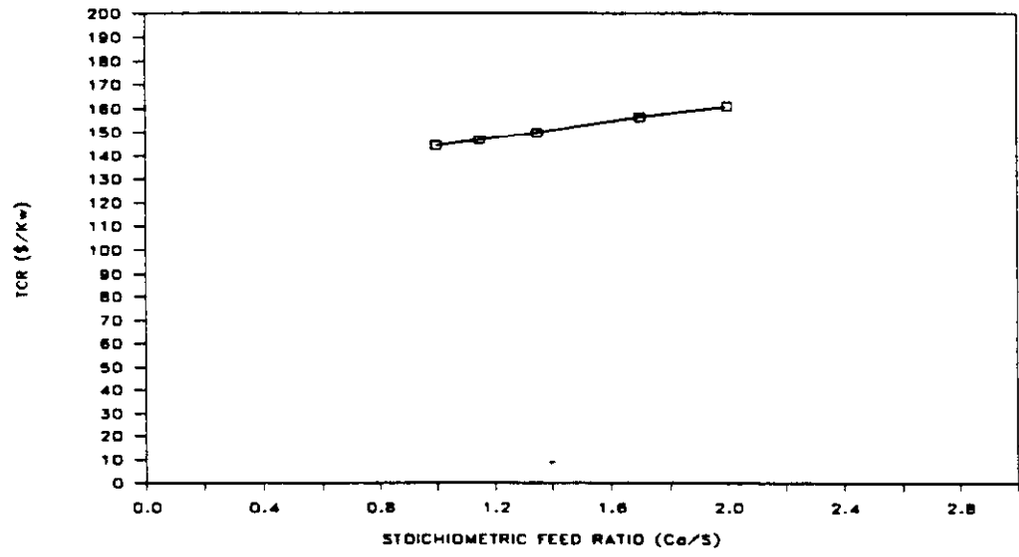
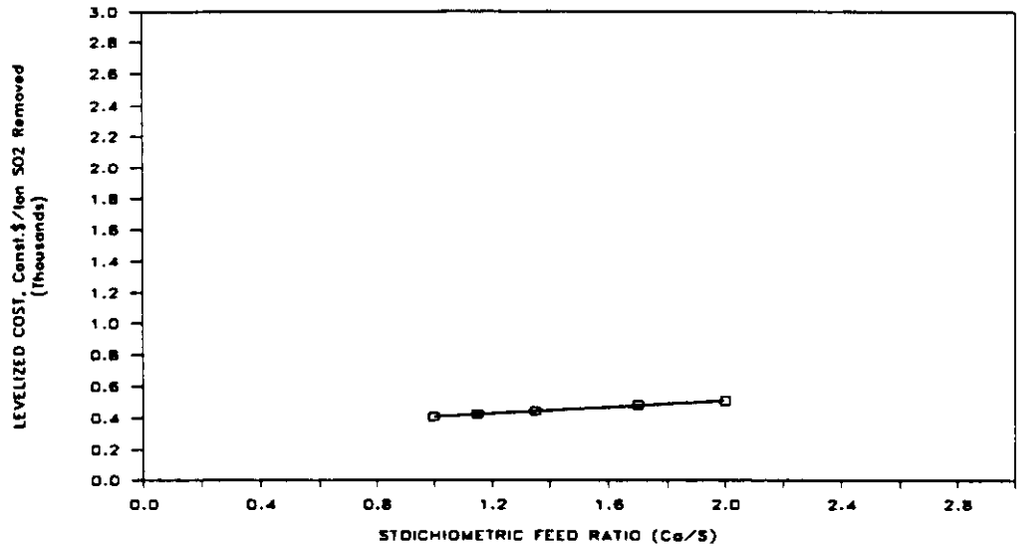
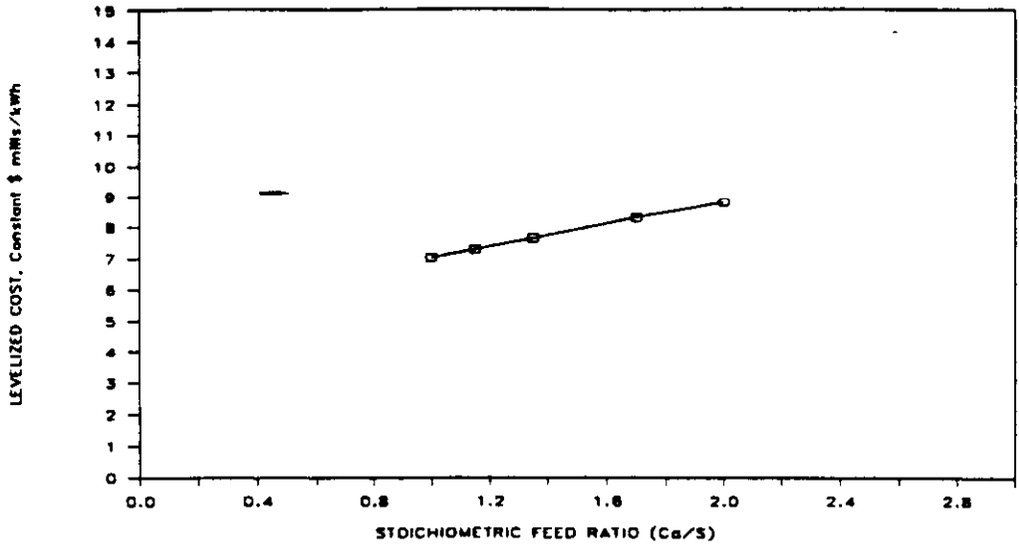


Figure 20-4. AirPol GSA Sensitivity Curves

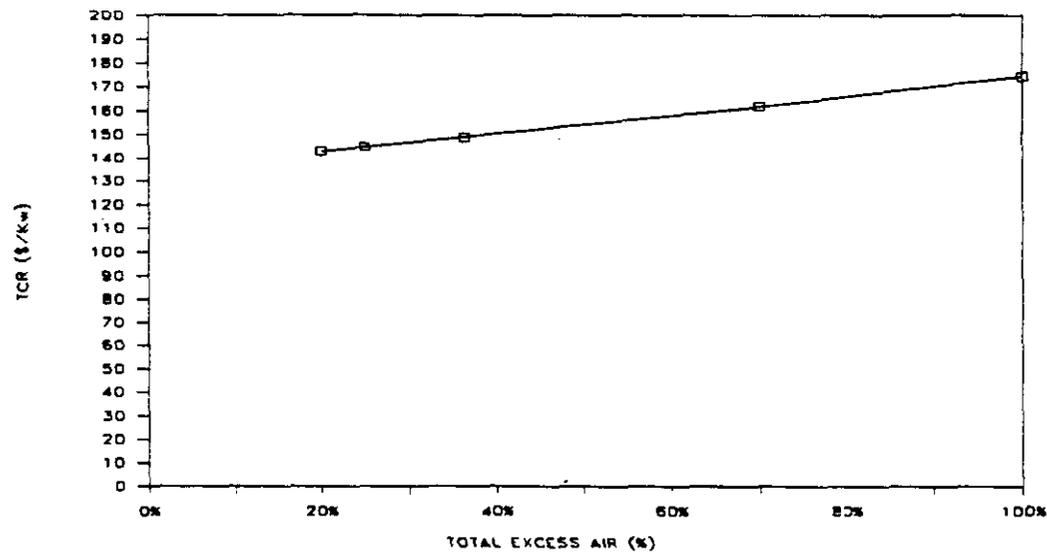
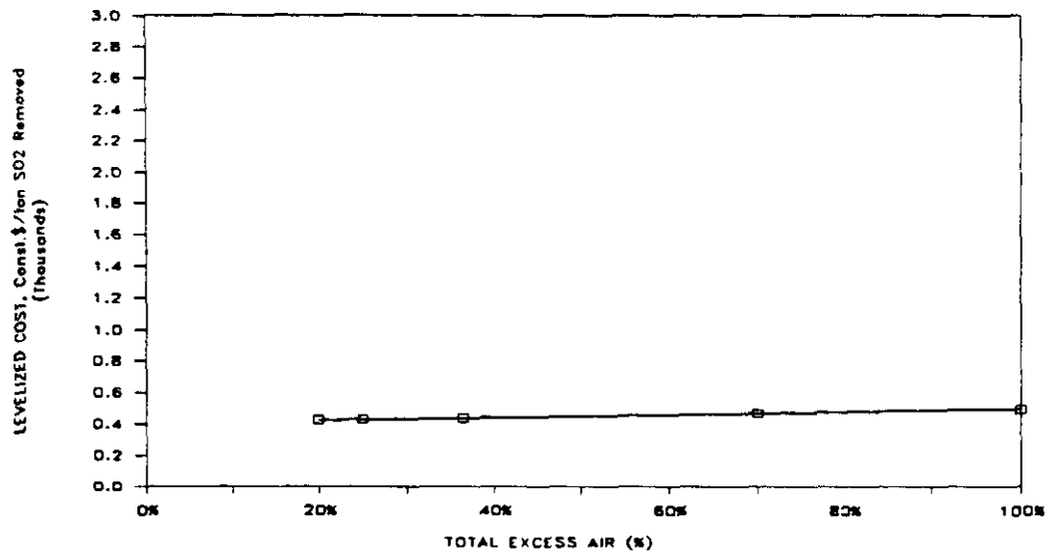
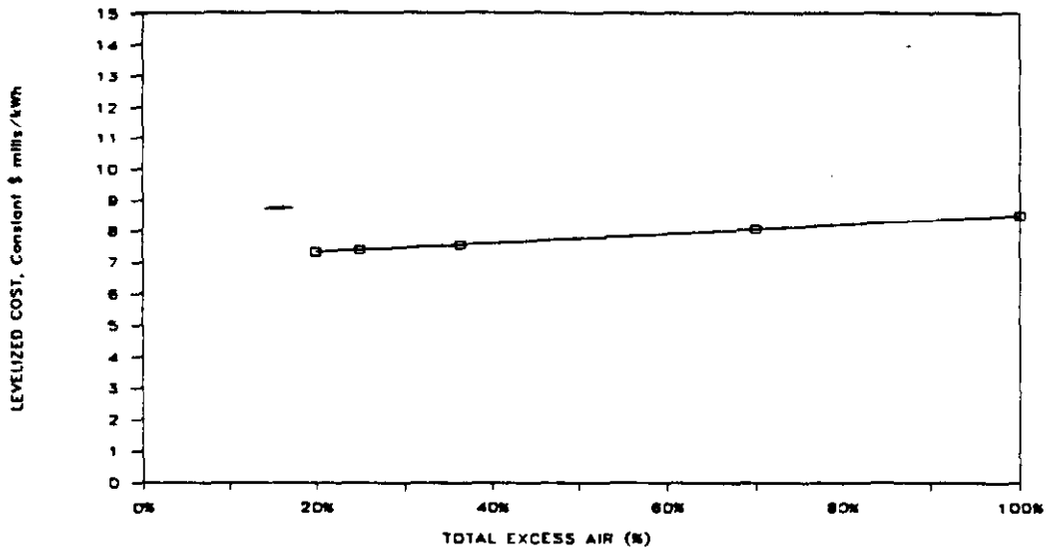


Figure 20-5. Airpol GSA Sensitivity Curves

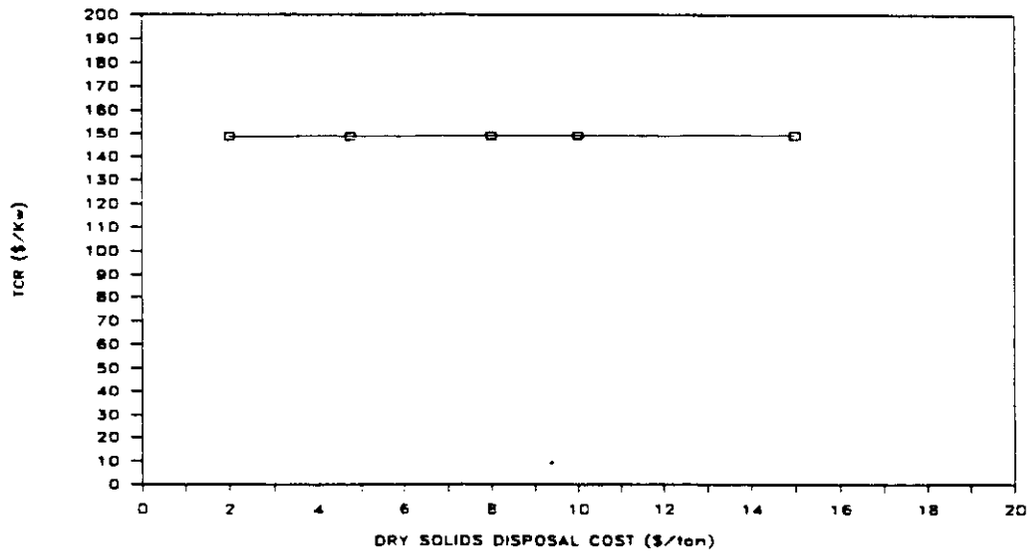
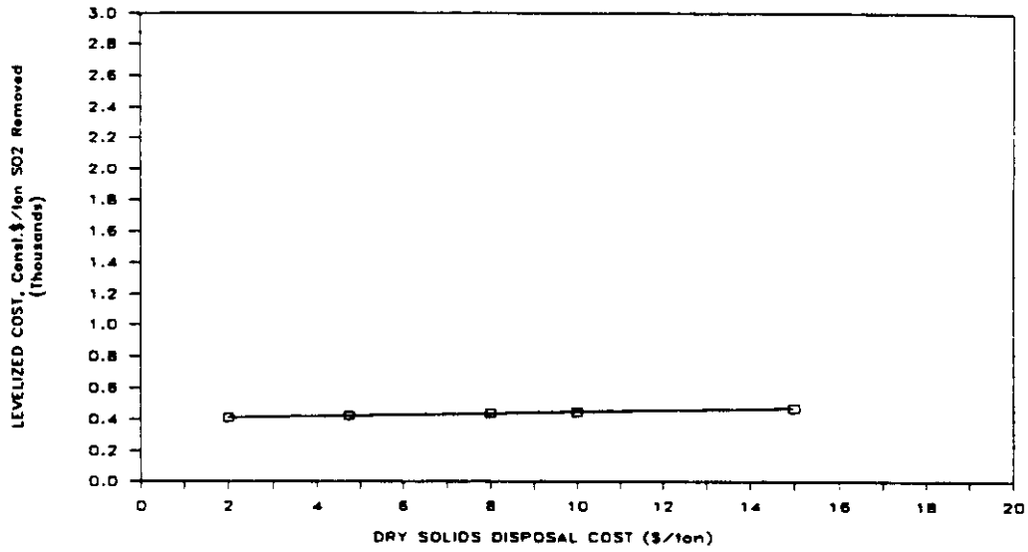
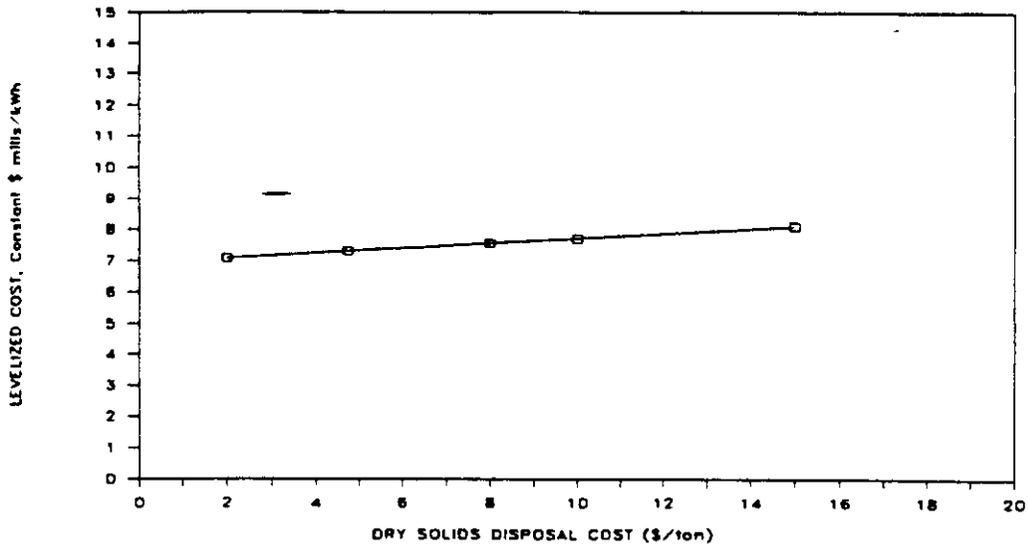


Figure 20-6. AirPol GSA Sensitivity Curves

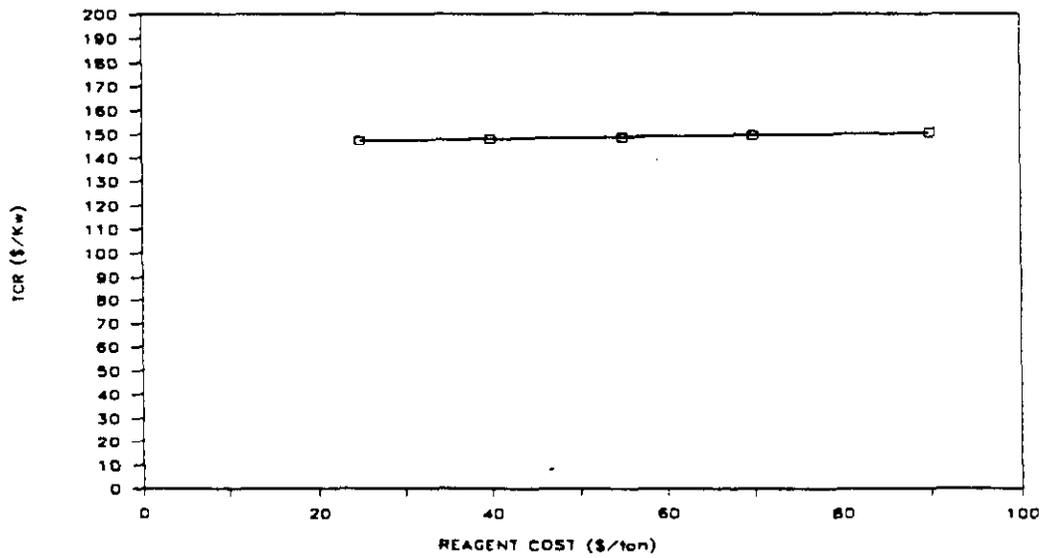
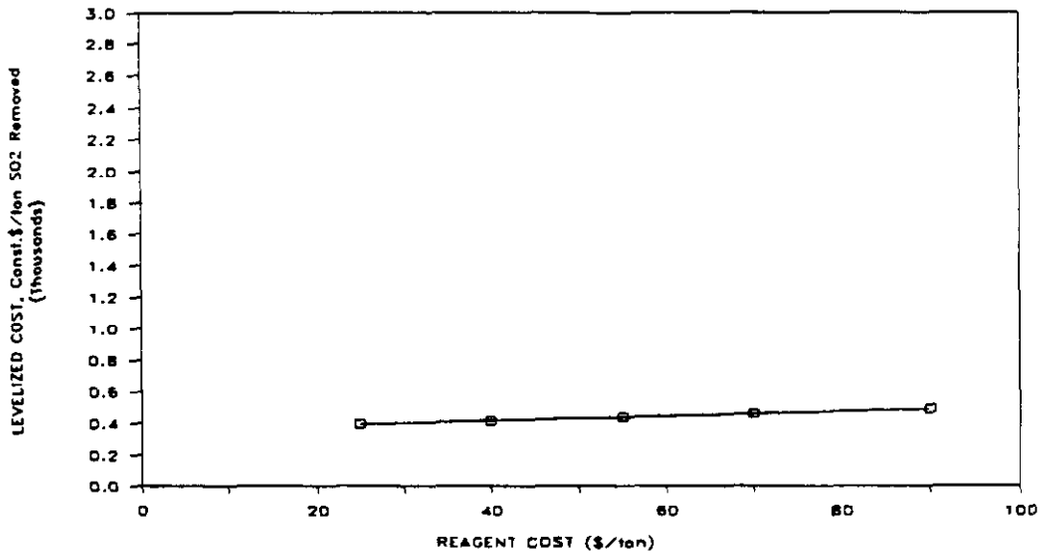
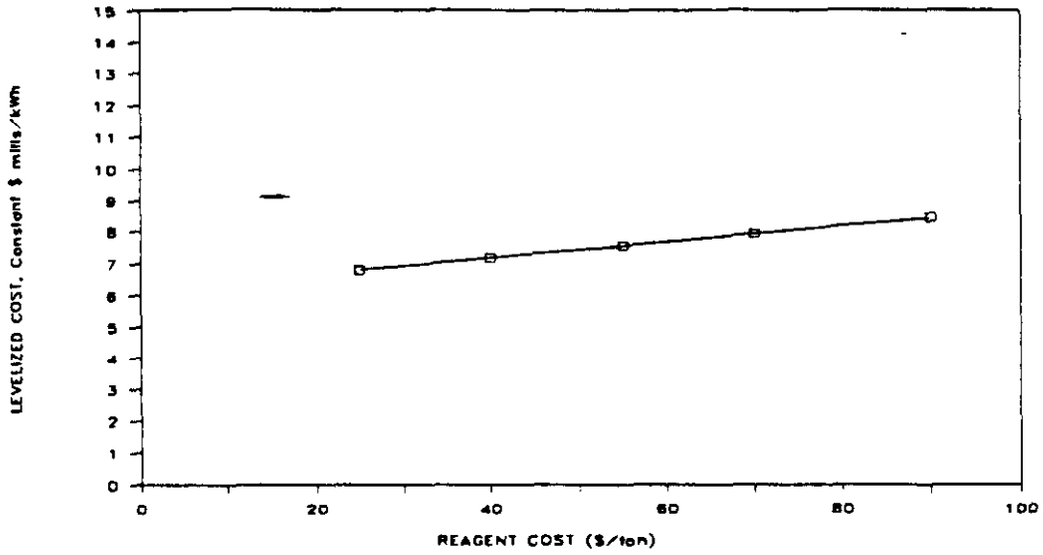


Figure 20-7. AirPol GSA Sensitivity Curves

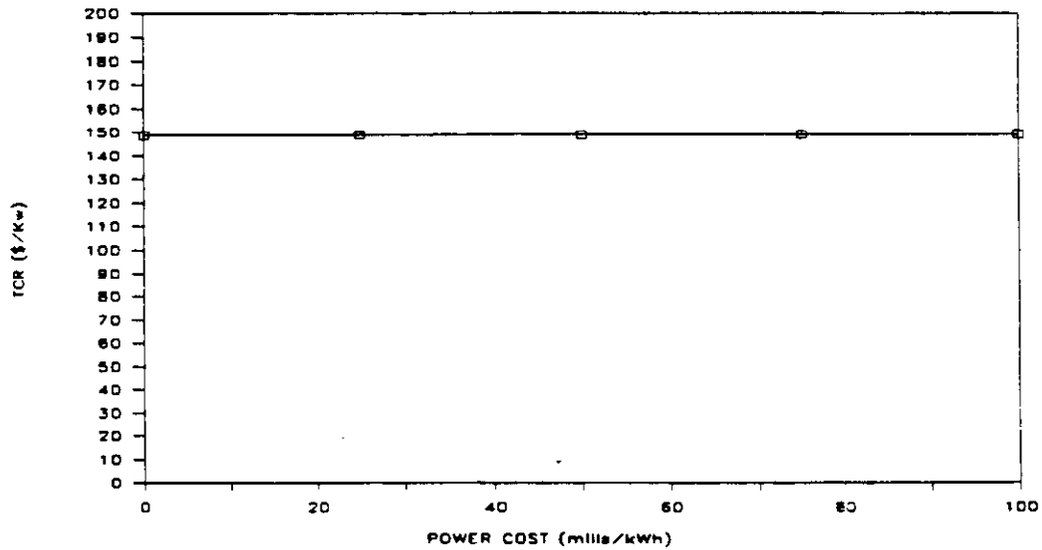
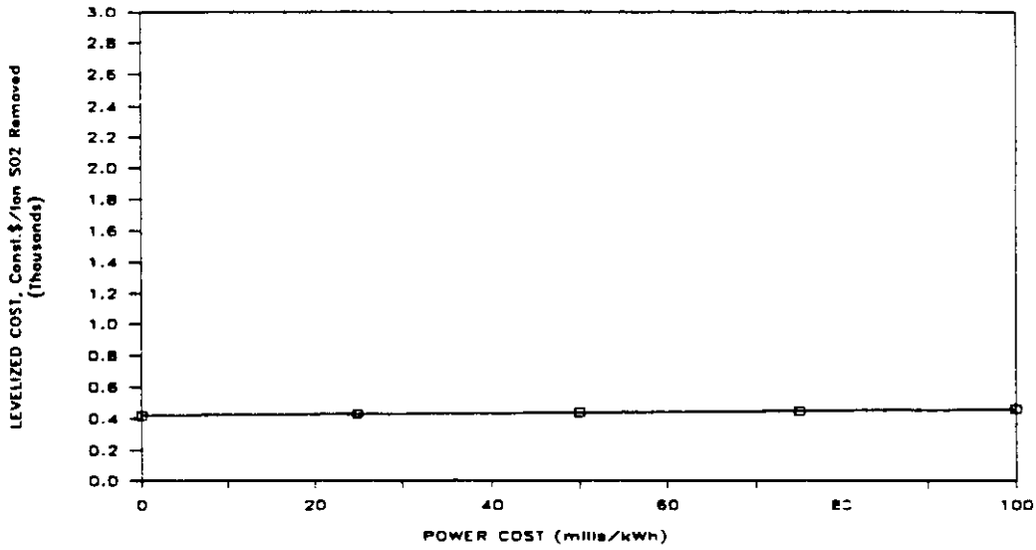
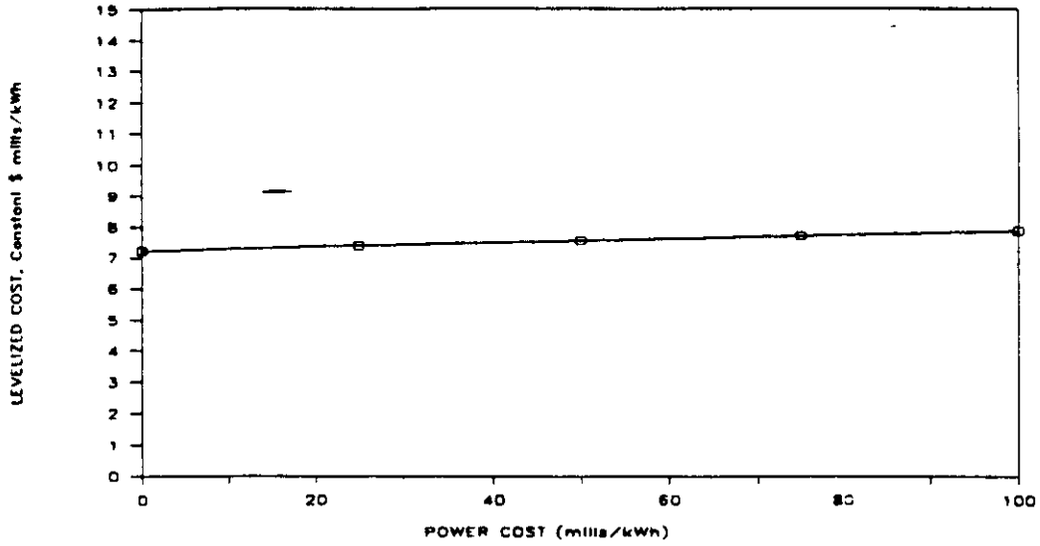


Figure 20-8. AirPol GSA Sensitivity Curves

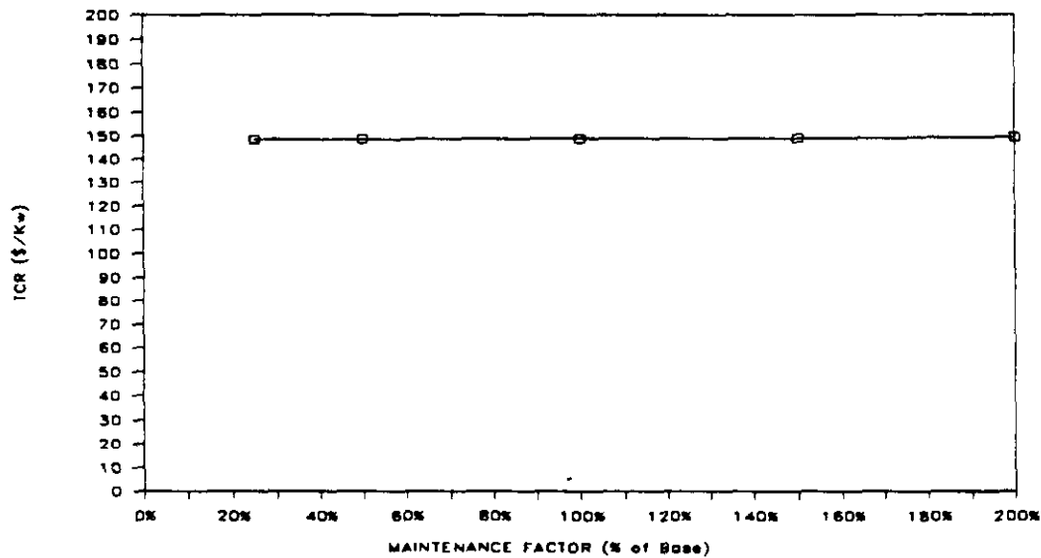
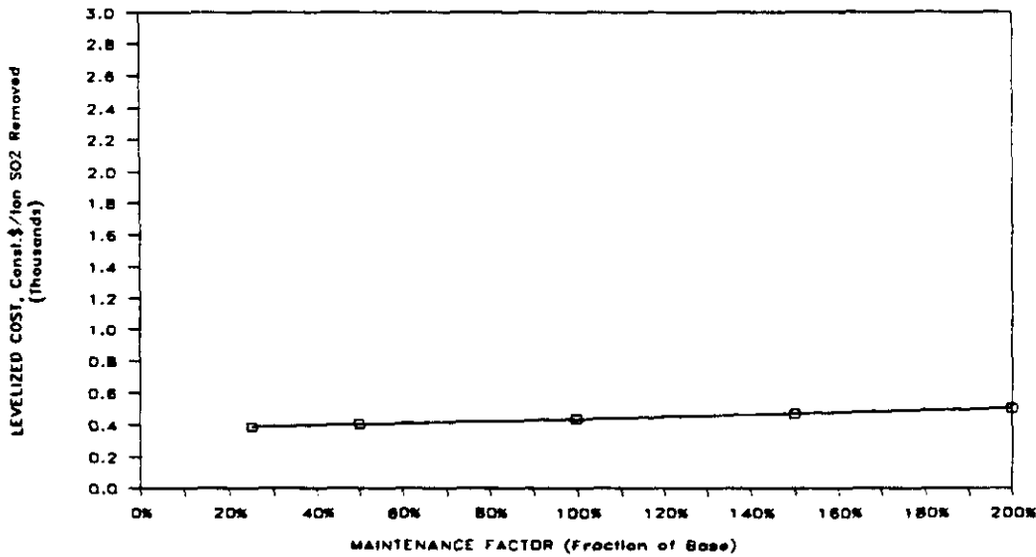
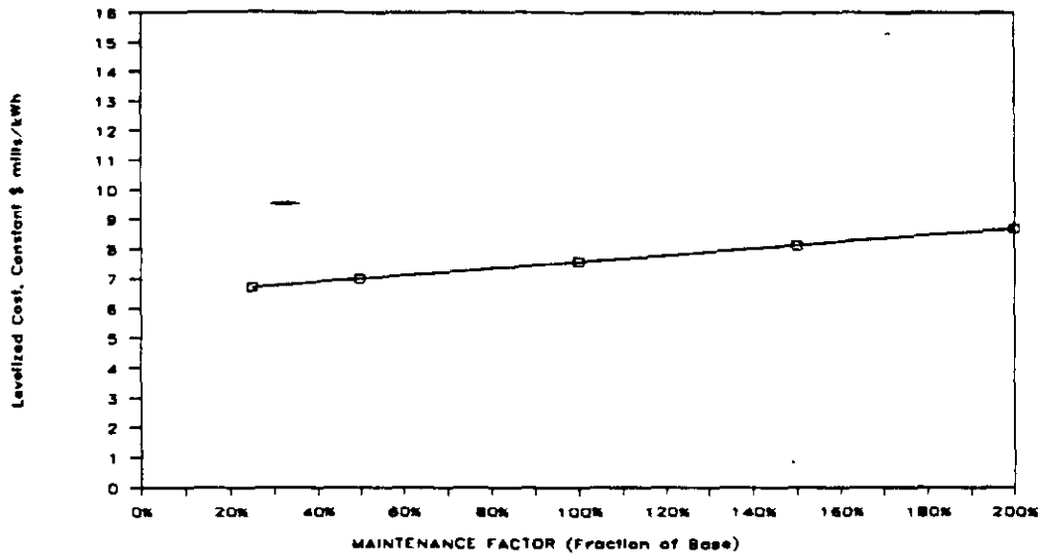


Figure 20-9. AirPoll GSA Sensitivity Curves

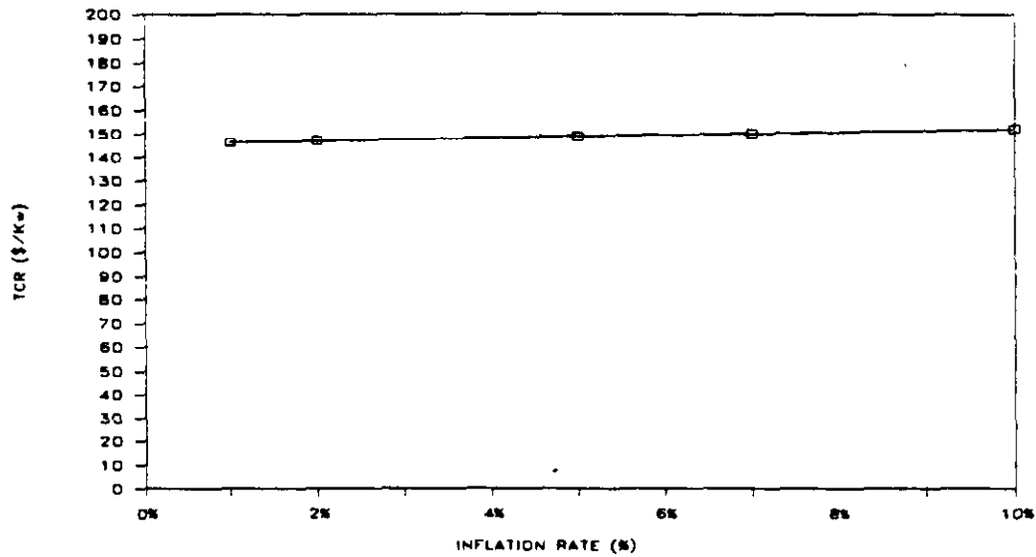
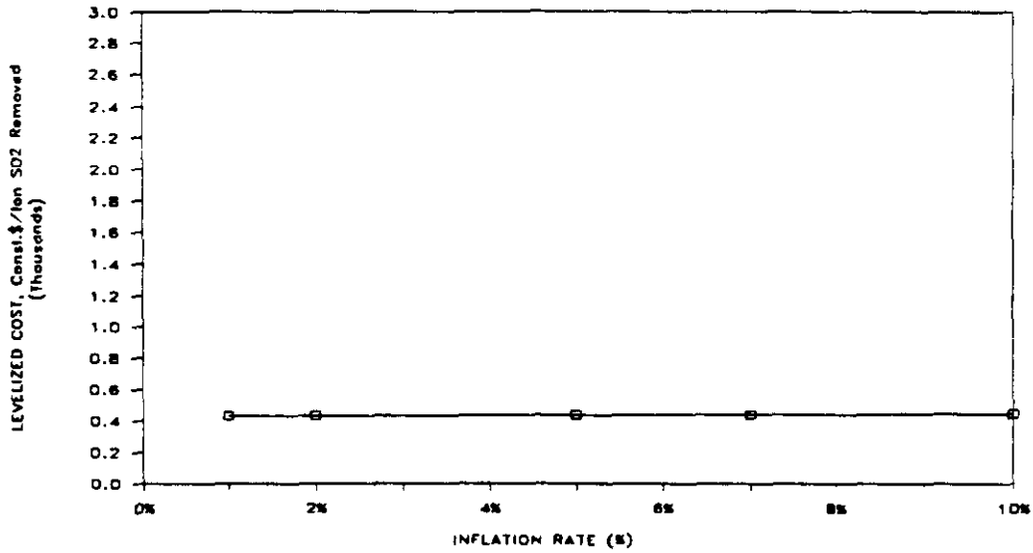
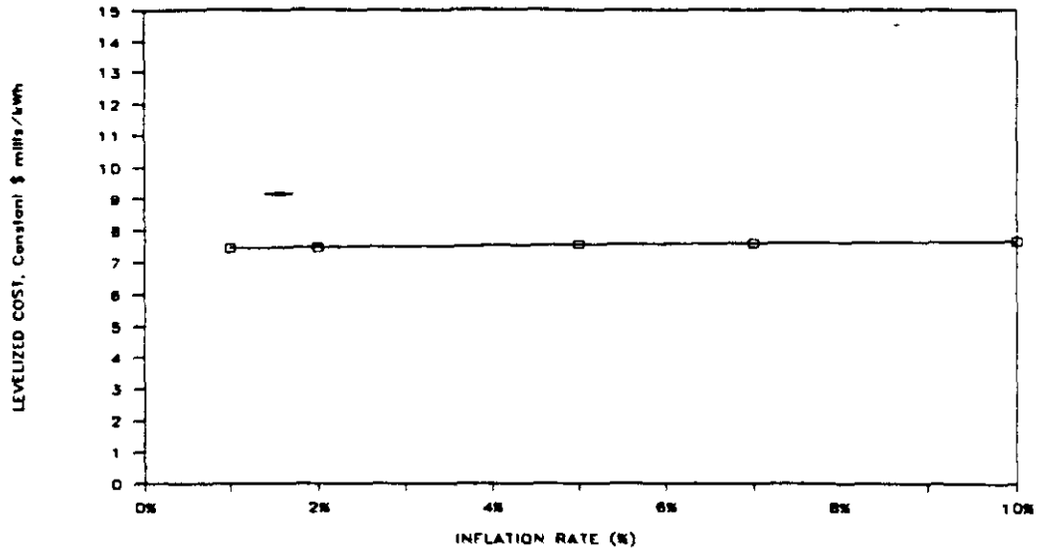


Figure 20-10. AirPol GSA Sensitivity Curves

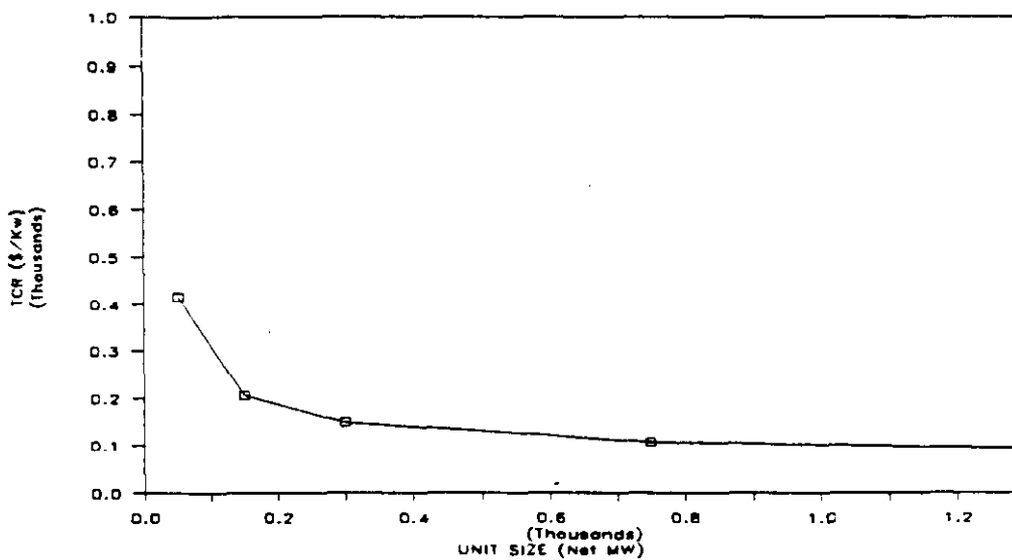
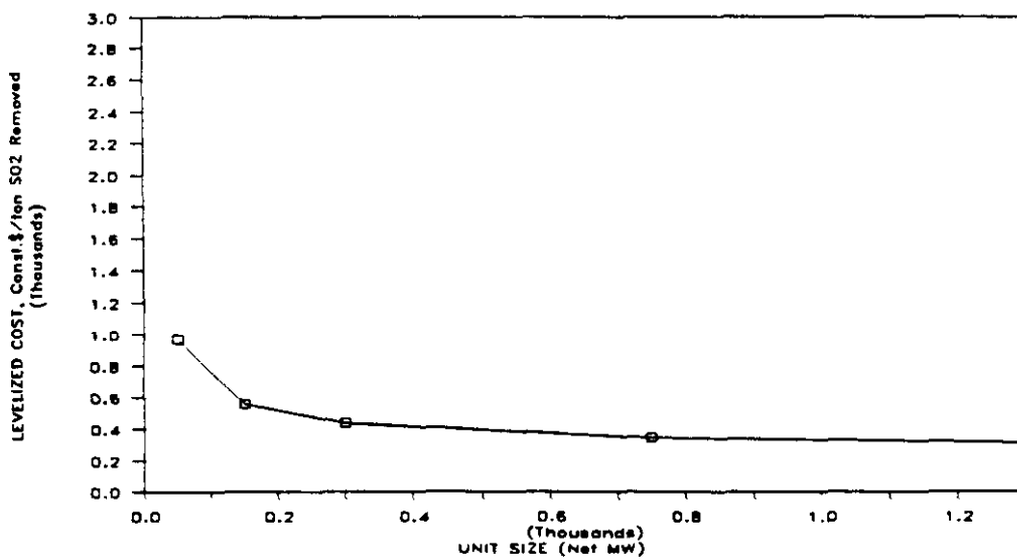
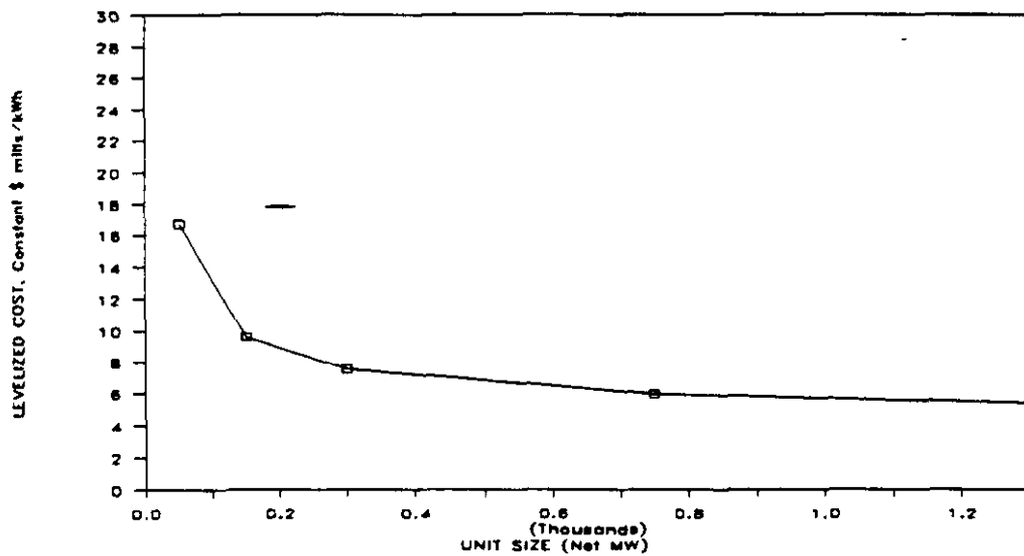


Figure 20-11. AirPol GSA Sensitivity Curves

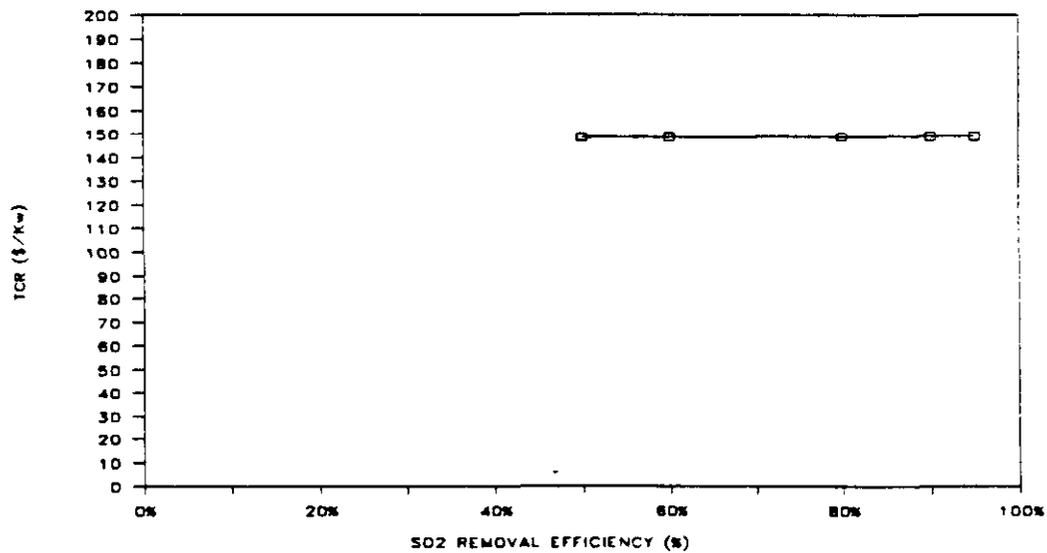
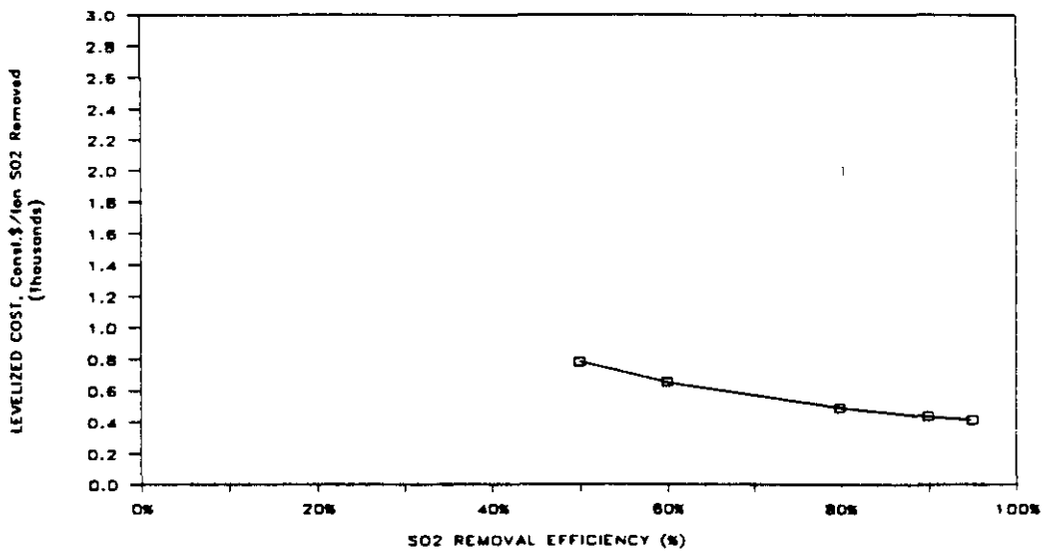
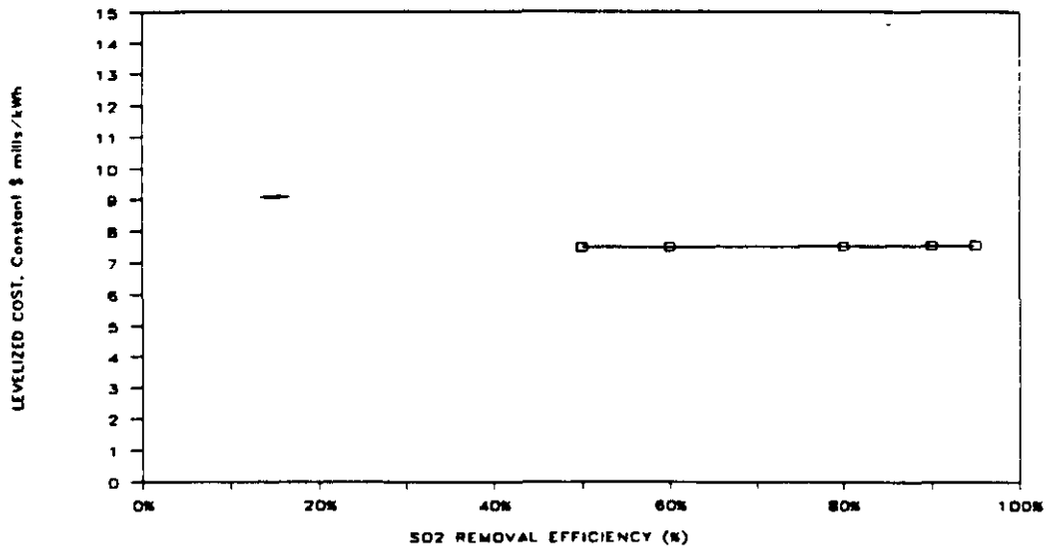


Figure 20-12. AirPol GSA Sensitivity Curves

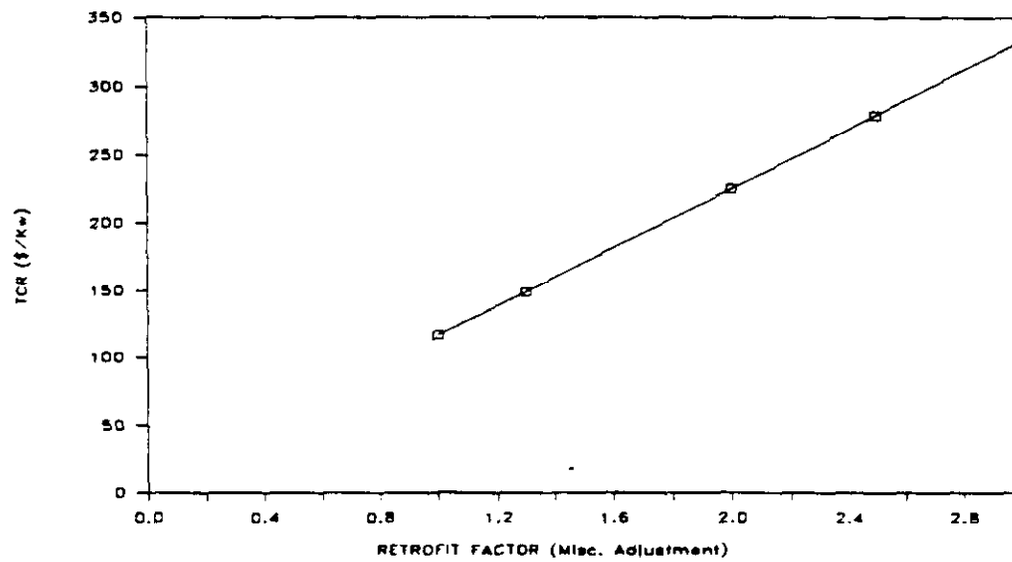
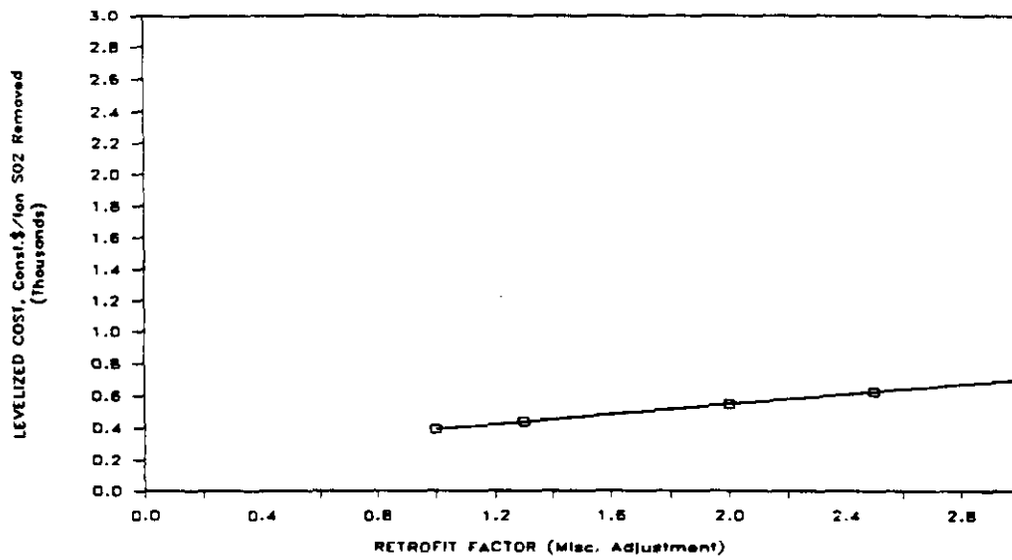
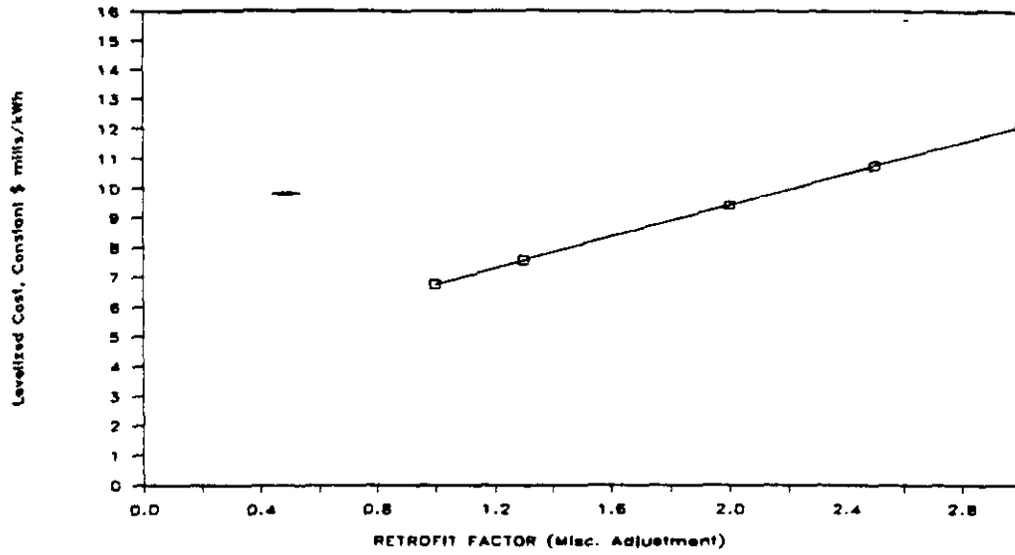


Figure 20-13. AirPol GSA Sensitivity Curves

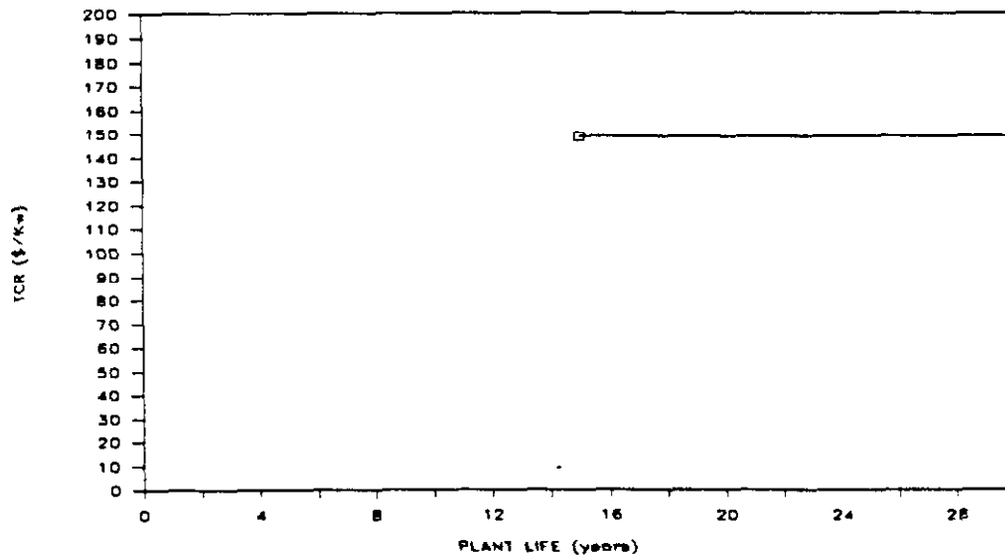
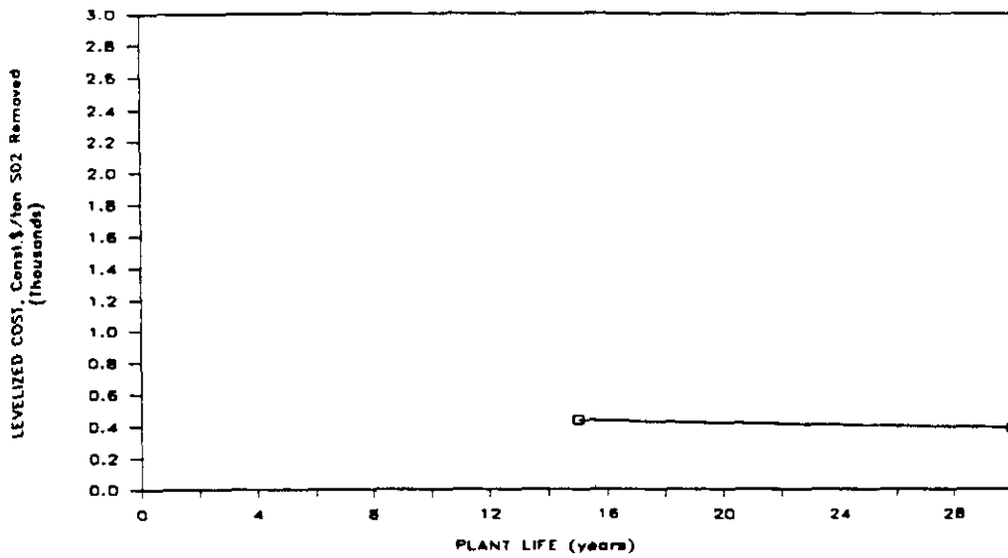
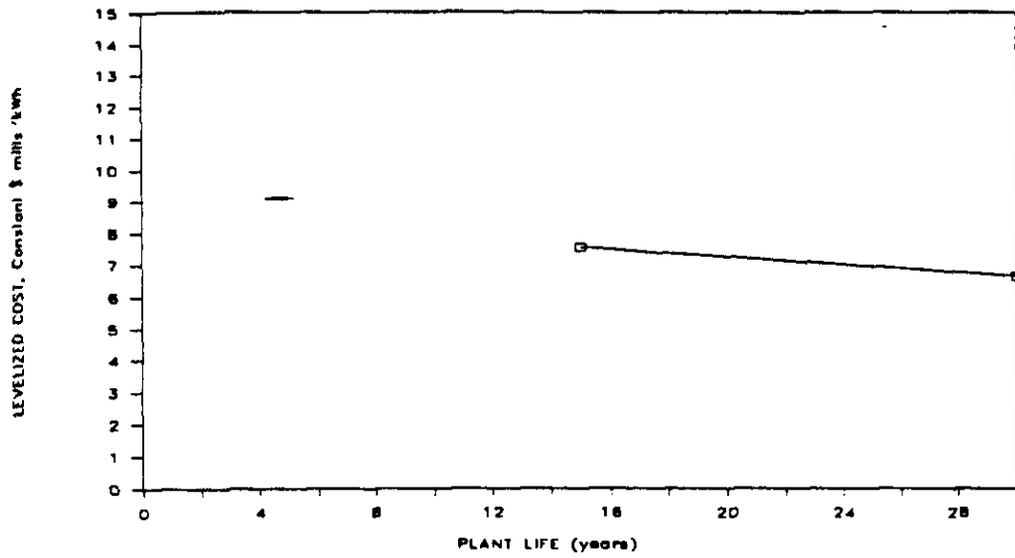


Figure 20-14. AirPol GSA Sensitivity Curves

REFERENCES

1. Hedenhag, J., "Technical and Economic Comparison of State of the Art Wet and Dry FGD Systems," presented at the Industrial Power Conference, Lakewood, Colorado, March 28-30, 1993.
2. "AirPol Gas Suspension Absorption Demonstration Plant," DOE Quarterly Project Summary, January 1993.
3. "TVA Begins Testing AirPol's Pollution Control Technology," (DOE) Fossil Energy Review, October-December, 1992.
4. Project specific transmittals of information from AirPol to Raytheon, June/July/October 1993.
5. Burnett, T. et.al. "10-MW Demonstration of the AirPol Gas Suspension Absorption Flue Gas Desulfurization Process," presented at the 1993 SO₂ Control Symposium, Boston, Massachusetts, August 24-27, 1993.
6. Burnett, T. et.al. "Recent Results from the 10-MW Demonstration of the Gas Suspension Absorption Process at TVA's Center for Emissions Research," to be presented at the 1994 International Joint Power Generation Conference, October 1994.

VENDOR COMMENTS

AirPol, Inc. (Mr. Frank Hsu)

- The first 18 months of the GSA demonstration at TVA's CER were successful in proving the GSA's capability to effectively remove SO₂ from the flue gas. TVA plans to continue the development of the GSA with the following activities:
 1. Continue to monitor and evaluate the performance of the ESP to ensure that the GSA process will not have an adverse impact on this particulate control device;
 2. Conduct lower SCA tests by deenergizing one or more fields in the ESP to determine the resulting effect on particulate emissions;
 3. Evaluate the effect of other limes on the performance of the GSA system; and
 4. Evaluate the potential for using the by-product material from the atmospheric fluidized bed combustion unit as a source of lime to displace some of the fresh lime feed to the system.

INDUSTRY COMMENTS

Tennessee Valley Authority (Mr. Tom Burnett)

- We assume that the request for detention slaker use is based on the need to remove the lime grit before feeding the lime slurry to the two-fluid nozzle since the current nozzle is not designed to be abrasion-resistant. (The lime grit is very abrasive even though it is ground to a very small size in the ballmill slaker and remains in the lime slurry that is fed to the two-fluid nozzle.) The utility industry is likely to have a strong preference for a ballmill slaker to prevent the production of an additional wastewater stream that must be handled and disposed of in an acceptable manner. We would suggest that AirPol either develop an abrasion-resistant nozzle or

develop and include a means for handling and disposing of the grit material to alleviate the utility concerns over this issue.

- The use of cooling tower blowdown for slaking the lime will similarly meet significant resistance in the utility industry. Previous work by the Electric Power Research Institute (among others) has indicated that the dissolved salts in the cooling tower blowdown adversely affect the lime slaking reaction leading to lower lime utilization rates in these dry, lime-based systems. Since the cost of lime is the major annual operating cost in a utility GSA system, any change that reduces the lime utilization rate will have severe repercussions for the process economics. We would suggest that AirPol undertake a test program to confirm that cooling tower blowdown can be used to slake the pebble lime without affecting the lime utilization before recommending its use in an actual application.
- We do not believe that the conversion of the pneumatic system for the overflow solids line to a mechanical screw conveyor and chute would be the best way to move this material from the recycle feeder box to the storage silo. At the 10-MW plant, where we had an existing bucket elevator to receive the overflow material and lift it to the top of the silo, a mechanical screw conveyor may have been appropriate. However, in a full-scale, new GSA installation, we believe that a pneumatic system will be used to move the by-product material to a silo. (There may also be a screw conveyor to remove the by-product material from the recycle feeder box and into a chute with a rotary valve that feeds the pneumatic transfer system, but a pneumatic system will still be necessary.)

AirPol GAS SUSPENSION ABSORBER PROCESS EQUIPMENT LIST
 (For One 300 MW Unit)

AREA 10 - REAGENT FEED SYSTEM

<u>Equipment Item and Description</u>	<u>Qty.</u>	<u>Material Unit Cost</u> *	<u>Material Total Cost</u> *
<u>Railspur.</u>	1	\$101,000	\$101,000
Service:	Provide FGD plant access for reagent delivery		
<u>Lime Receiving System.</u>	1 Lot	\$107,100	\$107,100
Service:	Pebble Lime		
Includes:	1 - two railcar capacity weather enclosure 2 - 50-T below rail hoppers 4 - hopper pressure feeders, 12" x 9" inlet tee, 12" isolation gate, 50 tph ea. 1 - 4000 ACFM baghouse, 10 Hp, 1600 ft ² 1 - concrete pit 1 - car shaker w/trolley hoist		
<u>Reagent Pneumatic Unloading Conveying System.</u>	1 Lot (1 spare blower included)	\$242,800	\$242,800
Type:	Pneumatic		
Service:	Quick lime		
Solids Rate:	50 tph		
Transfer Lines:	8" x 600'		
Blower Motor:	200 Hp ea.		
<u>Lime Storage Silo.</u>	2 op.	\$499,800	\$999,600
Dimensions:	43' dia. x 129' straight side, 60° conical bottom		
Capacity:	60 days (total)		
Dust Collector:	Baghouse, 1000 ft ²		
Material:	Concrete		
<u>Silo Pressure Feeders.</u>	20 op.	\$ 3,050	\$ 61,000
Type:	Slide gate or wafer, 12" inlet/outlet pneumatically operated		
Service:	Lime		
Capacity:	10 tph		
Material:	Carbon steel		
Includes:	12" x 9" inlet tee, 12" isolation gate		

*All costs listed are uninstalled equipment costs unless otherwise noted.
 Installed costs do not include an adjustment for construction labor.

AirPol

AREA 10 - REAGENT FEED SYSTEM (Continued)

<u>Equipment Item and Description</u>	<u>Qty.</u>	<u>Material Unit Cost</u>	<u>Material Total Cost</u>
<u>Reagent Pneumatic Transfer Conveying System.</u>	1 Lot (1 spare blower included)	\$179,700	\$179,700
Type: Pneumatic			
Service: Quick lime			
Solids Rate: 25 tph			
Transfer Lines: 6" x 600'			
Blower Motor: 125 Hp			
<u>Lime Day Bin.</u>	2 op.	\$ 41,650	\$ 83,300
Dimensions: 13' dia. x 39' straight side, 60° conical bottom			
Retention: 30 hrs			
Material: Carbon steel with cover			
Dust Collector: Baghouse, 400 ft ²			
<u>Detention Slaker.</u>	2 op. 1 spare	\$ 57,200	\$171,600
Service: Pebble lime, 1"			
Feed Rate: 3.6 tph			
Motor: 2.5 Hp			
Material: Carbon steel with rubber liner			
Accessories: Feeder, grit remover, pumps, piping, and controls			
<u>Lime Slurry Feed Tank.</u>	1 op.	\$59,100	\$118,200
Dimensions: 22' dia. x 22' straight side			
Service: 30% hydrated lime slurry			
Material: Carbon steel neoprene lined, open top, 4 baffles, agitator support			
Normal Level: 20'			
<u>Miscellaneous Process Equipment.</u>			\$103,200
(5 percent of total area capital requirement)			

AirPol

AREA 20 - SO₂ REMOVAL SYSTEM

<u>Equipment Item and Description</u>	<u>Qty.</u>	<u>Material Unit Cost</u>	<u>Material Total Cost</u>
<u>Gas Suspension Absorber.</u>	2 op. 1 spare	\$154,600	\$463,800
Type: Circulating fluid bed			
Service: Flue gas/hydrated lime			
Dimensions: 21' I.D. x 76' straight side, 164' overall height			
Capacity: 510 K AFCM @ 277°F inlet temperature			
Material: Carbon steel with 4" insulation			
Includes: Dry gas venturi on lower portion of vessel			
<u>Lime Slurry Feed Pump.</u>	1 op. 1 spare	\$ 2,300	\$ 4,600
Type: Horizontal centrifugal			
Flow: 150 gpm			
Head: 100'			
Material: Cast steel, rubber-lined			
Motor: 7.5 Hp			
<u>Slurry Atomizer.</u>	2 op. 1 spare	\$ 5,000	\$ 15,000
Service: Lime Slurry			
Nozzles: Single dual-fluid			
Flow: 70 gpm			
Piping: 4" dia transfer piping			
<u>Slurry Injection Air Compressor.</u>	1 op. 1 spare	\$122,150	\$244,300
Type: Multi-stage centrifugal air compressor unit complete with control unit, motor driver, prefilter, and after-filter			
Service: Air			
Motor: 300 Hp			

AirPol

AREA 20 - SO₂ REMOVAL SYSTEM (Continued)

<u>Equipment Item and Description</u>	<u>Qty.</u>	<u>Material Unit Cost</u>	<u>Material Total Cost</u>
<u>Cyclone Collector.</u>	4 op. 2 spare	\$203,100	\$1,218,600
Type: Mechanical tubular collector			
Service: Flue gas with spent sorbent/fly ash			
Capacity: 220 K AFCM @ 152°F			
Size: 24' dia. x 75' overall height			
Material: Carbon Steel			
Removal: 99%			
Efficiency:			
<u>Recirculation Solids Feeder Box.</u>	4 op. 2 spare	\$138,200	\$829,200
Service: Product solids and fly ash			
Material: Carbon steel			
Flow: 519 tph			
Includes: Return and overflow solids conveyors			
Size: 8'9" W x 8'9" L x 13'H			
<u>ESP Modifications.</u>	1 Lot	\$115,200	\$115,200
Service: Flue gas, 145°F			
Modifications: Remove existing external ESP insulation and add 6" of new insulation			
<u>Miscellaneous Process Equipment.</u>			\$144,500
(5 percent of total area capital requirement)			

AirPol

AREA 30 - FLUE GAS SYSTEM

<u>Equipment Item and Description</u>	<u>Qty.</u>	<u>Material Unit Cost</u>	<u>Material Total Cost</u>
<u>GSA Reactor Inlet Isolation Damper.</u>	3	\$155,000	\$465,000
Type: Double blade guillotine w/purge air			
Service: Flue gas			
Material: Carbon steel with nickel base metal for wetted parts			
Duct Size: 9' x 16'			
<u>GSA Reactor Outlet Isolation Damper.</u>	3	\$132,200	\$396,600
Type: Double guillotine w/purge air			
Service: Treated flue gas			
Material: Incoloy 825			
Duct Size: 9' x 14'			
<u>FGD System By-pass Damper.</u>	1	\$268,800	\$268,800
Type: Double-louver, parallel blade w/purge air			
Service: Flue gas, 277°F			
Material: Carbon steel with nickel base metal for wetted parts			
Duct Size: 14' x 20'			
<u>Bypass Ducting.</u>	1 Lot	\$551,300	\$551,300
Type: Rectangular with external stiffeners			
Service: Flue gas			
Material: Carbon steel w/4" stiffeners			
Surface Area: 6,730 ft ²			
<u>Inlet Reactor Ducting, including Manifold.</u>	1 Lot	\$363,900	\$363,900
Type: Rectangular w/external stiffeners			
Service: Flue gas			
Material: Carbon steel w/4" stiffeners			
Surface Area: 4,400 ft ²			

AirPol

AREA 30 - FLUE GAS SYSTEM (Continued)

<u>Equipment Item and Description</u>	<u>Qty.</u>	<u>Material Unit Cost</u>	<u>Material Total Cost</u>
<u>Outlet Reactor/Cyclone Ducting, including Manifold.</u>	1 Lot	\$720,500	\$720,500
Type: Rectangular w/external stiffeners			
Service: Treated flue gas			
Material: Carbon steel, reinforced polyester lined, insulated			
Surface Area: 5,340 ft ²			
<u>Duct Lining Outlet from ESP to ID Fan.</u>	1 Lot	\$104,300	\$104,300
Service: Scrubbed flue gas			
Materials: Reinforced polyester lining with insulation			
Surface Area: 9,960 ft ²			
<u>Duct Lining from ID Fan Outlet, including Chimney Transition.</u>	1 Lot	\$ 56,300	\$ 56,300
Service: Flue gas			
Materials: Reinforced polyester lining with insulation			
Surface Area: 5,370 ft ²			
<u>ID/Booster Fan Modification.</u>	2 op.	\$381,700	\$763,400
Type: Centrifugal fan			
Operating Conditions: 448 K AFCM @ 10.0" WC and 145°F			
BHP: 880 Hp			
Test Conditions: 538 K AFCM @ 14.0" WC and 145°F			
Motor Test			
Block Rating: 1500 Hp			

AirPol

AREA 60 - SOLIDS HANDLING SYSTEM

<u>Equipment Item and Description</u>	<u>Qty.</u>	<u>Material Unit Cost</u>	<u>Material Total Cost</u>
<u>Solids Pneumatic Conveying System.</u>	1 Lot (1 spare blower included)	\$179,700	\$179,700
Type: Pneumatic			
Service: Fly ash/calcium sulfite/ calcium sulfate mixture			
Capacity: 25 tph			
Transfer Lines: 6" x 600'			
Blower Motor: 150 Hp			
<u>Solids Silo.</u>	1	\$268,800	\$268,800
(Sized for spent sorbent quantity only assuming capacity exists for flyash collected)			
Dimensions: 28' dia. x 84' straight side, 60° Cone			
Capacity: 3 day storage			
Service: Spent sorbent/fly ash mixture			
Material: Carbon steel w/stainless steel cone			
Accessories: Baghouse, 1000 ft ²			
<u>Pug Mill.</u>	2 op.	\$ 33,950	\$ 67,900
Service: Spent sorbent/fly ash/water			
Discharge			
Capacity: 30 tph			
Motor: 40 Hp			
<u>Demolition of Ash Piping.</u>	1 Lot	\$ 57,750	\$ 57,750
Includes: Removal of all existing pneumatic conveying piping, blowers, and all associated equipment			
<u>Miscellaneous Process Equipment.</u>			\$ 28,700
(5 percent of total area capital requirement)			

AirPol

AREA 70 - GENERAL SUPPORT EQUIPMENT

<u>Equipment Item and Description</u>	<u>Qty.</u>	<u>Material Unit Cost</u>	<u>Material Total Cost</u>
<u>Instrument/Plant Air Compressor System.</u>	2 op.	\$ 48,100	\$ 96,200
Type: Multi-stage, rotary screw air compressor unit complete with control unit, motor driver, instrument air-dryer, pre-filter, after-filter, after-cooler, and air receivers Service: Air Air-dryer Power Consumption: 6.6 kW Motor: 85 Hp			
<u>Makeup Blowdown Water Tank.</u>	1 op.	\$ 8,200	\$ 8,200
Size: 10' dia. x 10' straight side Service: Cooling tower blowdown Material: Carbon steel Normal Level: 9'			
<u>Miscellaneous Process Equipment.</u>			\$ 5,200
(5 percent of total area capital requirement)			

AREA 80 - ADDITIONAL EQUIPMENT

<u>Onsite Electric Power.</u>	1 Lot	\$886,000	\$886,000
Type: Miscellaneous electrical equipment Includes: Grounding, transformers			

APPENDIX "B"

**10 MW DEMONSTRATION OF
THE GAS SUSPENSION ABSORPTION PROCESS
AT TVA'S CENTER FOR EMISSIONS RESEARCH**

By Tennessee Valley Authority

**Prepared for the United States Department of Energy
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**Cleared by Office of Patent Counsel
Chicago Operations Office
U.S. Department of Energy**