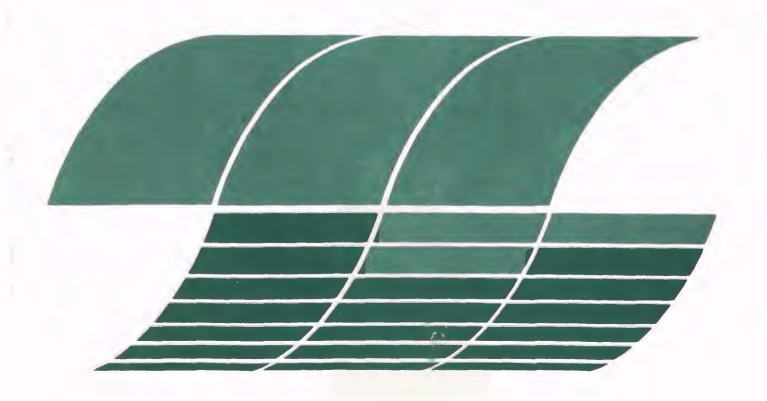
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Technology Assessment Report for Industrial Boiler Applications: NO_X Flue Gas Treatment

Interagency Energy/Environment R&D Program Report



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Technology Assessment Report for Industrial Boiler Applications: NO_x Flue Gas Treatment

by

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ABSTRACT

This study assesses the applicability of NO, flue gas treatment technology to industrial boilers and is one of a series of technology assessment reports to aid in determining the technological basis for a New Source Performance Standard for Industrial Boilers. The status of development and performance of alternative NO_x flue gas treatment control techniques were assessed and the cost, energy, and environmental impacts of the most promising processes were identified. It was found that processes utilizing selective catalytic reduction (SCR) of NO, with ammonia can achieve 90 percent reduction of NO, emissions, and that these processes are the nearest to commercialization in the U.S. In Japan, SCR processes have been successfully operated on commercial scale gas-and oil-fired sources and are being installed on coal-fired sources. Cost estimates of applying SCR processes in the U.S. indicated that the cost effectiveness varies significantly depending on the fuel fired, boiler size, and control level. However, boiler size is the most significant factor affecting cost effectiveness with the economy of scale causing control of large sources to be the most effective. The energy impact of applying SCR processes averaged about 0.5 percent of boiler capacity. No adverse environmental impacts were apparent although there are emissions, liquid effluents, and solid wastes that must be controlled. For regulatory purposes this assessment must be viewed as preliminary, pending the results of the more extensive examination of impacts called for under Section 111 of the Clean Air Act.

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PREFACE

The 1977 Amendments to the Clean Air Act required that emission standards be developed for fossil-fuel-fired steam generators. Accordingly, the U.S. Environmental Protection Agency (EPA) recently promulgated revisions to the 1971 new source performance standard (NSPS) for electric utility steam generating units. Further, EPA has undertaken a study of industrial boilers with the intent of proposing an NSPS for this category of sources. The study is being directed by EPA's Office of Air Quality Planning and Standards, and technical support is being provided by EPA's Office of Research and Development. As part of this support, the Industrial Environmental Research Laboratory at Research Triangle Park, NC, prepared a series of technology assessment reports to aid in determining the technological basis for the NSPS for industrial boilers. This report is part of that series. The complete report series is listed below:

Title	Report No.
The Population and Characteristics of Industrial/ Commercial Boilers	EPA-600/7/79-178a
Technology Assessment Report for Industrial Boiler Applications: Oil Cleaning	EPA-600/7-79-178b
Technology Assessment Report for Industrial Boiler Applications: Coal Cleaning and Low Sulfur Coal	EPA-600/7-79-178c
Technology Assessment Report for Industrial Boiler Applications: Synthetic Fuels	EPA-600/7-79-178d
Technology Assessment Report for Industrial Boiler Applications: Fluidized-Bed Combustion	EPA-600/7-79-178e
Technology Assessment Report for Industrial Boiler Applications: $\mathrm{NO}_{ imes}$ Combustion Modification	EPA-600/7/79-178f

Title	Report No.
Technology Assessment Report for Industrial Boiler Applications: NO _X Flue Gas Treatment	EPA-600/7-79-178g
Technology Assessment Report for Industrial Boiler Applications: Particulate Collection	EPA-600/7-79-178h
Technology Assessment Report for Industrial Boiler Applications: Flue Gas Desulfurization	EPA-600/7-79-178i

These reports will be integrated along with other information in the document, "Industrial Boilers - Background Information for Proposed Standards," which will be issued by the Office of Air Quality Planning and Standards.

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SECTION 1

EXECUTIVE SUMMARY

1.1 INTRODUCTION

1.1.1 Background and Objectives

The Clean Air Act Amendments of 1977 require the Environmental Protection Agency to coordinate and lead the development and implementation of regulations on air pollution. These include standards of performance for new and modified sources of pollution. Fossil fired steam generators are specifically mentioned in the act and EPA has undertaken a study of industrial boilers with intent to propose emission control levels based upon the results of this and other studies.

This specific report examines the im_Pacts of application of flue gas treatment (FGT) for NO_X control on industrial boilers. The overall objective is to provide a background document that quantifies the economic, energy and environmental impacts as well as establish whether or not the technology is demonstrated and available to the U.S. market. All potential FGT technologies are considered and detailed analyses are performed on those which are the most promising.

1.1.2 Report Organization and Approach

Several boiler/FGT combinations are considered in the detailed analyses that follow. In Section 2, all NO_X control processes that have been developed to treat boiler flue gas are discussed in moderate detail. The section

is divided into three subsections based on fuel--coal, oil, and gas. This is done for two reasons: 1) to make this report consistent in format with other Individual Technology Assessment Reports (ITAR's, see preface), and 2) to examine the effect of fuel type on the various technologies considered. In the case of FGT, the majority of the technologies can be applied to the majority of the fuels. As a result, much of the material in the three subsections is very similar, especially with regard to the technical descriptions of the systems.

A decision was made early in the development of this report to produce essentially three stand-alone sections; one for each fuel type. This allows one or more fuel types to be eliminated from consideration without impacting the quality of the data in the remaining sections. As a result, there is a significant amount of repetition in the three subsections. The subsection dealing with applications to oil-fired boilers (Section 2.3) contains descriptions of all of the FGT technologies considered and has the greatest amount of information on specific systems. Therefore, for most readers, it is necessary to read only subsections 2.1 and 2.3 for a complete description of all FGT technologies considered. Subsections 2.2 and 2.4, dealing with coal- and gas-fired applications, do contain unique information on the status of development and number of applications and can be consulted if this specific information is desired. The Executive Summary is organized differently than the body of the report in that all fuel types are discussed together instead of separately. This is done to allow the reader to directly examine the effect of fuel type on the economic, energy, and environmental impacts. The summary discusses each of the impacts separately and also separates NO_{γ} -only systems from NO_{γ}/SO_{γ} systems.

The large number of potential fuel/boiler/FGT system combinations requires the combinations be reduced to those systems with a high potential for commercial application and successful operation. This is done in Section 3. The data used to make these preliminary evaluations is derived from Section 2. The combinations selected in Section 3 are then analyzed in detail in subsequent sections.

Section 4 presents the economic impacts of these FGT processes on ten industrial boilers. Standard costing techniques are used to develop annualized costs which are plotted to show the effect of several parameters on the total system costs. The process specifications used in the economic analyses are developed in Sections 5 and 6. These two Sections present the results of material and energy balances performed for each case to quantify the energy and environmental impacts of FGT systems. These balances are also used to size most of the individual pieces of equipment.

In Sections 4, 5 and 6 results are not presented for all possible control levels. That is, while some systems have data presented for three levels of control, others have data presented for only two or one control level(s). The curves, however, are plotted over the range of the three control levels (70, 80 and 90 percent). When the individual Sections were initially prepared, data was calculated for all three control levels. During the interim period prior to the compilation of these Sections into the final report, new cases were added and several economic premises were changed. In order to meet budget and time constraints, it was necessary to reduce the number of analyses.

It was observed that curve shapes were all very similar and that new curves could be drawn accurately without a complete set of points. In cases with a lot of similarity, i.e. among the coal-fired boilers, a curve shape was established for one case by using three points. For the other cases, an analysis was made to determine the midpoint of the curve. Curves, similar in shape to that developed by a three-point analysis, were then passed through these midpoints. In cases where there was little similarity, a two-point analysis was performed to determine the end-points of the curve. A curve was then drawn through these points using the original curve (determined by three-point analysis in the initial case analysis) to determine the shape. It should be noted that, even if straight lines were used, the interpolated and extrapolated results would not be changed

significantly. This is why a limited set of analyses are used to determine a complete set of cost and energy data.

The final section, Section 7, deals with test data that have been determined for operating FGT systems. These types of data do not exist for U.S. applications since FGT has yet to be applied in this country. A Japanese consultant with contacts among FGT system users was retained to obtain test data from industrial boilers in Japan. The test data presented in Section 7 represents the most complete set of data of this type available.

For the reader interested in the details involved with the analyses presented in this report, the Appendices present an example calculation as well as complete sets of process selection criteria material and energy balances, and cost breakdowns.

1.1.3 Scope of Study

Several variables are considered in order to make the study as comprehensive as possible, these being

- Fuel
- Boiler, type and size
- Control level
- FGT process type
- Pollutants controlled.

As mentioned previously, three fuel types are considered: coal, oil and natural gas. Coal and oil are further divided as shown below.

Coal

High Sulfur Eastern (3.5% S) Low Sulfur Eastern (0.9% S) Low Sulfur Western (0.6% S)

0il

Distillate Residual

One boiler type is considered for natural gas, distillate oil and residual oil. However, four boiler types are considered for the coal fuels. The combinations of fuel and boiler type considered at the beginning of the study are shown in Table 1.1.3-1. These boilers are termed "standard boilers" because they apply to all of the ITAR's. The NO_{\times} emissions from these boilers are shown in Table 1.1.3-2.

In the ensuing discussion of emission control technologies, candidate technologies were compared using three emission control levels labeled "moderate, intermediate, and stringent." These control levels were chosen only to encompass all candidate technologies and form bases for comparison of technologies for control of specific pollutants considering performance, costs, energy, and non-air environmental effects.

From these comparisons, candidate "best" technologies for control of individual pollutants are recommended by the contractor for consideration in subsequent industrial boiler studies. These "best technology" recommendations do not consider combinations of technologies to remove all pollutants and have not undergone the detailed environmental, cost, and energy impact assessments necessary for regulatory action. Therefore, the levels of "moderate, intermediate, and stringent" and the recommendation of "best technology" for individual pollutants are not to be construed as indicative of the regulations that will be developed for industrial boilers. EPA will

Туре	Fuel*	Rating MW _t (MBtu/hr)	Gas Flow Rate Nm ³ /hr
Package, Firetube	Distillate Oil	4.4 (15)	5,400
Package, Firetube	Natural Gas	4.4 (15)	5,600
Package, Watertube	Residual Oil	8.8 (30)	9,500
Package, Watertube Underfeed Stoker	HSE LSE LSW	8.8 (30) 8.8 (30) 8.8 (30)	12,500 12,600 12,900
Package, Watertube Chaingrate Stoker	HSE LSE LSW	22 (75) 22 (75) 22 (75)	31,300 31,000 32,400
Package, Watertube	Natural Gas	44 (150)	52,800
Package Watertube	Distillate Oil	44 (150)	51,900
Package, Watertube	Residual Oil	44 (150)	47,800
Field Erected, Watertube Spreader Stoker	HSE LSE LSW	44 (150) 44 (150) 44 (150)	62,900 62,700 64,700
Field Erected, Watertube Pulverized Coal	HSE LSE LSW	58.6 (200) 58.6 (200) 58.6 (200)	72,600 72,800 75,500

TABLE 1.1.3-1. CHARACTERISTICS OF THE STANDARD BOILERS

*Coal types: HSE = High Sulfur Eastern (3.5% S)

LSE = Low Sulfur Eastern (0.9% S)

LSW = Low Sulfur Western (0.6% S)

Boiler	Fuel*	NO _× Emissions				
		g/s	(1b/hr)	ng/J	(1b/10 ⁶ Btu)	ppm
Package, Firetube	Distillate Oil	0.300	(2.38)	68.8	(0.16)	97
Package, Firetube	Natural Gas	0.332	(2.63)	77.4	(0.18)	104
Package, Watertube	Residual Oil	2.02	(16.0)	228	(0.53)	373
Package, Watertube Underfeed Stoker	HSE LSE LSW	2.40 2.06 2.95	(10.05) (16.35) (23.40)	275 237 335	(0.64) (0.55) (0.78)	335 288 402
Package, Watertube Chaingrate	HSE LSE LSW	6.02 5.15 7.40	(47.70) (40.80) (58.65)	275 232 335	(0.64) (0.54) (0.78)	336 290 401
Package, Watertube	Natural Gas	3.31	(26.26)	75.3	(0.18)	110
Package, Watertube	Distillate Oil	2.99	(23.76)	68.0	(0.16)	101
Package, Watertube	Residual Oil	7.47	(60.00)	172	(0.40)	292
Field Erected, Watertube Spreader Stoker	HSE LSE LSW	12.0 10.3 14.8	(95.40) (81.45) (117.15)	275 232 335	(0.64) (0.54) (0.78)	337 288 400
Field Erected, Watertube Pulverized Coal	HSE LSE LSW	19.2 16.5 23.7	(152.46) (130.50) (187.56)	327 280 404	(0.76) (0.65) (0.94)	466 396 550

TABLE 1.1.3-2. NO_{\times} EMISSION RATES FOR THE STANDARD BOILERS

*Coal types: HSE = High Sulfur Eastern (3.5% S)

LSE = Low Sulfur Eastern (0.9% S)

LSW = Low Sulfur Western (0.6% S)

perform rigorous examination of several comprehensive regulatory options before any decisions are made regarding the standards for emission from industrial boilers. The control levels are defined in Table 1.1.1-3.

The types of FGT systems considered are different for each fuel type and these are discussed in subsequent sections for each specific fuel. The project schedule required that the number of potential combinations of boiler, fuel, and control level be reduced in order to keep the number of required analyses manageable. Detailed analyses were performed on the cases shown in Table 1.1.3-4. Note that these are the cases for FGT processes which remove only NO_{χ} . For FGT processes which remove both NO_{χ} and SO_{χ} a separate set of cases was developed and is shown in Table 1.1.3-5.

Only one coal is considered for the NO_{\times} -only cases. This is due to the fact that FGT process designs and impacts are not significantly affected by fuel sulfur content and therefore analyzing each coal type would not yield any additional information. The flue gas flow rates and NO_{\times} concentrations vary somewhat among the coal types considered, but not enough to cause much difference in the size of the necessary FGT process. With all of the FGT systems analyzed, the equipment size is primarily a function of the flue gas flow rate and secondarily a function of NO_{\times} concentration. However, since coal sulfur level can affect the environmental impact, two coal types are considered in this section.

For the processes which remove both NO_{\times} and SO_{\times} , two coals are analyzed to show the effect of coal sulfur level on the various impacts. High sulfur eastern and low sulfur western were selected in order to have the widest range of coal sulfur levels. Also, NO_{\times}/SO_{\times} processes for oil-fired boiler application are considered only for the case of residual oil since this oil has the most significant SO_{\times} emissions. NO_{\times}/SO_{\times} processes are examined for application to these boilers to enable comparison between a simultaneous NO_{\times}/SO_{\times} system and a combination of a NO_{\times} -only system and an FGD system. This comparison will be made during a future phase of the industrial boiler evaluation, but not in this report.

Baseline NO _X Emissions			Mode	Moderate, 70%		Level of Control Intermediate, 80%		Stringent, 90%	
Fuel	ng/J	(1b/MBtu)	ng/J	(1b/MBtu)	ng/J	(1b/MBtu)	ng/J	(1b/MBtu)	
Pulverized Coal	404	(0.94)	121	(0.28)	80.8	(0.19)	40.4	(0.094)	
Stoker Coal	335	(0.78)	101	(0.23)	67.0	(0.16)	33.5	(0.078)	
Residual Oil	172	(0.40)	51.6	(0.12)	34.4	(0.080)	17.2	(0.040	
Distillate Oil	68	(0.16)	20.4	(0.047)	13.6	(0.032)	6.8	(0.016)	
Natural Gas	75	(0.18)	22.6	(0.053)	15.1	(0.035)	7.5	(0.018)	

TABLE 1.1.1-3. NO_× CONTROL LEVELS

Where emissions are dependent on boiler size, the largest boiler is shown.

~

Boiler	Fuel*	Size ^{MW} t	Control Level %
Package, Firetube	Distillate Oil	4.4	70, 90
Package, Firetube	Natural Gas	4.4	70, 90
Package, Watertube	Residual Oil	8.8	70, 90
Package, Watertube Underfeed Stoker	LSW	8.8	80
Package, Watertube Chaingrate Stoker	LSW	22	70, 80, 90
Package, Watertube	Natural Gas	44	70, 90
Package, Watertube	Distillate Oil	44	70, 90
Package, Watertube	Residual Oil	44	70, 90
Field Erected, Watertube Spreader Stoker	LSW	44	80
Field Erected, Watertube Pulverized Coal	LSW	58.6	70, 90

TABLE 1.1.3-4. CASES SELECTED FOR DETAILED ANALYSIS - NOx-ONLY FGT PROCESSES

*LSW = Low Sulfur Western Coal (0.6%S)

TABLE 1.1.3-5.	CASES SELECTED	FOR DETAILED	ANALYSIS -
	SO_{\times}/NO_{\times} FGT PR	OCESSES	

Boiler	Fuel*	Boiler Size, ^{MW} t	<u>Control Level</u> % NO _X % SO _X	
Package, Watertube	Residual Oil	44	80	85
Package, Watertube Underfeed Scoker	HSE LSW	8.8	80	85
Field Erected, Watertube Pulverized Coal	HSE LSW	58.6	80	85

*HSE = High Sulfur Eastern Coal (3.5% S)

LSW = Low Sulfur Western Coal (0.6% S)

It should be noted that FGT technology for NO_× control has not yet been commercially applied to coal-fired boilers. However, pilot units have been tested and two full scale systems are scheduled. Coal-fired applications are considered here since they are currently being offered in the U.S. and it is felt that they will be demonstrated commercially in the near future.

1.2 FLUE GAS TREATMENT FOR CONTROL OF NO $_{\times}$ ONLY

The systems of emission reduction considered in this study for applications to coal-fired boilers are divided into two general categories: those which remove only NO_{χ} and those which remove both NO_{χ} and SO_{χ} . Here and throughout the study these two types of systems are considered separately to avoid confusion.

1.2.1 System Descriptions

The $\mathrm{NO}_{\times}\text{-}\mathrm{only}$ systems considered are as follows:

- Fixed Packed Bed Selective Catalytic Reduction (SCR)
- Moving Bed SCR
- Parallel Flow SCR
- Absorption-Oxidation

From the comparison evaluation of these systems, the candidates for "best" emission control systems were selected. These candidate systems are shown, along with a brief description, in Table 1.2.1-1.

SCR systems utilize ammonia to selectively reduce nitrogen oxides. The chemical mechanisms can be summarized by the following gas-phase reactions.

 $4NO + 4NH_3 + O_2 \stackrel{\rightarrow}{\leftarrow} 4N_2 + 6H_2O$ (1-1)

 $2NO_2 + 4NH_3 + O_2 \stackrel{\Rightarrow}{\leftarrow} 3N_2 + 6H_2 0$ (1-2)

Process	Description	Fuel Application
Moving Bed SCR	Utilizes NH_3 to selectively reduce NO_x to N_2 ; capable of achieving stringent NO_x control level; catalyst (rings or pellets) gravity-bed, mechani- cally-screened, and returned to reactor.	Residual Oil
Parallel Flow SCR	Utilizes NH_3 to selectively reduce NO_X to N_2 ; capable of achieving stringent NO_X control level; special catalyst arrangement (honeycomb, parallel plate or tubes) greatly reduces particulate impac- tion as gas flow is parallel to catalyst surface.	Coal Residual Oil
Fixed Packed Bed SCR	Utilizes NH ₃ to selectively reduce NO_X to N_2 ; capable of achieving stringent NO_X control level; ring shaped catalyst pellets packed in fixed bed.	Distillate Oil Natural Gas

TABLE 1.2.1-1. CANDIDATES FOR BEST EMISSION CONTROL SYSTEM

The first reaction predominates as flue gas NO_{\times} consists primarily of NO. Oxygen is in large excess in the flue gas and does not limit the extent of reaction. A process flow diagram is shown for an SCR system in Figure 1.2.1-1. Flue gas is taken from the boiler between the economizer and air preheater. Ammonia, taken from a liquid storage tank and vaporized, is injected and mixed with the flue gas prior to the reactor. The flue gas passes through the catalyst bed where NO_{\times} is reduced to N_2 . The flue gas then exits the reactor and is sent to the air preheater and, if necessary, further treatment equipment.

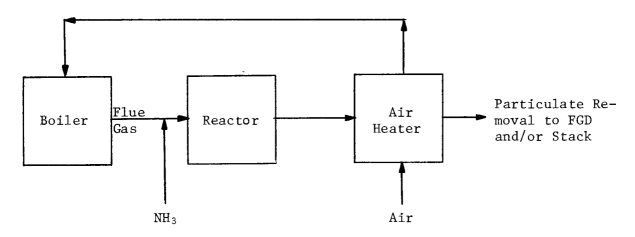


Figure 1.2.1-1. Flow diagram for typical NO_x-only SCR process.⁵

With this and all SCR systems it is desirable to treat flue gas exiting the economizer at 350-400°C prior to any air preheater since it is at this temperature range than the catalysts show the optimum combination of activity and selectivity. The analyses conducted in this study assumed that the boilers were operated constantly at full load and, therefore, had constant flue gas temperatures. However, it is possible that the boiler may experience large and frequent load swings which result in a variable flue gas temperature. FGT systems in this service will require flue gas heating in order to maintain sufficiently high temperatures. Temperature control can be accomplished by either a heater or a slipstream around the economizer. The heater will effectively decouple the FGT system from the boiler and does not require flow control of a flue gas slipstream. The economizer

bypass will not derate the boiler since it will only be required during low load situations. In each of these approaches, much of the heat added to the flue gas will be captured in the air preheater. Both alternatives do, however, present an additional economic impact.

SCR systems can generally be applied to all boiler sizes and types, although with existing boilers there may be problems with spacial limitations. All of the catalysts considered here for use in treating flue gas containing SO₂ and SO₃ are resistant to poisoning by these compounds. Long term tests of these catalysts in the presence of SO_x have shown very little or no decrease in activity or selectivity. Reactor size is proportional to flue gas flow rate, and this will determine the size and cost of the SCR system while the particulate concentration will determine the necessary catalyst/reactor combination.

The reactor itself is usually sized on the basis of a space velocity which is defined as the gas flow rate divided by the catalyst volume. A typical space velocity for a parallel flow system is about 6000 hr⁻¹ compared to 8000 hr⁻¹ for a moving bed or fixed, packed bed SCR system. The pressure drop through parallel flow systems is typically on the order of 100 mm H₂O, which is somewhat higher than moving or fixed, packed bed systems. The pressure drop is being reduced as this technology develops.

Parallel flow, moving bed and fixed, packed bed SCR systems are all capable of attaining the stringent level of NO_{χ} control. Greater than 90 percent NO_{χ} reduction is achieved at $NH_3:NO_{\chi}$ mole ratios of 1:1 on commercial systems applied to industrial boilers in Japan. All of these systems have been applied to a variety of oil-fired industrial boilers in Japan and appear to be viable techniques of attaining up to 90 percent NO_{χ} control.

There are some potential problems downstream of the FGT systems due to the presence of the unreacted ammonia in the flue gas. Two things can happen: 1) the NH_3 can react with SO_2 or SO_3 to form ammonium bisulfate or ammonium sulfate or 2) the NH₃ can enter the downstream equipment unreacted. The bisulfate has been shown to cause air preheater pluggage and this is the subject of ongoing research both at the EPA and the Electric Power Research Institute (EPRI). Both the bisulfate and sulfate exist as a particulate, but may be difficult to collect if the particles are submicron in size. Unreacted NH₃ is not likely to present any operational problems. A recent study has shown that if an ESP exists downstream, then most of the NH₃ will exit with the ash. NH₃ can actually improve the performance of an FGD system.¹⁶

1.2.2 Economic Impacts

The costs of NO_{χ} FGT systems applied to the industrial boilers are presented in this section. Two types of data are presented. First the capital and annual costs are shown as a function of boiler size. Then the cost effectiveness in terms of $k \log NO_{\chi}$ removed is evaluated. Tables 1.2.2-1 through 1.2.2-3 show the range of annual cost for the moderate to stringent level of control for the various boiler/size/control system combinations.

The annual costs in terms of \$/MBtu/hr are plotted against boiler size in Figures 1.2.2-1 through 1.2.2-4. In all cases, there is clearly an economy of scale with the larger units. An interesting result is that for the small residual oil-fired boiler, the parallel flow system is somewhat less expensive, but with the larger boiler, the moving bed system is less expensive. This is a result of the labor cost, which is a fixed cost, and is higher for moving bed systems than for parallel flow systems. Therefore, with small systems, the labor component has a significant effect on the annual cost of these systems. This result is the primary reason why it is not possible to choose a best system for residual oil applications. The effect of fuel type on annual cost is shown in Figure 1.2.2-5 when costs for the 44 MW, (150 MBtu/hr) boilers are compared for each fuel type. Systems applied to coal-fired boilers are the most expensive while those

Boiler	Size, MBtu/hr	Control	System	<u>Annual Cos</u> Moderate	t, \$1000/yr Stringent		
Underfeed Stoker	30	Parallel	Flow SCR	108	130		
Chaingrate	75	Paralle1	Flow SCR	153	197		
Spreader Stoker	150	Paralle1	Flow SCR	221	291		
Pulverized Coal	200	Parallel	Flow SCR	254	351		

TABLE 1.2.2-1.	ANNUAL COST	OF NO_{\times}	CONTROL	SYSTEMS	APPLIED
	TO COAL-FIRM	ED BOILI	ERS		

TABLE 1.2.2-2. ANNUAL COST OF NO_{\times} CONTROL SYSTEMS APPLIED TO OIL FIRED-BOILERS

Boiler	Size, MBtu/hr	Control System	<u>Annual Cos</u> Moderate	t, \$1000/yr Stringent
Distillate Oil	15	Fixed Packed Bed SCR	64	67
Distillate Oil	150	Fixed Packed Bed SCR	137	176
Residual Oil	30	Parallel Flow SCR	96	108
Residual Oil	30	Moving Bed SCR	120	130
Residual Oil	150	Parallel Flow SCR	181	223
Residual Oil	150	Moving Bed SCR	168	204

TABLE 1.2.2-3. ANNUAL COST OF NO $_{\times}$ CONTROL SYSTEMS APPLIED TO NATURAL GAS-FIRED BOILERS

Boiler	Size, MBtu/hr	Control System	<u>Annual Cos</u> Moderate	t, \$1000/yr Stringent
Package, Firetube	15	Fixed Packed Bed SCR	64.4	67.6
Package, Watertube	150	Fixed Packed Bed SCR	129	175



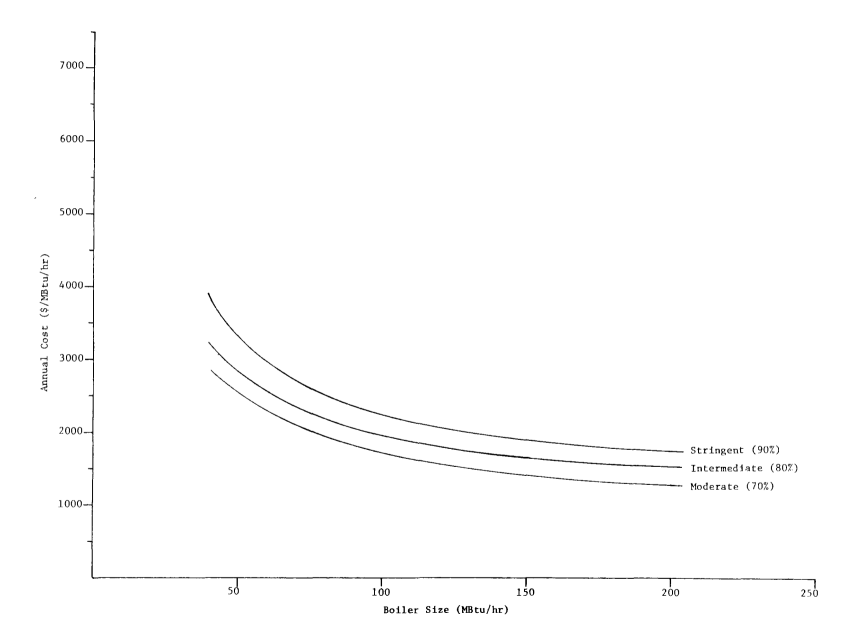


Figure 1.2.2-1. Annual cost of parallel flow SCR NO_{\times} FGT systems for coal-fired boilers.

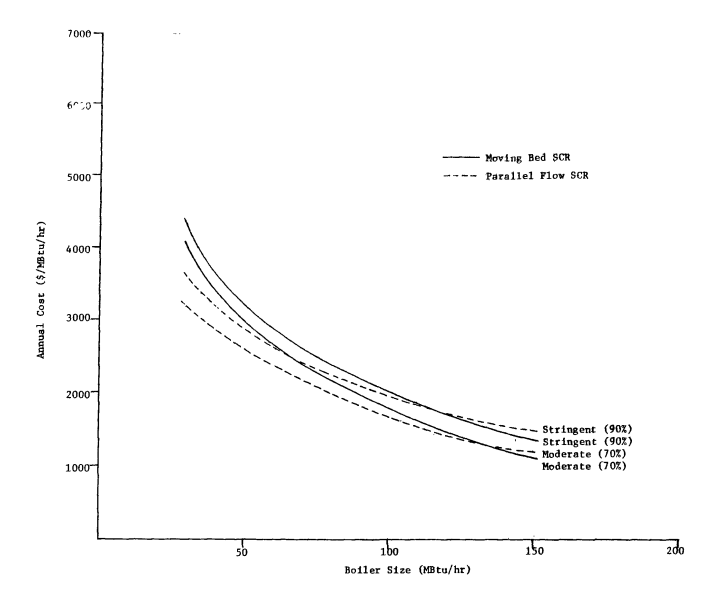


Figure 1.2.2-2. Annual cost comparison of NO_{χ} FGT systems for residual oil boilers.

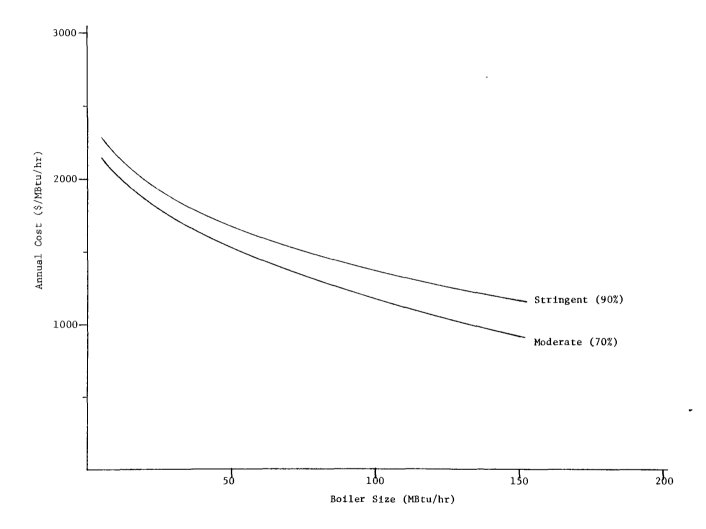


Figure 1.2.2-3. Annual cost of fixed packed bed SCR NO_{X} FGT systems for distillate oil boilers.

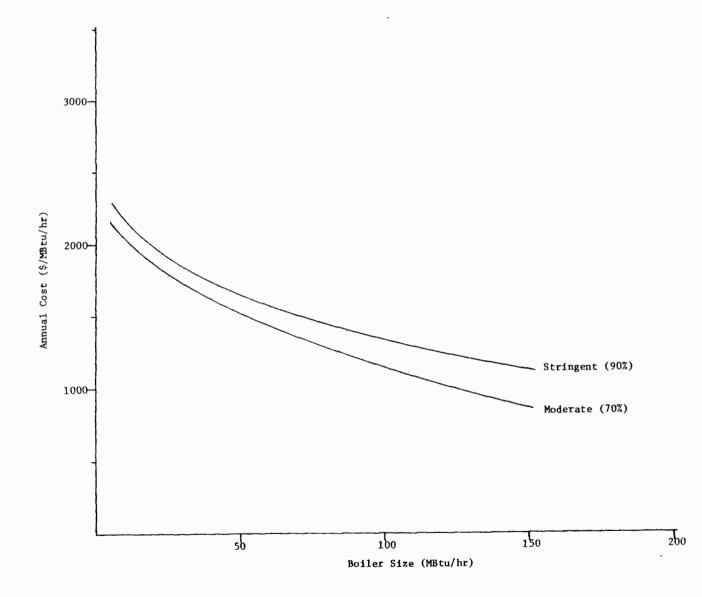


Figure 1.2.2-4. Annual cost of fixed packed bed SCR NO $_{\times}$ FGT systems for natural gas boilers.

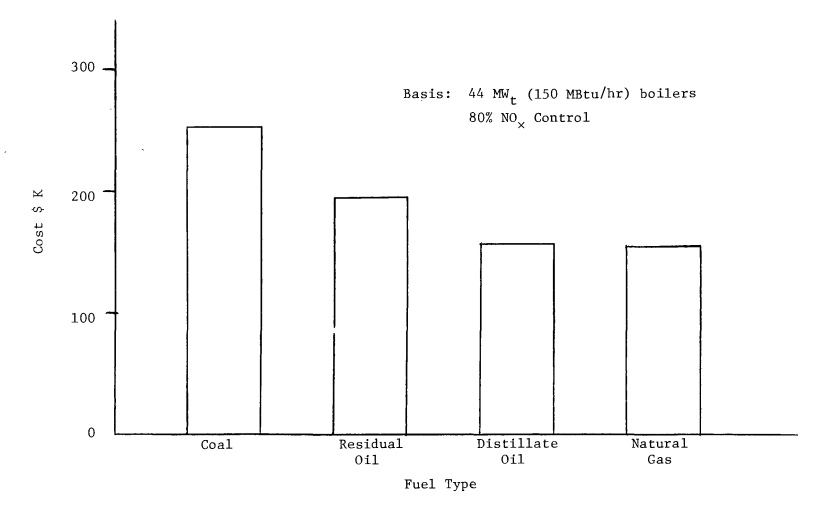


Figure 1.2.2-5. Comparison of annual costs of NO_{\times} FGT systems applied to 150 MBtu/hr boilers. (Average costs used where two systems apply.)

applied to distillate oil- and natural gas-fired boilers are the least expensive. Annual costs for residual oil-fired boilers lie in between these two extremes. The higher costs for systems which treat flue gas from coal-and residual oil-fired boilers are a result of the use of systems that will handle high particulate loadings, higher baseline NO_{χ} emissions with these fuels and, in the case of coal, higher flue gas flow rates.

Capital costs in terms of \$/MBtu/hr are presented as a function of boiler size in Figures 1.2.2-6 through 1.2.2-9. These figures also show larger systems to be less expensive in terms of cost per unit of capacity. This is due to the fact that the equipment costs used in this study were either constant for all sizes or varied exponentially with size.

In addition to determining the annual and capital costs, the study also examines the cost effectiveness of the various combinations. Cost effectiveness is defined as $\frac{1}{\text{kg NO}_{\times}}$ removed. Comparing the systems in this manner shows which combinations provide the largest environmental benefit for the lowest cost. Cost effectiveness if plotted against the level of NO_{\times} control in Figures 1.2.2-10 through 1.2.2-13.

For coal-fired boilers, the increased annual cost over an uncontrolled boiler for NO_x -only parallel flow SCR systems ranges approximately 6-12 percent, depending on the boiler and level of control. The figure plainly shows economy of scale as the largest coal-fired standard boiler, pulverized coal, has the most cost effective NO_x control system. Annual costs for the small boilers are labor cost-dominant, hence the maximum cost effectiveness at 90 percent NO_x control. The large boiler's costs are catalyst costdominant, hence the maximum cost effectiveness at 70 percent NO_x control (additional catalyst is required to remove the additional NO_x). Similar effects occur with the other fuels as well. In all cases it is apparent that the system size has a significantly larger effect on the cost effectiveness than does the control level.

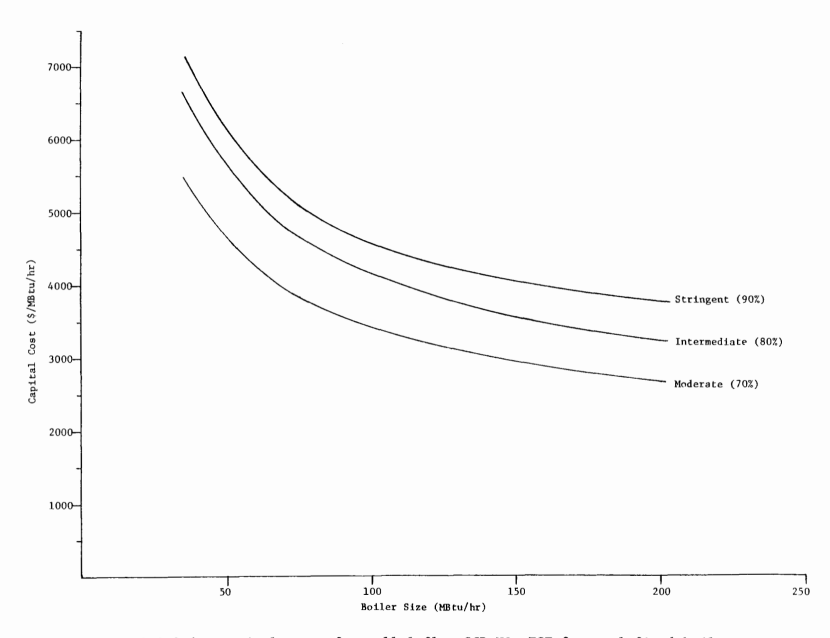


Figure 1.2.2-6. Capital cost of parallel flow SCR $\rm NO_{X}$ FGT for coal-fired boilers.

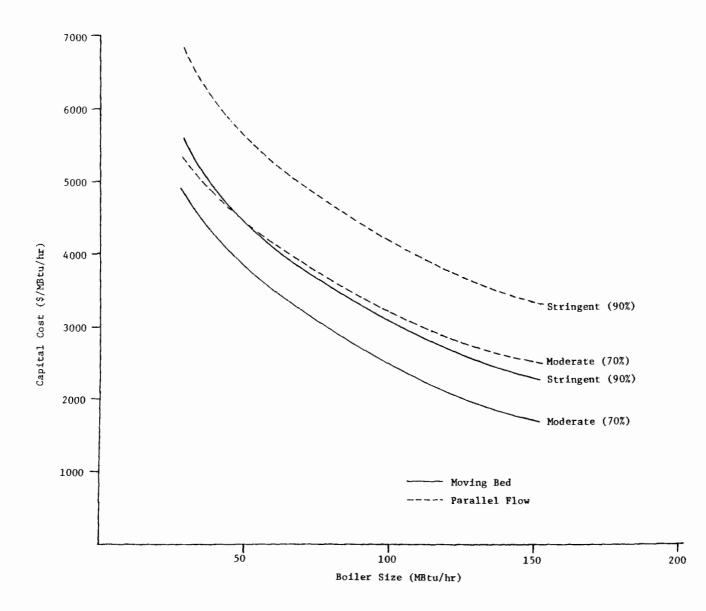


Figure 1.2.2-7. Capital cost comparison of NO_{\times} FGT systems for residual oil boilers.

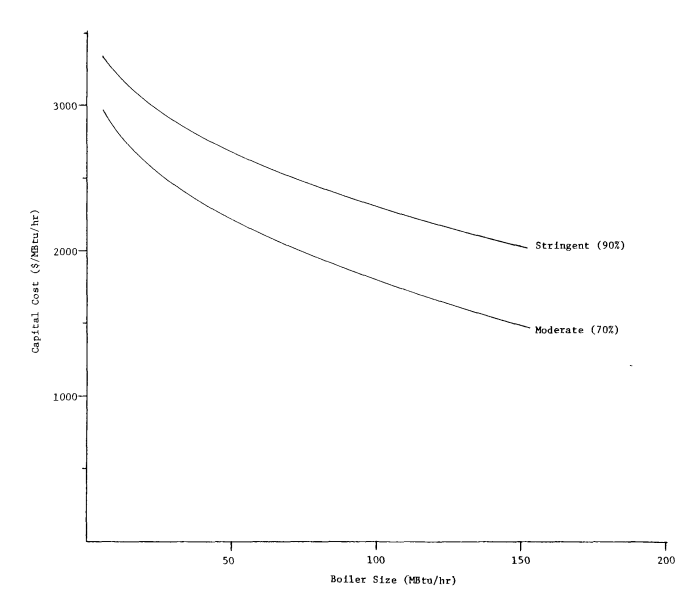


Figure 1.2.2-8. Capital cost of fixed packed bed SCR NO_{χ} FGT systems for distillate oil boilers.

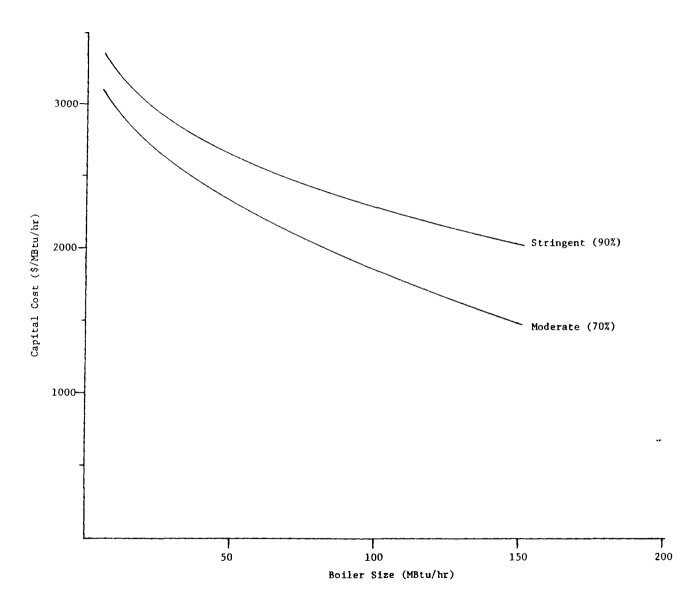


Figure 1.2.2-9. Capital cost of fixed packed bed SCR NO_{χ} FGT systems for natural gas boilers.

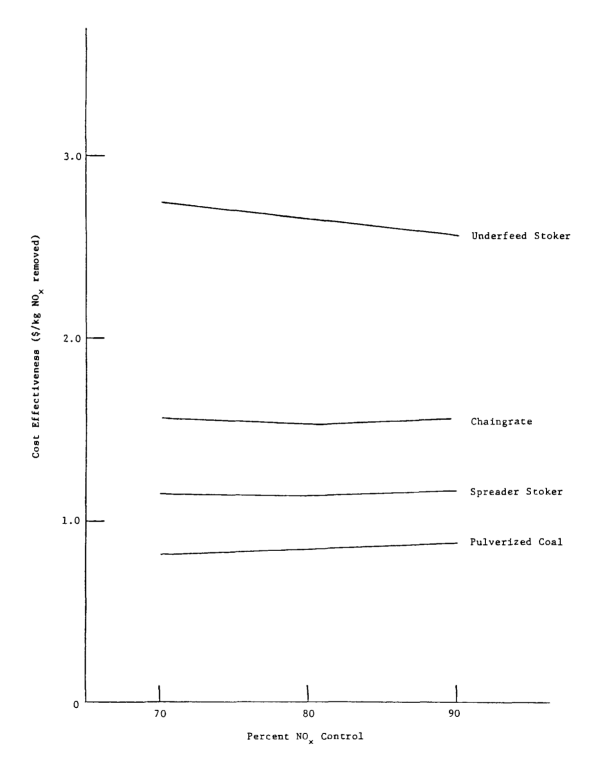
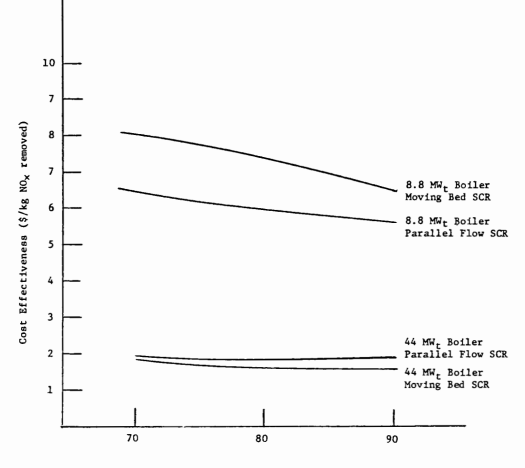
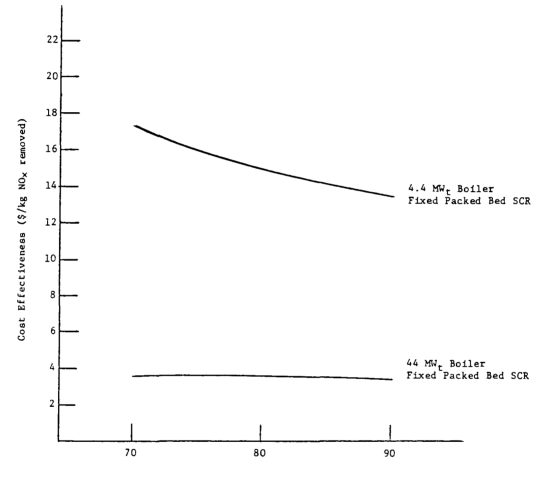


Figure 1.2.2-10. Cost effectiveness of parallel flow $\rm NO_{\times}$ control systems for coal-fired boilers.



Percent NO_X Control

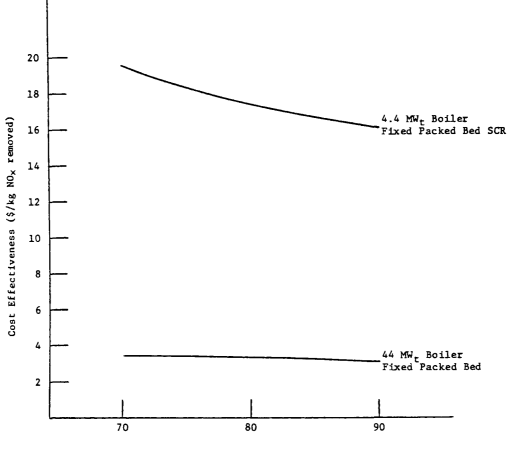
Figure 1.2.2-11. Cost effectiveness of FGT systems applied to residual oil-fired boilers.



Percent NO_X Control

Figure 1.2.2-12. C

Cost effectiveness of FGT systems applied to distillate oil-fired boilers.



Percent NO_X Control

Figure 1.2.2-13. Cost effectiveness of FGT systems applied to natural gas-fired boilers.

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The costs of SCR applications to modified or reconstructed facilities will be higher than those shown here. It is estimated that these costs will range from 25 to 120 percent more than applications to new boilers.

1.2.3 Energy Impacts

In calculating energy usage for each of the cases, all sources of energy consumption were considered. These sources are shown in Table 1.2.3-1.

NO _x FGT System	Energy Consumption Step (equipment)	Type of Energy Consumed
Parallel Flow SCR	Reactor Draft Loss (Fan) Liquid NH ₃ Transfer (Pump) NH ₃ Vaporization (Vaporizer) NH ₃ Dilution	Electrical Electrical Steam Steam
Moving Bed SCR	Reactor Draft Loss (Fan) Liquid NH ₃ Transfer (Pump) Catalyst Screening & Transfer (Elevator) Baghouse Draft Loss (Blower) NH ₃ Vaporization (Vaporizer) NH ₃ Dilution	Electrical Electrical Electrical Electrical Steam Steam
Fixed Packed Bed SCR	Reactor Draft (Fan) Liquid NH ₃ Transfer (Pump) NH ₃ Vaporization (Vaporizer) NH ₃ Dilution Soot Blowing-Distillate Oil Boiler Only	Electrical Electrical Steam Steam Steam

TABLE 1.2.3-1. AREAS OF ENERGY CONSUMPTION IN NO, FGT SYSTEMS

The energy impacts are presented in two forms. In the first, energy consumption in terms of MBtu/hr is plotted as a function of boiler size. These data are shown in Figures 1.2.3-1 through 1.2.3-4. Essentially, in all of the cases energy consumption is less than 1 MBtu/hr and represents a small amount of energy. The relative amount of energy consumed is shown in Figures 1.2.3-5 through 1.2.3-8 where usage is shown as a percent of the boiler heat input.

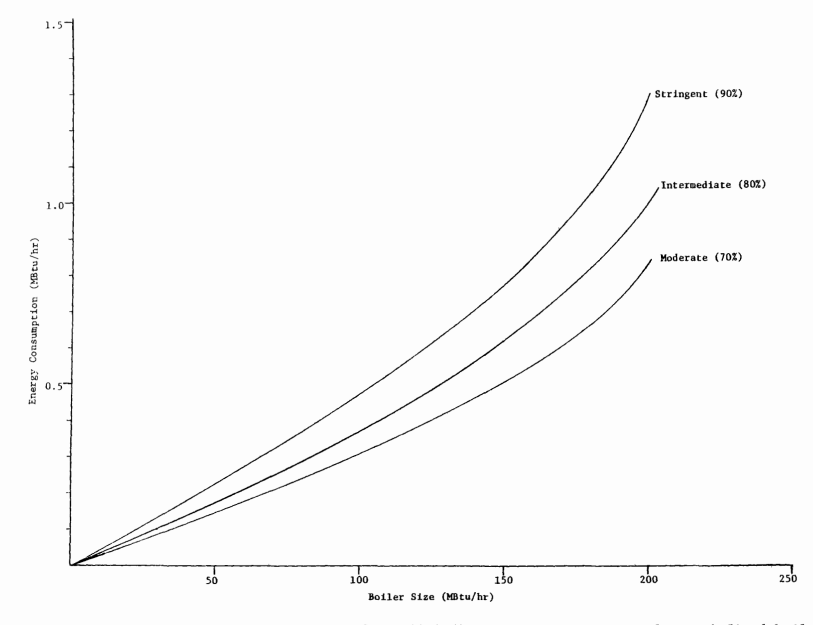


Figure 1.2.3-1. Energy consumption of parallel flow SCR NO_× FGT systems for coal-fired boilers.

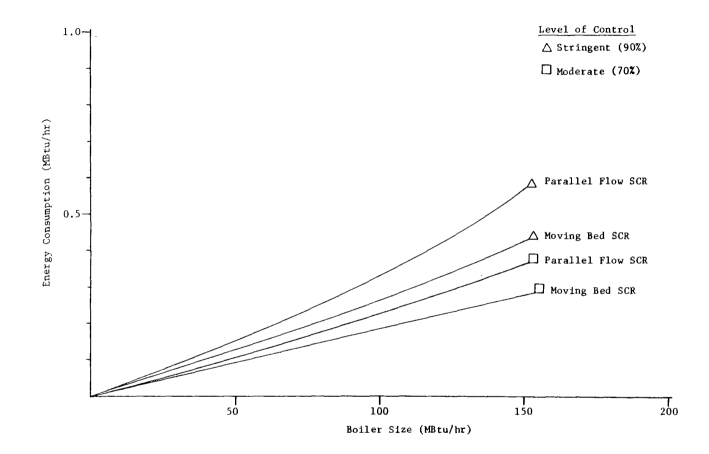


Figure 1.2.3-2. Energy consumption of NO_{\times} FGT systems for residual oil boilers.

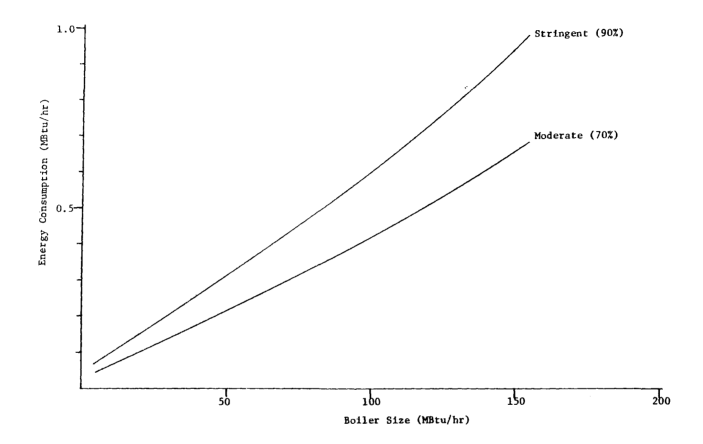


Figure 1.2.3-3. Energy consumption of fixed packed bed SCR NO_{χ} FGT systems for distillate oil boilers.



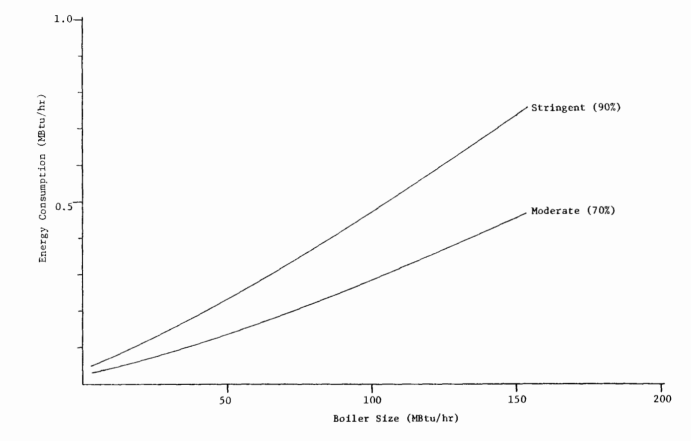


Figure 1.2.3-4. Energy consumption of fixed packed bed SCR NO_{χ} FGT systems for natural gas boilers.

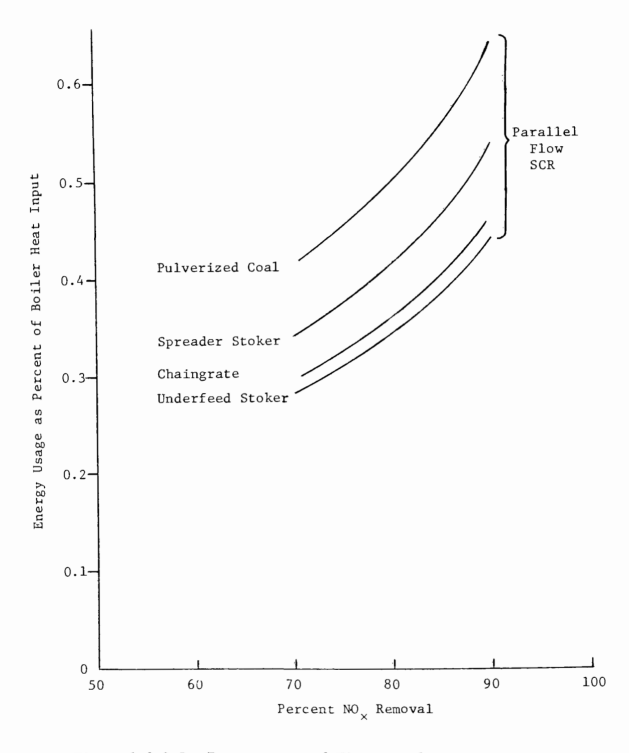


Figure 1.2.3-5. Energy usage of $\rm NO_X$ control systems as percent of boiler heat input. Coal-fired boilers.

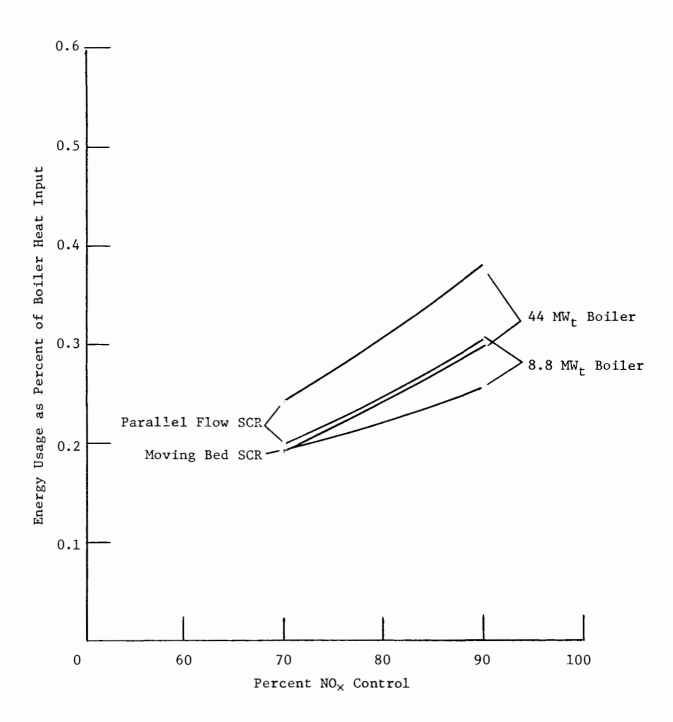
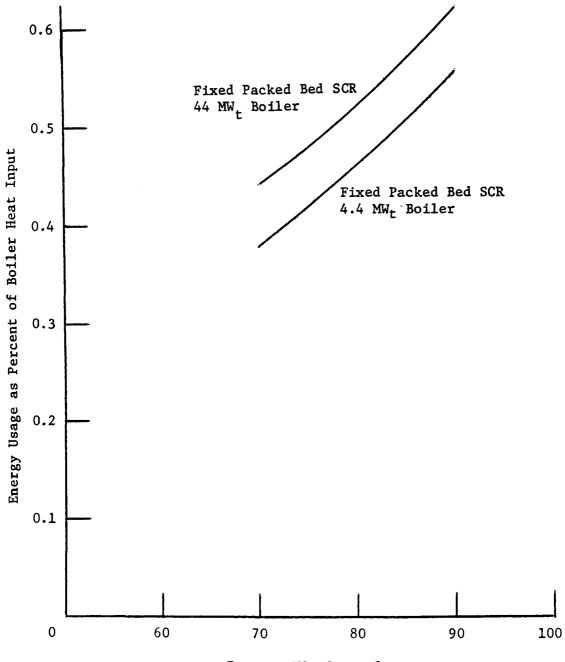


Figure 1.2.3-6. Energy usage of NO_× control systems as percent of boiler heat input. Residual oil-fired boilers.



Percent NO_× Control

Figure 1.2.3-7. Energy usage of NO_x control systems as a function of boiler heat input. Distillate oil-fired boilers.

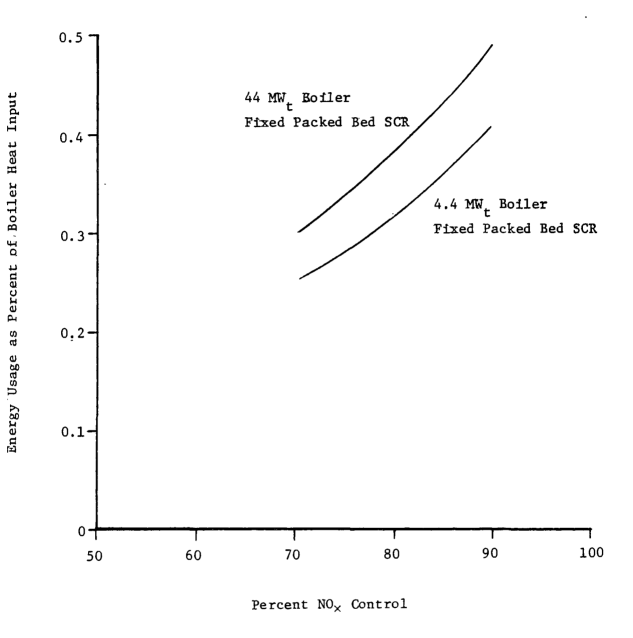


Figure 1.2.3-8. Energy usage of $\rm NO_{\times}$ control systems as percent of boiler heat input. Natural gas-fired boilers.

For the coal-fired boilers, the systems have a range of 0.27 to 0.64 percent. Energy usage is more a function of NO_{χ} control level than boiler size. In this analysis, energy usage is higher for larger boilers as a result of the reactor design method which allowed the reactor pressure drop to vary. Also, for the pulverized coal case, higher inlet NO_{χ} concentrations lead to higher energy usage.

With the oil-fired boilers, the parallel flow SCR systems have a range of 0.20 to 0.38 percent (from moderate to stringent level of control) and the moving bed systems from 0.19 to 0.29. One can see that the moving bed systems require less energy than the parallel flow systems. This is due to the greater pressure drop across a parallel flow reactor, which is larger than the ΔP across a moving bed reactor. Again, as in economics, the two candidate systems are considered to have similar energy impacts. For the distillate oil-fired boiler, the energy consumption ranged from 0.33 to 0.62 percent of the boiler heat input. For NO_X FGT applied to gas-fired boilers, the small fixed packed bed SCR system has a range of 0.27 to 0.42 percent, whereas the large system varies from 0.30 to 0.49 percent.

In all cases, energy usage was less than 0.64 percent of the boiler heat input, and in most cases it was less than half of this amount.

1.2.4 Environmental Impacts

There are some potential adverse environmental impacts of SCR systems. The use of NH₃ as the gaseous reducing agent introduces the possibility of ammonia emissions. The level of NH₃ emissions experienced by commercial SCR operations range from 1 to 10 ppm depending on the control level. Even at elevated NH₃:NO_× ratios (>1.0), the NH₃ emissions are reported to be less than 20 ppm. It is possible that NH₃ emissions will increase as the catalyst ages; however, commercial applications have not operated long enough to show this effect. Ten ppm of NH₃ may be an optimistic value, especially considering that currently there is no continuous monitoring technique for measuring

 NH_3 in the presence of SO_{\times} . The data, therefore, represent spot measurements and not continuous data. It seems reasonable to assume that 10 ppm represents a minimum level of NH_3 emissions. NH_3 emissions in terms of pounds per MBtu are presented in Figures 1.2.4-1 through 1.2.4-3 as a function of boiler size. While there is some variation, emission levels are essentially constant for all boiler sizes.

Another potential environmental problem is the formation of ammonium bisulfate, NH_4HSO_4 , or ammonium sulfate, $(NH_4)_2SO_4$. The presence of NH_3 , SO_3 , and H_2O in the hot flue gas leads to the formation of liquid NH_4HSO_4 upon cooling to approximately $180-220^{\circ}C$ by the following reaction.

$$NH_3(g) + SO_3(g) + H_2O(g) \neq NH_4HSO_4(1)$$
 (1-3)

This can create a plugging and corrosion problem in heat exchange equipment, particularly when medium- or high-sulfur fuels are fired. A beneficial effect is obtained by the tying up of SO_3 which is more hazardous than SO_2 and difficult to catch with FGD.¹² Further cooling to about 190°C precipitates the formation of solid ammonium sulfate by the following reaction.

$$NH_4HSO_4(1) + NH_3(g) \neq (NH_4)_2SO_4(s)$$
 (1-4)

It is speculated that minor, if any, amounts of these sulfates will be emitted to the atmosphere in situations where particulate control equipment exists downstream of the NO_{χ} control system. Sulfate formation is not a problem with gas-fired boilers since there is no sulfur present in the fuel.

Disposal of spent catalyst is the final environmental concern of the parallel flow SCR systems. Catalysts such as titanium dioxide (TiO_2) and vanadium pentoxide (V_2O_5) are probably recycled due to their high cost. This question is currently unanswered since all applications of this technology are very recent and none have yet required a catalyst change.

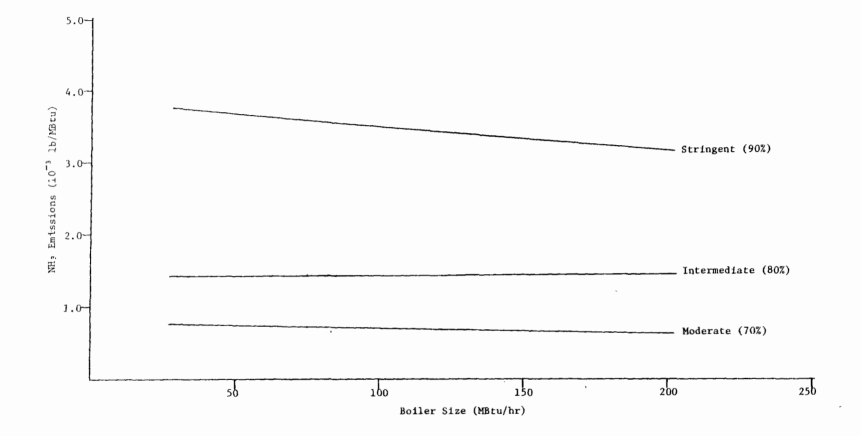


Figure 1.2.4-1. NH_3 emissions from SCR NO_{χ} FGT systems for coal-fired boilers.

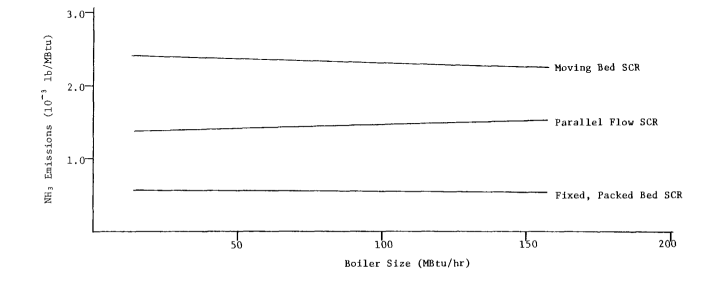


Figure 1.2.4-2. $\rm NH_3$ emissions from SCR $\rm NO_{\times}$ FGT systems for oil-fired boilers.

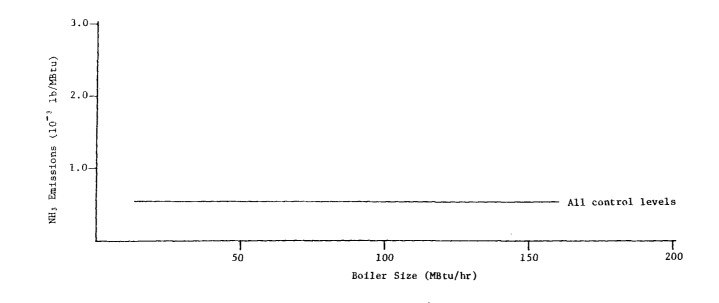


Figure 1.2.4-3. NH_3 emissions from SCR NO_{\times} FGT systems for natural gas-fired boilers.

Summarizing, FGT processes are relatively clean, possess minor potential air pollution and waste problems, and have no water, thermal, or noise pollution.

1.2.5 Development Status

Parallel flow SCR processes have been applied in Japan to several residual oil-fired industrial boilers. Oil-fired utility boilers and other sources with high particulate concentrations are also being treated. SCR processes have not yet been demonstrated commercially on coal-fired boilers. However, pilot units have been operated and some U.S. firms are offering SCR processes for use on coal-fired boilers. Two applications to coal-fired utility boilers are planned for 1980 (Table 1.2.5-1) although none exist at the present time. A coal-fired pilot unit demonstration of one parallel flow design is currently underway in the U.S. under EPA sponsor-ship¹⁴ and several have been conducted in Japan. The EPA facility should be operational by early 1980. Another U.S. demonstration of a NO_X -only SCR process will be performed in 1980 by the Electric Power Research Institute.

Location	User	Process Developer	Fuel	Capacity (Nm³/hr)	Completion Date
Takehara	Electric Power Development Co.	Has not been selected	Coal	800,000	July 1981
Tomato	Hokkaido Electric	Hitachi, Ltd.	Coal	280,000	October 1980

TABLE 1.2.5-1. PLANNED FGT INSTALLATIONS OF SCR COAL-FIRED UTILITY BOILERS¹³

Parallel flow and moving bed SCR processes have been applied in Japan to several oil-fired industrial and utility boilers. These operations are summarized in Tables 1.2.5-2 through 1.2.5-5. SCR systems are considered commercially available for oil-fired boilers at this time.

Location (Japan)	User	Process Developer	Fuel	Capacity (Nm³/hr)	Completion Date
Sodegaura	Fuji Oil	Mitsubishi H.I.	Resid	200,000	January 1978
Kawasaki	Ajinomoto	Ishikawajima H.I.	Resid	180,000	January 1978
Chiba	Ukishima Pet. Chem.	Mitsui Engineering	Resid	220,000	April 1978

TABLE 1.2.5-2. EXISTING FGT INSTALLATIONS OF SCR PARALLEL FLOW SYSTEMS OIL-FIRED INDUSTRIAL BOILERS¹⁷

TABLE 1.2.5-3. EXISTING FGT INSTALLATIONS OF SCR PARALLEL FLOW SYSTEMS OIL-FIRED UTILITY BOILERS¹⁸

Location (Japan)	User	Process Developer	Fuel	Capacity (Nm ³ /hr)	Completion Date
Yokosuka	Tokyo Electric	Mitsubishi H.I.	Resid	40,000	March 1977
Chita	Chubu Electric	Mitsubishi H.I.	Resid	1,920,000	February 1980
Kudamatsu	Chugoku Electric	Ishikawajima H.I.	Resid	1,900,000	July 1979
Niigata	Tohoku Electric	Ishikawajima H.I.	Resid	1,660,000	August 1981

Location (Japan)	User	Process Developer	Fuel	Capacity (Nm³/hr)	Completion Date
Kaizuka	Chiyoda Kenzai	Hitachi, Ltd.	Resid	15,000	October 1977
Amagasaki	Nippon Oils & Fats	Hitachi, Ltd.	Resid	20,000	April 1978
Sodegaura	Sumitomo Chemical	Mitsubishi H.I.	Resid	300,000	September 1976
Sodegaura	Sumitomo Chemical	Sumitomo Chemical, Mitsubishi H.I.	Resid	300,000	October 1976
Hirakatu	Kurabo	Kurabo	Resid	30,000	August 1975

TABLE 1.2.5-4. EXISTING FGT INSTALLATIONS OF SCR MOVING BED SYSTEMS OIL-FIRED INDUSTRIAL BOILERS¹⁸

Location (Japan)	User	Process Developer	Fuel	C apacity (Nm ³ /hr)	Completion Date
Amagasaki	Kansai Paint	Hitachi, Ltd.	Distillate	16,000	October 1977
Amagasaki	Nisshin Steel	Hitachi, Ltd.	Resid	20,000	August 1977
Amagasaki	Nisshin Steel	Hitachi, Ltd.	Resid	19,000	July 1977
Sakai	Nisshin Steel	Hitachi, Ltd.	Distillate	30,000	December 1978
Hokkaichi	Shindaikyowa P.C.	Hitachi Zosen	Resid	440,000	November 1975
Sodegaura	Sumitomo Chemical	Sumitomo Chemical Eng.	Resid	30,000	July 1973
Sodegaura	Sumitomo Chemical	Sumitomo Chemical Eng.	Resid	240,000	March 1976
Sorami	Toho Gas	Sumitomo Chemical Eng.	Distillate	62,000	October 1977
Sorami	Toho Gas	Sumitomo Chemical Eng.	Distillate	23,000	December 1977
Sorami	Toho Gas	Sumitomo Chemical Eng.	Distillate	23,000	June 1978
Sorami	Toho Gas	Sumitomo Chemical Eng.	Distillate	19,000	July 1978
Kawasaki	Nippon Yakin	Mitsubishi Kakoki	Resid	14,000	July 1978
Kawasaki	Toho Gas	Mitsubishi Kakoki	Distillate	30,000	October 1977
Chita	Toho Gas	Mitsubishi Kakoki	Distillate	30,000	October 1977

TABLE 1.2.5-5. EXISTING FGT INSTALLATIONS OF SCR FIXED BED SYSTEMS OIL-FIRED INDUSTRIAL BOILERS¹⁷

Table 1.2.5-6 shows the numerous industrial fixed packed bed SCR applications. Although gas-fired boilers, both industrial and utility, are numerous in Japan, few have been equipped with SCR units so far. This is due to the fact that less expensive NO_X reduction by combustion modifications on these boilers has been adequate to meet environmental regulations. Fixed packed bed SCR systems are considered to be commercially available for natural gas-fired boilers at this time.

Company	Site	Capacity (Nm ³ /hr)	Reactor type	Completion date			
Osaka Gas	Takaishi	15,000x2	FPB	December 1976			
Chubu Electric	Chita	1,910,000	FPB	April 1978			
Kyushu Electric	Kokura	1,610,000	FPB	July 1978			
Chubu Electric	Chita	1,910,000	FPB	September 1978			
Hyushu Electric	Kokura	1,610,000	FPB	December 1978			

TABLE 1.2.5-6. GAS-FIRED SCR PLANTS IN JAPAN¹⁷

1.3 FLUE GAS TREATMENT FOR CONTROL OF NO, AND SO,

Some FGT processes have the capability of removing SO_{\times} in addition to NO_{\times} . These processes are typically more complex and costly than those which remove just NO_{\times} ; however, this is offset by the simultaneous dual pollutant control capability. For this reason, these processes are considered separately from the NO_{\times} -only processes.

1.3.1 System Description

The following $\mathrm{NO}_{\times}/\mathrm{SO}_{\times}$ systems are considered for application to the coal-fired boilers:

- Parallel Flow SCR
- Adsorption
- Electron Beam Radiation
- Absorption-Reduction
- Oxidation-Absorption-Reduction
- Oxidation-Absorption.

Parallel flow SCR is selected as the only candidate for "best" NO_{χ}/SO_{χ} system. The choice here is a combination of no serious secondary environmental impacts, system performance, system reliability, and status of development. The process is described briefly in Table 1.3.1-1.

Process	Description
Parallel Flow SCR	Utilizes NH_3 to catalytically reduce NO_X after SO_X is adsorbed by and reacted with catalyst; capable of achieving >90 percent NO_X and SO_X reduction; SO_X saturated catalyst is regenerated while flue gas is diverted to alternate reactor.

TABLE 1.3.1-1. BEST NO_X/SO_X EMISSION CONTROL SYSTEM FOR COAL-FIRED BOILERS

The NO_x reduction reactions occurring in this process are the same as those described by reactions (1-1) and (1-2). The process utilizes an acceptor material to adsorb SO₂ and the product of the adsorption reaction then acts as an NO_x reduction catalyst. Elemental copper is converted to oxide form by flue gas oxygen.³

$$Cu(s) + {}^{1}_{2}O_{2}(g) + CuO(s)$$
 (1-5)

Sulfur dioxide reacts with the copper oxide, as described by:³

$$SO_2(g) + {}^{1}_{2}O_2(g) + CuO(s) + CuSO_4(s)$$
 (1-6)

 SO_3 in the flue gas is also removed:

$$SO_3(g) + CuO(s) \rightarrow CuSO_4(s) \tag{1-7}$$

It is this final copper sulfate $(CuSO_4)$ reaction product that acts as the primary catalyst for NO_X reduction by ammonia in the parallel flow SCR NO_X/SO_X system. After the spent reactor is isolated from flue gas flow, the reactor is purged with steam. A reducing gas, usually hydrogen, is then added which reacts with the copper sulfate in the following manner:⁴

$$CuSO_{4}(s) + 2H_{2}(g) \rightarrow Cu(s) + SO_{2}(g) + 2H_{2}O(g)$$
 (1-8)

The off-gas of this reaction is cooled to condense out the steam, reducing the gas volume and thus concentrating the SO_2 . The concentrated SO_2 is compressed and sent to a workup section to produce either elemental sulfur, liquid SO_2 , or sulfuric acid. (Sulfuric acid is produced in the cases studied in detail for this report.)

Besides the catalyst regeneration and sulfur treatment sections of the parallel flow SCR simultaneous NO_X/SO_X system, the NO_X -only and NO_X/SO_X parallel flow SCR systems are quite similar and the technical description contained in the previous section (1.2.1) applies here also.

No continuous coal-fired NO_x removal test data for NO_x/SO_x systems are available. Continuous oil-fired NO_x removal test data for the parallel flow $SCR NO_x/SO_x$ system have been obtained from a 40 MW_e unit in Japan. These data show the process capable of achieving up to 80 percent NO_x control (average = 65 percent) and 90 percent SO_x control. However, the process developers believe the process capable of achieving NO_x control levels of greater than 90 percent as a result of testing on the 40 MW_e unit. SO_2 removal results from coal-fired tests in the U.S. also show the process capable of attaining greater than 90 percent SO_2 reduction.

1.3.2 Economic Impacts

The annual and capital costs of this system applied to two coal-fired boilers and a residual oil-fired boiler are presented in this section. The costs are based on using a sulfuric acid plant for SO₂ workup and a compressor/gasholder for flow smoothing. Table 1.3.2-1 shows the annual costs of applications to coal-fired boilers. Two boiler types and two coals are presented and these data are plotted in Figure 1.3.2-1. Once again, economy of scale with large systems is evident; however, the effect is most significant for the high sulfur coal cases.

TABLE 1.3.2-1. ANNUAL COST OF PARALLEL FLOW NO_X/SO_X CONTROL SYSTEMS

Boiler	Fuel	Annual Cost, \$1000/yr
Pulverized Coal	High Sulfur Eastern Low Sulfur Western	1805 894
Underfeed Stoker Coal	High Sulfur Eastern Low Sulfur Western	711 462

Capital costs are shown in Figure 1.3.2-2 and the significant effect of boiler size on costs can again be seen.

Table 1.3.2-2 shows the annual cost of the single case studied. Only one case was analyzed for reasons described earlier and, as a result, it is not possible to plot the results or show trends.

Boiler	Fuel	Annual Cost, \$1000/yr	Capital Cost, \$1000	
Package, Watertube	Residual Oil	1092	3801	

TABLE 1.3.2-2. COSTS OF PARALLEL FLOW NO₂/SO₂ CONTROL SYSTEM

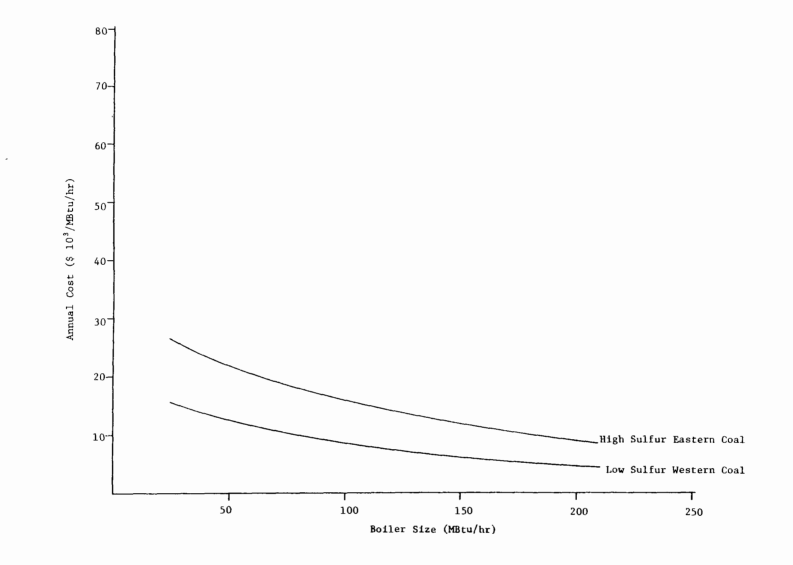


Figure 1.3.2-1. Annual cost of parallel flow SCR NO_X/SO_X FGT for coal-fired boilers at intermediate level of control.

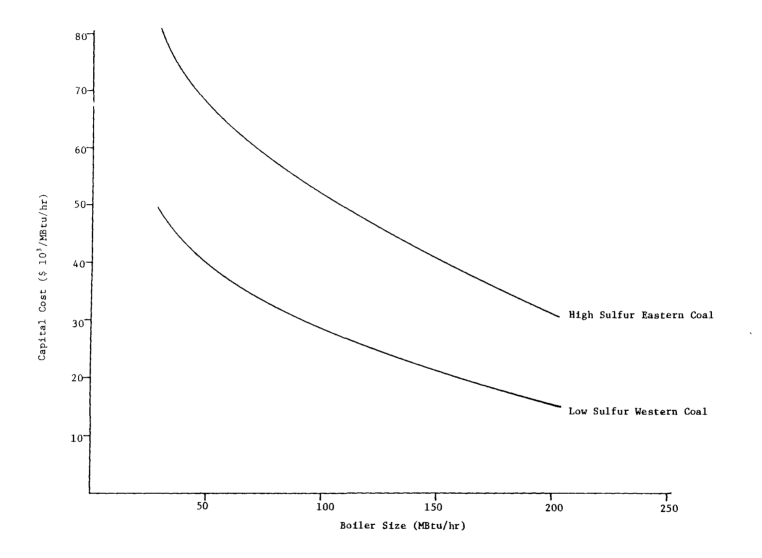


Figure 1.3.2-2. Capital cost of parallel flow SCR NO_X/SO_X FGT for coal-fired boilers at intermediate level of control.

1.3.3 Energy Impacts

Energy usage for these cases is summarized in Table 1.3.3-1 and plotted in Figure 1.3.3-1 where energy consumption is plotted against flue gas flow rate. The curves are essentially linear with the high sulfur case having a significantly greater impact. With high sulfur coal, the energy usage is 7.7 percent of the boiler heat input for both boiler types. With the low sulfur coal, this figure drops to 2.2 percent of the boiler heat input.

TABLE 1.3.3-1. ENERGY CONSUMPTION OF NO_{\times}/SO_{\times} CONTROL PROCESSES APPLIED TO COAL FIRED BOILERS

Boiler	Fuel	Energy Consumption, MW _t (MBtu/hr)
Pulverized Coal	High Sulfur Eastern Low Sulfur Western	$\begin{array}{c} 4.5 & (15) \\ 1.2 & (4.1) \end{array}$
Underfeed Stoker	High Sulfur Eastern Low Sulfur Western	0.68 (2.3) 0.20 (0.68)

Energy use for the oil-fired boiler is shown in Table 1.3.3-2. Here again, it is not possible to plot the result.

TABLE 1.3.3-2. ENERGY CONSUMPTION OF PARALLEL FLOW NO×/SO× CONTROL SYSTEM

		Energy Consumption				
Boiler	Fuel	MWt	(MBtu/hr)	% of Boiler Heat Input		
Package, Watertube	Residual Oil	2.0	(6.6)	4.4		

1.3.4 Environmental Impacts

The environmental impacts of this NO_{\times}/SO_{\times} process are similar to those of the NO_{\times} -only processes. The primary adverse environmental impact is from NH_3 emissions. The process developers claim that these emissions are low

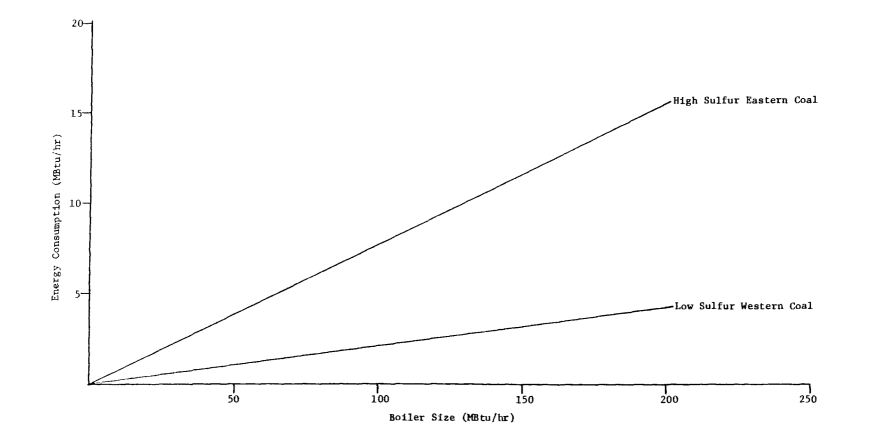


Figure 1.3.3-1. Energy consumption of parallel flow SCR NO_x/SO_x FGT systems for coal-fired boilers.

(<10 ppm); however, a continuous NH_3 monitoring method for use with gases containing sulfur oxides will be necessary before NH_3 emissions can be accurately quantified. The potential problems with ammonium bisulfate and sulfate formation should be much less with the NO_X/SO_X process since much of the SO_X has been removed from the flue gas.

1.3.5 Development Status

The integrated process has been tested on oil but not coal; however, the NO_x and SO_x removal capabilities have been demonstrated separately. The SO_2 capabilities have been demonstrated on a pilot unit treating coalfired flue gas. An EPA-sponsored U.S. test of the integrated process on flue gas from a coal-fired boiler is scheduled for 1980. Pilot and demonstration units of Shell/UOP process are summarized in Table 1.3.5-1 and commercial applications are summarized in Table 1.3.5-2.

Location/ Company	Designed By	Fuel/ Application	Size, Nm ³ /hr	Type of Operation	Dates
Shell Ref. at Pernis	Shell	Residual Fuel Oil- Proc. Heater	600-1000	SO _x -only	1967-1972
Rotterdam Utility	Shell	Coal- Steam Boiler	-	Heavy Fly Ash Loading	1971
Tampa Elec. Big Bend	UOP	Coal- Wet-Bottom Utility Boiler	1200-2000	SO _x -only SO _x -NO _x Simultaneous	1 974-197 6
JGC Yokohama Yokohama	JGC	Fuel 0il	250-700	NO _x -only	1974-
Nippon Steel	JGC	Sintering Furnace	2000	NO _x -only	1976-1978
	JGC	Coke Oven	400	NO _× -only	1976-1977

Unit	Designed By	Fuel/ Application	Size, Nm ³ /hr	Type of Operation	Dates
SYS* Yokkaichi	Shell	Residual Fuel Oil- Ref. Boiler	125,000	SO _x -only; NO _x -SO _x Simultaneous	1973–1975 1975–
Kashima Oil Co. Ltd.	JGC	Fuel Oil- Process Unit Heater	50,000	NO _x -only	1975-
Fuji Oil Co. Ltd.	JGC	CO Boiler	70,000	NO _× -only	1976-
Nippon Steel Corp.	JGC	Sintering Furnace	150,000	NO _x -only	1978-

TABLE 1.3.5-2. SHELL/UOP PROCESS COMMERCIAL APPLICATIONS

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SECTION 2

EMISSION CONTROL TECHNIQUES

This section presents descriptions of all control techniques for NO_{\times} control by flue gas treatment (FGT). Each control technique is described separately, however, there may be several vendors offering processes that are similar. Where this occurs, an effort has been made to generalize the various processes into a single technique within a single category. This is usually, but not always, possible. Where significant differences exist, they are discussed separately.

A distinction has been made between those processes which remove only NO_{\times} and those which remove both NO_{\times} and SO_2 . This is necessary because when final process comparisons are made it will be necessary to compare the cost of a NO_{\times} only process <u>plus</u> an FGD system versus the cost of a NO_{\times}/SO_{\times} process. In the subsections which follow, all NO_{\times} only processes are grouped together and presented first and the NO_{\times}/SO_{\times} process are presented second.

Economics for the various NO_× control processes are presented only for comparison and use in Section III for process selection. These economic figures do not necessarily represent costs for application of these systems to industrial boilers, in fact, most were developed with utility applications in mind. However, at this time they are the only published economic data available. Detailed cost estimates for several systems as applied to industrial boilers will be developed for this study in Section IV.

2.1 Principles of Control

The FGT systems are examined on the basis of application to industrial boilers. These boilers are generally smaller than those used for utility applications and produce steam for purposes such as electrical power generation, process heating and space heating. They range in size from small package units to large field erected units. The demand on the boilers may be constant, such as with process heating, or cyclic, such as with space heating.¹ Industrial boilers generally have fewer burners than utility boilers and, therefore, taking just one burner out of service can have a significant effect on the flue gas characteristics.² Also, the stoker units typically run with higher excess air. These characteristics of industrial boilers indicate that typical flue gases can have a wide variety of characteristics.

This study considers seven standard boilers as selected for a variety of reasons in a separate study.³ These boilers are described in Table 2.1-1. Three coals, two oils and natural gas are included as well as four sizes of coal-fired boilers. The coals considered are low sulfur western (0.6%S), low sulfur eastern (0.9%S) and high sulfur eastern (3.5%S). The two oils are distillate oil (#2) and residual oil (#6).

 NO_{χ} is formed in boilers by two mechanisms. In one mechanism, thermal fixation, N_2 and O_2 present in the combustion air react to form NO. This reaction requires the high temperatures that are present in the burner flame and is dependent also on the O_2 concentration in the flame. The reaction does not reach equilibrium and therefore the amount of NO_{χ} formed by this mechanism is governed by reaction kinetics.⁴ The second mechanism, fuel nitrogen conversion, involves the reaction of nitrogen contained in the molecular structure of the fuel with O_2 in the combustion air. The rate of reaction is a function of fuel nitrogen conversion and O_2 concentration. A more detailed description of the NO_{χ} formation mechanisms is contained in the Technology Assessment Report on NO_{χ} control by combustion modifications.

Boiler	Fuel*	Size ^{MW} t	Control Level %
	NO _X -Only FGT Systems		
Package, Firetube	Distillate Oil	4.4	70, 90
Package, Firetube	Natural Gas	4.4	70, 90
Package, Watertube	Residual Oil	8.8	70, 90
Package, Watertube Underfeed Stoker	LSŴ	8.8	80
Package, Watertube Chaingrate Stoker	ĽSW	22	70, 80, 90
Package, Watertube	Natural Gas	44	70, 90
Package, Watertube	Distillate Oil	44	70, 90
Package, Watertube	Residual Oil	44	70, 90
Field Erected, Watertube Spreader Stoker	LSW	44	80
Field Erected, Watertube Pulverized Coal	LSW	58.6	70, 90
	NO _X /SO _X FGT Systems		
Package, Watertube	Residual Oil	44	80 NO_{\times} , 85 SO_{\times}
Package, Watertube Underfeed Stoker	HSE LSW	8.8	80 NO _X , 85 SO _X
Field Erected, Watertube Pulverized Coal	HSE LSW	58.6	80 NO _× , 85 SO _×

TABLE 2.1-1.CHARACTERISTICS OF THE STANDARD BOILERS CONSIDERED
FOR ANALYSIS IN THIS REPORT

*HSE = High Sulfur Eastern Coal (3.5% S)
LSW = Low Sulfur Western Coal (0.6% S)

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The NO_{\times} emissions for the various coals considered are different, presumably due to different fuel nitrogen concentrations. However, the emissions from the stoker boilers, on a ppm and mass per energy input basis, do not change from boiler to boiler. The mass rates do change due to differences in the flue gas flow rates for the various boilers. Emission rates for the standard boilers are shown in Table 2.1-2. The emission rates are based on AP-42 calculations.

In the sections which follow Section II, it is shown that NO_{χ} FGT system designs are not significantly affected by NO_{χ} concentration. The most significant design variables are flue gas flow rate and control level. For this reason, it is possible to generate information over the entire boiler size range while considering only one coal type. The coal chosen for analysis is low sulfur western since this coal has both the highest flue gas flow rates and NO_{χ} emissions.

FGT systems utilize either a gas phase reaction or liquid absorption to treat the flue gas. In most cases the gas phase reaction is between NO_{χ} and NH_3 in the presence of a solid phase catalyst. The catalyst is contained within a reactor and may be either fixed or moving bed. The NO_{χ} is converted to N_2 which exits with the flue gas.

Systems utilizing a liquid absorption technique contact flue gas and absorbent in conventional scrubbers. The absorbed NO_{χ} either remains in the scrubbing liquor and is treated in the liquid phase or reacts with a solute to form N_2 which degasses and leaves with the flue gas.

The NO_{\times} FGT systems discussed in the following subsections are divided into two categories. Those which remove only NO_{\times} are presented first and the simultaneous NO_{\times}/SO_{\times} processes are discussed second. The distinction is made since the two process types cannot be accurately compared unless FGD flue gas desulfurization (FGD) is included with the NO_{\times} -only processes. This comparison will be made, but only in the Comprehensive Technology Assessment

			N	IO× Emissio	ns	
Boiler	Fuel*	g/s	(1b/hr)	g/MJ	(1b/10 ⁶ Btu)	ррт
Package, Firetube	Distillate Oil	0.300	(2.38)	0.0688	(0.16)	97
Package, Firetube	Natural Gas	0.332	(2.63)	0.0774	(0.18)	104
Package, Watertube	Residual Oil	2.02	(16.0)	0.228	(0.53)	373
Package, Watertube Underfeed Stoker	HSE LSE LSW	2.40 2.06 2.95	(19.05) (16.35) (23.40)	0.275 0.237 0.335	(0.64) (0.55) (0.78)	335 288 402
Package, Watertube Chaingrate	HSE LSE LSW	6.02 5.15 7.40	(47.70) (40.80) (58.65)	0.275 0.232 0.335	(0.64) (0.54) (0.78)	336 290 401
Package, Watertube	Natural Gas	3.31	(26.26)	0.0753	(0.175)	110
Package, Watertube	Distillate Oil	2.99	(23.76)	0.0680	(0.158)	101
Package, Watertube	Residual Oil	7.47	(60.00)	0.172	(0.40)	292
Field Erected, Watertube Spreader Stoker	HSE LSE LSW	12.0 10.3 14.8	(95.40) (81.45) (117.15)	0.275 0.232 0.335	(0.64) (0.54) (0.78)	337 288 400
Field Erected, Watertube Pulverized Coal	HSE LSE LSW	19.2 16.5 23.7	(152.46) (130.50) (187.56)	0.327 0.280 0.404	(0.76) (0.65) (0.94)	466 396 550

TABLE 2.1-2. NO $_{\times}$ EMISSION RATES FOR THE STANDARD BOILERS

*Coal types: HSE = High Sulfur Eastern (3.5%S) LSE = Low Sulfur Eastern (0.9%S) LSW = Low Sulfur Western (0.6%S) Report (CTAR) which follows completion of the Individual Technology Assessment Reports (ITAR's). Therefore, in Section III of this ITAR, NO_{\times} only processes will only be compared with other NO_{\times} -only processes and $NO_{\times}/$ SO_{\times} processes will only be compared with other NO_{\times}/SO_{\times} processes. This distinction will be maintained throughout the other sections of the ITAR also. The NO_{\times} -only processes described are:

- Fixed Packed Bed Selective Catalytic Reduction (SCR)
- Moving Bed SCR
- Parallel Flow SCR
- Absorption-Oxidation

The NO_{χ}/SO_{χ} processes described are:

- Parallel Flow SCR
- Adsorption
- Electron Beam Radiation
- Absorption-Reduction
- Oxidation-Absorption-Reduction
- Oxidation-Absorption

2.2 CONTROLS FOR COAL-FIRED BOILERS

2.2.1 Selective Catalytic Reduction-Fixed Packed Bed Reactors

Fixed packed bed systems for selective catalytic reduction of NO_x are applicable only to flue gas streams containing particulate emissions of less than 20 mg/Nm³. Particulate emissions for all coals are higher, on the order of 1-5 grams per Nm³. Although it is possible to install a hot ESP to reduce the particulate level to 20 mg/Nm³ this is expensive and not always effective. For these reasons fixed packed bed SCR systems are not considered for application to coal-fired boilers by process vendors.⁵

2.2.2 Selective Catalytic Reduction-Moving Bed Reactors

Moving bed systems for selective catalytic reduction of NO_{\times} are applicable only to flue gas streams containing less than 1 g/Nm³. Particulate emissions for all coals are higher, on the order of 1-5 grams per Nm³. Although it is possible to install a hot ESP to reduce the particulate level to 1 g/Nm³ this is expensive and not always effective. For these reasons moving bed SCR systems are not considered for application to coal-fired boilers in this report.

2.2.3 Selective Catalytic Reduction-Parallel Flow Reactor

2.2.3.1 System Description--

The distinguishing aspect of this process is the catalyst shape which is produced in a variety of shapes. The catalysts are produced in either a honeycomb, pipe, or plate shape. Both metal and ceramic supports are employed. Several shapes are illustrated in Figure 2.2.3-1. The catalyst shapes allow particulate laden flue gas to pass through the reactor with no inertial impaction of the particles while the NO_× is transported to the catalyst surfaces by basic diffusion. The catalysts can handle all of the particulate levels emitted by the standard boilers. All of the catalysts considered here for use in treating flue gas containing SO₂ and SO₃ are resistant to poisoning by these compounds. Long term tests of these catalysts in the presence of SO_× have shown very little or no decrease in activity or selectivity.

The reactors used are similar to standard fixed packed bed units and an example is shown in Figure 2.2.3-2. The catalyst is usually prepared in small modules and manually stacked within the reactor. The specific arrangement will depend on the particular process under consideration.

A typical flow diagram for a parallel flow SCR system is shown in Figure 2.2.3-3. The arrangement is similar to the other SCR processes in that hot flue gas leaving the boiler economizer is injected with NH_3 and

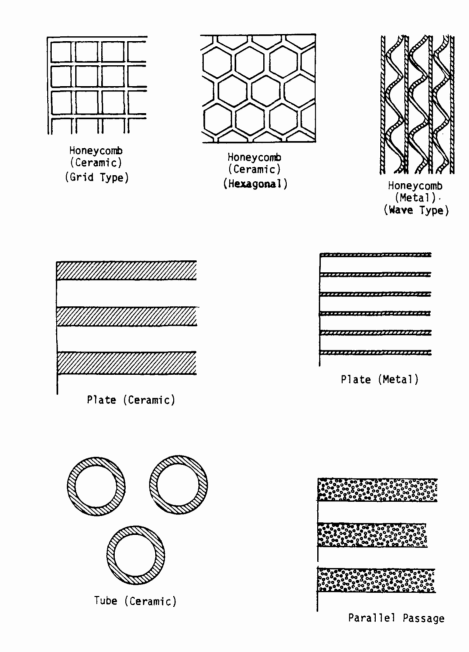


Figure 2.2.3-1. Shapes of parallel flow catalysts.²²

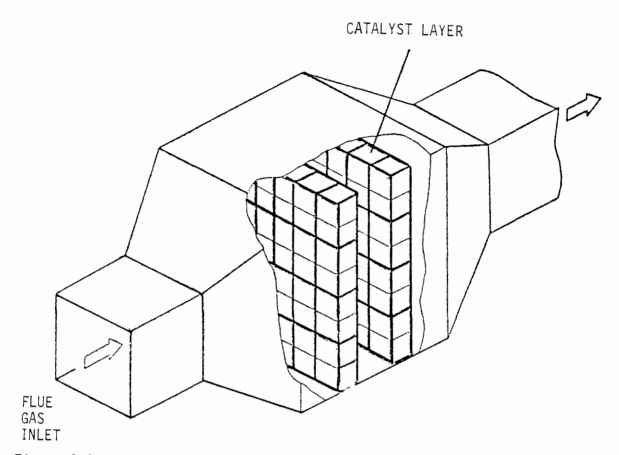


Figure 2.2.3-2. Typical reactor used with parallel flow SCR process.²³

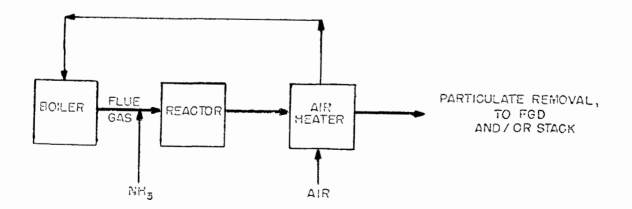


Figure 2.2.3-3. Flow diagram for parallel flow SCR process.²⁴

passed through a catalyst bed. Temperature control is important and can be accomplished with either a fired heater or an economizer bypass. NH_3 can be controlled using boiler operating condition inputs to conventional control components.

Within the reactor, NO_{\times} reacts with NH_3 to form N_2 and H_2O according to the following reactions. 12

$$4NH + 4NH_3 + O_2 \neq 4N_2 + 6H_2O$$
 (2-1)

$$2NO_2 + 4NH_3 + O_2 \neq 3N_2 + 6H_2O$$
 (2-2)

Reaction (2-1) is the primary reaction since flue gas NO_{\times} is typically 90-95 percent NO. O_2 is necessary for both reactions and is present in sufficient quantities (>3 percent) in all of the flue gases from the standard boilers.

The catalyst volume for a desired NO_{\times} removal can be determined by the fundamental design equation for a plug flow reactor.¹³

$$\frac{V}{F} = \int_{0}^{x} \frac{dx}{r}$$
(2-3)

The reaction rate, r, can be expressed as

$$r = k[NH_3]^a [NO]^b [O_2]^c$$
 (2-4)

The variables presented here have the same definitions as those presented in equations 2-3 and 2-4 of Section 2.3.2. Catalyst volume can also be determined by knowing the space velocity for a given catalyst and NO conversion level. The space velocity is defined as the flue gas flow rate divided by the catalyst volume. The reaction rate is different for each catalyst formulation since different catalysts will lower the activation energy by different amounts. The activation energy affects the reaction rate constant, k, according to the Arrhenius equation.

$$k = Ae^{-\frac{E}{RT}}$$
(2-5)

Example values of k, a, b, and c for two catalyst formulations are shown in Table 2.2.3-1.

An important design variable with catalytic systems is the space velocity which expresses the volume of catalyst required to treat one volume per hour of flue gas. Space velocity varies with catalyst formulation, catalyst shape, and control level. Typical values of space velocity for various catalyst shapes are shown in Table 2.2.3-2. Also shown are other catalyst design variables such as catalyst dimensions, gas velocities, bed depth and pressure drop. Ranges of values are used since specific values are different for each catalyst. The values shown pertain to 90 percent NO_{χ} removal and an NH_3/NO_{χ} mole ratio of 1:1.

Both NH_3/NO_{χ} ratio and space velocity will change with removal level. The NH_3/NO_{χ} mole ratio will range from 0.7-1.0 and the space velocity will range approximately as shown in the table for control levels of 70 to 90 percent.¹⁵

Variables associated with the boiler can also affect the performance of these systems. These are

- flue gas flow rate
- NO_{χ} concentration
- boiler load variability

TABLE 2.2.3-1. REACTION RATE DATA FOR TWO CATALYST FORMULATIONS¹¹ Catalyst: V₂O₅ on Al₂O₃ $k = 2.05 \times 10^{3} e^{-\frac{9650}{RT}}$ a = 0.30b = 0.22c = 0.05Catalyst: Fe-Cr on Al₂O₃ $k = 3.25 \times 10^{3} e^{-\frac{10,860}{RT}}$ a = 0.45b = 0.10c = 0.15

TABLE 2.2.3-2. CATALYST DESIGN VARIABLES FOR VARIOUS CATALYST SHAPES²⁵ (Basis: 90% NO_{\times} removal at NH₃/NO_{\times} ratio of 1:1, 350-400°C)

	Honeycomb Honeycomb,		Parallel Plate		
	(metallic)	tube (ceramic)	(Ceramic)	(Metallic)	
Catalyst size (mm)					
Thickness	1	2.3-5	8-10	1	
Opening	4-8	6-20	8-14	5-10	
Gas velocity (m/sec) ^a	2-6	5-10	5-10	4-8	
Bed depth (m)	1-2	1.5-5	4-6	2-5	
$SV (1,000 hr^{-1})^{b}$	5-8	4-8	1.5-3	2-4	
Pressure drop ($mmH_2^{\circ}0$)	40-80	40-160	80-160	60-120	

^aVelocity at 350-400°C in open column (superficial velocity).

^bGas volume $(Nm^{3}/hr)/catalyst$ bed volume (m^{3}) .

The flue gas flow rate and control level determine the catalyst volume (hence reactor size) necessary. Increases in either also increase the reactor size. The NO_{\times} concentration is primarily a function of fuel type used in the standard boilers. Higher concentrations require larger NH_3 storage and vaporization equipment; reactor size is not affected. Boiler load can affect several things including flue gas temperature, flow rate, and NO_{\times} concentration. It is necessary to maintain reaction temperatures of 350 to 400°C. Temperature control equipment may be necessary to accomodate large boiler load variations which may lower the flue gas temperature. Where these variations are present, some equipment overdesign may be warranted to insure a constant control level. These variables are discussed in more detail in the section on moving bed SCR systems for coal-fired boilers, Section 2.2.2.

Parallel flow SCR processes have been applied in Japan to several residual oil-fired industrial boilers. Oil-fired utility boilers and other sources with high particulate concentrations are also being treated. Two applications to coal-fired utility boilers are planned for 1980 (Table 2.2.3-3) although none exist at the present time. A coal-fired pilot unit demonstration of one NO_-only parallel flow design is currently underway in the U.S. under EPA sponsorship and several have been conducted in Japan. The EPA facility should be operational by mid-1979. Also, a parallel flow pilot system will be applied to flue gas from a coal-fired boiler in a study sponsored by the Electric Power Research Institute (EPRI). The unit is expected to be operational by 1980. A list of vendors of parallel flow SCR systems is presented in Table 2.2.3-4. The number of pilot unit demonstrations indicates that application of parallel flow SCR processes to coal-fired industrial boilers is feasible.

2.2.3.2 System Performance--

Performance data based on pilot plant testing were not found in the literature, however, data do exist for oil-fired applications. Since many of the flue gas characteristics are similar for oil and coal-fired boilers,

Location	User	Process Developer	Fuel	Capacity (Nm ³ /hr)	Completion Date
Takehara	Electric Power C.	Not yet announced	Coal	800,000	July 1981
Tomato	Hokkaido Electric	Hitachi, Ltd.	Coal	88,000	October 1980

TABLE 2.2.3-3. PLANNED FGT INSTALLATIONS OF SCR COAL-FIRED UTILITY BOILERS²⁶

TABLE 2.2.3-4. PROCESS VENDORS OF PARALLEL FLOW SCR PROCESSES²⁸

	Demonstrated on Coal		
Vendor	Yes/No	Scale	
Hitachi Zosen	yes .	pilot	
Hitachi, Ltd.	yes	pilot	
Japan Gasoline Corp.	no		
Mitsui Engineering & Shipbuilding	no		
Mitsubishi Heavy Industries	yes	pilot	
Ishikawajima-Harima Heavy Industries	yes	pilot	
Kobe Steel	no		
Kawasaki Heavy Industries	yes	pilot	
Shell/UOP	by 1979	pilot	

it is expected that the FGT performance will be roughly similar. Detailed data on oil-fired applications are contained in Section 2.3.

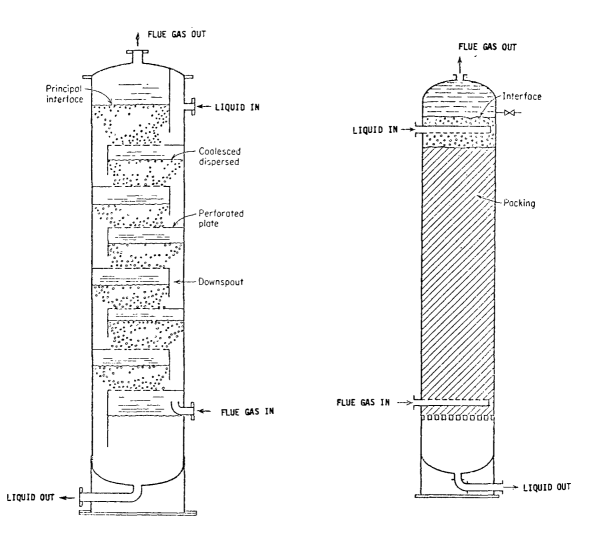
There are some potential problems downstream of the SCR systems due to the presence of the unreacted ammonia in the flue gas. Two things can happen: 1) the NH₃ can react with SO_2 or SO_3 to form ammonium bisulfate or ammonium bisulfate or 2) the NH₃ can enter the downstream equipment unreacted The bisulfate has been shown to cause air preheater pluggage and this is the subject of ongoing research both at the EPA and the Electric Power Research Institute (EPRI). Both the bisulfate and sulfate exist as a particulate, but may be difficult to collect if the particles are submicron in size. Unreacted NH₃ is not likely to present any operational problems. A recent study has shown that if an ESP exists downstream, then most of the NH₃ will exit with the ash. NH₃ can actually improve the performance of an FGD system.¹²⁹

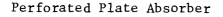
2.2.4 Absorption-Oxidation

2.2.4.1 System Description--

Absorption-oxidation processes remove NO_{\times} from flue gas by absorbing the NO or NO_{\times} into a solution containing an oxidant which converts the NO_{\times} to a nitrate salt. Two types of gas/liquid contactors can be used and examples of each type are shown in Figure 2.2.4-1. Both perforated plate and packed towers accomplish NO_{\times} absorption by generating high gas/liquid interfacial areas. The choice of one type of contactor is a design decision made to achieve a given removal for the least cost.

A generalized process flow diagram is shown in Figure 2.2.4-2. Flue gas is taken from the boiler after the air preheater. Before the gas can be sent to the NO_{χ} absorber, it must be SO_2 -free since SO_2 consumes prohibitive amounts of the costly liquid-phase oxidant. In most cases, the oxidant is permanganate (MnO_{4}) , but $Ca(ClO)_2$ can also be used. Therefore, a conventional FGD unit is required ahead of the NO_{χ} absorber. A prescrubber to cool





Packed Absorber

Figure 2.2.4-1. Gas/liquid contactor options for Absorption-Oxidation Processes.²⁹

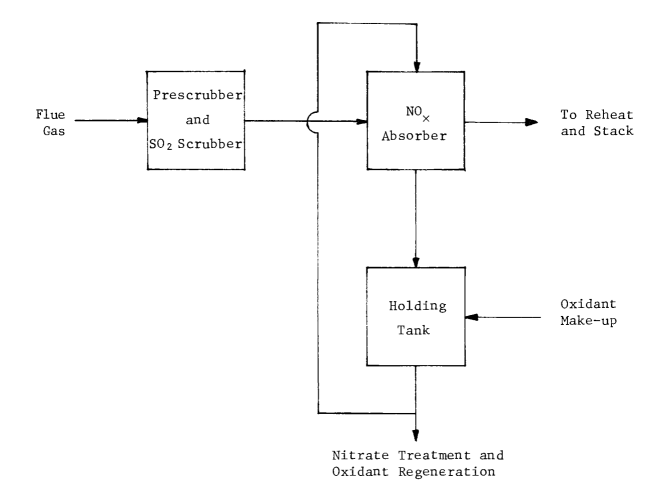


Figure 2.2.4-2. Process flow diagram for Absorption-Oxidation Process.³⁰

the gas and remove both particulates and Cl⁻ prior to FGD is also necessary. After having passed through these two scrubbing sections, the flue gas enters the distributing space at the bottom of the NO_{\times} absorber, below the packing or plates. The gas passes upward through the column, countercurrent to the flow of the liquid absorbent/oxidant (usually a KOH solution containing KMnO₄). NO_{\times} is absorbed and then oxidized over the length of the column according to the following reactions.³¹

NO (g)
$$\rightarrow$$
 NO (aq) (2-6)

$$NO(aq) + KMnO_4(aq) \rightarrow KNO_3(aq) + MnO_2(s)$$
(2-7)

$$2NO_2(g) \rightarrow N_2O_4(g)$$
 (2-8)

$$N_2O_4(g) \rightarrow N_2O_4(aq) \tag{2-9}$$

$$N_2O_4(aq) + 2K_2MnO_4(aq) \rightarrow 2KMnO_4(aq) + 2KNO_2(aq)$$
(2-10)

Since most of the NO_{χ} from combustion processes occurs as NO,³² reactions 2-6 and 2-7 predominate. The clean gas passes out of the top of the absorber to a heater for plume buoyancy and is sent to the stack. The absorbing solution drops to a holding tank where makeup KOH and/or KMnO₄ are added. This solution flows to a centrifuge to separate the solid MnO₂ which is then electrolytically oxidized to MnO₄. The remaining solution is either concentrated in an evaporator to form a weak KNO₃ solution or is electrochemically treated to produce a weak HNO₃ solution and a mixed stream of KOH and KNO₃.

The fundamental design equation used for gas absorption column design is

$$\int_{Y_{a}}^{Y_{b}} \frac{dy}{(y-y^{*})} = \left(\frac{K_{y}a}{G_{y}}\right) z \qquad (2-11)$$

- where y = bulk NO_x concentration (mole fraction) of gas phase at any given point in column
 - y-y* = overall driving force for absorption (y* being the NO_{χ} concentration of a gas in equilibrium with a given liquid NO_{χ} concentration)
 - $Y_{b} = inlet NO_{x}$ concentration
 - $Y_{2} = outlet NO_{2}$ concentration
 - K_y = overall gas-phase mass transfer coefficient, lb-moles NO_x/
 (ft²)(hr)(mole fraction)
 - a = area of gas-liquid interface per unit packed volume, ft^2/ft^3
 - $G_y = molal gas mass velocity, lb-moles flue gas/(ft²)(hr)$
 - Z = length of packed section of column, ft

In a column containing a given packing or plate configuration and being irrigated with a certain liquid flow, there is an upper limit to the gas flow rate. This limit's superficial gas velocity (volumetric gas flow rate/ cross-sectional area of column) is called the flooding velocity. At this point, the gas flow completely impedes the downward motion of the liquid and blows the liquid out of the top of the column. The gas velocity, obviously, must be lower than the flooding velocity. How much lower is a design decision. Often, it is an economic tradeoff between power costs and equipment costs. A low gas velocity will lower the pressure drop and, hence, the power costs but the absorber will have a larger diameter and cost more. High gas velocities have an opposite effect. Usually the optimum gas velocity is about one-half the flooding velocity.34 The height of the column depends on the desired level of removal and on the rate of mass transfer. The latter is a major problem for these systems trying to achieve large NO_{x} reductions since NO is relatively insoluble in water. This can be seen in Table 2.2.4-1.

	Boiling Point, °C	Solubility in Cold Water (O°C), cm ³	Solubility in Hot Water (60°C), cm ³
NO	-151.8	7.34/100 cc H ₂ 0	2.37/100 cc H ₂ 0
NO ₂	21.2	soluble, decomposes	

TABLE	2.2.4-1.	NITROGEN	OXIDES	CHARACTERISTICS ³⁵

One can see that NO has a very limited solubility in water and, since most NO_{\times} is present as NO, the rate of mass transfer (absorption) is going to be relatively slow. This means that the absorber must be tall with a high liquid flow rate. Table 2.2.4-2 presents the effects of boiler/flue gas variables on the design of absorption-oxidation systems.

Variable	Design Effect
Presence of particulates	Requires prescrubber
Presence of SO ₂	Requires FGD pretreatment
Increased gas flow	Requires larger column diameter; increased liquid flow rate
Increased NO_{χ} concentration	Requires larger column height; increased oxidant concentration

TABLE 2.2.4-2. SYSTEM DESIGN CONSIDERATIONS

Both flue gas flow rate and NO_{\times} concentration can be affected by boiler operating conditions. Therefore a change in load on an industrial boiler may alter these variables markedly. The absorber must be designed to accommodate any anticipated load changes. The column size and the liquid and oxidant flows must be designed for each application after examining the boiler operating history and establishing ranges of variation.

None of the sources consulted for this study could supply typical ranges for operating variables such as liquid/gas ratio, reagent concentrations or pressure drops and, as a result, none are presented here. Economic data were not presented either. One source did estimate the removal for absorptionoxidation processes to be 85 percent.³⁶

Presently, absorption-oxidation processes are still in the pilot unit stage of development. Table 2.2.4-3 presents a list of absorptionoxidation process vendors and the status of development of their projects. One can see from the table that no coal-fired flue gas tests have been performed.

Vendor	Status of Development
Hodogaya	No information available; stopped development on process
Kobe Steel	1974: 1000 Nm ³ /hr gas from iron-ore sintering furnace; stopped development on process
MON (Mitsubishi Metal, MKK, Nikon Chemical)	1974: 4000 Nm ³ /hr flue gas from oil-fired boiler
Nissan Engineering	1972: 4 pilot plants, 100-2000 Nm ³ /hr tail gas from HNO ₃ plant

TABLE 2.2.4-3. PROCESS VENDORS OF ABSORPTION-OXIDATION PROCESSES 37,38

2.2.4.2 System Performance--

No coal-fired tests have been made. No information has been published on tests conducted with other fuels. The relative insolubility of NO in water may present a major obstacle to achieving the stringent level of control (90 percent NO, reduction) by absorption-oxidation processes. Another primary drawback of absorption-oxidation systems is the production of nitrate salts (see Equation 4-2), a secondary pollutant. These processes probably could not be applied on a large scale as wastewater treatment systems (chemical or biological) do not remove nitrogen compounds from the wastewater.³⁹ Trying to recover the nitrates as nitric acid for industrial use or potassium nitrate for fertilizer does not seem promising as the by-products are of low quality. Also, the use of an expensive, liquid-phase oxidant requires stainless steel and other corrosion resistant materials of construction. High sulfur coals require an FGD system prior to the NO_{v} absorber to prevent excessive oxidant consumption by SO_2 . The process steps of several absorber columns in series (large fan requirements), oxidant regeneration (electrolysis), and flue gas reheat (inline heater) are all energy intensive and present technical and economic disadvantages.

2.2.5 Selective Catalytic Reduction-NO_X/SO₂ Removal

2.2.5.1 System Description--

From a NO_{χ} removal standpoint, this process is very similar to those discussed in Sections 2.2.2 and 2.2.3. The primary difference is the additional equipment necessary to collect and process the SO₂. The main feature of the process is the reactor and catalyst which remove both NO_{χ} and SO₂. This process was developed by Shell although the U.S. licensor, UOP, is currently marketing and developing the process. The NO_{χ}/SO_2 version of the process is commonly called the SFGT process which stands for the Shell Flue Gas Treatment process.

A uniquely designed parallel flow type of reactor is used to avoid problems with particulates. The reactor consists of a series of packages containing catalyst material, arranged in a parallel configuration which allows flue gas flow between the packages. Each package consists of catalyst material placed between two layers of wire gauze. Figure 2.2.5-1 illustrates the internals of the parallel passage reactor. The flue gas flows between the catalyst packages and not directly through the catalyst material. This prevents plugging of the catalyst with particulate matter in the flue gas.

For convenient fabrication and handling, catalyst packages of a standard size are appropriately spaced and placed in a container to form a unit cell or module. SO_2 removal efficiency and capacity are determined by the number of unit cells placed in series in a cell stack. For a given level of SO_2 removal, a greater number of cells in the stack increases the capacity and reduces the frequency of regeneration. The number of stacks is determined largely by the flue gas rate and the flue gas velocity through a single stack is generally not a design variable. For most design situations, 4 to 5 unit cells in a stack are adequate to achieve high SO_2 removal, however, if a high level of denitrification is required, more unit cells per stack may be necessary.

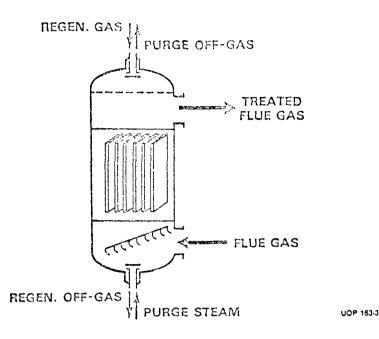


Figure 2.2.5-1. The SFGT parallel flow reactor. 40

The SFGT process is a dry process with two or more reactors operating in a cyclic manner. The desulfurization aspect of the process is regenerable, while NO_x removal is accomplished by catalytic reduction with ammonia. The catalyst material is commonly called an acceptor since SO₂ removal involves adsorption or "acceptance" of SO₂. The desulfurization cycle consists of the following steps:

- 1) oxidation of acceptor bed/acceptance of SO₂,
- 2) purge reactor,
- 3) regeneration with reducing gas, and
- 4) purge reactor.

The products of the oxidation and acceptance reactions in step 1 above catalyze the reaction of NO_{\times} with ammonia to form nitrogen and water. NO_{\times} removal is accomplished by metering ammonia into the untreated flue gas upstream of the reactors. The catalytic reaction takes place across the partially spent acceptor beds.

Also associated with the SFGT process are facilities for generating reducing gas and for the processing of SO_2 in regeneration off gases into sulfur by-products. Figure 2.2.5-2 illustrates the process flow for a typical SFGT system.

Boiler flue gas is withdrawn upstream of the air preheater and particulate removal device by the SFGT system fan and discharged to the reactor inlets. The flue gas then flows through fixed bed reactors in open channels alongside and in contact with the acceptor material. Ammonia is added to the flue gas upstream of the SFGT system fan to insure complete mixing before the flue gas enters the reactor.

Fresh acceptor material is elemental copper on an alumina support. This is converted to the oxide form by flue gas oxygen shortly after initiation of the acceptance cycle. SO_2 is removed by reaction with the copper oxide and oxygen as the flue gas flows through the channels, converting the acceptor material to copper sulfate. Simultaneous with the desulfurization process, the reduction of flue gas NO_x by ammonia is selectively catalyzed by copper oxide and copper sulfate in the acceptor bed. As the flue gas leaves the SFGT system reactors it is returned to the boiler flue gas duct downstream of SFGT fan suction.

Flue gas is fed to a reactor until an unacceptable amount of SO₂ begins to pass through the reactor. This occurs when a large fraction of the acceptor has been converted to the sulfate form. Flue gas flow is then diverted to another reactor and the spent reactor is isolated. Any flue gas remaining in the spent reactor is purged with an inert gas such as steam, and the regeneration cycle is initiated.

Regeneration is accomplished by passing a reducing gas through the bed countercurrent to the direction of the flue gas flow. The reducing gas, which is primarily hydrogen, reacts with the copper sulfate in the spent reactor to convert it to elemental copper. An off gas of SO₂ and water is

2-24

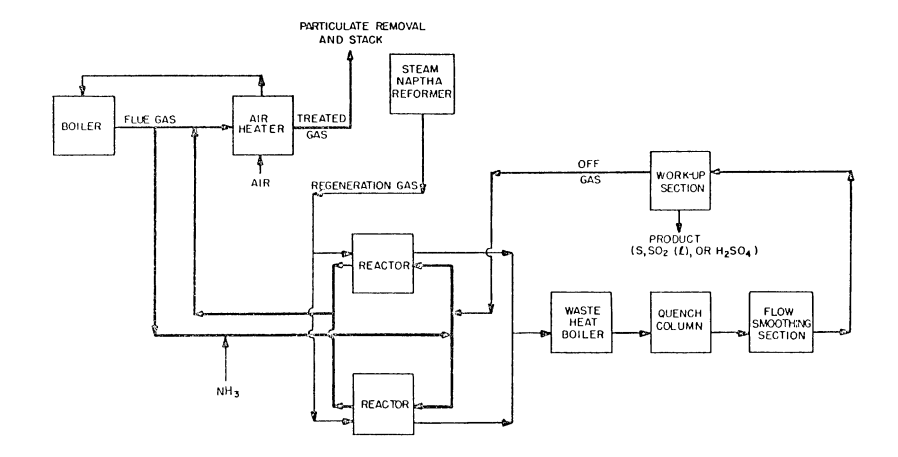


Figure 2.2.5-2. Flow diagram of the SFGT process.⁴¹

produced by the reaction. After regeneration is complete, the reactor is again purged with steam and is ready for another acceptance cycle. Regeneration gas can be produced from a number of sources, but steam-naphtha reform-ing is proposed by UOP as being the most economical.⁴²

The regeneration off-gas treatment section consists of flow smoothing equipment and SO_2 workup equipment. Typically, the regeneration off-gas is cooled and most of the steam condensed, raising the SO_2 concentration from 10 percent to 80 percent by volume. The concentrated SO_2 is then compressed into an intermediate holding vessel to provide a smooth flow rate to the workup section. The workup section may be a modified Claus unit which produces an elemental sulfur by-product, a fractionation unit which produces liquid SO_2 , or a sulfuric acid plant.

Each process step consists of different chemical reactions. The is converted to the oxide form by the following reaction:

$$Cu + \frac{1}{2}O_2 + CuO$$
 (2-12)

This oxide readily reacts with flue gas SO2 and oxygen, as described by:

$$Cu0 + \frac{1}{2}O_2 + SO_2 \rightarrow CuSO_4$$
 (2-13)

SO3 in the flue gas is also removed by the following reaction:

$$CuO + SO_3 \rightarrow CuSO_4 \tag{2-14}$$

The reaction scheme for reduction of NO_{X} is described by the following: $^{1\,2}$

$$4NO + 4NH_3 + O_2 \neq 4N_2 + 6H_2O$$
 (2-1)

$$2NO_2 + 4NH_3 + O_2 \neq 3N_2 + 6H_2O$$
 (2-2)

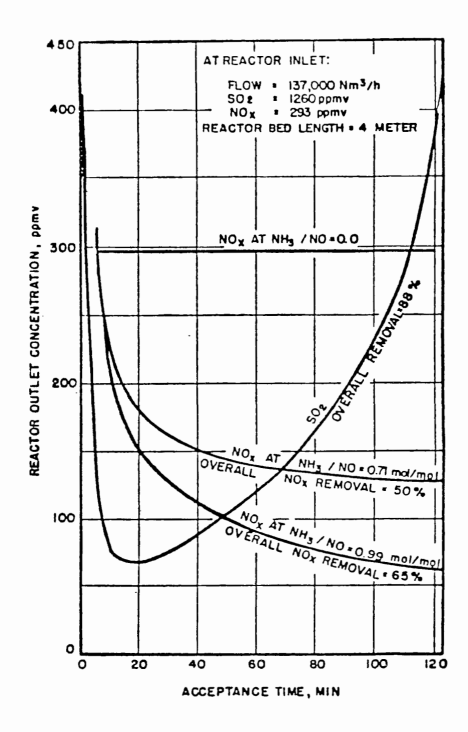


Figure 2.2.5-3. SFGT reactor performance *versus* acceptance time.⁴⁴

Excess ammonia which is not consumed in reactions 2-1 and 2-2 may be catalytically oxidized to nitrogen and water by reaction with flue gas oxygen, as described by:

$$4NH_3 + 3O_2 \neq 2N_2 + 6H_2O$$
 (2-15)

Maximum NO_{\times} removal efficiency is achieved at the point of SO_2 breakthrough, where conversion of the acceptor material from the oxide to the sulfate form is essentially complete. Figure 2.2.5-3 illustrates reactor outlet SO_2 and NO concentrations during a typical SFGT acceptance cycle.

Copper sulfate is reduced to the elemental copper form by reducing gas hydrogen as described by the following reaction:

$$CuSO_4 + 2H_2 \rightarrow Cu + SO_2 + 2H_2O$$
 (2-16)

Any acceptor material present in the reactor as the oxide will also be reduced, according to the following reaction:

$$CuO + H_2 \rightarrow Cu + H_2O$$
 (2-17)

The regeneration step occurs at the same temperature as the acceptance step, $400^{\circ}C$ (750°F).

The general reactor design equation is the same as that described in earlier sections for SCR processes. The primary variables are the gas rate, reaction rate, and control level. Reaction rate data have not been released for this process except that the NO_v reduction is first order.

The gas flow rate and control level will determine the reactor size. Increases in either variable will increase the reactor volume. The effect of control level can be seen in Figure 2.2.5-4. It is necessary for the flue gas to enter the reactor at 400°C and therefore it must be taken from

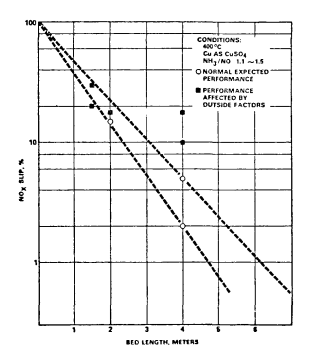


Figure 2.2.5-4. Unconverted NO_X as a function of catalyst bed length.⁴⁵

an appropriate point in the boiler, most likely from between the economizer and air preheater. Alternatively, a cooler gas can be heated to 400°C by an inline heater.

The removal efficiency of NO_{\times} for a given reactor size is determined by the amount of NH_3 injected as shown in Figure 2.2.5-5. Since the reaction is first order in NO_{\times} , control level is not affected by NO_{\times} concentration.⁴⁷ The SO₂ control efficiency is primarily a function of the acceptance time of the reactor (Figure 2.2.5-3). Typical ranges of operating variables are shown in Table 2.2.5-1.

Since the SFGT system can handle full particulate loading ($\geq 10 \text{ gr/sft}^3$) it is not dependent on any pretreatment facilities. Also, the SFGT system operation is independent of boiler operation. The system fan takes suction from the flue gas duct between the economizer and air preheater and the reactor discharge returns to the boiler flue gas duct just downstream of the

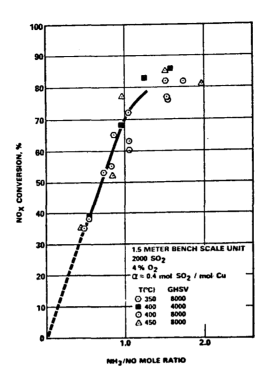


Figure 2.2.5-5. NO_{\times} reduction with NH_3 over commercial SFGT acceptor.46

Variable	Typical Range
Space Velocity	$5,000 - 8,000 \text{ hr}^{-1}*$
NH₃:NO _× Mole Ratio	1.0:1.0 to 1.2:1.0*
Flue Gas Temperature	400°C
Pressure Drop	5-6 in. H ₂ 0*
Maximum Particulate Loading	≥23 g/Nm

TABLE 2.2.5-1. DESIGN AND OPERATING VARIABLES FOR SFGT SYSTEM^{4 8}

*Actual value will depend on required removal level.

suction point, with no valves between the two points. The system fan provides a constant flow rate through the SFGT system. If the boiler flue gas rate is greater than the fan rate, flue gas will bypass the system through the open duct. If the boiler flue gas rate is lower than that of the system fan, treated gas will recycle back to the system fan suction. Recycle of treated gas to the reactor inlet with "open bypass" arrangement presents no operating problems. This is due to the fact that both the level of desulfurization and denitrification are independent of inlet concentrations, and the system does not humidify the flue gas.

Tables 2.2.5-2 and 2.2.5-3 present test and commercial applications of the SFGT process. The development history of the process can also be seen in these tables.

In the U.S., from 1974 to 1976 a pilot scale unit at Tampa Electric Company (TECO) was operated using flue gas from a coal-fired boiler. Testing was for SO₂ removal only, NO_x control was not attempted during this period. The process developer is currently modifying the TECO pilot unit to accommodate 7 meters of bed height, up from the previous maximum of 5 meters. This should permit simultaneous removal of NO_x and SO_x to the 90 percent level. Also, provisions are being made for injection of a CO/CO₂ gas mixture into the regeneration gas in order to simulate medium-Btu gas from a coal gasifier.

The costs for an industrial size boiler have not been estimated. However, costs for a 500 MW utility boiler application are available and are shown in Tables 2.2.5-4, 5, and 6. Also shown are the estimated energy and raw material requirements.

2-31

Location/ Company	Designed By	Fuel/ Application	Size, Nm³/hr	Type of Operation	Dates	Comments
Shell Ref. at Pernis	Shell	Residual Fuel Oil- Proc. Heater	600-1000	SO _x -only	1967-1972	SO _x reduction - approx. 90%
Rotterdam Utility	Shell	Coal- Steam Boiler	-	Heavy Fly Ash Loading	1971	Particulate mat- ter - loadings to 20 Gr/Nm ³
Tampa Elec. Big Bend	UOP	Coal- Wet-Bottom Utility Boiler	1200-2000	SO _× -only; SO _× -NO _× Simultaneous	1974–1976 1979–	SO - 90%; SO _x -NO _x - 90/90% fly ash to 25 Gr/Nm ³
JGC Yokohama	JGC*	Fuel Oil	250-700	NO _x -only	1974-	NO _x reduction - 90-99%
Nippon Steel	JGC	Sintering Furnace	2000	NO _x -only	1976-1978	NO _× reduction - 90-97%
	JGC	Coke Oven Gas	400	NO _x -only	1976-1977	NO _x reduction - 90%; special low temp. cat. evalua- tion

TABLE 2.2.5-2. SFGT PROCESS, PILOT AND DEMONSTRATION UNITS

*JGC Corporation, licensing agent in Japan.

Unit	Designed By	Fuel/ Application	Size, Nm ³ /hr	Type of Operation	Dates	Comments
SYS* Yokkaichi	Shell	Residual Fuel Oil- Ref. Boiler	125,000	SO _x -only; NO _x -SO _x Simultaneous	1973–1975 1975–	SO _x reduction - 90%; Simultaneous - 90/50%
Kashima Oil Co. Ltd.	JGC	Fuel Oil- Process Unit Heater	50,000	NO _x -only	1975-	95–98%
Fuji Oil Co. Ltd.	JGC	CO Boiler	70,000	NO _× -only	1976-	93-96%
Nippon Steel Corp.	JGC	Sintering Furnace	150,000	NO _× -only	1978-	∿95% (low temp. cata- lyst)

TABLE 2.2.5-3. SFGT PROCESS, COMMERCIAL UNITS

*Showa Yokkaichi Sekiyu

	BASIS:
Incorporated Units:	Steam-Naphtha Reformer SFGD Reactor Section Compressor/Gasholder Flow Smooth Section Modified Claus Unit
	Modified Claus Unit
Power Plant Size	500 MW
Fuel	Coal
S-Content, Wt-%	
C as e 1	3.5
Case 2	2.5
Case 3	0.8
HHV	10,500 Btu/1b
Heat Rate	9,000 Btu/kWh
Excess Air	20%
Air Preheater Leakage	13%
Flue Gas Rate	1,582,000 Nm ³ /h (983,000 SCFM)
SO ₂ Content, ppmv	
Case 1	2,580
Case 2	1,850
Case 3	590
Mid-1977, Gulf Coast Location	
Load Factor	7,000 h/a
Capital Charges	15%/a
Cost of:	
Naphtha	\$0.35/gal
Steam (40 psi, SAT.)	\$1.50/M 1b
Electricity	\$0.018/kWh
Labor	\$10.00/hr
Heat Credits	\$2.50/MMBtu
Sulfur	\$45.00/ton

TABLE 2.2.5-4. ECONOMICS OF SFGT SYSTEM⁴⁹

- - - - -

		SFGD Section	Flow Smooth Section	Mod. Claus Section	Reformer Section	Total
Case 1						
Electricity	kW	5,770	850	115	480	7215
Steam**	kmo1/h	1,820	-380*	-740*	-600*	100
Naphtha***	Gcal/h				90.92	90.92
Heat Credits	G cal/ h					42.53
S° Produced	kg/h			5250		5250
Case 2						
Electricity	kW	5,800	570	82	300	6782
Steam**	kmol/h	1,300	-270*	-530*	-415*	85
Naphtha***	Gcal/h				62.75	62.75
Heat Credits	Gcal/h					32.48
S° Produced	kg/h			3760		3760
Case 3						
Electricity	kW	5,120	180	30	110	5440
Steam**	kmol/h	480	-95*	-170*	-140*	75
Naphtha***	Gcal/h				21.01	21.01
Heat Credits	Gcal/h					18.46
S° Produced	kg/h			1200		1200

TABLE 2.2.5-5. ECONOMICS OF SFGT SYSTEM ESTIMATED CHEMICALS AND UTILITY REQUIREMENTS⁵⁰

*Produced

**40 psig, Saturated

***5.175 MMBtu/Bb1 produces 11,500 SCF Hydrogen/Bb1

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	L AND OFERALL		
	Case 1	Case 2	Case 3
EEC. (MM\$)			
SFGD Reactor Section	28.95	28.53	22.94
Compressor/Gasholder	7.82	6.10 /	2.65
Modified Claus	2.76	2.26	1.14
Steam-Naphtha Reformer	8.81	7.14	4.17
Estimated Annual Revenue Requirements (M\$/a)			
Capital Charges	7251	6604	4634
Maintenance	967	881	618
Labor	123	123	123
Acceptor	1479	1053	411
Electricity	909	855	685
Steam	42	35	31
Naphtha	7174	4951	1658
Heat Credits	-2977	-2273	-1292
Sulfur Credits	-1570	-1126	-359
Capital Cost, Operating Cost, Energy Requirement			
Capital Cost, \$/kW	97	88	62
Operating Cost, ¢/kWh	0.38	0.32	0.19
Energy Requirement, Btu/kWh*	525	371	124
*Defined as the sum of;			<u></u>
Electricity at 9000 Btu/kWh Steam at 40000 Btu/kmol Naphtha at 4 Btu/kcal Heat Credits at 4 Btu/kcal			

TABLE 2.2.5-6. ECONOMICS OF SFGT SYSTEM ESTIMATED CAPITAL AND OPERATING COST⁵¹

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2.2.5.2 System Performance--

 NO_{x} control by the SFGT process is shown graphically in Figure 2.2.5-5. As can be seen, at a space velocity of 8000 hr⁻¹, NO_{x} control of >80 percent can be achieved. Figure 2.2.5-4 indicates that the process developers feel the process to be capable of NO_{y} control levels of >90 percent.

Several different test series were conducted using the TECO pilot plant and the operating conditions for these tests are shown in Table 2.2.5-7. The SO₂ removal efficiency for several of these runs is shown in Figure 2.2.5-6 plotted against the number of cycles, which can be converted to time. No data of this type are available for NO_× control using coal-fired flue gas, however, these data should be available in about one year.

As mentioned earlier, the system is not impacted by changes in the boiler gas rate or particulate concentrations. Changes in the NO_{χ} concentration due to boiler load changes can be compensated for by a conventional control system used in conjunction with the NH₃ injection equipment. This control system will be developed during the upcoming pilot tests at the TECO pilot plant.

2.2.6 Adsorption

2.2.6.1 System Description--

The adsorption process removes NO_{\times} and SO_2 from flue gas by adsorbing them onto a special activated char. Adsorbed NO_{\times} is reduced to N_2 while SO_2 is reduced and condensed to an elemental S by-product.

A process flow diagram is shown in Figure 2.2.6-1. Flue gas is taken from the boiler air preheater and passed through a particulate removal device to prevent blinding of the adsorption bed. The flue gas then enters the adsorber, a vertical column with parallel louver beds containing the char in

TABLE 2.2.5-7.	SUMMARY OF PLANT AT T		ERATING C	ONDITIONS ON	THE SFGT	PILOT
Run No.	1	2	3	4	5	6
Duration, Months	5	2 ¹ 2	1^{1}_{2}	2	5	3
Cycles	2488	1520	1292	1412	4328	2210
Cumulative Cycles	2488	4008	5300	6712	11040	13250
Flow Rate, SCFM	1090	1090	1090	1090/1420	1090	1090
Acc. Time, Min.	20	20	20	20	20	20
Reg. Time, Min.	20	20	20	20	20	20
Flue Gas Source*	1	1	2	1	1	3
Eff. SOR		92	95	95	95	93
Eff. EOR		82	95	80	92	93

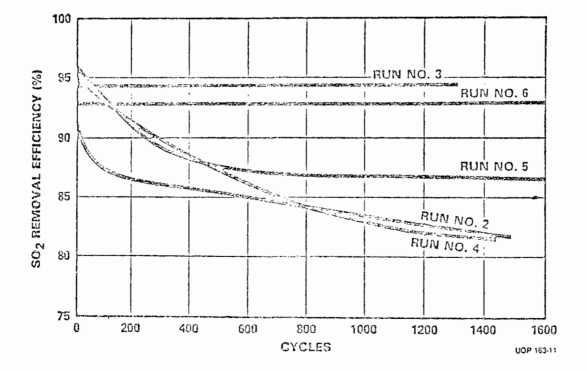


Figure 2.2.5-6. SO₂ removal efficiency vs. cycles.⁵³

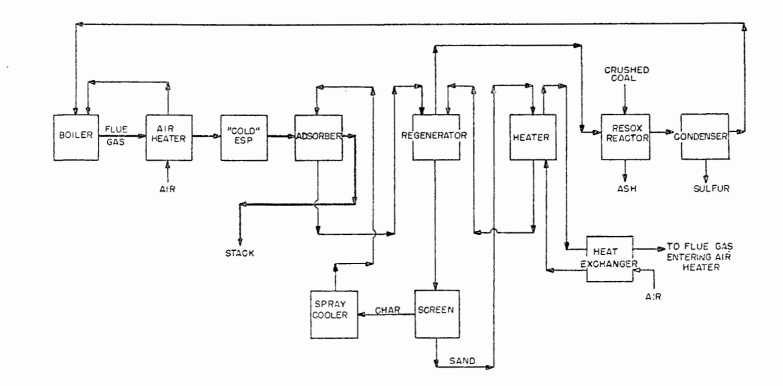


Figure 2.2.6-1. Flow diagram of Foster Wheeler-Bergbau Forschung Dry Adsorption Process.⁵⁴

pellet form. NO_X and SO_2 are adsorbed on the char which slowly moves downward through the bed. The NO_X adsorption mechanism is unknown but SO_2 undergoes the following reaction.

$$SO_2(g) + H_2O(g) + \frac{1}{2}O_2(g) + H_2SO_4(1)$$
 (2-18)

The reaction product is held in the pores of the char pellets. The flue gas exits the adsorber and passes to the stack. The saturated char leaves the bottom of the adsorber and is screened to remove any fly ash deposits. It is then conveyed to a regenerator where it is mixed with hot sand (650°C) and the following reactions take place.^{55,56}

$$2H_2SO_4(1) + C(s) + CO_2(g) + 2H_2O(g) + 2SO_2(g)$$
 (2-19)

$$2NO(g) + C(s) \rightarrow CO_2(g) + N_2(g)$$
 (2-20)

This SO_2 -rich gas product stream is sent to an off-gas treatment reactor containing hot, crushed coal (650-820°C) and the following reactions take place.⁵⁶

$$SO_2(g) \rightarrow S(g) + O_2(g)$$
 (2-21)

$$C(s) + O_2(g) + CO_2(g)$$
 (2-22)

The gas then passes to a condenser where the S vapor forms molten S. The char/sand mixture from the regenerator is screened to separate the two solids. The char is recycled to the adsorber via a spray cooler and the sand is recycled to the regenerator after passing through a heater.

This process operates at 120-150°C, however, typical values for other operating variables were not found. NO_{χ} and SO_2 control levels were reported to be 40-60 percent and 80-95 percent, respectively.⁵⁷ The economics of the process vary with the fuel sulfur level. For fuel sulfur levels of 0.9-4.3

percent, the capital costs range from 40-90/kW and the operating costs range from 1.0-2.3 mills/kWh.⁵⁸ The costs were based on applying the process to a utility boiler of >200 MW capacity.

Presently, the adsorption process is in the prototype unit stage of development. The one reported process developer in the field, Foster Wheeler-Bergbau Forschung has a 20 MW prototype unit and several small pilot plants treating coal-fired flue gas.

2.2.6.2 System Performance--

Tests have shown the adsorption process to be primarily a SO₂ reduction process as NO_{χ} removal efficiency averages 40-60 percent while SO₂ removal had a range of 80-95 percent.⁵⁹

The primary drawback of this process, besides the low NO_{\times} removal level, is its complexity: numerous process steps involving hot solids handling. Solids flow can be difficult to control and high maintenance requirements could be expected. The vendor has reported several mechanical problems during testing which included control of adsorber-bed levels, poor char distribution, char-sand separation, hot sand conveying, and char cooling and feed. Some corrosion-resistant material is needed in the high temperature zones of the process. The ash waste stream from the off-gas treatment reactor appears to be the sole secondary pollutant associated with the process. The overall complexity and low NO_{\times} removal of the process present definite technical disadvantages.

2.2.7 Electron Beam Radiation

2.2.7.1 System Description--

This dry process utilizes an electron beam to bombard the flue gas, removing NO_{χ} and SO_2 in the process. A block flow diagram for the process is shown in Figure 2.2.7-1.

2-41

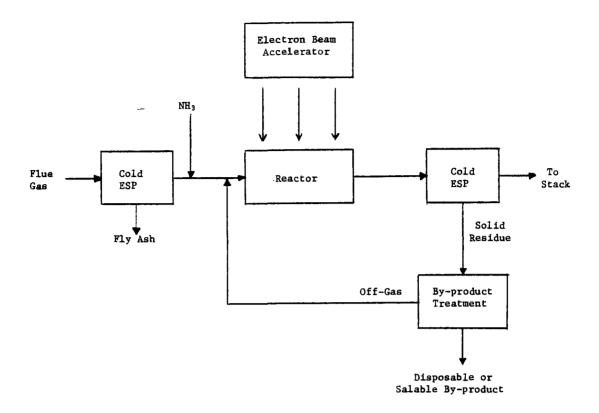


Figure 2.2.7-1. Process flow diagram for Ebara-JAERI electron beam process.⁶⁰

Flue gas is taken from the boiler air preheater and passed through a cold ESP to remove particulates. After a small amount of ammonia is added, the gas enters a reactor where it is bombarded with an electron beam. (The penetration of the gas stream by the beam will require a unique discharge pattern or other special design considerations.) A powder containing both ammonium nitrate and sulfate is generated by an unknown reaction mechanism. The gas then exits the reactor, passes through a second ESP to remove the solid by-product, and is sent to the stack. The by-product treatment system is still being developed. Various methods investigated include thermal decomposition in the presence of an inert gas, steam roasting with CaO, or steam roasting with H₂O. The by-product may eventually be useful as a fertilizer.⁶¹

The key subsystem of this process is the electron beam accelerator. Control of this unit's power supply is based upon inlet composition, flow rate, and temperature of the flue gas.

Some of the important variables and typical ranges are listed in Table 2.2.7-1.

	Typical Value
Temperature	∿100°C
Reactor residence time	1-20 sec
Radiation rate	10^5 – 10^6 rad*/sec
Total radiation absorbed	1-3 Mrad*

TABLE 2.2.7-1. SYSTEM VARIABLES⁶²

*Rad is the radiation dose absorbed
1 rad = .01 J/Kg

The operating cost with NO_{χ} removal only (low sulfur coals) is lower due to lower radiation levels, but the capital cost would be just as high as for simultaneous NO_{χ}/SO_{χ} removal. Capital costs are quite high for this process as the 2 ESP's and the accelerator are expensive. The cost for a 1000 Nm³/hr test unit are reported to be \$1000/kW, however, the cost of a full scale system is expected to be lower. Operating costs are not available.

No coal-fired tests have been performed at this time. The Ebara Manufacturing Company in conjunction with Japan Atomic Energy Research Institute (JAERI) has operated a 1000 Nm³/hr pilot plant treating flue gas from an oil-fired boiler. In 1976, a 3000 Nm³/hr pilot plant began treating off-gas from an iron ore sintering furnace at Nippon Steel. By-product treatment technology needs to be more fully developed before this process can be applied commercially.

2-43

In the U.S., the Department of Energy (DOE) is funding development of an electron beam process offered by Research-Cottrell. Pilot unit tests with flue gas are scheduled, however, the details of the program are not yet available.

2.2.7.2 System Performance--

No coal-fired testing has been done.

A summary of the oil-fired pilot tests is shown in Figure 2.2.7-2.

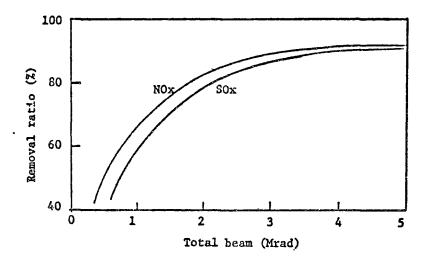


Figure 2.2.7-2. Oil-fired pilot plant test results.⁶⁴

One can see that NO_{\times}/SO_2 removal drops off drastically at a total radiation dose below 1 Mrad while the maximum removal is obtained at about 3 Mrad. The removal efficiencies decrease as the concentrations of NO_{\times} and SO_2 increase as can be seen in Figure 2.2.7-3.

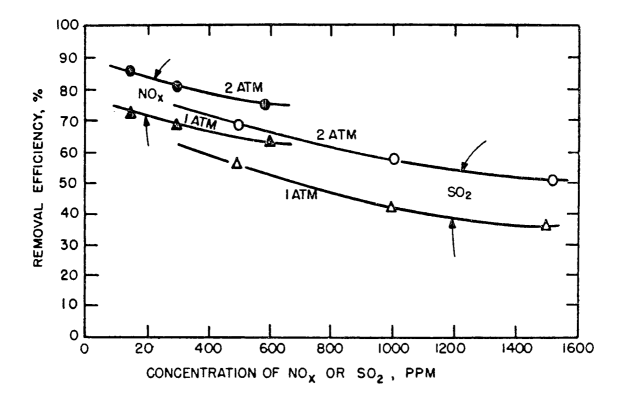


Figure 2.2.7-3. Effect of pollutant concentration on removal efficiency.⁶⁵

2.2.8 Absorption-Reduction

2.2.8.1 System Description--

Absorption-reduction processes simultaneously remove NO_{χ} and SO_2 from flue gas by absorbing them into a scrubbing solution. The processes are based on the use of chelating compounds, such as ethylenediamine tetraacetic acid (EDTA) complexed with iron, to "catalyze" the absorption of NO_{χ} . Most process vendors prefer a perforated-plate type of gas-liquid contactor. The advantages of a perforated-plate absorber over a packed bed absorber include easier cleaning when solids are present, wider operating ranges, and more economical handling of high liquid rates.⁶⁶ An example of a perforated plate absorber is shown in Figure 2.2.8-1. The most common design of a perforated plate is one that employs liquid crossflow over the face of the plate with the gas passing upward through the plate perforations. A schematic of the operation of a crossflow perforated plate is shown in Figure 2.2.8-2. The liquid is prevented from flowing through the plates by the upward flow of the gas. However, during periods of low gas flow (such as load changes on industrial boilers) liquid can drain through the openings in the plates. This reduces the liquid's time of contact with the gas on each plate and may decrease the overall operating efficiency of the absorber. To prevent this problem, there are two other types of dispersers utilized besides the basic sieve-plate: the valve-plate and the bubble cap, depicted in Figure 2.2.8-3. As the gas flow lowers, the valve or cap settles, sealing off the perforation so liquid cannot drain through. This design feature allows the perforated plate absorber to operate more efficiently at widely fluctuating gas rates.

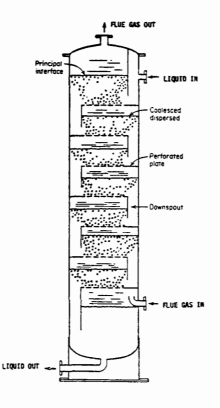


Figure 2.2.8-1. Perforated plate absorber option for Absorption-Reduction Processes.²⁸

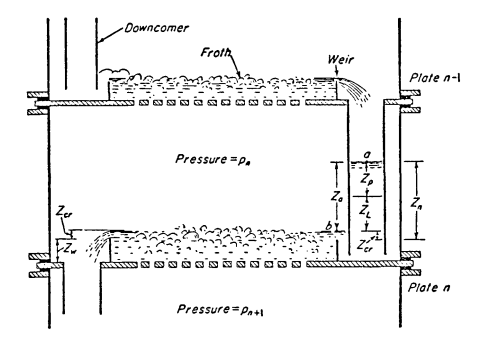
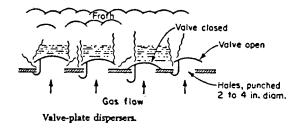
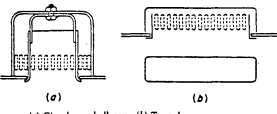


Figure 2.2.8-2. Normal operation of sieve plate. Z_{α} , height of station a above datum. Z_{CP} , weir crest. Z_L , liquid-friction head. Z_p , pressure head across plate. Z_n , net head in down pipe. Z_{ω} , weir height.⁶⁷





(a) Circular or bell cap. (b) Tunnel cap.

Bubble cap dispersers

Figure 2.2.8-3. Other gas dispersers.⁶⁸

While most all absorption-reduction processes utilize ferrous chelating compounds to enhance NO absorption, the scrubbing solutions, the by-product treatment and sorbent regeneration chemistry differ from process to process. For this reason, one of the simpler absorption-reduction processes, that of Kureha Chemical Industry Company, is examined here in detail.

A block flow diagram of the Kureha absorption-reduction process is shown in Figure 2.2.8-4. Flue gas is taken from the boiler after the air preheater. It passes through a prescrubber to adiabatically cool the gas and remove both particulates and chlorides. The flue gas then enters the distributing space at the bottom of the NO_{χ}/SO_2 absorber, below the plates or packing. The gas flows upward, countercurrent to a sodium acetate (CH₃COONa) scrubbing solution ($\sim 60^{\circ}$ C) containing ferrous iron and EDTA and a few seed crystals of gypsum (to prevent scaling). Most of the SO₂ is rapidly absorbed at the bottom of the absorber according to the following reactions.⁷¹

$$SO_2(g) \rightarrow SO_2(aq)$$
 (2-23)

 $SO_2(aq) + 2CH_3COONa(aq) + H_2O \rightarrow Na_2SO_3(aq) + 2CH_3COOH(aq)$ (2-24)

The NO_{χ} (which consists mainly of NO) is relatively insoluble; therefore, it is absorbed gradually over the length of the column. The ferrous chelating compounds effect on NO absorption is described in Figure 2.2.8-5. The NO_{χ} is absorbed and undergoes the following reactions.⁷³

$$NO(g) \rightarrow NO(aq)$$
 (2-6)

$$2NO_2(g) \rightarrow N_2O_4(aq)$$
 (2-25)

$$N_2O_4(g) \rightarrow N_2O_4(aq)$$
 (2-9)

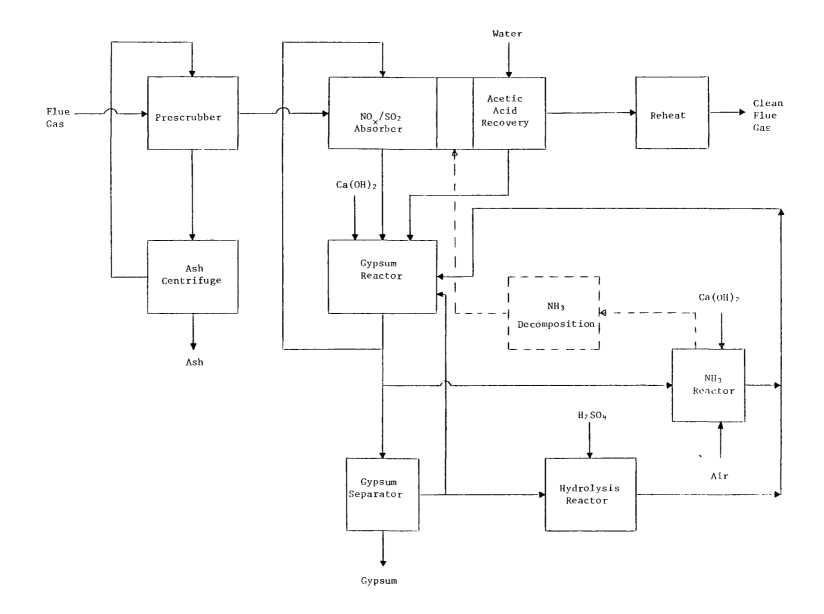


Figure 2.2.8-4. Process flow diagram of Kureha absorption-reduction process. 69,70

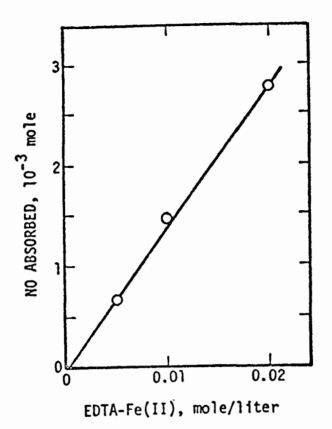


Figure 2.2.8-5. EDTA-Fe(II) concentration and NO absorption at 50°C.72

 $2NO(aq) + 5Na_2SO_3(aq) + 4CH_3COOH(aq) \rightarrow 2NH(SO_3Na)_2(aq) + Na_2SO_4(aq) + 4CH_3COONa(aq) + H_2O \quad (2-26)$

 $2N_{2}O_{4}(aq) + 7Na_{2}SO_{3}(aq) + 4CH_{3}COOH(aq) \rightarrow 2NH(SO_{3}Na)_{2}(aq) + 3Na_{2}SO_{4}(aq) + 4CH_{3}COONa(aq) + H_{2}O \quad (2-27)$

Some of the acetic acid (CH₃COOH) formed at the bottom of the absorber via reaction (2-24) is vaporized. It must be captured and is done so by water scrubbing at the very top of the absorber. From the top of the absorber column the clean flue gas passes to a heater for plume buoyancy and is then sent to the stack. The liquid effluent drops from the bottom of the absorber to a gypsum, CaSO₄·2H₂O, production reactor. Here, the solution is mixed with with the purge stream from the acetic acid recovery section and a lime slurry stream. The lime, Ca(OH)2, treatment involves the following reactions.⁷⁴

$$2CH_{3}COOH(aq) + Ca(OH)_{2}(aq) \rightarrow (CH_{3}COO)_{2}Ca(aq) + 2H_{2}O$$
 (2-28)

 $(CH_{3}COO)_{2}Ca(aq) + Na_{2}SO_{4}(aq) + 2H_{2}O \rightarrow CaSO_{4} \cdot 2H_{2}O(s) \neq + 2CH_{3}COONa(aq)$ (2-29)

The gypsum formed by reaction 2-29 is centrifuged. Most of the liquor discharged is returned to the gypsum reactor and on to the absorber. The remaining liquor is sent to a reactor where sulfuric acid (H_2SO_4) is added to hydrolyze the imidodisulfonate, $NH(SO_3Na)_2$, by the following reaction.⁷⁵

$$H^{+}$$

NH(SO₃Na)₂(aq) + 2H₂O $\stackrel{\rightarrow}{\rightarrow}$ NH₄HSO₄(aq) + Na₂SO₄(aq) (2-30)

The effluent from this reactor is then recycled to the gypsum production reactor. A small purge stream is taken from the gypsum reactor to another reactor where the ammonium bisulfate (NH₄HSO₄) formed in the hydrolysis reaction is treated with lime to yield gypsum and NH₃ off-gas by the following reaction.⁷⁶

$$\mathrm{NH}_{4}\mathrm{HSO}_{4}(\mathrm{aq}) + \mathrm{Ca}(\mathrm{OH})_{2}(\mathrm{s}) \rightarrow \mathrm{CaSO}_{4} \cdot \mathrm{2H}_{2}\mathrm{O}(\mathrm{s}) + \mathrm{NH}_{3}(\mathrm{g})^{\uparrow} \qquad (2-31)$$

The gaseous ammonia is stripped from the solution by an air stream. If no use for the ammonia can be found, the gas mixture is sent to a catalytic reactor where ammonia is oxidized by the following reaction.⁷⁷

$$4NH_{3}(g) + 30_{2}(g) \xrightarrow{\text{catalyst}} 2N_{2}(g) + 6H_{2}O(g) \qquad (2-32)$$

350°C

The product stream is then sent to the deacetating section of the absorber column.

The fundamental design equation used for gas absorption column design is $^{\rm 32}$

$$\int_{Y_{a}}^{Y_{b}} \frac{dy}{(y-y^{*})} = \left(\frac{K_{a}}{G_{y}}\right) Z \qquad (2-11)$$

- where y = bulk NO_x concentration (mole fraction of gas phase at any given point in column
 - y-y* = overall driving force for absorption (y* being the NO_x concentration of a gas in equilibrium with given liquid NO_x concentration)
 - $Y_{\rm b}$ = inlet NO_x concentration
 - $Y_a = outlet NO_{\times} concentration$
 - Ky = overall gas-phase mass transfer coefficient, lb-moles NO_x/
 (ft²)(hr)(mole fraction)
 - a = area of gas-liquid interface per unit packed volume, ft^2/ft^3
 - $G_y = molal gas mass velocity, lb-moles flue gas/(ft²)(hr)$
 - Z = length of packed section of column, ft

In a column containing a given plate or packing configuration and being irrigated with a certain liquid flow, there is an upper limit to the gas flow rate. This limit's superficial gas velocity (volumetric gas flow rate/ cross-sectional area of column) is called the flooding velocity. At this point, the gas flow completely impedes the downward motion of the liquid and blows the liquid out of the top of the column. The gas velocity, obviously, must be lower than the flooding velocity. How much lower is a design decision. Often it is an economic tradeoff between power costs and equipment costs. A low gas velocity will lower the pressure drop and, hence, the power costs but the absorber will have a larger diameter and cost more. High gas velocities have an opposite effect. Usually the optimum gas velocity is about one-half the flooding velocity.33 The height of the column depends on the desired level of removal and on the rate of mass The latter consideration is the reason why a chelating compound transfer. is used in absorption-reduction processes to aid in NO_{χ} absorption. Table 2.2.8-1 presents the effects of boiler/flue gas variables on the design of

2 - 52

absorption-reduction systems. Both flue gas flow rate and NO_{\times} concentration can be affected by boiler operating conditions. Therefore a change in load on an industrial boiler may alter these variables markedly. The absorber must be designed to accommodate any anticipated load change. The column size and the liquid flows must be designed for each application after examining the boiler operating history and establishing ranges of variation.

Variable	Design Effect
Presence of particulates	Requires prescrubber
Presence of SO ₂	Requires SO ₂ :NO mole ratio of at least 3-5 ⁹ (depending on process) for absorption- reduction to be effective.
Increased gas flow	Requires larger column diameter; increased liquid flow rate
Increased NO_{χ} concentration	Requires larger column height; increased catalyst concentration

TABLE 2.2.8-1. SYSTEM DESIGN CONSIDERATIONS

The process vendors have not released much information on the operating conditions of these processes. This is primarily due to the competitive status of these similar processes at this early stage of development. Typical values for some of the process variables are shown in Table 2.2.8-2.

Variable	Range
Liquid/Gas ratio, 1/Nm ³	10-30
$\mathrm{SO}_{\times}/\mathrm{NO}_{\times}$ mole ratio	2.5-3.0
Superficial Gas Velocity, m/s	1-3

TABLE 2.2.8-2.TYPICAL VALUES FOR PROCESS VARIABLES
OF ABSORPTION-REDUCTION PROCESSES

Cost estimates for this type of process cover a large range, presumably due to the differences in sorbent regeneration technique. Capital costs for utility applications are reported to range from \$65-127/kW and operating costs from 4.8-7.4 mills/kWh.⁷⁹

Presently, absorption-reduction processes are in the pilot-unit stage of development. Table 2.2.8-3 presents a list of absorption-reduction process vendors and the status of development of their projects. One can see from the table that only one coal-fired flue gas test has been performed.

Vendor	Status of Development
Asahi	1974: 600 Nm ³ /hr flue gas from residual oil- fired boiler (1000 hours continuous).
Chisso	1975: 300 Nm ³ /hr flue gas from oil-fired boiler (335 hours continuous)
Kureba	1976: 5000 Nm ³ /hr flue gas from heavy oil-fired boiler (3000 hours continuous)
Mitsui Engineering and Shipbuilding	1974: 150 Nm^3/hr flue gas from oil-fired boiler
Pittsburgh Environmental	1976: 3000 Nm ³ /hr flue gas from coal-fired boiler (52 hours continuous, absorption section only)

TABLE 2.2.8-3. PROCESS VENDORS OF ABSORPTION-REDUCTION PROCESSES⁸⁰

2.2.8.2 System Performance--

The one coal-fired test showed 60-70 percent NO_{χ} and 90 percent SO_2 reductions are possible.¹¹ The longest continuous operation was for 52 hours and the absorption section was the only part of the process tested. Pilot-plant testing was discontinued after two months. Plans are being made for further coal-fired pilot tests on the integrated system.

Absorption-reduction processes are readily applicable only to high sulfur coals as a $SO_2:NO_{\times}$ mole ratio in the flue gas of at least 3-5 is required for maximum performance. This can easily be shown by observing reactions 2-24 and 2-26 reprinted below.

$$SO_2(aq) + 2CH_3COONa(aq) + H_2O \rightarrow Na_2SO_3(aq) + 2CH_3COOH(aq)$$

One can see that 1 mole of SO_2 absorbed in solution reacts to form 1 mole of sodium sulfite (Na₂SO₃). Then, 5 moles of sodium sulfite are required to reduce 2 moles of NO. So, the minimum stoichiometric $SO_2:NO_x$ mole ratio required is $\frac{5}{2}$ or 2.5. Also, some of the sodium sulfite is oxidized to sodium sulfate by oxygen present in the flue gas according to:

$$Na_2SO_3(aq) + \frac{1}{2}O_2(aq) \rightarrow Na_2SO_4(aq)$$
 (2-33)

and is not available for NO_{\times} reduction. Low-sulfur coals would require SO_2 to be added to the flue gas for these processes to perform; therefore, they should be considered applicable to high sulfur coals only.

Absorption-reduction processes require large absorbers with high liquid rates due to relative insolubility of NO, even when the absorption catalyst is used. Also, the regeneration of the absorption catalyst and the flue gas reheat for plume buoyancy are energy intensive. Some corrosion-resistant material is necessary due to the corrosive nature of the absorbing solution. However, absorption-reduction appears to be the most promising of the "wet" NO_X/SO_2 removal processes. This is due primarily to its not utilizing oxidants which require much corrosion-resistant material and, more importantly, create serious secondary pollution problems. Also, the primary by-products of absorption-reduction processes, gypsum, can be used as landfill material or in building materials. For all the above reasons, absorption-reduction processes appear, at this preliminary stage, competitive with other wet $NO_{\rm x}/SO_2$ removal processes.

2.2.9 Oxidation-Absorption-Reduction

2.2.9.1 System Description--

Oxidation-absorption-reduction processes simultaneously remove NO_{χ} and SO_2 from flue gas by oxidizing relatively insoluble NO to relatively soluble NO_2 and then absorbing both NO_2 and SO_2 into a scrubbing solution. The processes are based on the use of gas-phase oxidants, either ozone (O₃) or chlorine dioxide (ClO₂), to selectively oxidize NO to NO_2 . Both perforated-plate and packed bed absorption columns are utilized by various process vendors.

Most of the oxidation-absorption-reduction processes are similar in that they consist of five major sections:

- prescrubbing
- gas-phase oxidation
- NO₂/SO₂ absorption
- reduction of absorbed NO₂ and oxidation of SO_3^-
- wastewater treatment

The areas where processes differ are gas-phase oxidation - O_3 or ClO_2 ; absorption solutions - limestone slurry (CaCO₃), H₂SO₄, or NaOH; and the amount and type of waste treatment required. Thermal decomposition, biological denitrification, or wastewater evaporation wastewater treatment systems can be used. Because of these differences, only one of the oxidationabsorption-reduction processes, that of Mitsubishi Heavy Industries, is examined here in detail. A block flow diagram of the MHI oxidation-absorption-reduction process is shown in Figure 2.2.9-1.

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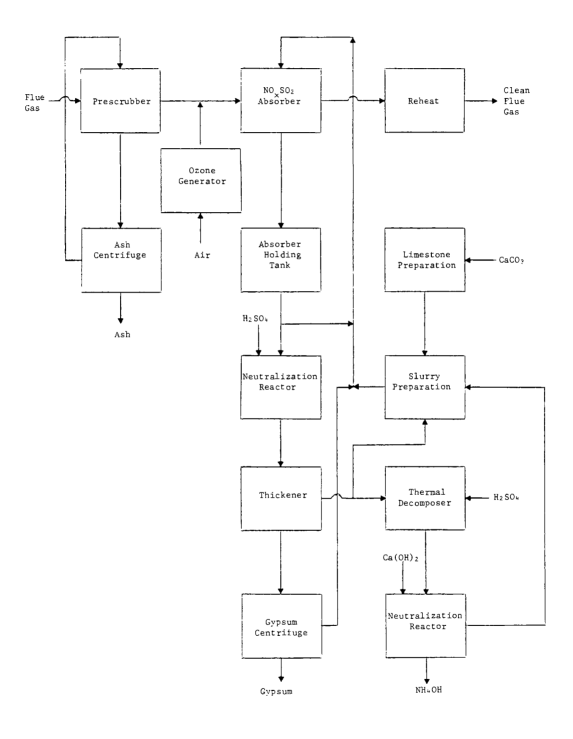


Figure 2.2.9-1. Process flow diagram for MHI oxidationabsorption-reduction process.⁸¹

Flue gas is taken from the boiler after the air preheater and passed through a prescrubber to cool the gas and remove particulates and chlorides. The flue gas then enters a duct where it is injected with ozone (about 1 percent by weight in air)⁸² such that the 03:NO ratio is 1:1. Ozone selectively oxidizes NO by the following reatcion.⁸³

$$NO(g) + O_3(g) \rightarrow NO_2(g) + O_2(g)$$
 (2-34)

After injection, the flue gas passes countercurrent to a lime/limestone slurry in a grid-packed absorption column. A water-soluble catalyst is added to the slurry to enhance NO_2 absorption (even though NO_2 is more soluble than NO, it is still less soluble than SO_2). SO_2 is absorbed quickly at the bottom of the column and undergoes the following reactions.¹⁵

$$SO_2(g) \rightarrow SO_2(aq)$$
 (2-23)

$$SO_2(aq) + CaCO_3(s) + {}_{2}H_20 \rightarrow CaSO_3 \cdot {}_{2}H_2O(s) + CO_2(g)$$
 (2-35)

$$SO_2(aq) + CaSO_3(aq) + H_2O \rightarrow Ca(HSO_3)_2(aq)$$
 (2-36)

 $\rm NO_2$ is absorbed gradually over the length of the column and reacts as follows. $^{\rm 16}$

$$2NO_{2}(g) + Ca(OH)_{2}(s) + CaSO_{3} \cdot \frac{1}{2}H_{2}O(s) + \frac{1}{2}H_{2}O \rightarrow Ca(NO_{2})_{2}(aq) + CaSO_{4} 2H_{2}O(s)$$

$$(2-37)$$

Once both the NO₂ and SO₂ are absorbed, the nitrite ion formed by reaction 2-37 is reduced by the bisulfate ion formed by reaction $2-36.^{84}$

$$Ca(NO_2)_2(aq) + 3Ca(HSO_3)_2(aq) \rightarrow 2Ca[NOH(SO_3)_2](aq) + 2CaSO_3 \cdot \frac{1}{2}H_2O(s) + H_2O$$

(2-38)

These hydroxylamine $[NOH(SO_3)_2^{=}]$ compounds are reduced further by the sulfite ion⁸⁵

$$Ca[NOH(SO_3)_2](aq) + CaSO_3 \cdot \frac{1}{2}H_2O(s) + \frac{3}{2}H_2O \rightarrow Ca[NH(SO_3)_2](aq) + CaSO_4 \cdot 2H_2O(s) + (2-39)$$

Upon leaving the top of the absorber, the clean flue gas is reheated for plume buoyancy and sent to the stack. The slurry solution drops to a holding tank from which most of the solution is returned to the top of the absorber. A small stream passes to a neutralization reactor where sulfuric acid is added to convert the sulfite solid to soluble bisulfite and solid gypsum:⁸⁶

$$2\text{CaSO}_{3} \cdot \frac{1}{2}\text{H}_{2}O(s) + \text{H}_{2}SO_{4}(aq) + \text{H}_{2}O \rightarrow \text{CaSO}_{4} \cdot 2\text{H}_{2}O(s) + \text{Ca}(\text{HSO}_{3})_{2}(aq)$$
(2-40)

This stream passes to a thickener from which the bottoms are sent to a centrifuge to separate the solid gypsum by-product from the liquor which is returned to the absorber. The overflow from the thickener is primarily recycled to the limestone slurry preparation tank. The remainder is sent to a thermal decomposer where sulfuric acid is added to hydrolyze the N-S compounds.¹⁸

$$2Ca[NH(SO_3)_2](aq) + 2H_2O \xrightarrow{H^+} Ca(NH_2SO_3)_2(aq) + Ca(HSO_4)_2(aq)$$
 (2-41)

 $Ca(NH_2SO_3)_2(aq) + Ca(HSO_4)_2(aq) + 6H_2O \xrightarrow{H^+} 2NH_4HSO_4(aq) + 2CaSO_4 \cdot 2H_2O(s) \downarrow$ (2-42)

The ammonium bisulfate solution is pumped to another neutralization reactor where lime is added.⁸⁷

$$\mathrm{NH_4HSO_4(aq)} + \mathrm{Ca(OH)_2} + \mathrm{H_2O} \rightarrow \mathrm{CaSO_4 \cdot 2H_2O(s)} + \mathrm{NH_4OH(aq)} \qquad (2-43)$$

MHI has three possible methods of removing this ammonium hydroxide:

- decompose by increasing pH
- decompose thermally
- strip out with makeup H₂O

The remaining gypsum slurry is pumped to the limestone slurry preparation tank.

The fundamental design equation used for gas absorption column design is 32

$$\int_{Y_{a}}^{Y_{b}} \frac{dy}{(y-y^{\star})} = \left(\frac{K_{a}}{G_{y}}\right) Z$$
(2-11)

- where y = bulk NO_x concentration (mole fraction) of gas phase at any given point in column
 - y-y* = overall driving force for absorption (y* being the NO_x concentration of a gas in equilibrium with a given liquid NO_x concentration)
 - $Y_{\rm b}$ = inlet NO_x concentration

- Ky = overall gas-phase mass transfer coefficient, lb-moles NO_x/
 (ft²)(hr)(mole fraction)
- a = area of gas-liquid interface per unit packed volume, ft^2/ft^3
- G_y = molal gas mass velocity, lb-moles flue gas/(ft²)(hr)
- Z = length of packed section of column, ft

In a column containing a given plate or packing configuration and being irrigated with a certain liquid flow, there is an upper limit to the gas flow rate. This limit's superficial gas velocity (volumetric gas flow rate/cross-sectional area of column) is called the flooding velocity. At this point, the gas flow completely impedes the downward motion of the

liquid and blows the liquid out of the top of the column. The gas velocity obviously, must be lower than the flooding velocity. How much lower is a design decision. Often, it is an economic tradeoff between power costs and equipment costs. A low gas velocity will lower the pressure drop and, hence, the power costs but the absorber will have a larger diameter and cost more. High gas velocities have an opposite effect. Usually the optimum gas velocity is about one-half the flooding velocity.³³ The height of the column depends on the desired level of removal and on the rate of mass transfer. The latter consideration is why oxidation-absorption-reduction processes oxidize NO to more soluble NO₂ prior to the absorber and why some processes add water soluble catalysts to the scrubbing solution to aid NO2 absorption. The oxidation step enables these processes to use shorter absorbers with lower liquid rates than either the absorption-oxidation or absorption-reduction processes. Table 2.2.9-1 presents the effects of boiler/flue gas variables on the design of oxidation-absorption-reduction systems. Both flue gas flow rate and NO, concentration can be affected by boiler operating conditions. Therefore a change in load on an industrial boiler may alter these variables markedly. The absorber must be designed to accommodate any anticipated load change. The column size and the liquid, oxidant, and catalyst flows must be designed for each application after examining the boiler operating history and establishing ranges of variation.

Typical ranges for several operating parameters for this type of process are shown in Table 2.2.9-2. Reagent concentrations were not available. Economics for the various processes cover a wide range presumably due to different techniques for oxidant generation and treatment of the scrubbing solution. Costs are reported to range from \$84-134/kW for utility applications' capital expense and 6.7-9 mills/kWh for operating expense.⁹¹

Presently, some of the oxidation-absorption-reduction processes have reached the prototype stage of development. Table 2.2.9-3 presents a list of oxidation-absorption-reduction process vendors and the status of development of their projects. One can see from the table that no coal-fired flue gas tests have been made as of yet.

Variable	Design Effect
Presence of particulates	Requires prescrubber
Presence of SO ₂	Depends on individual process: if NO_2 is com- pletely reduced to N_2 or NH_3 by SO_3^- (as does MHI), then at least the stoichiometric $SO_2:NO_{\chi}$ mole ratio of 3:1 is required ⁸⁸ [see equation (9-6)]; if NO_2^- is not reduced completely, then a different ratio will be necessary
Increased gas flow	Requires larger column diameter; increased liquid flow rate
Increased NO _× concentration	Requires larger column height; increased gas- phase oxidant flow rate; increased liquid- phase catalyst concentration

TABLE 2.2.9-1. SYSTEM DESIGN CONSIDERATIONS

TABLE 2.2.9-2. TYPICAL RANGES OF OPERATING VARIABLES FOR OXIDATION-ABSORPTION-REDUCTION PROCESSES^{89,90}

Variable		Range	
Liquid/Gas Ratio, 1/Nm ³		2–12	
Oxidant/NO Mole Ratio	O ₃ systems	0.6-1.0	
	C10 ₂ systems	0.55	
SO_2/NO_{\times} Mole Ratio		2.5-5.0	
Superficial Gas Velocity	v, m/s	3-5	
Pressure Drop, mmH_2O		200-500	

Vendor	Status of Development
Chiyoda	1975: 1000 Nm ³ /hr flue gas from heavy oil- fired boiler
Ishikawajima-Harima Heavy Industries	1975: 5000 Nm ³ /hr flue gas from oil-fired boiler (3000 hours continuous)
Mitsubishi Heavy Industries	1975: 2000 Nm ³ /hr flue gas from heavy oil- fired boiler (700 hours continuous)
Osaka Soda	1976: 60,000 Nm ³ /hr flue gas from oil-fired boiler
Shirogane	1974: 48,000 Nm ³ /hr flue gas from oil-fired boiler
Sumitomo Metal-Fujikasui: Calcium Process	1976: 25,000 Nm ³ /hr flue gas from sintering furnace
Sumitomo Metal-Fujikasui: Sodium Process	1973: 62,000 Nm ³ /hr flue gas from heavy oil- fired boiler (5 others)

TABLE 2.2.9-3. PROCESS VENDORS OF OXIDATION-ABSORPTION-REDUCTION PROCESSES^{92,93}

2.2.9.2 System Performance--

No coal-fired testing has been performed. Results of oil-fired tests show up to 90 percent NO $_{\rm v}$ reduction and >95 percent SO $_2$ reduction.

The primary disadvantage of these processes is the utilization of costly gas-phase oxidants which create secondary wastewater pollution problems. Both ozone and chlorine dioxide are highly unstable so they cannot be stored and must be generated onsite. O_3 , the more expensive of the two, is generated by a high energy corona discharge in air. This instantaneous process requires significantly large amounts of electricity. ClO_2 is generated by a slower chemical reaction (requires about 20 minutes to respond to a change in demand) which could make it less responsive to boiler load changes. The use of ClO_2 introduces an additional secondary pollutant, chlorides, besides the nitrite salt problem. Significant amounts of corrosion-resistant material are required for oxidation-absorption-reduction processes, regardless of which oxidant is utilized. Some of the processes would not be applicable to low sulfur coals as they require large amounts of SO₂ to obtain NO₂(aq) or NO₂ reduction.

2.2.10 Oxidation-Absorption

2.2.10.1 System Description--

As a group, oxidation-absorption processes include those oxidation processes which do not qualify for the oxidation-absorption-reduction category. Basically, there are two types of oxidation-absorption processes. One is a simplified version of the oxidation-absorption-reduction process and uses an excess of ozone to selectively oxidize NO_{χ} to N_2O_5 which is absorbed into aqueous solution and concentrated to form a 60 percent nitric acid (HNO₃) by-product. There is no reduction of $NO_{\chi}(NO_2)$ by the absorption of SO_2 (as SO_3^{-}) and no wastewater treatment facility. The other type of oxidation-absorption process is based on equimolar NO-NO₂ absorption: absorbing N_2O_3 which is formed by the gas-phase reaction of NO and NO₂.

A flow diagram of the Kawasaki Heavy Industries oxidation-absorption process is shown in Figure 2.2.10-1. Flue gas is taken from the boiler after the air preheater. It passes countercurrent to a magnesium hydroxide $[Mg(OH)_2]$ slurry in the first section of the absorber. There, SO₂ is absorbed and undergoes the following reactions.⁹⁵

$$SO_2(g) \rightarrow SO_2(aq)$$
 (2-23)

$$Mg(OH)_{2}(s) + SO_{2}(aq) + 5H_{2}O \rightarrow MgSO_{3} \cdot 6H_{2}O(s) \downarrow$$
(2-44)

The SO_2 -free flue gas passes to the first denitrification section of the absorber while the liquid effluent drops to a holding tank. A recycle NO_2 stream is added to the flue gas to bring the $NO:NO_2$ mole ratio to 1:1. The

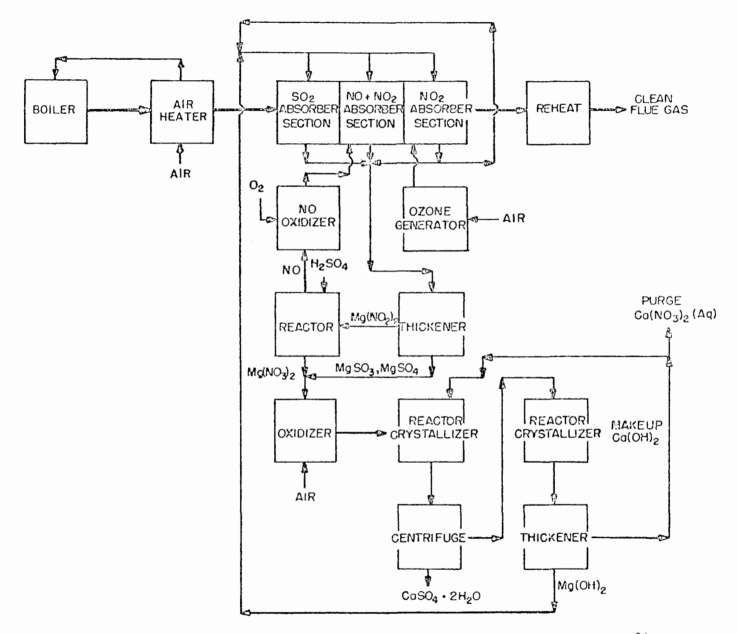


Figure 2.2.10-1. Flow diagram of Kawasaki Heavy Industries process.94

resulting mixture then passes countercurrent to a $Mg(OH)_2$ slurry. Equimolar amounts of NO and NO₂ react and are absorbed in the following manner.⁹⁶

$$NO(g) + NO_2(g) \rightarrow N_2O_3(g)$$
 (2-45)

$$N_2O_3(g) \rightarrow N_2O_3(aq)$$
 (2-46)

$$Mg(OH)_2(aq) + N_2O_3(aq) \rightarrow Mg(NO_2)_2(aq) + H_2O$$
 (2-47)

The flue gas passes out of the top of this absorption section while the liquid effluent drops to the holding tank. Because the rate of reaction 2-45 decreases with NO_{χ} concentration (below 200 ppm it becomes negligible), it is necessary to further reduce NO_{χ} by injecting ozone to oxidize the remaining NO to NO_2 . The mixture then passes to the final denitrification section of the absorber and is passed countercurrent to a Mg(OH)₂ slurry. This section of the absorber is described by the following reactions.⁹⁷

$$2NO_2(g) \rightarrow N_2O_4(g)$$
 (2-8)

$$N_2O_4(g) \rightarrow N_2O_4(aq)$$
 (2-9)

$$2N_2O_4(aq) + 2Mg(OH)_2(s) \rightarrow Mg(NO_3)_2(aq) + Mg(NO_2)_2(aq) + 2H_2O$$
 (2-48)

The clean flue gas leaves the top of this absorber section, is passed to a reheater for plume buoyancy and sent to the stack. Part of the liquid effluent from this section is recycled to the tops of the absorber sections while the rest drops to the holding tank. The slurry solution is pumped to a thickener which separates the soluble nitrite (NO_2) and nitrate (NO_3) salts from the solid magnesium sulfite. The overflow from the thickener passes to a NO_2 decomposition reactor where sulfuric acid is added.⁹⁸

$$3Mg(NO_2)_2(aq) + 2H_2SO_4(aq) \rightarrow 2MgSO_4(aq) + Mg(NO_3)_2(aq) + 4NO(g) \uparrow + 2H_2O$$

(2-49)

The NO off-gas passes through an oxidizer where it is oxidized by air to NO_2 and sent to the first denitrification section of the absorber. The effluent from the decomposition reactor is mixed with the thickener bottoms and pumped to a second oxidizer.⁹⁹

$$MgSO_3 \cdot 6H_2O(s) + \frac{1}{2}O_2(g) \rightarrow MgSO_4(aq) + 6H_2O$$
 (2-50)

The magnesium sulfate formed in the oxidizer is treated with calcium nitrate in a gypsum production reactor.²⁵

$$Ca(NO_3)_2(aq) + MgSO_4(aq) + 2H_2O \rightarrow CaSO_4 \cdot 2H_2O(s) \downarrow + Mg(NO_3)_2(aq)$$

(2-51)

The products of this reaction are sent to a centrifuge to remove the solid gypsum by-product. The liquid from the centrifuge goes to another decomposition reactor where makeup lime slurry is added.¹⁰⁰

$$Mg(NO_3)_2(aq) + Ca(OH)_2(s) \rightarrow Ca(NO_3)_2(aq) + Mg(OH)_2(s) \qquad (2-52)$$

The magnesium hydroxide product is separated in a thickener and recycled to the absorbers. The thickener overflow stream is split and part is recycled to the gypsum production reactor and the rest is concentrated to form low-grade liquid fertilizer by-product, $Ca(NO_3)_2$.

Since the processes in this category are all very different, especially with respect to chemistry, generalization of typical ranges of operating variables is not meaningful and, therefore, not presented. No published economics for these processes were found.

Presently, the equimolar absorption-type oxidation-absorption processes are still in the pilot-unit stage of development. Table 2.2.10-1 presents a list of all oxidation-absorption process vendors and their project's status of development.

Vendor	Status of Development
Kawasaki Heavy Industries	1975: 5000 Nm ³ /hr flue gas from coal- fired boiler
Tokyo Electric-MHI (NO _× only)	1974: 100,000 Nm ³ /hr flue gas from natural gas-fired boiler
Ube Industries	No information available

TABLE 2.2.10-1. PROCESS VENDORS OF OXIDATION-ABSORPTION PROCESSES¹⁰⁰

2.2.10.2 System Performance--

Only one coal-fired test has been performed. No information has been published on any of the tests conducted.

The production of nitrate salts poses a potential secondary pollution problem. The plan for reclaiming and concentrating the nitrates as $Ca(NO_3)_2(aq)$ for liquid fertilizer is questionable as the by-product is of low quality and may not be easily marketable in the U.S. Also, the gypsum by-product would be contaminated with various nitrate and sulfite salts, and therefore, would probably be useful only as landfill material. Much corrosion-resistant material is necessary due to the utilization of ozone and circulating magnesium slurries. The three absorber sections, with their respective operating conditions, and ozone generation present complex process control problems. The process steps of several absorber sections in series (large fan requirements), ozone generation (corona discharge), flue gas reheat (inline heater), and by-product and wastewater treatment are all energy intensive and present technical and economic disadvantages when compared to other simpler FGT processes.

2.3 CONTROLS FOR OIL-FIRED BOILERS

2.3.1 Selective Catalytic Reduction-Fixed Packed Bed Reactors

2.3.1.1 System Description--

Fixed packed bed systems are applicable only to flue gas streams containing less than 20 mg/Nm³ of particulates. As such, they are applicable to distillate oil-fired boilers (19 mg/Nm³) but not to residual oil-fired boilers (330 mg/Nm³).

The primary feature of these systems is the reactor which contains the catalyst. As the name implies, the granular catalyst is randomly packed in a stationary bed. An example of a typical fixed bed reactor is shown in Figure 2.3.1-1. The important features of the reactor are:

- the catalyst
- the catalyst support
- the gas distributor

The catalyst can be either spherical or cylindrical in shape. Spherical granules typically range in size from 4-10 mm in diameter.¹⁰³ The composition varies from process to process and most formulations are proprietary. All of the catalysts considered here for use in treating flue gas containing SO_2 and SO_3 are resistant to poisoning by these compounds. Long term tests of these catalysts in the presence of SO_{\times} have shown very little or no decrease in activity or selectivity. The catalyst is supported either by inert packing (as shown in Figure 2.3.1-1) or by a perforated support plate (Figure 2.3.1-2).

The catalyst supports hold the catalyst fixed in place in order to prevent both mobilization of the particles by the gas stream and catalyst rearrangement which would allow channelling of the flue gas. The gas distributor can be a perforated plate or similar device which spreads the gas flow across the entire cross-section of the catalyst bed.

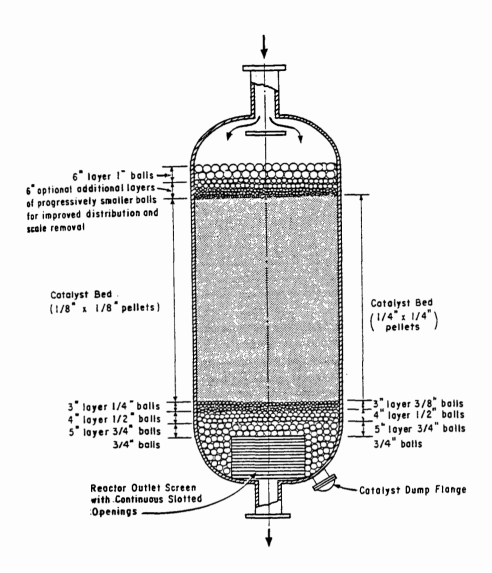


Figure 2.3.1-1. Example of typical fixed packed bed reactor.¹⁰¹

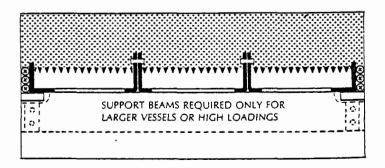


Figure 2.3.1-2. Example of catalyst support plate.¹⁰²

A typical fixed bed SCR process layout is presented in Figure 2.3.1-3. Several arrangements are possible, however, for application to new boilers this arrangement is the most desirable.⁸

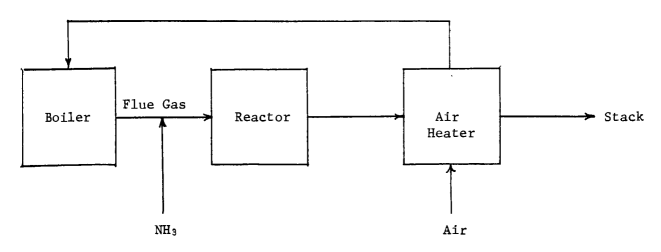


Figure 2.3.1-3. Process layout for fixed bed SCR process.

The principle of operation of these systems involves a gas phase reaction between ammonia (NH₃) and NO_{\times} (NO and NO₂). These reactions are presented most accurately by¹²

$$4NO + 4NH_3 + O_2 \neq 4N_2 + 6H_2O$$
 (2-1)

$$2NO_2 + 4NH_3 + O_2 \neq 3N_2 + 6H_2O$$
 (2-2)

The first reaction predominates since flue gas NO_{\times} is typically 90-95 percent NO. As shown, the NO_{\times} is reduced to molecular nitrogen (N₂) which exits with the flue gas stream.

The primary design equation used with these processes is the standard equation for reactor design, 13

$$\frac{V}{F} = \int_{0}^{x} \frac{dx}{r}$$
(2-3)

where V is the catalyst volume F is the mass (or molar) flow rate x is the conversion of NO_x to N_2 r is the reaction rate <u>mass (or moles)</u> volume of catalyst x time

The reaction rate, r, for each NO reduction reaction can be represented by

$$r = k[NH_3]^{a}[NO]^{b}[O_2]^{c}$$
(2-4)

where k is the reaction rate constant

[NH₃], [NO], [O₂] are the reactant concentrations, and a, b, c are empirically determined exponents

The catalyst volume can also be determined if the space velocity is known for the catalyst and removal level of interest. The space velocity is defined as the gas flow rate divided by the catalyst volume. The reaction rate is different for each catalyst formulation and therefore, values for k, a, b, and c must be determined for the particular catalyst to be used before any design can be performed. The reaction rate constant is usually described by the Arrhenius equation.

$$k = Ae^{-\frac{E}{RT}}$$
(2-5)

where A is the frequency factor E is the activation energy R is the universal gas constant, and T is the temperature Values for k, a, b and c for two catalyst formulations are shown in Table 2.3.1-1. Values for other catalyst formulations will be different. The most important design and operating variables are similar to those for moving bed systems using granular catalysts. These are listed, along with typical ranges, in Table 2.3.1-2.

Other variables that affect the process are

- flue gas flow rate
- NO_v control level
- NO_v concentration
- boiler load variation

The flue gas flow rate and control level determine the catalyst volume (hence reactor size) necessary. Increases in either also increases the reactor size. The NO_x concentration is a function of fuel type used in the standard boilers. Higher concentrations require larger NH_3 storage and vaporization equipment; reactor size is not affected. Boiler load can affect several things including flue gas temperature, flow rate and NO_x concentration. It is necessary to maintain reactions temperatures of 350 to 400°C and temperature control equipment may be necessary if the boiler experiences large load variations. Where these variations are present, some equipment overdesign may be warranted to insure a constant control level. These variables are discussed in more detail in the section on moving bed SCR systems for coal-fired boilers, Section 2.3.2. Costs of fixed packed bed systems range from 16-49/kW (capital) and 1.2-1.8 mills/kWh (operating). These costs are based on utility applications as well as a variety of processes and operating conditions.

There are vendors of fixed packed bed SCR systems and all are Japanese. Vendors are listed in Table 2.3.1-3 and the scale of development is also noted. Fixed packed systems have been applied to industrial but not utility boilers in Japan. Existing and planned installations are shown in Table 2.3.1-4. Currently, there are no installations in the U.S.

TABLE 2.3.1-1. REACTION RATE DATA FOR TWO CATALYST FORMULATIONS¹¹

Catalyst: V_2O_5 on Al_2O_3 $k = 2.05 \times 10^3 e^{-\frac{9650}{RT}}$ a = 0.30 b = 0.22 c = 0.05Catalyst: Fe-Cr on Al_2O_3 $k = 3.25 \times 10^3 e^{-\frac{10,860}{RT}}$ a = 0.45 b' = 0.10c = 0.15

TABLE 2.3.1-2. DESIGN AND OPERATING VARIABLES FOR FIXED PACKED BED SYSTEMS¹⁴

Variable	Typical Range
Gas Velocity, m/s	1 - 1.5
Bed Depth, m	0.2 - 0.6
Space Velocity, hr ⁻¹	6,000 - 10,000
Pressure Drop, mmH_2O	40 - 80
Temperature, °C	350 - 400

Vendor	Notes
Sumitomo Chemical	Tested on commercial scale equipment
Hitachi Zosen	Tested on commercial scale equipment
Hitachi, Ltd.	Tested on commercial scale equipment
Mitsubishi Heavy Industries	Tested on commercial scale equipment
Ishikawjima-Harima Heavy Industries	Tested on commercial scale equipment
Mitsui Toatsu Chemical	Has not been applied to boilers
Kawasaki Heavy Industries	Tested on pilot scale equipment
Mitsubishi Kakoki Kaisha	Tested on commercial scale equipment

TABLE 2.3.1-3. VENDORS OF SCR FIXED BED SYSTEMS FOR OIL-FIRED APPLICATIONS²¹

2.3.1.2 System Performance--

Typical performance data for packed fixed bed SCR systems are shown in Figures 2.3.1-4 and 2.3.1-5 and Tables 2.3.1-5 through 2.3.1-7. These data indicate that NO_{χ} removals up to 90 percent are achievable with these systems. This allows them to be considered for all control levels of interest in this study.

There are some potential problems downstream of the SCR systems (fixed packed bed, moving, and parallel flow) due to the presence of the unreacted ammonia in the flue gas. Two things can happen: 1) the NH₃ can react with SO_2 or SO_3 to form ammonium bisulfate or ammonium sulfate or 2) the NH₃ can enter the downstream equipment unreacted. The bisulfate has been shown to cause air preheater pluggage and this is the subject of ongoing research both at the EPA and the Electric Power Research Institute (EPRI). Both the bisulfate and sulfate exist as a particulate, but may be difficult to collect if the particles are submicron in size. Unreacted NH₃ is not likely to present any operational problems. A recent study has shown that if an ESP exists downstream, then most of the NH₃ will exit with the ash. NH₃ can actually improve the performance of an FGD system.¹²⁹

Location (Japan)	User	Process Developer	Fue1	Capacity (Nm ³ /hr)	Completion Date
Amagasaki	Kansai Paint	Hitachi, Ltd.	Distillate	16,000	October 1977
Amagasaki	Nisshin Steel	Hitachi, Ltd.	Resid	20,000	August 1977
Amagasaki	Nisshin Steel	Hitachi, Ltd.	Resid	19,000	July 1977
Sakai	Nisshin Steel	Hitachi, Ltd.	Distillate	30,000	December 1978
Hokkaichi	Shindaikyowa P.C.	Hitachi Zosen	Resid	440,000	November 1975
Sodegaura	Sumitomo Chemical	Sumitomo Chemical Eng.	Resid	30,000	July 1973
Sodegaura	Sumitomo Chemical	Sumitomo Chemical Eng.	Resid	240,000	March 1976
Sorami	Toho Gas	Sumitomo Chemical Eng.	Distillate	62,000	October 1977
Sorami	Toho Gas	Sumitomo Chemical Eng.	Distillate	23,000	December 1977
Sorami	Toho Gas	Sumitomo Chemical Eng.	Distillate	23,000	June 1978
Sorami	Toho Gas	Sumitomo Chemical Eng.	Distillate	19,000	July 1978
Kawasaki	Nippon Yakin	Mitsubishi Kakoki	Resid	14,000	July 1978
Kawasaki	Toho Gas	Mitsubishi Kakoki	Distillate	30,000	October 1977
Chita	Toho Gas	Mitsubishi Kakoki	Distillate	30,000	October 1977

TABLE 2.3.1-4. EXISTING FGT INSTALLATIONS OF SCR FIXED BED SYSTEMS OIL-FIRED INDUSTRIAL BOILERS²¹

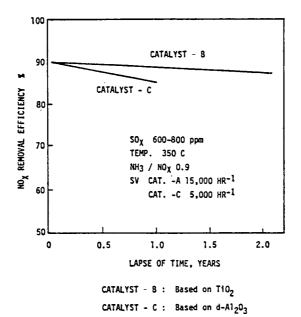
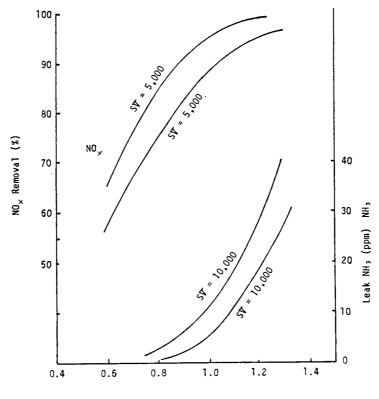


Figure 2.3.1-4. Performance of experimental catalyst of Sumitomo Chemical.¹⁰⁴



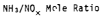


Figure 2.3.1-5. Typical example of operation data (oil-fired boiler, 350-400°C, granular or honeycomb catalyst).¹⁰⁵

	Idemitsu Kosan	Shindaikyowa Petrochemical	Kawasaki Steel
Completed	Oct. 1975	Nov. 1975	Nov. 1976
Plant site	Chiba	Yokkaichi	Chiba
Gas source	FCC-CO Boiler and furnace	Oil-fired Boiler	Iron-ore Sintering machine
Capacity (Nm ³ /hr)	350,000	440,000	762,000
Load factor (%)	50-100	50-100	70-100
Pretreatment of gas	Heating	EP*, FGD, Heating	EP, FGD WEP [†] , Heating
Reactor inlet			
NO _× (ppm)	230	150	200-300
SO _× (ppm)	50-80	80-130	5-20
Dust (mg/Nm ³)	20-50	30-100	3-10
02 (%)	2.3	3.2	11.2
Reactor type	Fixed bed	Fixed bed	Fixed bed
Reaction temp.	400	420	
NO _× /NH ₃ ratio	1.0	1.0	1.0
Catalyst No.	204	304	304
$SV (hr^{-1})$	5,000	10,000	4,000
NO _x removal (%)	93	80 [‡]	95
Pressure drop by SCR reactor (mmH ₂ O)	170	160	50
Catalyst life	1 year	l year	l year

TABLE 2.3.1-5. OPERATION PARAMETERS OF MAJOR PLANTS CONSTRUCTED BY HITACHI ZOSEN¹⁰⁶

*Electrostatic precipitator

 † Wet electrostatic precipitator

[‡]Including leakage in heat exchanger

	Mitsui Petro- Chemical Co.
Capacity (Nm ³ /hr)	200,000
Gas composition	
NO _× (ppm)	190
SO _x (ppm)	None
Dust (mg/Nm ³)	20-50
Catalyst and reactor	
Catalyst carrier	$A1_20_3$
Catalyst shape	Granule
$SV (hr^{-1})$	2,600
Temperature (°C)	350
$\rm NH_3/NO_{\times}$ mole ratio	1.0
NO _x removal (%)	Above 90
Total pressure drop (mmH_20)	
Leak NH3 (ppm)	
Operation start	Oct. 1975
Plant cost (10 ⁶ yen)	
Denitrification cost	
(yen/kWhr)*	

TABLE 2.3.1-6. SCR PLANT BY MITSUI ENGINEERING & SHIPBUILDING CO.¹⁰⁷

*Including 7 years depreciation.

	Pilot	Commercial
Gas for SCR (Nm ³ /hr)	30,000	240,000
Fuel	0i1(S=0.7%)	0i1(S=0.7%)
Load fluctuation	60-100%	60-100%
Stack height (m)	70	140
Inlet gas composition		
02 (%)	6	6
SO _× (ppm)	400	400
NO _× (ppm)	200	200
Particulates after EP (mg/Nm^3)	5-20	5-10
FGD unit	None	None
$SV (hr^{-1})$	5,000	5,000
Temperature (°C)	320	320
NO _× removal (%)	Over 90	Over 90
NH ₃ /NO mole ratio	1.0	1.0
Leak ammonia (ppm)	10-20	10-20
Type of reactor	Fixed bed	Fixed bed
Pressure drop (mmH_2O)		
Reactor		200
Total system		500
Plant completed	July 1973	March 1976

TABLE 2.3.1-7. OPERATION DATA OF SCR PLANTS FOR DIRTY GAS¹⁰⁸

2.3.2 Selective Catalytic Reduction-Moving Bed Reactor

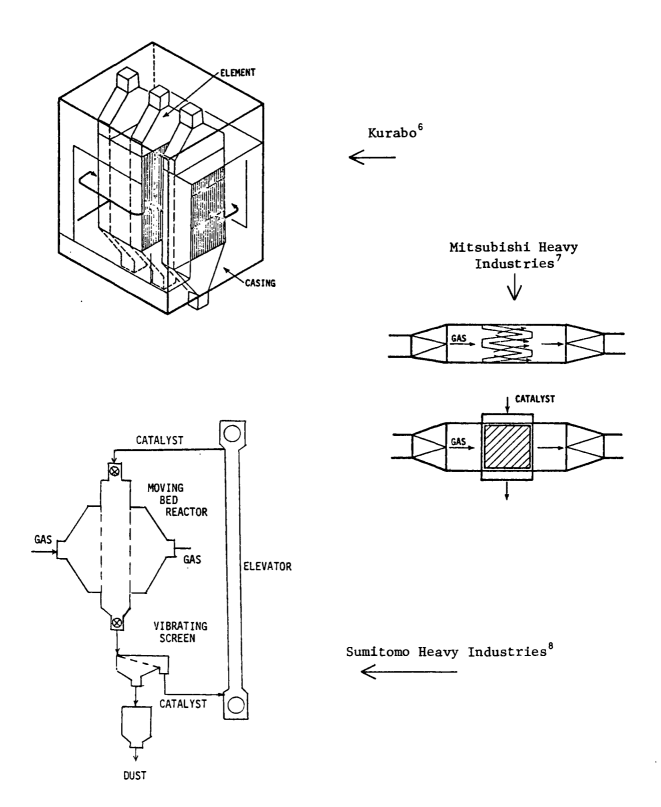
2.3.2.1 System Description--

The primary feature of this and other selective catalytic reduction (SCR) processes is the reactor. The reactor contains the catalyst which allows the reduction reaction to proceed at 350-400°C. In this case the reactor is a moving-bed type in which a portion of the catalyst is either continuously or intermittently removed from service in order to remove the accumulated particulates. Some moving bed reactors are shown in Figure 2.3.2-1. The actual reactor arrangement is highly process specific, however, the principle of operation is the same for all of the processes.

For moderate particulate loadings the bed is moved intermittently and operated as a fixed on bed system most of the time. High particulate loadings require that the bed be moved continuously. Moving bed reactors are reportedly capable of handling up to 2 g/Nm³ of particulates. However, this is more a theoretical than a practical particulate load limit.⁹ If possible, this would be sufficient to handle the residual oil-fired boilers (0.33 g/Nm³).

Entrained particulates are generally removed from the catalyst bed by vibrating the bed and screening the catalyst or some other mechanical means. Particulate removal by the bed can be as high as 70 percent.¹⁰

An example flow diagram for a moving bed SCR process is shown in Figure 2.3.2-2. Flue gas is taken from the boiler between the boiler and the air preheater. An economizer bypass is utilized for temperature control. Ammonia is injected and mixed with the flue gas stream just prior to the reactor. The flue gas passes through the catalyst bed where NO_{χ} is reduced to N_2 and is then sent to the stack or further treatment facilities. The catalyst circulates through the reactor and is screened to remove particulates. The particulates are blown to a small baghouse where they are collected.





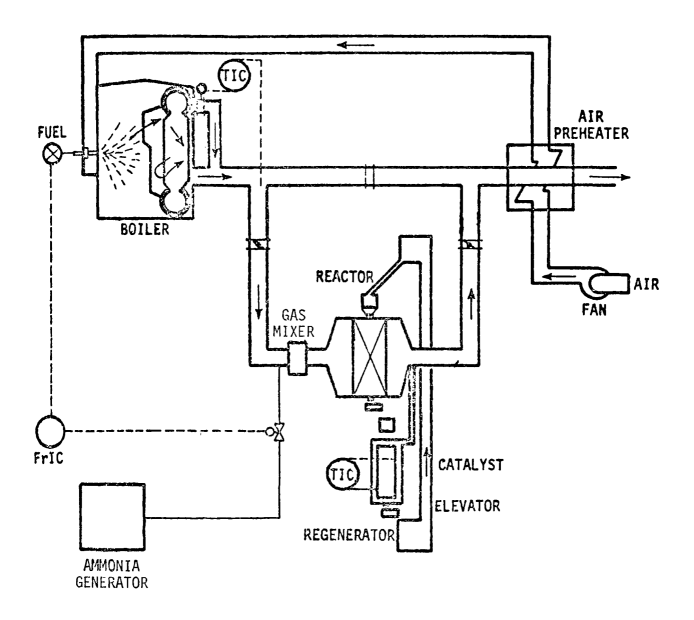


Figure 2.3.2-2. Process flow diagram for moving bed SCR process.¹¹

The NO reduction reactions are represented most accurately by¹²

$$4NO + 4NH_3 + O_2 \stackrel{2}{\leftarrow} 4N_2 + 6H_2O$$
 (2-1)

$$2NO_2 + 4NH_3 + O_2 \stackrel{\rightarrow}{\leftarrow} 3N_2 + 6H_2O$$
 (2-2)

The first reaction predominates since flue gas NO_{χ} typically consists of 90-95 percent NO. As shown, the NO_{χ} is reduced to molecular nitrogen, N_2 , which exits with the flue gas. O_2 is also a reactant, but is in large excess (>3 percent) in the flue gas.

The fundamental design equation used for catalytic reactor design is¹³

$$\frac{V}{F} = \int_{0}^{x} \frac{dx}{r}$$
(2-3)

where V = catalyst volume
F = mass flow rate (or molar flow rate)
x = conversion of NO_x to N₂
r = reaction rate, mass (or moles)
volume of catalyst x time

The reaction rate, r, for each of the NO_{\times} reduction reactions can be represented by

$$r = k [NH_3]^a [NO]^b [O_2]^c$$
 (2-4)

where k = reaction rate constant
[NH3], [NO], [O2] = reactant concentrations, and
a, b, c = empirically determined exponents

The reaction rate is different for each catalyst formulation and, therefore, values for k, a, b, and c must be determined for each particular catalyst. The catalyst volume can also be determined if the space velocity is known for the catalyst and removal level of interest. The space velocity is defined as the gas flow rate divided by the catalyst volume.

The most important design and operating variables are presented in Table 2.3.2-1 and typical values for these are also shown. Although some of the data used in developing this table are for utility applications, the values should not be different for industrial applications. There are other variables that are important, but must be determined for each individual case. These are:

- flue gas flow rate
- NO_v control level
- NO, concentration

Variable	Typical Range
Space Velocity	$6000 - 10,000 \text{ hr}^{-1} \star$
$\rm NH_3:NO_{\times}$ Mole Ratio	0.7 - 1.0*
Flue Gas Temperature	350 - 400°C*
Pressure Drop	40 - 80 mm Hg
Catalyst Diameter (ring)	4 – 8 mm

TABLE 2.3.2-1. DESIGN AND OPERATING VARIABLES FOR MOVING BED SCR SYSTEMS^{14,15}

*Actual value will depend on required removal level.

The first two of these variables are the most important since they determine the size of the reactor. Higher flow rates or removal levels require larger reactors. Pressure drop for FGT systems does not usually change for a particular process. To accommodate the higher flow rates, the reactor cross-sectional area is increased to provide a constant linear velocity. The NO_{\times} concentration will affect the NH₃ and dilution steam requirements, but will not affect reactor size. Both flow rate and NO_{\times} concentration can be affected by boiler operating conditions. Since industrial boilers have fewer burners than utility boilers,² one burner represents a more significant fraction of the total boiler capacity. Therefore, a change in load on an industrial boiler may change these variables substantially if a burner is taken out of service

The FGT system will have to be designed to accommodate load changes. The necessary design accomodations will have to be determined for each application after examining operating data and establishing ranges of variation. Most likely this will involve a slight overdesign of the reactor and other equipment. The process control loops used with utility boiler applications should be capable of following load changes in industrial boilers.

Space velocity is usually defined as the volume of catalyst or reactor required to treat a given flow rate of flue gas.¹⁶ The magnitude of the space velocity is dependent entirely on the particular catalyst being considered. As can be seen in Table 2.3.2-1, the range for moving bed processes is $6000-10,000 \text{ hr}^{-1}$. These values are typically reported for 90 percent removal. For lower control levels, the value will be proportionally greater.

Almost all SCR processes require temperatures in the 350-400°C range in order to achieve good reaction rates. The temperature can vary with such things as boiler load, excess air, and ambient air temperatures. To control temperature two techniques are possible. The first involves bypassing a portion of the hot flue gas around the economizer and mixing with the economizer outlet gas so that the desired temperature is maintained.¹¹ The other technique uses inline heaters to obtain the desired temperature.¹⁷

The NH₃:NO_x mole ratio is also a function of the necessary removal level and, to a lesser extent, space velocity.¹⁸ For the three levels of removal considered in this study, 70, 80 and 90 percent, NH₃:NO_x mole ratios of 0.7, 0.8, and 0.9 are required, respectively. These data are for oil-fired boilers.

The catalyst shape and size is determined by the process vendor and is simply a design decision. Ring shapes (shaped like Raschig rings) are the most resistant to particulate plugging and, for this reason, were selected for this study.

The most recently published cost estimates for SCR systems are those of the Japanese Environment Agency which were published in Dr. Jumpei Ando's most recent report on Japanese NO_x control technology. Values taken from this study for two gas flow rates are shown below. The smaller gas flow rate is typical of industrial sized units while the larger flow rate is typical of utility installations.¹⁹

Gas Flow Rate	<u>Capital Cost</u>	Operating Cost
50,000 Nm ³ /hr	\$0.5 x 10 ⁶	\$0.2 x 10 ⁶
1,200,000 Nm ³ /hr	\$5.0 x 10 ⁶	\$3.5 x 10 ⁶

The cost for the large unit was included for comparison with other costs for large units that were reported for other process types in Section II. The reactor and catalyst type (fixed packed bed, moving bed or parallel flow) were not disclosed and, as a result, those costs are assumed to apply to all NO_y-only SCR systems.

Vendors of moving bed SCR systems are listed in Table 2.3.2-2 and the relative levels of application are noted. Although there are seven vendors, only four have applied their process to boilers. Of these, three have been applied to commercial scale equipment. Table 2.3.2-3 lists the moving bed systems that have been applied to oil-fired industrial boilers in Japan. Moving bed systems have not been applied to utility boilers in that country. Presently, there are no moving bed systems operating in the U.S. The Japaneese installations all treat gas from residual oil-fired boilers, implying that the technology is not necessary for distillate oil-fired applications which can use fixed packed beds.

2.3.2.2 System Performance--

The performance of several moving bed catalysts and plants is illustrated in Figures 2.2.3-2 through 2.3.2-8. The data presented indicate that NO_{χ} control greater than 90 percent is possible through the correct selection of process design variables. Outlet NH_3 concentrations are also shown. These are discussed in detail in Section VI. Table 2.3.2-4 shows several operating values from a commercial installation.

Vendor	Notes					
Sumitomo Chemical & Mitsubishi Heavy Industries	Tested on commercial scale equipment					
Hitachi, Ltd.	Tested on commercial scale equipment					
Ishikawajima-Harima Heavy Industries	Tested on pilot scale equipment					
Kurabo	Tested on commercial scale equipment					
Kobe Steel	Has not been applied to boilers					
Sumitomo Heavy Industries	Has not been applied to boilers					
Asahi Glass Company	Has not been applied to boilers					

TABLE 2.3.2-2. VENDORS OF SCR MOVING BED SYSTEMS FOR OIL-FIRED APPLICATIONS²¹

TABLE 2.3.2-3. EXISTING FGT INSTALLATIONS OF SCR MOVING BED SYSTEMS OIL-FIRED INDUSTRIAL BOILERS²¹

Location (Japan)	User	Process Developer	Fuel	Capacity (Nm ³ /hr)	Completion Date	
Kaizuka	Chiyoda Kenzai	Hitachi, Ltd.	Resid	15,000	Oct	1977
Amagasaki	Nipp on Oils & Fats	Hítachi, Ltd.	Resid	20,000	Apr	1978
Sodegaura	Sumitomo Chemical	Mitsubishi H.I.	Resid	300,000	Sept	1976
Sodegaura	Sumitomo Chemical	Sumitomo Chemical/ Mitsubishi H.I.	Resid	300,000	0ct	1976
Hirakatu	ľurabo	Kurabo	Resid	30,000	Aug	1975

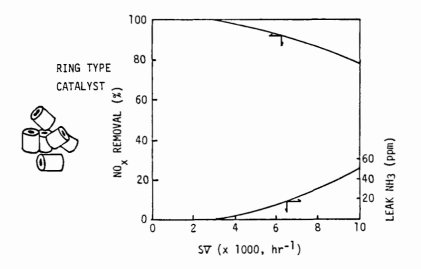


Figure 2.3.2-3. SV vs. NO_X removal and NH₃ leak (ring type catalyst, 15 mm diameter, 350°C NH₃/NO 1.0, inlet NO_X 250 ppm).¹⁰⁹

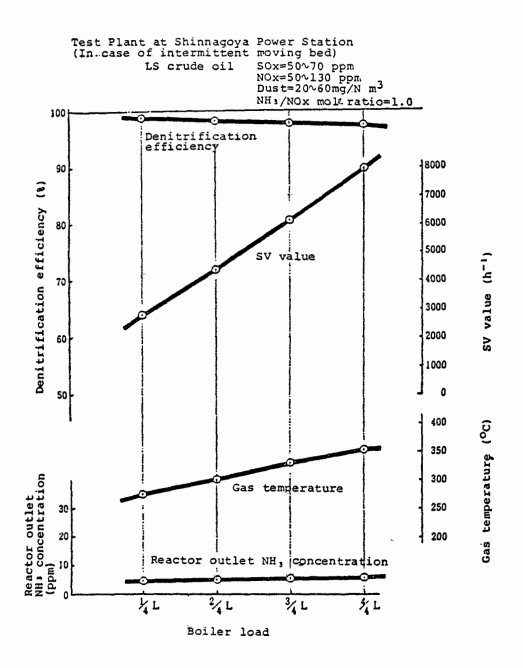
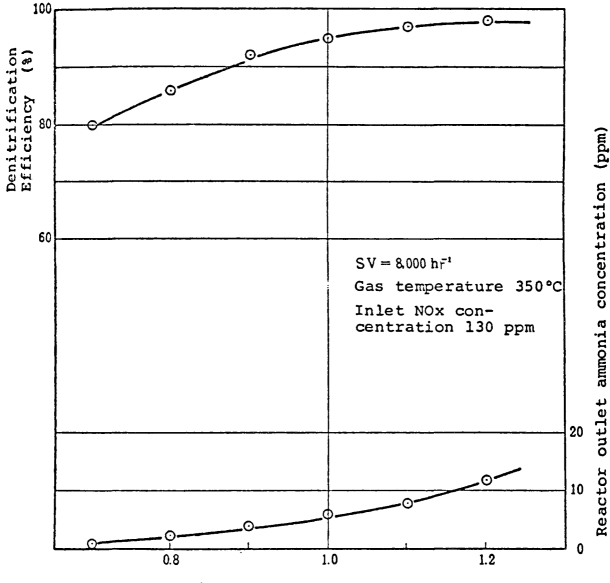


Figure 2.3.2-4. Relation between boiler load and denitrification efficiency (one example).¹¹⁰



NH3/NOx mol ratio (mol/mol)

Figure 2.3.2-5. NH_3/NO_{\times} mole ratio vs. denitrification efficiency and reactor outlet ammonia concentration.

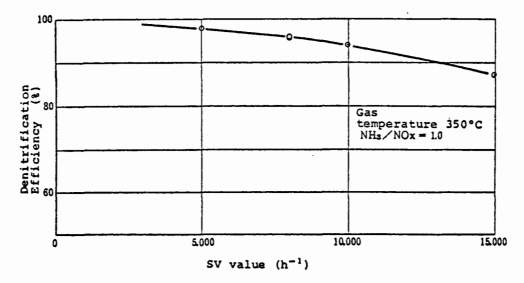


Figure 2.3.2-6. SV value and denitrification efficiency ¹¹² (for small, <1 mm, diameter particles).

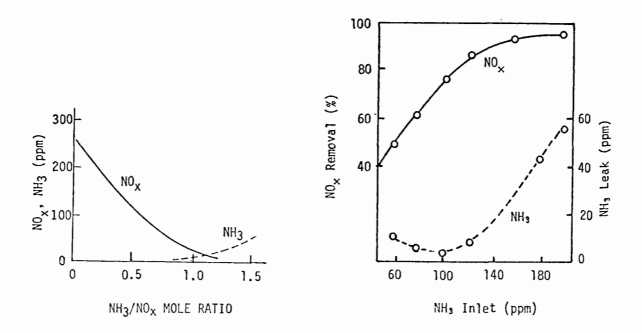


Figure 2.3.2-7. Relationship of NH₃/NO_× ratio to outlet NO_×¹¹³ NH₃ concentrations.

Figure 2.3.2-8. At 300°C.¹¹⁴

Gas for SCR (Nm ³ /hr)	300,000
Fuel	0il (S=0.7%)
Load fluctuation	60-100%
Stack height (m)	140
Inlet gas composition	
02 (%)	6
SO _x (ppm)	400
NO _× (ppm)	200
Particulates after EP (mg/Nm ³)	10-20
FGD unit	Scheduled
$SV (hr^{-1})$	5,000
Temperature (°C)	320
NO _x removal (%)	Over 90
NH3/NO mole ratio	1.0
Leak ammonia (ppm)	10-20
Type of reactor	Moving bed
Plant completed	Oct. 1976

TABLE 2.3.2-4. OPERATION DATA OF A COMMERCIAL SCR PLANT FOR DIRTY GAS¹⁰⁸

2.3.3 Selective Catalytic Reduction-Parallel Flow Reactor

2.3.3.1 System Description--

The distinguishing aspect of this process is the catalyst shape which is produced in a variety of shapes. The catalysts are produced in either a honeycomb, pipe, or plate shape. Both metal and ceramic supports are employed Several shapes are illustrated in Figure 2.3.3-1. The catalyst shapes allow particulate laden flue gas to pass through the reactor with no inertial impaction of the particles while the NO_x is transported to the catalyst surfaces by basic diffusion. The catalysts can handle all of the particulate levels emitted by the standard boilers.

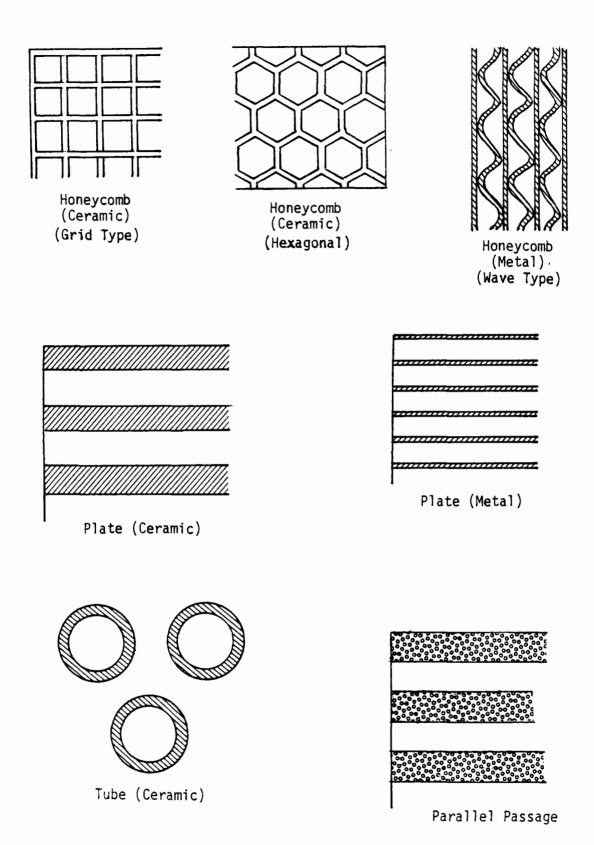


Figure 2.3.3-1. Shapes of parallel flow catalysts.²²

The reactors used are similar to standard fixed bed units and an example is shown in Figure 2.3.3-2. The catalyst is usually prepared in small modules and manually stacked within the reactor. The specific arrangement will depend on the particular process under consideration.

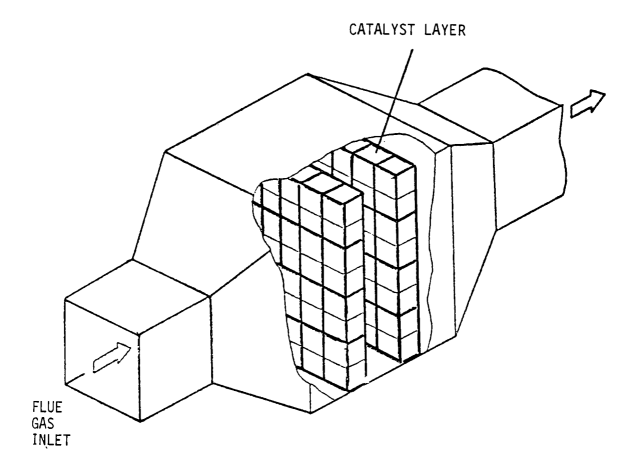


Figure 2.3.3-2. Typical reactor used with parallel flow SCR process.²³

A typical flow diagram for a parallel flow SCR system is shown in Figure 2.3.3-3. The arrangement is similar to the other SCR processes in that hot flue gas leaving the boiler economizer is injected with NH_3 and passed through a catalyst bed. Temperature control is important and can be accomplished with either a fired heater or an economizer bypass. NH_3 can be controlled using boiler operating condition inputs to conventional control components.

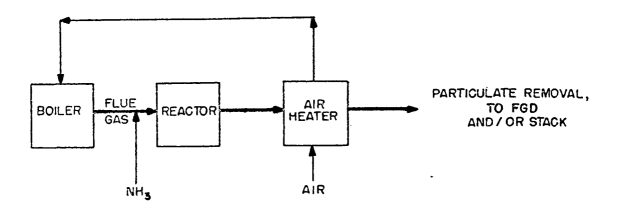


Figure 2.3.3-3. Flow diagram for parallel flow SCR process.²⁴

Within the reactor, NO_{χ} reacts with NH_3 to form N_2 and H_2O according to the following reactions. 12

$$4NO + 4NH_3 + O_2 \neq 4N_2 + 6H_2O \qquad (2-1)$$

$$2NO_2 + 4NH_3 + O_2 \neq 3N_2 + 6H_2O$$
 (2-2)

Reaction (2-1) is the primary reaction since flue gas NO_{\times} is typically 90-95 percent NO. O_2 is necessary for both reactions and is present in sufficient quantities (>3 percent) in all of the flue gases from the standard boilers.

The catalyst volume for a desired NO_{\times} removal can be determined by the fundamental design equation for a plug flow reactor.¹³

$$\frac{V}{F} = \int_{0}^{x} \frac{dx}{r}$$
(2-3)

The reaction rate, r, can be expressed as

$$r = k[NH_3]^a [NO]^b [O_2]^c$$
 (2-4)

The variables presented here have the same definitions as those presented in equations 2-3 and 2-4 of Section 2.2.2. The catalyst volume can also be determined if the space velocity is known for the catalyst and removal level of interest. The space velocity is defined as the gas flow rate divided by the catalyst volume.

The reaction rate is different for each catalyst formulation since different catalysts will lower the activation energy by different amounts. The activation energy affects the reaction rate constant, k, according to the Arrhenius equation.

$$k = Ae^{-\frac{E}{RT}}$$
(2-5)

An important design variable with catalytic systems is the space velocity which expresses the volume of catalyst required to treat one volume per hour of flue gas. Space velocity varies with catalyst formulation, catalyst shape, and control level. Typical values of space velocity for various catalyst shapes are shown in Table 2.3.3-1. Also shown are other catalyst design variables such as catalyst dimensions, gas velocities, bed depth and pressure drop. Ranges of values are used since specific values are different for each catalyst. The values shown pertain to 90 percent NO_{χ} removal and an NH_3/NO_{χ} mole ratio of 1:1.

Both NH_3/NO_{\times} ratio and space velocity will change with removal level. The NH_3/NO_{\times} mole ratio will range from 0.7-1.0 and the space velocity will range approximately as shown in the table for control levels of 70 to 90 percent.¹⁵

Variables associated with the boiler can also affect the performance of these systems. These are

- flue gas flow rate
- NO₂ concentration
- boiler load variability

2-97

TABLE 2.3.3-1. CATALYST DESIGN VARIABLES FOR VARIOUS CATALYST SHAPES (Basis: 90% NO_{\times} removal at NH_3/NO_{\times} ratio of 1:1, 350-400°C)

	Honeycomb Honeycomb		Paralle	el Plate	
	(metallic)	tube (ceramic)	(Ceramic)	(Metallic)	
Catalyst size (mm)					
Thickness	0.5-1	1.5-3	8-10	1	
Opening	4-8	6-20	8-14	5-10	
Gas velocity (m/sec) ^a	2-6	5-10	5-10	4-8	
Bed depth (m)	1-2	1.5-5	4-6	2-5	
SV (1,000 hr ⁻¹) ^b	5-8	4-8	1.5-3	2-5	
Pressure drop (mmH ₂ 0)	30-80	40-160	80-160	60-120	

^aVelocity at 350-400°C in open column (superficial velocity). ^bGas volume (Nm³/hr)/catalyst bed volume (m³).

The flue gas flow rate and control level determine the catalyst volume (hence reactor size) necessary. Increases in either also increase the reactor size. The NO_{χ} concentration is a function only of fuel type used in the standard boilers. Higher concentrations require larger NH_3 storage and vaporization equipment; reactor size is not affected. Boiler load can affect several things including flue gas temperature, flow rate and NO_{χ} concentration. It is necessary to maintain reactions temperatures of 350 to 400° C and temperature control equipment may be necessary if the boiler experiences large load variations. Where these variations are present, some equipment overdesign may be warranted to insure a constant control level. These variables are discussed in more detail in the section on moving bed SCR systems for oil-fired boilers, Section 2.3.2.

Parallel flow SCR processes have been applied in Japan to several residual oil-fired industrial boilers. Oil-fired utility boilers and other sources with high particulate concentrations are also being treated. A list of vendors of parallel flow SCR systems is presented in Table 2.3.3-2. Notes on the relative level of application are also shown. Four of the eight vendors have applied their systems to oil-fired boilers indicating that application of this technology to industrial boilers is technically feasible. Parallel flow SCR systems have been applied to both industrial and utility boilers. Specific applications are listed in Tables 2.3.3-3 and 2.3.3-4. There have been no applications in the U.S. The tables indicate that the parallel flow technology is designed primarily for residual oils and not distillate oils.

TABLE 2.3.3-2. VENDORS OF SCR PARALLEL FLOW SYSTEMS FOR OIL-FIRED APPLICATIONS²¹

Vendor	Notes		
Hitachi Zosen	Tested on pilot scale equipment		
Hitachi, Ltd.	Tested on commercial scale equipment		
JGC	Has not been tested on boilers		
Mitsui Engineering & Shipbuilding	Tested on commercial scale equipment		
Mitsubishi Heavy Industries	Tested on commercial scale equipment		
Ishikawajima-Harima Heavy Industries	Tested on commercial scale equipment		
Kobe Steel	Has not been tested on boilers		
Kawasaki Heavy Industries	Tested on pilot scale equipment		

TABLE 2.3.3-3. EXISTING FGT INSTALLATIONS OF SCR PARALLEL FLOW SYSTEMS OIL-FIRED INDUSTRIAL BOILERS²¹

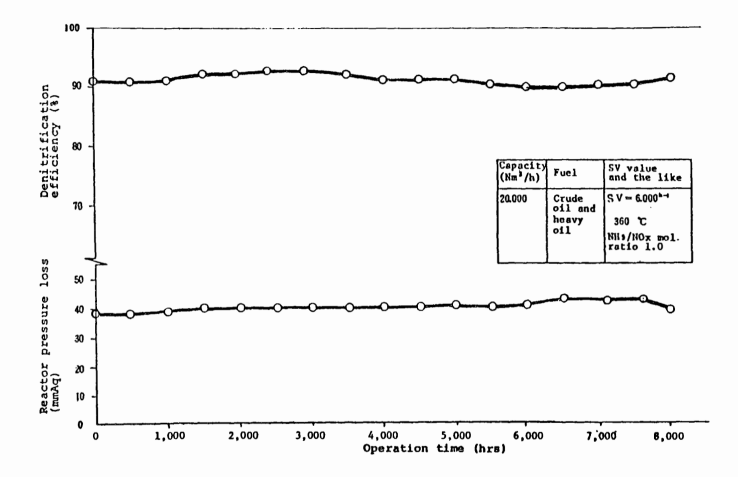
Location (Japan)	User	Process Developer	Fuel	Capacity (Nm³/hr)	Completion Date
Sodegaura	Fuji Oil	Mitsubishi H.I.	Resid	200,000	January 1978
Kawasaki	Ajinomoto	Ishikawajima H.I.	Resid	180,000	January 1978
Chiba	Ukishima Pet. Chem.	Mitsui Engineering	Resid	220,000	April 1978

Location (Japan)	User	Process Developer	Fuel	Capacity (Nm³/hr)	Completion Date
Yokosuka	Tokyo Electric	Mitsubishi H.I.	Resid	40,000	March 1977
Chita	Chubu Electric	Mitsubishi H.I.	Resid	1,920,000	February 1980
Kudamatsu	Chugoku Electric	Ishikawajima H.I.	Resid	1,900,000	July 1979
Niigata	Tohoku Electric	Ishikawajima H.I.	Resid	1,660,000	August 1981

TABLE 2.3.3-4. EXISTING FGT INSTALLATIONS OF SCR PARALLEL FLOW SYSTEMS OIL-FIRED UTILITY BOILERS²¹

2.3.3.2 System Performance--

The performance of several parallel flow catalysts is illustrated in Figures 2.3.3-4 through 2.3.3-9. Table 2.3.3-5 shows several operating data for a single parallel flow SCR installation. The data presented indicated that NO_{\times} control levels of greater than 90 percent are obtainable through selection of the appropriate process design variables. Other data are also presented and these are discussed in subsequent sections.



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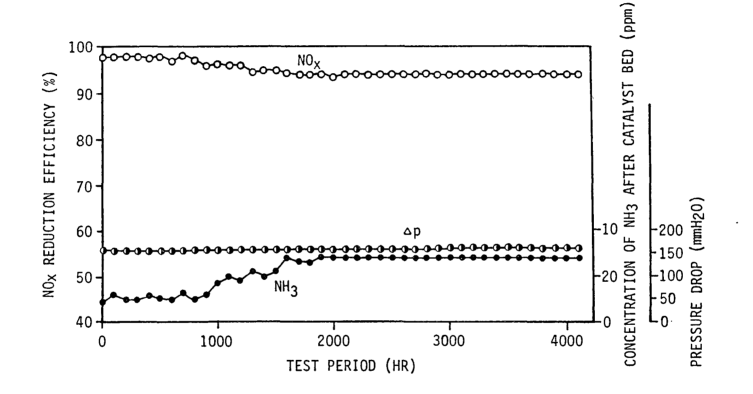
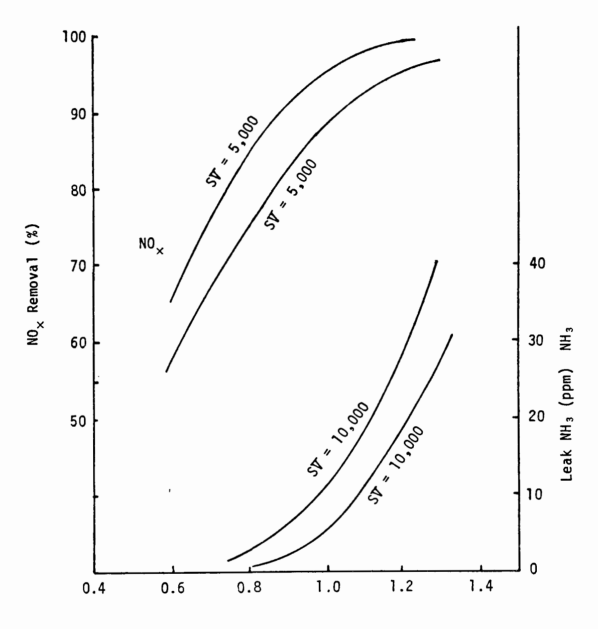


Figure 2.3.3-5. Durability of NO_x removal catalyst for exhaust gas of high sulfur oil burning boiler.¹¹⁶



 $\rm NH_3/NO_{ imes}$ Mole Ratio

Figure 2.3.3-6. Typical example of operation data (oil-fired boiler, 350-400°C, granular or honeycomb catalyst).¹⁰⁵

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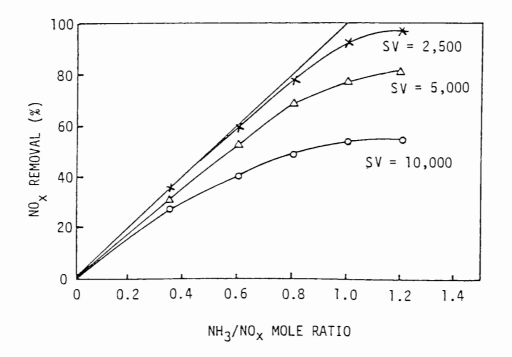


Figure 2.3.3-7. NH_3/NO_x mole ratio vs. NO removal (plate catalyst; 350°C, LV 5.9 m/sec).¹¹⁷

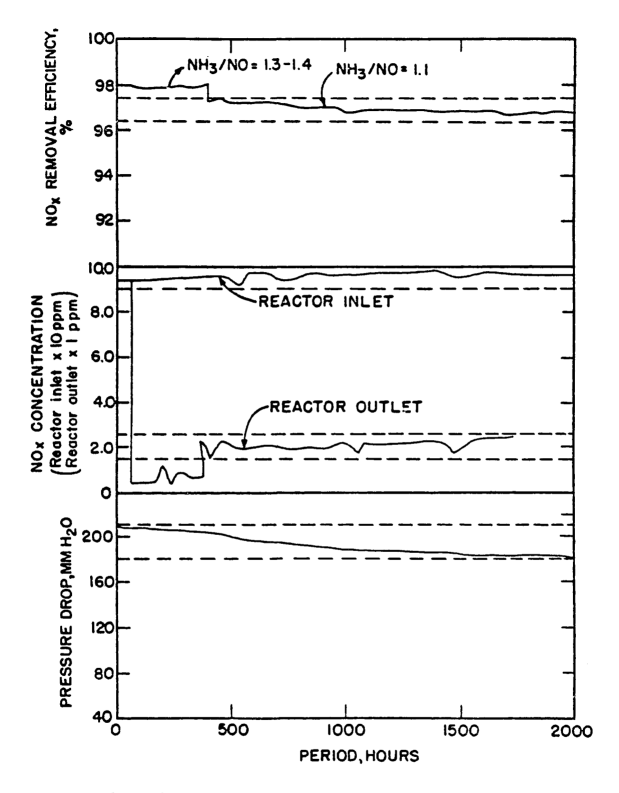


Figure 2.3.3-8.

 $\rm NO_{\chi}$ removal efficiency, $\rm NO_{\chi}$ concentration, and pressure loss over 2,000 hr test period for JGC Paranox Process. 118

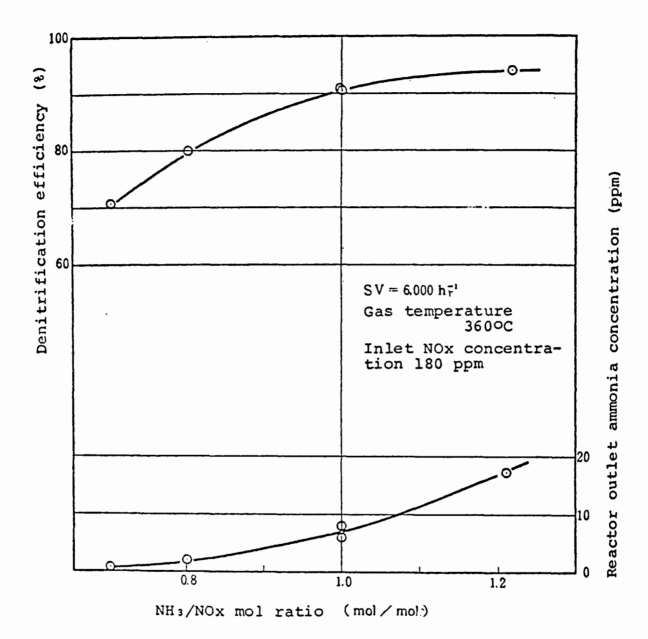


Figure 2.3.3-9. NH_3/NO_{\times} mole ratio and denitrification efficiency and reactor outlet ammonia concentration.¹¹⁹

Capacity (Nm ³ /hr)	220,000
Gas composition	
NO _× (ppm)	150
SO _× (ppm)	300
Dust (mg/Nm ³)	100-150
Catalyst and reactor	
Catalyst carrier	TiO
Catalyst shape	PP
$SV (hr^{-1})$	4,000
Temperature (°C)	350-400
NH₃/NO _× mole ratio	1.0
NO _× removal (%)	Above 90
Total pressure drop (mmH_2O)	180
Leak NH3 (ppm)	Below 10
Operation start	July 1977
Plant cost (10 ⁶ yen)	260

TABLE 2.3.3-5. SCR PLANT BY MITSUI ENGINEERING AND SHIPBUILDING CO.²⁰

2.3.4 Absorption-Oxidation

2.3.4.1 System Description--

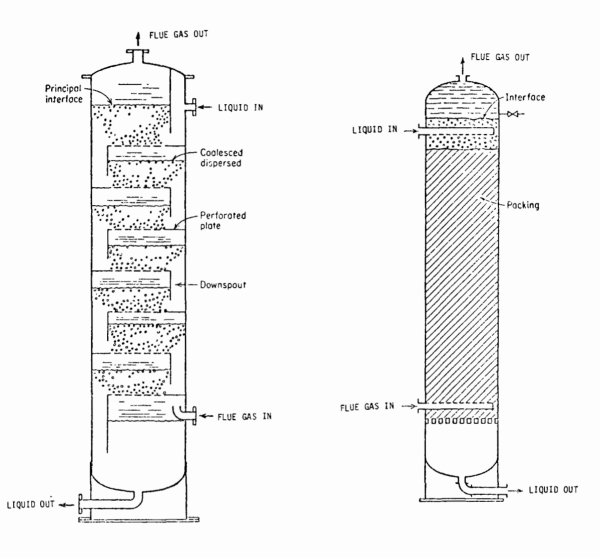
Absorption-oxidation processes remove NO_{χ} from flue gas by absorbing the NO or NO_2 into a solution containing an oxidant which converts the NO_{χ} to a nitrate salt. Two types of gas/liquid contactors can be used and examples of each type are shown in Figure 2.3.4-1. Both perforated plate and packed towers accomplish NO_{χ} absorption by generating high gas/liquid interfacial areas. The choice of one type of contactor is a design decision made to achieve a given removal for the least cost.

A generalized process flow diagram is shown in Figure 2.3.4-2. Flue gas is taken from the boiler after the air preheater. Before the gas can be sent to the NO_X absorber, it must be SO₂-free since SO₂ consumes prohibitive amounts of the costly liquid-phase oxidant. In most cases, the oxidant is permanganate (MnO_{4}) ; however, Ca(ClO)₂ can also be used. Therefore, a conventional FGD unit is required ahead of the NO_X absorber. A prescrubber to cool the gas and remove both particulates and Cl⁻ prior to FGD is also necessary. After having passed through these two scrubbing sections, the flue gas enters the distributing space at the bottom of the NO_X absorber, below the packing or plates. The gas passes upward through the column, countercurrent to the flow of the liquid absorbent/oxidant (usually a KOH solution containing KMnO₄). NO_X is absorbed and then oxidized over the length of the column according to the following reactions.³¹

$$NO(g) \rightarrow NO(aq)$$
 (2-6)

$$NO(aq) + KMnO_4(aq) \rightarrow KNO_3(aq) + MnO_2(s)$$
(2-7)

$$2NO_2(g) \rightarrow N_2O_4(g)$$
 (2-8)



Perforated Plate Absorber

Packed Absorber

Figure 2.3.4-1. Gas/liquid contactor options for Absorption-Oxidation Processes.²⁹

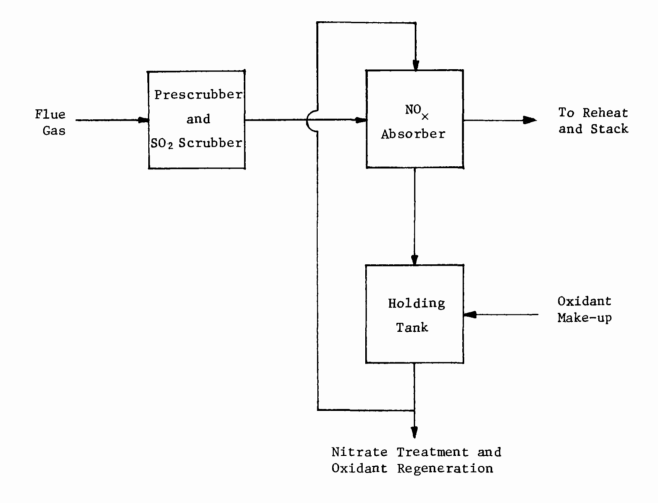


Figure 2.3.4-2. Process flow diagram for absorptionoxidation process.³⁰

$$N_2O_4(g) \rightarrow N_2O_4(aq)$$
 (2-9)

$$N_2O_4(aq) + 2K_2MnO_4(aq) + 2KMnO_4(aq) + 2KNO_2(aq)$$
 (2-10)

Since most of the NO_{χ} from combustion processes occurs as NO_{Λ}^{32} reactions 2-6 and 2-7 predominate. The clean gas passes out of the top of the absorber to a heater for plume buoyancy and is sent to the stack. The absorbing solution drops to a holding tank where makeup KOH and/or KMnO₄ are added. This solution flows to a centrifuge to separate the solid MnO₂ which is then electrolytically oxidized to MnO₄. The remaining solution is either concentrated in an evaporator to form a weak KNO₃ solution or is electrochemically treated to produce a weak HNO₃ solution and a mixed stream of KOH and KNO₃.

The fundamental design equation used for gas absorption column design is

$$\int_{Y_{a}}^{Y_{b}} \frac{dy}{(y-y^{\star})} = \left(\frac{K_{a}a}{G_{y}}\right) Z \qquad (2-11)$$

- where y = bulk NO_x concentration (mole fraction) of gas phase at any given point in column
 - y-y* = overall driving force for absorption (y* being the NO_x concentration of a gas in equilibrium with a given liquid NO_x concentration)
 - $Y_{b} = inlet NO_{x}$ concentration
 - $Y_{2} = outlet NO_{x}$ concentration
 - Ky = overall gas-phase mass transfer coefficient, lb-moles NO_x/
 (ft²)(hr)(mole fraction)

a = area of gas-liquid interface per unit packed volume, ft^2/ft^3 G_y = molal gas mass velocity, lb-moles flue gas/(ft²)(hr) Z = length of packed section of column, ft

In a column containing a given packing or plate configuration and being irrigated with a certain liquid flow, there is an upper limit to the gas flow rate. This limit's superficial gas velocity (volumetric gas flow rate/ cross-sectional area of column) is called the flooding velocity. At this point, the gas flow completely impedes the downward motion of the liquid and blows the liquid out of the top of the column. The gas velocity, obviously, must be lower than the flooding velocity. How much lower is a design decision. Often, it is an economic tradeoff between power costs and equipment costs. A low gas velocity will lower the pressure drop and, hence, the power costs but the absorber will have a larger diameter and cost more. High gas velocities have an opposite effect. Usually the optimum gas velocity is about one-half the flooding velocity.³⁴ The height of the column depends on the desired level of removal and on the rate of mass transfer. The latter is a major problem for these systems trying to achieve large NO_x reductions since NO is relatively insoluble in water. This can be seen in Table 2.3.4-1.

	Boiling P oint, °C	Solubility in Cold Water (0°C), cm ³	Solubility in Hot Water (60°C), cm ³
NO	-151.8	7.34/100 cc H ₂ 0	$2.37/100 \text{ cc } H_20$
NO ₂	21.2	soluble, decomposes	

TABLE 2.3.4-1. NITROGEN OXIDES CHARACTERISTICS³⁵

One can see that NO has a very limited solubility in water and, since most NO_{\times} is present as NO, the rate of mass transfer (absorption) is going to be relatively slow. This means that the absorber must be tall with a high

liquid flow rate. Table 2.3.4-2 presents the effects of boiler/flue gas variables on the design of absorption-oxidation systems.

Variable	Design Effect
Presence of particulates	Requires prescrubber
Presence of SO ₂	Requires FGD pretreatment
Increased gas flow	Requires larger column diameter; increased liquid flow rate
Increased NO_{χ} concentration	Requires larger column height; increased oxidant concentration

TABLE 2.3.4-2. SYSTEM DESIGN CONSIDERATIONS

Both flue gas flow rate and NO_{\times} concentration can be affected by boiler operating conditions. Therefore a change in load on an industrial boiler may alter these variables markedly. The absorber must be designed to accommodate any anticipated load changes. The column size and the liquid and oxidant flows must be designed for each application after examining the boiler operating history and establishing ranges of variation.

None of the sources consulted for this study could supply typical ranges for operating variables such as liquid/gas ratio, reagent concentrations or pressure drops and, as a result, none are presented here. Economic data were not presented either. One source did estimate the removal for absorptionoxidation processes to be 85 percent.³⁶

Presently, absorption-oxidation processes are still in the pilot unit stage of development. Table 2.3.4-3 presents a list of absorptionoxidation process vendors and the status of development of their projects.

Vendor	Status of Development		
Hodogaya	No information available; stopped development on process		
Kobe Steel	1974: 1000 Nm ³ /hr gas from iron-ore sintering furnace; stopped development on process		
MON (Mitsubishi Metal, MKK, Nikon Chemical)	1974: 4000 Nm ³ /hr flue gas from oil-fired boiler		
Nissan Engineering	1972: 4 pilot plants, 100-2000 Nm³/hr tail gas from HNO3 plant		

TABLE 2.3.4-3. PROCESS VENDORS OF ABSORPTION-OXIDATION PROCESSES^{37,38}

Only one of the process vendors has piloted this process on flue gas from an oil-fired boiler and there have been no commercial applications either in Japan or the U.S.

2.3.4.2 System Performance--

No information has been published on tests conducted with flue gas from oil-fired boilers. The relative insolubility of NO in water may present a major obstacle to achieving the stringent level of control (90 percent NO, reduction) by absorption-oxidation processes. Another primary drawback of absorption-oxidation systems is the production of nitrate salts (see Equation 2-7), a secondary pollutant. These processes probably could not be applied on a large scale as wastewater treatment systems (chemical or biological) do not remove nitrogen compounds from the wastewater.³⁹ Trying to recover the nitrates as nitric acid for industrial use or potassium nitrate for fertilizer does not seem promising as the by-products are of low quality. Also, the use of an expensive, liquid-phase oxidant requires stainless steel and other corrosion resistant materials of construction. High sulfur coals require an FGD system prior to the $\mathrm{NO}_{\scriptscriptstyle\!\!\!\!\!\!\!\!\!}$ absorber to prevent excessive oxidant consumption by SO2. The process steps of several absorber columns in series (large fan requirements), oxidant regeneration (electrolysis). and flue gas reheat (inline heater) are all energy intensive and present technical and economic disadvantages.

2.3.5.1 System Description--

From a NO_{χ} removal standpoint, this process is very similar to those discussed in Sections 2.3.1 through 2.3.3. The primary difference is the additional equipment necessary to collect and process the SO₂. The main feature of the process is the reactor and catalyst which remove both NO and SO₂. This process was developed by Shell although the U.S. licensor, UOP, is currently marketing and developing the process. The NO_{χ}/SO_2 version of the process is commonly called the SFGT process which stands for the Shell Flue Gas Treatment Process.

A uniquely designed parallel flow type of reactor is used to avoid problems with particulates. This design is necessary only with flue gas from residual oil-fired boilers. The reactor consists of a series of packages containing catalyst material, arranged in a parallel configuration which allows flue gas flow between the packages. Each package consists of catalyst material placed between two layers of wire gauze. Figure 2.3.5-1 illustrates the internals of the parallel passage reactor. The flue gas flows between the catalyst packages and not directly through the catalyst material. This prevents plugging of the catalyst with particulate matter in the flue gas.

For convenient fabrication and handling, catalyst packages of a standard size are appropriately spaced and placed in a container to form a unit cell or module. SO_2 removal efficiency and capacity are determined by the number of unit cells placed in series in a cell stack. For a given level of SO_2 removal, a greater number of cells in the stack increases the capacity and reduces the frequency of regeneration. The number of stacks is determined largely by the flue gas rate and the flue gas velocity through a single stack is generally not a design variable. For most design situations, 4 to 5 unit cells in a stack are adequate to achieve high SO_2 removal, however, if a high level of denitrification is required, more unit cells per stack may be necessary.

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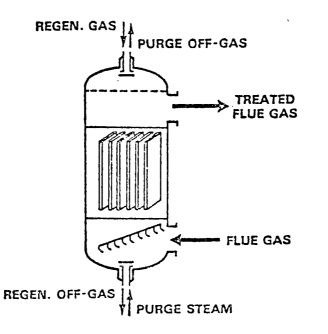


Figure 2.3.5-1. The SFGT parallel flow reactor. 40

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The SFGT process is a dry process with two or more reactors operating in a cyclic manner. The desulfurization aspect of the process is regenerable, while NO_{χ} removal is accomplished by catalytic reduction with ammonia. The catalyst material is commonly called an acceptor since SO_2 removal involves adsorption or "acceptance" of SO_2 . The desulfurization cycle consists of the following steps:

- 1) oxidation of acceptor bed/acceptance of SO₂,
- 2) purge reactor,
- 3) regeneration with reducing gas, and
- 4) purge reactor.

The products of the oxidation and acceptance reactions in step 1 above catalyze the reaction of NO_x with ammonia to form nitrogen and water. NO_x removal is accomplished by metering ammonia into the untreated flue gas upstream of the reactors. The catalytic reaction takes place across the partially spent acceptor beds.

Also associated with the SFGT process are facilities for generating reducing gas and for the processing of SO_2 in regeneration off gases into sulfur by-products. Figure 2.3.5-2 illustrates the process flow for a typical SFGT system.

Boiler flue gas is withdrawn upstream of the air preheater and particulate removal device by the SFGT system fan and discharged to the reactor inlets. The flue gas then flows through fixed bed reactors in open channels alongside and in contact with the acceptor material. Ammonia is added to the flue gas upstream of the SFGT system fan to insure complete mixing before the flue gas enters the reactor.

Fresh acceptor material is elemental copper on an alumina support. This is converted to the oxide form by flue gas oxygen shortly after initiation of the acceptance cycle. SO_2 is removed by reaction with the copper oxide and oxygen as the flue gas flows through the channels, converting the acceptor material to copper sulfate. Simultaneous with the desulfurization process, the reduction of flue gas NO_x by ammonia is selectively catalyzed by copper oxide and copper sulfate in the acceptor bed. As the flue gas leaves the SFGT system reactors it is returned to the boiler flue gas duct downstream of SFGT fan suction.

Flue gas is fed to a reactor until an unacceptable amount of SO_2 begins to pass through the reactor. This occurs when a large fraction of the acceptor has been converted to the sulfate form. Flue gas flow is then diverted to another reactor and the spent reactor is isolated. Any flue gas remaining in the spent reactor is purged with an inert gas such as steam, and the regeneration cycle is initiated.

Regeneration is accomplished by passing a reducing gas through the bed countercurrent to the direction of the flue gas flow. The reducing gas, which is primarily hydrogen, reacts with the copper sulfate in the spent reactor to convert it to elemental copper. An off gas of SO_2 and water is

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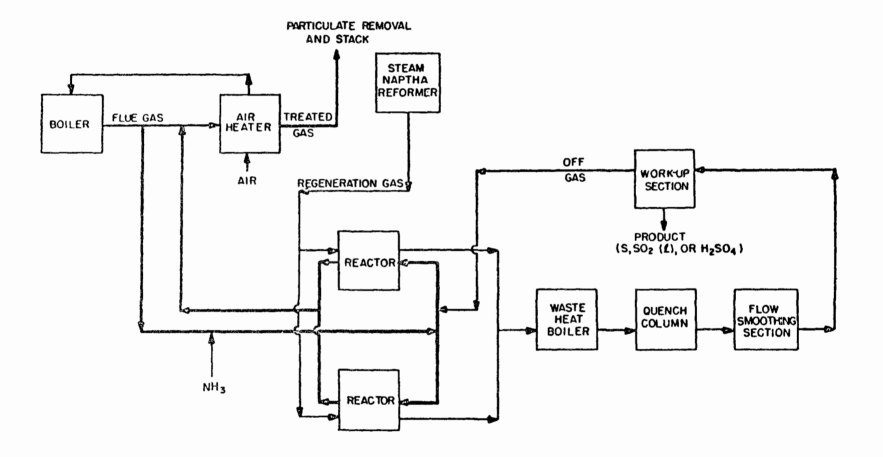


Figure 2.3.5-2. Flow diagram of the SFGT process,⁴¹

produced by the reaction. After regeneration is complete, the reactor is again purged with steam and is ready for another acceptance cycle. Regeneration gas can be produced from a number of sources, but steam-naphtha reforming is proposed by UOP as being the most economical.⁴²

The regeneration off-gas treatment section consists of flow smoothing equipment and SO_2 workup equipment. Typically, the regeneration off-gas is cooled and most of the steam condensed, raising the SO_2 concentration from 10 percent to 80 percent by volume. The concentrated SO_2 is then compressed into an intermediate holding vessel to provide a smooth flow rate to the workup section. The workup section may be a modified Claus unit which produces an elemental sulfur by-product, a fractionation unit which produces liquid SO_2 , or a sulfuric acid plant.

Each process step consists of different chemical reactions. The acceptor material is converted to the oxide form by the following reaction:

$$Cu + \frac{1}{2}O_2 \rightarrow CuO$$
 (2-12)

This oxide readily reacts with flue gas SO_2 and oxygen, as described by:

$$Cu0 + \frac{1}{2}O_2 + SO_2 + CuSO_4$$
 (2-13)

 SO_3 in the flue gas is also removed by the following reaction:

$$Cu0 + SO_3 \rightarrow CuSO_4$$
 (2-14)

The reaction scheme for reduction of NO_{\times} is described by the following:¹²

$$4NO + 4NH_3 + O_2 \neq 4N_2 + 6H_2O$$
 (2-1)

$$2NO_2 + 4NH_3 + O_2 \neq 3N_2 + 6H_2O \qquad (2-2)$$

Excess ammonia which is not consumed in reactions 2-1 and 2-2 may be catalytically oxidized to nitrogen and water by reaction with flue gas oxygen, as described by:

$$4NH_3 + 3O_2 \rightarrow 2N_2 + 6H_2O \qquad (2-15)$$

Maximum NO_{\times} removal efficiency is achieved at the point of SO_2 breakthrough, where conversion of the acceptor material from the oxide to the sulfate form is essentially complete. Figure 2.3.5-3 illustrates reactor outlet SO_2 and NO concentrations during a typical SFGT acceptance cycle.

A different set of reactions is involved during regeneration of the catalyst bed.

Copper sulfate is reduced to the elemental copper form by reducing gas hydrogen as described by the following reaction:

$$CuSO_4 + 2H_2 + Cu + SO_2 + 2H_2O$$
 (2-16)

Any acceptor material present in the reactor as the oxide will also be reduced, according to the following reaction:

$$CuO + H_2 + Cu + H_2O$$
 (2-17)

The regeneration step occurs at the same temperature as the acceptance step, $400^{\circ}C$ (750°F).

The general reactor design equation is the same as that described in earlier sections for SCR processes. The primary variables are the gas rate, reaction rate, and control level. Reaction rate data have not been released for this process except that the NO_x reduction is first order.

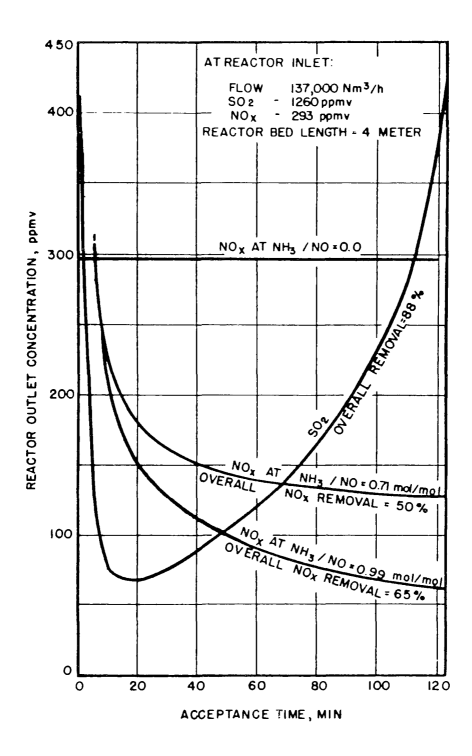


Figure 2.3.5-3. SFGT reactor performance versus acceptance time.44

The gas flow rate and control level will determine the reactor size. Increases in either variable will increase the reactor volume. The effect of control level can be seen in Figure 2.3.5-4. It is necessary for the flue gas to enter the reactor at 400°C and therefore it must be taken from an appropriate point in the boiler, most likely from between the economizer and air preheater. Alternatively, a cooler gas can be heated to 400°C by an inline heater.

The removal efficiency of NO_{x} for a given reactor size is determined by the amount of NH_{3} injected as shown in Figure 2.3.5-5. Since the reaction is first order in NO_{x} , control level is not affected by NO_{x} concentration.⁴⁷ The SO₂ control efficiency is primarily a function of the acceptance time of the reactor (Figure 2.3.5-3). Typical ranges of operating variables are shown in Table 2.3.5-1.

Since the SFGT system can handle full particulate loading ($\geq 10 \text{ gr/sft}^3$) it is not dependent on any pretreatment facilities. Also, the SFGT system operation is independent of boiler operation. The system fan takes suction from the flue gas duct between the economizer and air preheater and the reactor discharge returns to the boiler flue gas duct just downstream of the suction point, with no valves between the two points. The system fan provides a constant flow rate through the SFGT system. If the boiler flue gas rate is greater than the fan rate, flue gas will bypass the system through the open duct. If the boiler flue gas rate is lower than that of the system fan, treated gas will recycle back to the system fan suction. Recycle of treated gas to the reactor inlet with "open bypass" arrangement presents no operating problems. This is due to the fact that both the level of desulfurization and denitrification are independent of inlet concentrations, and the system does not humidify the flue gas.

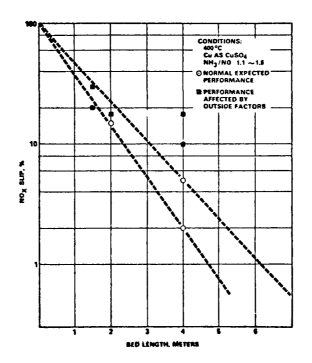


Figure 2.3.5-4. Unconverted NO_{\times} as a function of catalyst bed length.⁴⁵

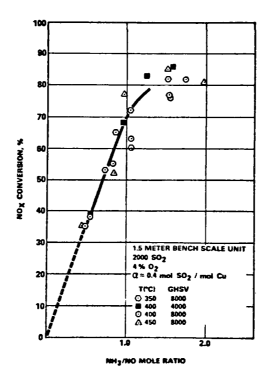


Figure 2.3.5-5. $\rm NO_{x}$ reduction with $\rm NH_{3}$ over commercial SFGT acceptor.46

Variable	Typical Range
Space Velocity	$5,000 - 8,000 \text{ hr}^{-1}$ *
$NH_3:NO_X$ Mole Ratio	1.0:1.0 to 1.2:1.0*
Flue Gas Temperature	400°C
Pressure Drop	5-6 in. H ₂ 0*
Maximum Particulate Loading	≥23 g/Nm

TABLE 2.3.5-1. DESIGN AND OPERATING VARIABLES FOR SFGT SYSTEM⁴⁸

*Actual value will depend on required removal level.

Tables 2.3.5-2 and 2.3.5-3 present the test and commercial applications of the SFGT process. The development history of the process can also be seen in these tables. In the U.S., from 1974 to 1976 a pilot scale unit at Tampa Electric Company (TECO) was operated using flue gas from a coal-fired boiler. Testing was for SO₂ removal only, NO_x control was not attempted during this period. The process developer is currently modifying the TECO pilot unit to accommodate 7 meters of bed height, up from the previous maximum of 5 meters. This should permit simultaneous removal of NO_x and SO_x to the 90 percent level. Also, provisions are being made for injection of a CO/CO₂ gas mixture into the regeneration gas in order to simulate medium-Btu gas from a coal gasifier.

The costs for an industrial size boiler have not been estimated. The only detailed cost estimates currently available are for coal-fired utility boilers. These are shown in Tables 2.3.5-4 through 2.3.5-6. Also shown are the estimated energy and raw material requirements.

Location/ Company	Designed By	Fuel/ Application	Size, Nm ³ /hr	Type of Operation	Dates	Comments
Shell Ref. at Pernis	Shell	Residual Fuel Oil- Proc. Heater	600-1000	SO _x -only	1967-1972	SO _x reduction - approx. 90%
Rotterdam Utility	Shell	Coal- Steam Boiler	-	Heavy Fly Ash Loading	1971	Particulate mat- ter - loadings to 20 Gr/Nm ³
Tampa Elec. Big Bend	UOP	Coal- Wet-Bottom Utility Boiler	1200-2000	SO _x -only; SO _X -NO _X Simultaneous	1974-1976 1979-	SO _x - 90%; SO _x -NO _x - 90/90% fly ash to 25 Gr/Nm ³
JGC Yokohama	JGC*	Fuel Oil	250-700	NO _× -only	1974-	NO _x reduction - 90-99%
Nippon Steel	JGC	Sintering Furnace	2000	NO _× -only	1976-1978	NO _x reduction - 90-97%
	JGC	Coke Oven Gas	400	NO _x -only	1976-1977	NO _x reduction - 90%; special low temp. cat. evalua- tion

TABLE 2.3.5-2. SFGT PROCESS, PILOT AND DEMONSTRATION UNITS

*JGC Corporation, licensing agent in Japan.

Unit	Designed By	Fuel/ Application	Size, Nm ³ /hr	Type of Operation	Dates	Comments
SYS* Yokkaichi	Shell	Residual Fuel Oil- Ref. Boiler	125,000	SO _x -only; NO _x -SO _x Simultaneous	1973–1975 1975–	SO _x reduction - 90%; Simultaneous - 90/50%
Kashima Oil Co. Ltd.	JGC	Fuel Oil- Process Unit Heater	50 , 000	NO _x -only	1975-	95–98%
Fuji Oil Co. Ltd.	JGC	CO Boiler	70,000	NO _× -only	1976-	93–96%
Nippon Steel Corp.	JGC	Sintering Furnace	150,000	NO _x -only	1978-	∿95% (low temp. cata- lyst)

TABLE 2.3.5-3.	SFGT PROCESS,	COMMERCIAL UNITS	
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*Showa Yokkaichi Sekiyu

	BASIS:
Incorporated Units:	Steam-Naphtha Reformer SFGD Reactor Section Compressor/Gasholder Flow Smooth Section Modified Claus Unit
Power Plant Size	500 MW
Fuel	Coal
S-Content, Wt-%	
C a se l	3.5
Case 2	2.5
Case 3	0.8
HHV	10,500 Btu/1b
Heat Rate	9,000 Btu/kWh
Excess Air	20%
Air Preleater Leakage	13%
Flue Gas Rate	1,582,000 Nm ³ /h (983,000 SCFM)
SO ₂ Content, ppmv	
Case 1	2,580
Case 2	1,850
Case 3	590
Mid-1977, Gulf Coast Location	
Load Factor	7,000 h/a
Capital Charges	15%/a
Cost of:	
Naphtha	\$0.35/gal
Steam (40 psí, SAT.)	\$1.50/M 1b
Electricity	\$0.018/kWh
Labor	\$10.00/hr
Heat Credits	\$2.50/MMBtu
Sulfur	\$45.00/ton

TABLE 2.3.5-4. ECONOMICS OF SFGT SYSTEM⁴⁹

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		SFGD Section	Flow Smooth Section	Mod. Claus Section	Reformer Section	Total
Case 1		* * * * * * * * * *				
Electricity	kW	5,770	850	115	480	7215
Steam**	kmo1/h	1,820	-380*	-740*	-600*	100
Naphtha***	Gcal/h				90.92	90.92
Heat Credits	Gcal/h					42.53
S° Produced	kg/h			5250		5250
Case 2						
Electricity	kW	5,800	570	82	300	6782
Steam**	kmo1/h	1,300	-270*	530*	-415*	85
Naphtha***	Gcal/h				62.75	62.75
Heat Credits	Gcal/h					32.48
S° Produced	kg/h			3760		3760
Case 3						
Electricity	kW	5,120	180	30	110	5440
Steam**	kmo1/h	480	-95*	-170*	-140*	75
Naphtha***	Gcal/h				21.01	21.01
Heat Credits	Gcal/h					18.46
S° Produced	kg/h			1200		1200

TABLE 2.3.5-5. ECONOMICS OF SFGT SYSTEM ESTIMATED CHEMICALS AND UTILITY REQUIREMENTS⁵⁰

*Produced

**40 psig, Saturated

***5.175 MMBtu/Bb1 produces 11,500 SCF Hyárogen/Bb1

	Case 1	Case 2	Case 3
EEC. (MM\$)			
SFGD Reactor Section	28.95	28.53	22.94
Compressor/Gasholder	7.82	6.10	2.65
Modified Claus	2.76	2.26	1.14
Steam-Naphtha Reformer	8.81	7.14	4.17
Estimated Annual Revenue Requirements (M\$/a)			
Capital Charges	7251	6604	4634
Maintenance	967	881	618
Labor	123	123	123
Acceptor	1479	1053	411
Electricity	909	855	685
Steam	42	35	31
Naphtha	7174	4951	1658
Heat Credits	-2977	-2273	-1292
Sulfur Credits	-1570	-1126	-359
Capital Cost, Operating Cost, Energy Requirement			
Capital Cost, \$/kW	97	88	62
Operating Cost, ¢/kWh	0.38	0.32	0.19
Energy Requirement, Btu/kWh*	525	371	124

TABLE 2.3.5-6. ECONOMICS OF SFGT SYSTEM ESTIMATED CAPITAL AND OPERATING COST⁵¹

Electricity at9000 Btu/kWhSteam at40000 Btu/kmolNaphtha at4 Btu/kcalHeat Credits at4 Btu/kcal

2.3.5.2 System Performance--

 NO_{\times} control by the SFGT process is shown graphically in Figure 2.3.5-5. and in Figure 2.3.5-3 presented earlier. As can be seen, at a space velocity of 8000 hr⁻¹, NO_{\times} control of >80 percent can be achieved. Figure 2.3.5-4 indicates that the process developers feel the process to be capable of NO_{\times} control levels of >90 percent. Ando indicates that NO_{\times} and SO_{\times} removals of 70 percent and 90 percent, respectively are achievable at an NH₃/NO mole ratio of 0.99.¹²⁰ He also indicates that higher NO_{\times} control may be possible. But unless some process modifications are made, SO_2 control will decrease and NH₃ emissions will increase.

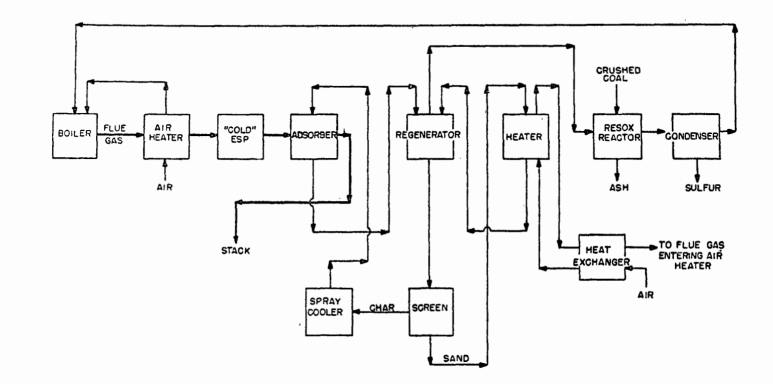
As mentioned earlier, the system is not impacted by changes in the boiler gas rate or particulate concentrations. Changes in the NO_{χ} concentration due to boiler load changes and be compensated for by conventional control system used in conjunction with the NH₃ injection equipment. This control system will be developed during the upcoming pilot tests at the TECO pilot plant.

2.3.6 Adsorption

2.3.6.1 System Description--

The adsorption process removes NO_{\times} and SO_2 from flue gas by adsorbing them onto a special activated char. Adsorbed NO_{\times} is reduced to N_2 while SO_2 is reduced and condensed to an elemental sulfur by-product.

A process flow diagram is shown in Figure 2.3.6-1. Flue gas is taken from the boiler air preheater and passed through a particulate removal device to prevent blinding of the adsorption bed. The flue gas then enters the adsorber, a vertical column with parallel louver beds containing the char in pellet form. NO_{χ} and SO_2 are adsorbed on the char which slowly moves downward through the bed. The NO_{χ} adsorption mechanism is unknown but SO_2 undergoes the following reaction.⁵⁵



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Figure 2.3.6-1. Flow diagram of Foster Wheeler-Bergbau Forschung Dry Adsorption Process.⁵⁴

$$SO_2(g) + H_2O(g) + \frac{1}{2}O_2(g) + H_2SO_4(1)$$
 (2-18)

The reaction product is held in the pores of the char pellets. The flue gas exits the adsorber and passes to the stack. The saturated char leaves the bottom of the adsorber and is screened to remove any fly ash deposits. It is then conveyed to a regenerator where it is mixed with hot sand (650°C) and the following reactions take place.^{55,56}

$$2H_2SO_4(1) + C(s) \rightarrow CO_2(g) + 2H_2O(g) + 2SO_2(g)$$
 (2-19)

$$2NO(g) + C(s) + CO_2(g) + N_2(g)$$
 (2-20)

This SO_2 -rich gas product stream is sent to an off-gas treatment reactor containing hot, crushed coal (650-820°C) and the following reactions take place.⁵⁶

$$SO_2(g) \rightarrow S(g) + O_2(g)$$
 (2-21)

$$C(s) + O_2(g) \rightarrow CO_2(g)$$
 (2-22)

The gas then passes to a condenser where the S vapor forms molten S. The char/sand mixture from the regenerator is screened to separate the two solids. The char is recycled to the adsorber via a spray cooler and the sand is recycled to the regenerator after passing through a heater.

This process operates at 120-150°C, however, typical values for other operating variables were not found. NO_{\times} and SO_2 control levels were reported to be 40-60 percent and 80-95 percent, respectively.⁵⁷ The economics of the process vary with the fuel sulfur level. For coal-fired boilers with fuel sulfur levels of 0.9-4.3 percent, the capital costs range from \$40-90/kW and the operating costs range from 1.0-2.3 mills/kWh.⁵⁸ The costs were based on applying the process to a utility boiler of >200 MW capacity. Costs for oil-fired applications were not found.

Presently, the adsorption process is in the prototype unit stage of development. The one reported process developer in the field, Foster Wheeler-Bergbau Forschung has a 20 MW prototype unit and several small pilot plants treating coal-fired flue gas. The process should also be applicable to oilfired boilers.

2.3.6.2 System Performance--

Tests have shown the adsorption process to be primarily a SO_2 reduction process as NO_{χ} removal efficiency averages 40-60 percent while SO_2 removal had a range of 80-95 percent.⁵⁹

The primary drawback of this process, besides the low NO_{χ} removal level, is its complexity: numerous process steps involving hot solids handling. Solids flow can be difficult to control and high maintenance requirements could be expected. The vendor has reported several mechanical problems during testing which included control of adsorber-bed levels, poor char distribution, char-sand separation, hot sand conveying, and char cooling and feed. Some corrosion-resistant material is needed in the high temperature zones of the process. The ash waste stream from the off-gas treatment reactor appears to be the sole secondary pollutant associated with the process. The overall complexity and low NO_{χ} removal of the process present definite technical disadvantages.

2.3.7 Electron Beam Radiation

2.3.7.1 System Description--

This dry process utilizes an electron beam to bombard the flue gas, removing NO_{\times} and SO_2 in the process. A block flow diagram for the process is shown in Figure 2.3.7-1.

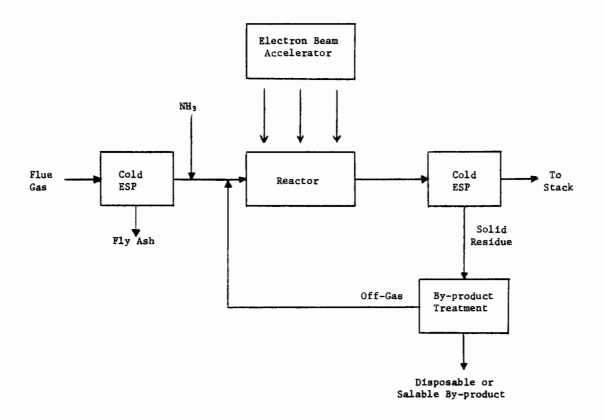


Figure 2.3.7-1. Process flow diagram for Ebara-JAERI electron beam process.⁶⁰

Flue gas is taken from the boiler air preheater and passed through a cold ESP to remove particulates. After a small amount of ammonia is added, the gas enters a reactor where it is bombarded with an electron beam. (The penetration of the gas stream by the beam will require a unique discharge pattern or other special design considerations.) A powder containing both ammonium nitrate and sulfate is generated by an unknown reaction mechanism. The gas then exits the reactor, passes through a second ESP to remove the solid by-product, and is sent to the stack. The by-product treatment system is still being developed. Various methods investigated include thermal decomposition in the presence of an inert gas, steam roasting with CaO, or steam roasting with H₂O. The by-product may eventually be useful as a fertilizer.⁶¹

The key subsystem of this process is the electron beam accelerator. Control of this unit's power supply is based upon inlet composition, flow rate, and temperature of the flue gas.

Some of the important variables and typical ranges are listed in Table 2.3.7-1.

	Typical Value
Temperature	∿100 [°] C
Reactor residence time	1-20 sec
Radiation rate	10 ⁵ -10 ⁶ rad*/sec
Total radiation absorbed	1-3 Mrad*
Total radiation absorbed	1-3 Mrad*

TABLE 2.3.7-1. SYSTEM VARIABLES⁶²

*Rad is the radiation dose absorbed
1 rad = .01 J/Kg

The operating cost with NO_{\times} removal only (low sulfur oils) is lower due to lower radiation levels, but the capital cost would be just as high as for simultaneous NO_{\times}/SO_{\times} removal. Capital costs are quite high for this process as the 2 ESP's and the accelerator are expensive. The costs for a 1000 Nm³/hr test unit are reported to be \$1000/kW; however, the costs of a full scale system are expected to be lower. Operating costs are not available.

The Ebara Manufacturing Company in conjunction with Japan Atomic Energy Research Institute (JAERI) has operated a 1000 Nm³/hr pilot plant treating flue gas from an oil-fired boiler. In 1976, a 3000 Nm³/hr pilot plant began treating of1-gas from an iron ore sintering furnace at Nippon Steel. Byproduct treatment technology needs to be more fully developed before this process can be applied commercially. In the U.S., the Department of Energy (DOE) is funding development of an electron beam process offered by Research-Cottrell. Pilot unit tests with flue gas are scheduled, however, the details of the program are not yet available.

2.3.7.2 System Performance--

A summary of the oil-fired pilot tests is shown in Figure 2.3.7-2.

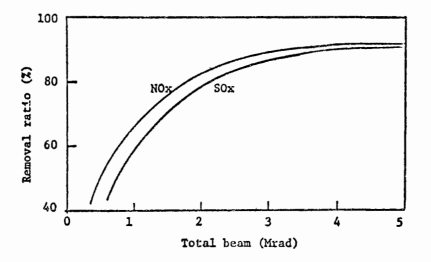


Figure 2.3.7-2. Oil-fired pilot plant test results.⁶⁴

One can see that NO_X/SO_X removal drops off drastically at a total radiation dose below 1 Mrad while the maximum removal is obtained at about 3 Mrad. The removal efficiencies decrease as the concentrations of NO_X and SO_X increase as can be seen in Figure 2.3.7-3.

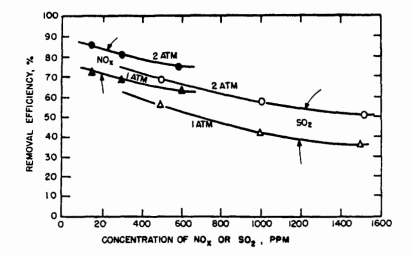


Figure 2.3.7-3. Effect of pollutant concentration on removal efficiency.65

2.3.8 Absorption-Reduction

2.3.8.1 System Description--

Absorption-reduction processes simultaneously remove $NO_{\rm v}$ and SO_2 from flue gas by absorbing them into a scrubbing solution. The processes are based on the use of chelating compounds, such as ethylenediamine tetraacetic acid (EDTA) complexed with iron, to "catalyze" the absorption of NO₂. Most process vendors prefer a perforated-plate type of gas-liquid contactor. The advantages of a perforated-plate absorber over a packed bed absorber include easier cleaning when solids are present, wider operating ranges, and more economical handling of high liquid rates.⁶⁶ An example of a perforated plate absorber is shown in Figure 2.3.8-1. The most common design of a perforated plate is one that employs liquid crossflow over the face of the plate with the gas passing upward through the plate perforations. A schematic of the operation of a crossflow perforated plate is shown in Figure 2.3.8-2. The liquid is prevented from flowing through the plates by the upward flow of the gas. However, during periods of low gas flow (such as load changes on industrial boilers) liquid can drain through the openings in the plates. This reduces the liquid's time of contact with the gas on each plate and may decrease the overall operating efficiency of the absorber. To prevent this problem, there are two other types of dispersers utilized besides the basic sieve-plate: the valve-plate and the bubble cap, depicted in Figure 2.3.8-3. As the gas flow lowers, the valve or cap settles, sealing off the perforation so liquid cannot drain through. This design feature allows the perforated plate absorber to operate more efficiently at widely fluctuating gas rates.

While most all absorption-reduction processes utilize ferrous chelating compounds to enhance NO absorption, the scrubbing solutions, the by-product treatment and sorbent regeneration chemistry differ from process to process. For this reason, one of the simpler absorption-reduction processes, that of Kureha Chemical Industry Company, is examined here in detail.

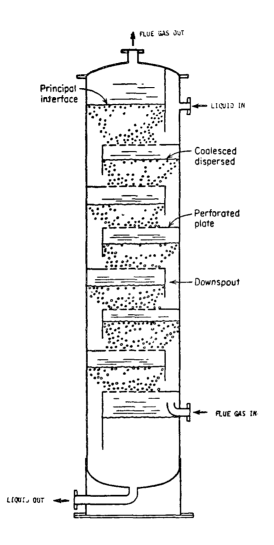


Figure 2.3.8-1. Perforated plate absorber option for Absorption-Reduction Processes.²⁸

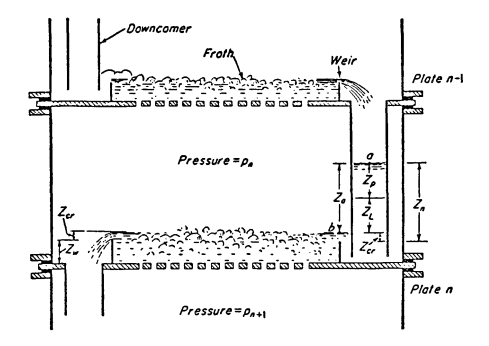
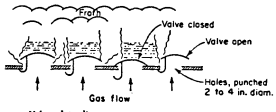
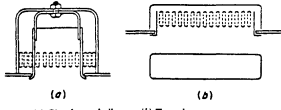


Figure 2.3.8-2. Normal operation of sieve plate. Z_{α} , height of station a above datum. $Z_{\mathcal{CP}}$, weir crest. Z_{L} , liquid-friction head. $Z_{\mathcal{p}}$, pressure head across plate. $Z_{\mathcal{N}}$, net head in down pipe. $Z_{\mathcal{W}}$, weir height.⁶



Valve-plate dispersers.



(a) Circular or bell cap. (b) Tunnel cap.

Bubble cap dispersers

Figure 2.3.8-3. Other gas dispersers.⁶⁸

A block flow diagram of the Kureha absorption-reduction process is shown in Figure 2.3.8-4. Flue gas is taken from the boiler after the air preheater. It passes through a prescrubber to adiabatically cool the gas and remove both particulates and chlorides. The flue gas then enters the distributing space at the bottom of the NO_{χ}/SO_2 absorber, below the plates or packing. The gas flows upward, countercurrent to a sodium acetate (CH₃COONa) scrubbing solution ($\sim 60^{\circ}$ C) containing ferrous iron and EDTA and a few seed crystals of gypsum (to prevent scaling). Most of the SO₂ is rapidly absorbed at the bottom of the absorber according to the following reactions.⁷¹

$$SO_2(g) \neq SO_2(aq)$$
 (2-23)

$$SO_2(aq) + 2CH_3COONa(aq) + H_2O \rightarrow Na_2SO_3(aq) + 2CH_3COOH(aq)$$
(2-24)

The NO_{χ} (which consists mainly of NO) is relatively insoluble; therefore, it is absorbed gradually over the length of the column. The ferrous chelating compounds effect on NO absorption is described in Figure 2.3.8-5. The NO_{χ} is absorbed and undergoes the following reactions.⁷³

$$NO(g) \rightarrow NO(aq)$$
 (2-6)

$$2NO_2(g) \rightarrow N_2O_4(aq)$$
 (2-25)

$$N_2O_4(g) \rightarrow N_2O_4(aq)$$
 (2-9)

 $2NO(aq) + 5Na_2SO_3(aq) + 4CH_3COOH(aq) \rightarrow 2NH(SO_3Na)_2(aq) + Na_2SO_4(aq) + 4CH_3COONa(aq) + H_2O \quad (2-26)$

 $2N_2O_4(aq) + 7Na_2SO_3(aq) + 4CH_3COOH(aq) \rightarrow 2NH(SO_3Na)_2(aq) + 3Na_2SO_4(aq) + 4CH_3COONa(aq) + H_2O$ (2-27)

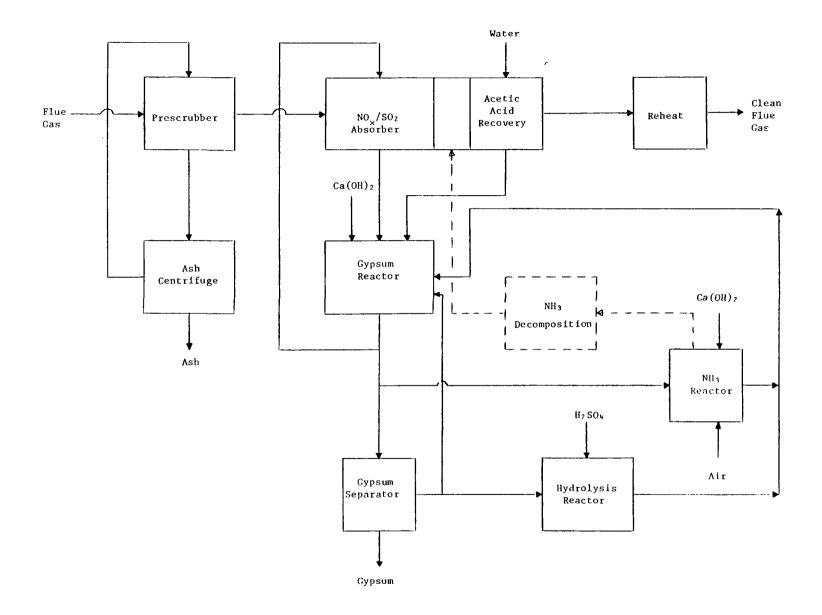


Figure 2.3.8-4. Process flow diagram of Kureha absorption-reduction process. 69,70

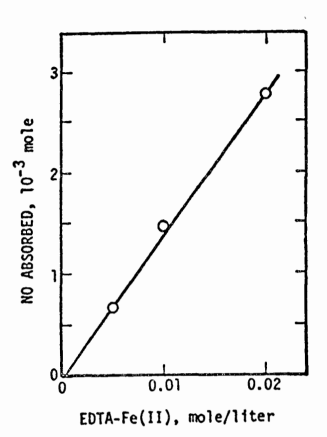


Figure 2.3.8-5. EDTA-Fe(II) concentration and NO absorption at 50°C.⁷²

Some of the acetic acid (CH₃COOH) formed at the bottom of the absorber via reaction (2-24) is vaporized. It must be captured and is done so by water scrubbing at the very top of the absorber. From the top of the absorber column the clean flue gas passes to a heater for plume buoyancy and is then sent to the stack. The liquid effluent drops from the bottom of the absorber to a gypsum, CaSO₄·2H₂O, production reactor. Here, the solution is mixed with with the purge stream from the acetic acid recovery section and a lime slurry stream. The lime, Ca(OH)₂, treatment involves the following reactions.⁷⁴

$$2CH_{3}COOH(aq) + Ca(OH)_{2}(aq) \rightarrow (CH_{3}COO)_{2}Ca(aq) + 2H_{2}O$$
 (2-28)

 $(CH_3COO)_2Ca(aq) + Na_2SO_4(aq) + 2H_2O \rightarrow CaSO_4 \cdot 2H_2O(s) + 2CH_3COONa(aq) (2-29)$

The gypsum formed by reaction 2-29 is centrifuged. Most of the liquor discharged is returned to the gypsum reactor and on to the absorber. The remaining liquor is sent to a reactor where sulfuric acid (H_2SO_4) is added to hydrolyze the imidodisulfonate, $NH(SO_3Na)_2$, by the following reaction.⁷⁵

$$H^{+}$$
NH(SO₃Na)₂(aq) + 2H₂O $\stackrel{\rightarrow}{\xrightarrow{}}$ NH₄HSO₄(aq) + Na₂SO₄(aq) (2-30) (2-30)

The effluent from this reactor is then recycled to the gypsum production reactor. A small purge stream is taken from the gypsum reactor to another reactor where the ammonium bisulfate (NH₄HSO₄) formed in the hydrolysis reaction is treated with lime to yield gypsum and NH₃ off-gas by the following reaction.⁷⁶

$$\mathrm{NH}_{4}\mathrm{HSO}_{4}(\mathrm{aq}) + \mathrm{Ca}(\mathrm{OH})_{2}(\mathrm{s}) \rightarrow \mathrm{CaSO}_{4} \cdot \mathrm{2H}_{2}\mathrm{O}(\mathrm{s}) \downarrow + \mathrm{NH}_{3}(\mathrm{g})^{\dagger} \qquad (2-31)$$

The gaseous ammonia is stripped from the solution by an air stream. If no use for the ammonia can be found, the gas mixture is sent to a catalytic reactor where ammonia is oxidized by the following reaction.⁷⁷

$$4NH_3(g) + 30_2(g) \xrightarrow{\text{catalyst}} 2N_2(g) + 6H_2O(g)$$
 (2-32)
350°C

The product stream is then sent to the deacetating section of the absorber column.

The fundamental design equation used for gas absorption column design is³²

$$\int_{Y_{a}}^{Y_{b}} \frac{dy}{(y-y^{*})} = \left(\frac{K_{y}^{a}}{G_{y}}\right) Z \qquad (2-11)$$

- where
- y = bulk NO_x concentration (mole fraction of gas phase at any given point in column
 - y-y* = overall driving force for absorption (y* being the NO_x concentration of a gas in equilibrium with given liquid NO_x concentration)
 - $Y_{L} = inlet NO_{x}$ concentration
 - $Y_a = outlet NO_x$ concentration
 - Ky = overall gas-phase mass transfer coefficient, lb-moles NO_x/
 (ft²)(hr)(mole fraction)
 - a = area of gas-liquid interface per unit packed volume, ft^2/ft^3
 - $G_y = molal gas mass velocity, lb-moles flue gas/(ft²)(hr)$
 - Z = length of packed section of column, ft

In a column containing a given plate or packing configuration and being irrigated with a certain liquid flow, there is an upper limit to the gas flow rate. This limit's superficial gas velocity (volumetric gas flow rate/ cross-sectional area of column) is called the flooding velocity. At this point, the gas flow completely impedes the downward motion of the liquid and blows the liquid out of the top of the column. The gas velocity, obviously, must be lower than the flooding velocity. How much lower is a design decision. Often it is an economic tradeoff between power costs and equipment costs. A low gas velocity will lower the pressure drop and, hence, the power costs but the absorber will have a larger diameter and cost more. High gas velocities have an opposite effect. Usually the optimum gas velocity is about one-half the flooding velocity.33 The height of the column depends on the desired level of removal and on the rate of mass transfer. The latter consideration is the reason why a chelating compound is used in absorption-reduction processes to aid in NO_x absorption. Table 2.3.8-1 presents the effects of boiler/flue gas variables on the design of absorption-reduction systems. Both flue gas flow rate and NO_x concentration can be affected by boiler operating conditions. Therefore a change in load on an industrial boiler may alter these variables markedly. The absorber

mest be designed to accommodate any anticipated load change. The column size and the liquid flows must be designed for each application after examining the boiler operating history and establishing ranges of variation.

Variable	Design Effect
Presence of particulates	Requires prescrubber
Presence of SO ₂	Requires SO ₂ :NO mole ratio of at least 3-5 ⁹ (depending on process) for absorption- reduction to be effective.
Increased gas flow	Requires larger column diameter; increased liquid flow rate
Increased NO_{χ} concentration	Requires larger column height; increased catalyst concentration

TABLE 2.3	.8-1. SYS	TEM DESIG	GN CONSIDERATIONS

The process vendors have not released much information on the operating conditions of these processes. This is primarily due to the competitive status of these similar processes at this early stage of development. Typical values for some of the process variables are shown in Table 2.3.8-2.

TABLE 2.3.8-2.TYPICAL VALUES FOR PROCESS VARIABLES
OF ABSORPTION-REDUCTION PROCESSES 78

Variable	Range	* * ~
Liquid/Gas ratio, 1/Nm ³	10-30	
SO _x /NO _x mole ratio	2.5-3.0	
Superficial Gas Velocity, m/s	1-3	

Cost estimates for this type of process cover a large range, presumably due to the differences in sorbent regeneration technique. Capital costs for an oil-fired system were not found, however, for coal-fired utility applications, capital costs ranged from \$65-127/kW. Operating costs for an oil-fired application are shown below for two flue gas flow rates.¹²¹

Gas Flow Rate, Nm ³ /hr	Operating Costs, mills/kWh
150,000	9.1
500,000	8.3

These costs are based on $\frac{200}{\$}$ and do not include flue gas reheat.

Presently, absorption-reduction processes are in the pilot-unit stage of development. Table 2.3.8-3 presents a list of absorption-reduction process vendors and the status of development of their projects. One can see from the table that several oil-fired flue gas tests have been performed.

Vendor	Status of Development
Asahi	1974: 600 Nm ³ /hr flue gas from residual oil- fired boiler (1000 hours continuous).
Chisso	1975: 300 Nm ³ /hr flue gas from oil-fired boiler (335 hours continuous)
Kureba	1976: 5000 Nm ³ /hr flue gas from heavy oil-fired boiler (3000 hours continuous)
Mitsui Engineering and Shipbuilding	1974: 150 Nm ³ /hr flue gas from oil-fired boiler
Pittsburgh invironmental	1976: 3000 Nm ³ /hr flue gas from coal-fired boiler (52 hours continuous, absorption section only)

TABLE 2.3.8-3. PROCESS VENDORS OF ABSORPTION-REDUCTION PROCESSES⁸⁰

2.3.8.2 System Performance--

Four of the vendors listed in Table 2.3.8-3 report NO_{\times} removals of at least 80 percent with oil-fired boiler flue gas. The Pittsburgh Environmental system, however, only achieves 60 percent with coal. SO_2 control for all of the systems is 90+ percent. No plots of system performance could be found for any of the oil-fired systems.

Absorption-reduction processes are readily applicable only to high sulfur oils as a $SO_2:NO_{\times}$ mole ratio in the flue gas of at least 3-5 is required for maximum performance. This can easily be shown by observing reactions 2-24 and 2-26 reprinted below.

 $SO_2(aq) + 2CH_3COONa(aq) + H_2O \rightarrow Na_2SO_3(aq) + 2CH_3COOH(aq)$

 $2NO(aq) + 5Na_2SO_3(aq) + 4CH_3COOH(aq) \rightarrow 2NH(SO_3Na)_2(aq) + Na_2SO_4(aq) + 4CH_3COONa(aq) + H_2O$

One can see that 1 mole of SO_2 absorbed in solution reacts to form 1 mole of sodium sulfite (Na₂SO₃). Then, 5 moles of sodium sulfite are required to reduce 2 moles of NO. So, the minimum stoichiometric $SO_2:NO_{\times}$ mole ratio required is $\frac{5}{2}$ or 2.5. Also, some of the sodium sulfite is oxidized to sodium sulfate by oxygen present in the flue gas according to:

$$Na_2SO_3(aq) + \frac{1}{2}O_2(aq) \rightarrow Na_2SO_4(aq)$$
 (2-33)

and is not available for NO_{\times} reduction. Low-sulfur oils would require SO_2 to be added to the flue gas for these processes to perform; therefore, they should be considered applicable to high sulfur oils only.

Absorption-reduction processes require large absorbers with high liquid rates due to relative insolubility of NO, even when the absorption catalyst is used. Also, the regeneration of the absorption catalyst and the flue gas reheat for plume buoyancy are energy intensive. Some corrosion-resistant material is necessary due to the corrosive nature of the absorbing solution. However, absorption-reduction appears to be the most promising of the "wet" NO_{χ}/SO_2 removal processes. This is due primarily to its not utilizing oxidants which require much corrosion-resistant material and, more importantly, create serious secondary pollution problems. Also, the primary by-product of absorption-reduction processes, gypsum, can be used as landfill material or in building materials. For all the above reasons, absorption-reduction processes appear, at this preliminary stage, to be competitive with other wet NO_{χ}/SO_{χ} removal processes.

2.3.9 Oxidation-Absorption-Reduction

2.3.9.1 System Description--

Oxidation-absorption-reduction processes simultaneously remove NO_{χ} and SO_2 from flue gas by oxidizing relatively insoluble NO to relatively soluble NO_2 and then absorbing both NO_2 and SO_2 into a scrubbing solution. The processes are based on the use of gas-phase oxidants, either ozone (O₃) or chlorine dioxide (ClO₂), to selectively oxidize NO to NO_2 . Both perforated-plate and packed bed absorption columns are utilized by various process vendors.

Most of the oxidation-absorption-reduction processes are similar in that they consist of five major sections:

- prescrubbing
- gas-phase oxidation
- NO_{χ}/SO_2 absorption
- reduction of absorbed NO_{γ} and oxidation of $SO_{3}^{=}$
- wastewater treatment

The areas where processes differ are gas-phase oxidation - O_3 or ClO_2 ; absorption solutions - limestone slurry (CaCO₃), H₂SO₄, or NaOH; and the amount and type of waste treatment required. Thermal decomposition, biological denitrification, or wastewater evaporation wastewater treatment systems can be used. Because of these differences, only one of the oxidationabsorption-reduction processes, that of Mitsubishi Heavy Industries, is examined here in detail.

A block flow diagram of the MHI oxidation-absorption-reduction process is shown in Figure 2.3.9-1.

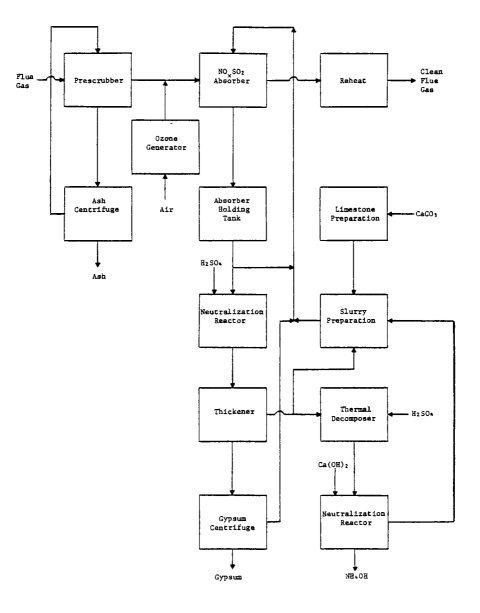


Figure 2.3.9-1. Process flow diagram for MHI oxidationabsorption-reduction process.⁸¹

Flue gas is taken from the boiler after the air preheater and passed through a prescrubber to cool the gas and remove particulates and chlorides. The flue gas then enters a duct where it is injected with ozone (about 1 percent by weight in air)⁸² such that the O3:NO ratio is 1:1. Ozone selectively oxidizes NO by the following reatcion.⁸³

$$NO(g) + O_3(g) \rightarrow NO_2(g) + O_2(g)$$
 (2-34)

After injection, the flue gas passes countercurrent to a lime/limestone slurry in a grid-packed absorption column. A water-soluble catalyst is added to the slurry to enhance NO_2 absorption (even though NO_2 is more soluble than NO, it is still less soluble than SO_2). SO_2 is absorbed quickly at the bottom of the column and undergoes the following reactions.¹⁵

$$SO_2(g) \rightarrow SO_2(aq)$$
 (2-23)

$$SO_2(aq) + CaCO_3(s) + {}^{\flat}H_2O \rightarrow CaSO_3 \cdot {}^{\flat}H_2O(s) + CO_2(g)$$

$$(2-35)$$

$$SO_2(aq) + CaSO_3(aq) + H_2O \rightarrow Ca(HSO_3)_2(aq)$$
(2-36)

 $\rm NO_2$ is absorbed gradually over the length of the column and reacts as follows. $^{1\,6}$

$$2NO_{2}(g) + Ca(OH)_{2}(s) + CaSO_{3} \cdot \frac{1}{2}H_{2}O(s) + \frac{1}{2}H_{2}O \rightarrow Ca(NO_{2})_{2}(aq) + CaSO_{4} 2H_{2}O(s)$$

$$(2-37)$$

Once both the NO₂ and SO₂ are absorbed, the nitrite ion formed by reaction 2-37 is reduced by the bisulfate ion formed by reaction $2-36.^{84}$

$$Ca(NO_{2})_{2}(a_{1}) + 3Ca(HSO_{3})_{2}(aq) \rightarrow 2Ca[NOH(SO_{3})_{2}](aq) + 2CaSO_{3} \cdot \frac{1}{2}H_{2}O(s) + H_{2}O(s) + H_{2}O($$

These hydroxylamine $[NOH(SO_3)_2^{-}]$ compounds are reduced further by the sulfite ion^{85}

$$Ca[NOH(SO_3)_2](aq) + CaSO_3 \cdot \frac{1}{2}H_2O(s) + \frac{3}{2}H_2O \rightarrow Ca[NH(SO_3)_2](aq) + CaSO_4 \cdot 2H_2O(s) \neq (2-39)$$

Upon leaving the top of the absorber, the clean flue gas is reheated for plume buoyancy and sent to the stack. The slurry solution drops to a holding tank from which most of the solution is returned to the top of the absorber. A small stream passes to a neutralization reactor where sulfuric acid is added to convert the sulfite solid to soluble bisulfite and solid gypsum:⁸⁶

$$2CaSO_{3} \cdot {}^{1}_{2}H_{2}O(s) + H_{2}SO_{4}(aq) + H_{2}O \rightarrow CaSO_{4} \cdot 2H_{2}O(s) + Ca(HSO_{3})_{2}(aq)$$
(2-40)

This stream passes to a thickener from which the bottoms are sent to a centrifuge to separate the solid gypsum by-product from the liquor which is returned to the absorber. The overflow from the thickener is primarily recycled to the limestone slurry preparation tank. The remainder is sent to a thermal decomposer where sulfuric acid is added to hydrolyze the N-S compounds.¹⁸

$$2Ca[NH(SO_3)_2](aq) + 2H_2O \stackrel{H^+}{\rightarrow} Ca(NH_2SO_3)_2(aq) + Ca(HSO_4)_2(aq) (2-41)$$

$$(NH_2SO_3)_2(aq) + Ca(HSO_4)_2(aq) + 6H_2O \stackrel{H^+}{\rightarrow} 2NH_4HSO_4(aq) + 2CaSO_4 \cdot 2H_2O(s) + (2-42)$$

The ammonium bisulfate solution is pumped to another neutralization reactor where lime is added.⁸⁷

$$NH_4HSO_4(aq) + Ca(OH)_2 + H_2O \rightarrow CaSO_4 \cdot 2H_2O(s) \downarrow + NH_4OH(aq) \qquad (2-43)$$

MHI has three possible methods of removing this ammonium hydroxide:

Ca

- decompose by increasing pH
- decompose thermally
- strip out with makeup H₂O

The remaining gypsum slurry is pumped to the limestone slurry preparation tank.

The fundamental design equation used for gas absorption column design is $^{\rm 32}$

$$\int_{Y_{a}}^{Y_{b}} \frac{dy}{(y-y^{\star})} = \left(\frac{K_{y}^{a}}{G_{y}}\right) Z \qquad (2-11)$$

- where y = bulk NO_X concentration (mole fraction) of gas phase at any given point in column
 - y-y* = overall driving force for absorption (y* being the NO_{χ} concentration of a gas in equilibrium with a given liquid NO_{χ} con-; centration)
 - $Y_{\rm b}$ = inlet NO_x concentration

$$Y_{2}$$
 = outlet NO₂ concentration

- a ^ Ky = overall gas-phase mass transfer coefficient, lb-moles NO_x/ (ft²)(hr)(mole fraction)
 - a = area of gas-liquid interface per unit packed volume, ft^2/ft^3
- G_y = molal gas mass velocity, 1b-moles flue gas/(ft²)(hr)
 - Z = length of packed section of column, ft

In a column containing a given plate or packing configuration and being irrigated with a certain liquid flow, there is an upper limit to the gas flow rate. This limit's superficial gas velocity (volumetric gas flow rate/cross-sectional area of column) is called the flooding velocity. At this point, the gas flow completely impedes the downward motion of the liquid and blows the liquid out of the top of the column. The gas velocity obviously, must be lower than the flooding velocity. How much lower is a design decision. Often, it is an economic tradeoff between power costs and equipment costs. A low gas velocity will lower the pressure drop and, hence, the power costs but the absorber will have a larger diameter and cost more. High gas velocities have an opposite effect. Usually the optimum gas velocity is about one-half the flooding velocity.³³ The height of the column depends on the desired level of removal and on the rate of mass transfer. The latter consideration is why oxidation-absorption-reduction processes oxidize NO to more soluble NO2 prior to the absorber and why some processes add water soluble catalysts to the scrubbing solution to aid NO2 absorption. The oxidation step enables these processes to use shorter absorbers with lower liquid rates than either the absorption-oxidation or absorption-reduction processes. Table 2.3.9-1 presents the effects of boiler/flue gas variables on the design of oxidation-absorption-reduction systems. Both flue gas flow rate and NO, concentration can be affected by boiler operating conditions. Therefore a change in load on an industrial boiler may alter these variables markedly. The absorber must be designed to accommodate any anticipated load change. The column size and the liquid, oxidant, and catalyst flows must be designed for each application after examining the boiler operating history and establishing ringes of variation.

Typical ranges for several operating parameters for this type of process are shown in Table 2.3.9-2. Reagent concentrations were not available. Economics for the various processes cover a wide range presumably due to different techniques for oxidant generation and treatment of the scrubbing solution. Costs are reported to range from \$84-134/kW for utility applications' capital expense and 6.7-9 mills/kWh for operating expense.⁹¹

Presently, some of the oxidation-absorption-reduction processes have reached the prototype stage of development. Table 2.3.9-3 presents a list of oxidation-absorption-reduction process vendors and the status of development of their projects. The applications of this process have been predominately to oil-fired boilers. Some of the applications treat flue gas flow

2-155

rates similar to those for the standard boilers of this study. Application to industrial boilers, therefore, is technically feasible.

Variable	Design Effect
Presence of particulates	Requires prescrubber
Presence of SO ₂	Depends on individual process: if NO_2 is com- pletely reduced to N_2 or NH_3 by SO_3^- (as does MHI), then at least the stoichiometric $SO_2:NO_{\times}$ mole ratio of 3:1 is required ⁸⁸ [see equation (9-6)]; if NO_2^- is not reduced completely, then a different ratio will be necessary
Increased gas flow	Requires larger column diameter; increased liquid flow rate
Increased NO_{χ} concentration	Requires larger column height; increased gas- phase oxidant flow rate; increased liquid- phase catalyst concentration

TABLE 2.3.9-1. SYSTEM DESIGN CONSIDERATIONS

TABLE 2.3.9-2. TYPICAL RANGES OF OPERATING VARIABLES FOR JXIDATION-ABSORPTION-REDUCTION PROCESSES^{89,90}

Variable		Range	
Liquid/Gas Ratio, 1/Nm ³		2–12	
Oxidant/NO Mole Ratio	0 ₃ systems	0.6-1.0	
	C1O ₂ systems	0.55	
SO_2/NO_{x} Mole Ratio		2.5-5.0	
Superficial Gas Velocity, m/s		3–5	
Pressure Drop, mmH_2O		200-500	

Vendor	Status of Development
Chiyoda	1975: 1000 Nm ³ /hr flue gas from heavy oil- fired boiler
Ishikawajima-Harima Heavy Industries	1975: 5000 Nm ³ /hr flue gas from oil-fired boiler (3000 hours continuous)
Mitsubishi Heavy Industries	1975: 2000 Nm ³ /hr flue gas from heavy oil- fired boiler (700 hours continuous)
Osaka Soda	1976: 60,000 Nm ³ /hr flue gas from oil-fired boiler
Shirogane	1974: 48,000 Nm ³ /hr flue gas from oil-fired boiler
Sumitomo Metal-Fujikasui: Calcium Process	1976: 25,000 Nm ³ /hr flue gas from sintering furnace
Sumitomo Metal-Fujikasui: Sodium Process	1973: 62,000 Nm ³ /hr flue gas from heavy oil- fired boiler (5 others)

TABLE 2.3.9-3. PROCESS VENDORS OF OXIDATION ABSORPTION-REDUCTION PROCESSES^{92,93}

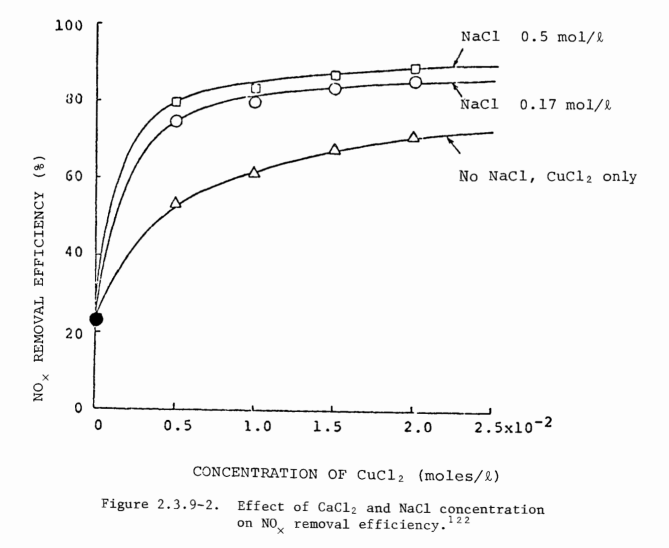
2.3.9.2 System Performance--

Results of oil-fired tests show up to 90 percent NO_{χ} reduction and >95 percent SO_2 reduction. Figures 2.3.9-2 and 2.3.9-3 illustrate NO_{χ} removals as a function of pH and additive concentrations.

The primary disadvantage of these processes is the utilization of costly gas-phase oxidants which create secondary wastewater pollution problems. Both ozone and chlorine dioxide are highly unstable so they cannot be stored and must be generated onsite. O_3 , the more expensive of the two, is generated by a high energy corona discharge in air. This instantaneous process requires significantly large amounts of electricity. ClO_2 is generated by a slower chemical reaction (requires about 20 minutes to respond to a change in demand) which could make it less responsive to boiler load changes. The use of ClO_2 introduces an additional secondary pollutant, chlorides, besides the nitrite salt problem. Significant amounts of corrosionresistant material are required for oxidation-absorption-reduction processes, regardless of which oxidant is utilized. Some of the processes would not be applicable to low sulfur oils as they require large amounts of SO_2 to obtain NO_2 (aq) or NO_2 reduction.

CONDITIONS

```
CONCENTRATION OF CaSO_3: 5 wt%
pH OF LIQUOR : 5.5
NO_2/(NO + NO_2) : 0.95
```



CONDITIONS

CONCENTRATION OF CaSO3	:	5 wt%
CONCENTRATION OF CuCl ₂	:	0.01 mole/l
CONCENTRATION OF NaCl	:	l wt%
$NO_2/(NO + NO_2)$:	0.95

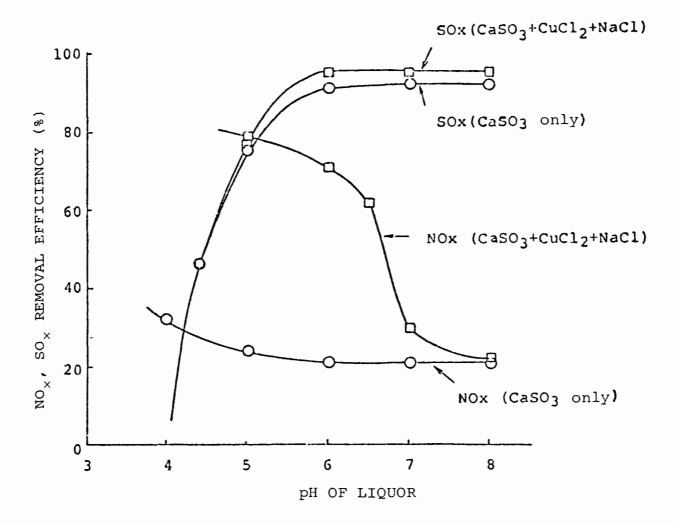


Figure 2.3.9-3. Effect of pH on \rm{SO}_{x} and \rm{NO}_{x} removal efficiency.¹²³

2.3.10 Oxidation-Absorption

2.3.10.1 System Description--

As a group, oxidation-absorption processes include those oxidation processes which do not qualify for the oxidation-absorption-reduction category. Basically, there are two types of oxidation-absorption processes. One is a simplified version of the oxidation-absorption-reduction process and uses an excess of ozone to selectively oxidize NO_x to N_2O_5 which is absorbed into aqueous solution and concentrated to form a 60 percent nitric acid (HNO₃) by-product. There is no reduction of $NO_x(NO_2)$ by the absorption of SO_2 (as SO_3^-) and no wastewater treatment facility. The other type of oxidation-absorption process is based on equimolar NO-NO₂ absorption: absorbing N_2O_3 which is formed by the gas-phase reaction of NO and NO₂.

A flow diagram of the Kawasaki Heavy Industries oxidation-absorption process is shown in Figure 2.3.10-1. Flue gas is taken from the boiler after the air preheater. It passes countercurrent to a magnesium hydroxide $[Mg(OH)_2]$ slurry in the first section of the absorber. There, SO₂ is absorbed and undergoes the following reactions.⁹⁵

$$SO_2(g) \rightarrow SO_2(aq)$$
 (2-23)

$$Mg(OH)_2(s) + SO_2(aq) + 5H_2O \rightarrow MgSO_3 \cdot 6H_2O(s) \downarrow$$

$$(2-44)$$

The SO_2 -free flue gas passes to the first denitrification section of the absorber while the liquid effluent drops to a holding tank. A recycle NO_2 stream is added to the flue gas to bring the $NO:NO_2$ mole ratio to 1:1. The resulting mixture then passes countercurrent to a $Mg(OH)_2$ slurry. Equimolar amounts of YO and NO_2 react and are reabsorbed in the following manner.⁹⁶

$$NO(g) + NO_2(g) + N_2O_3(g)$$
 (2-45)

$$N_2O_3(g) \rightarrow N_2O_3(aq)$$
 (2-46)

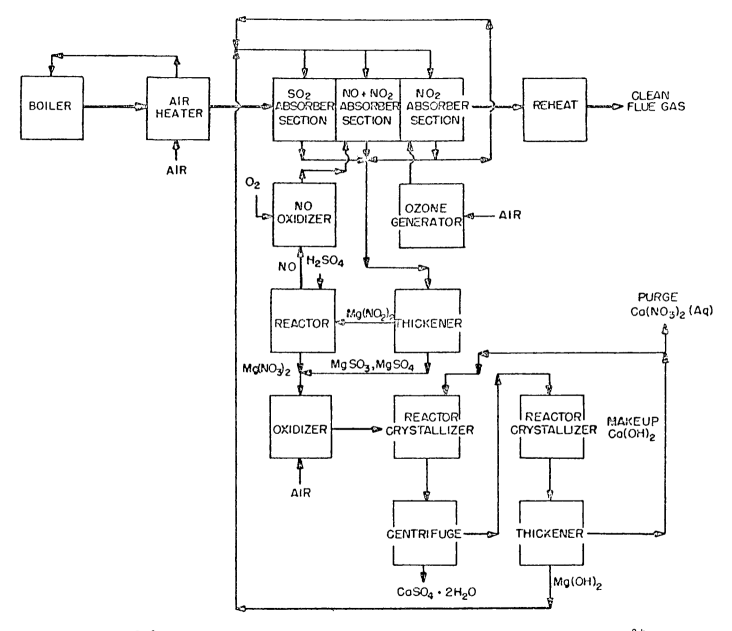


Figure 2.3.10-1. Flow diagram of Kawasaki Heavy Industries process.94

$$Mg(OH)_{2}(aq) + N_{2}O_{3}(aq) \rightarrow Mg(NO_{2})_{2}(aq) + H_{2}O \qquad (2-47)$$

The flue gas passes out of the top of this absorption section while the liquid effluent drops to the holding tank. Because the rate of reaction 2-45 decreases with NO_x concentration (below 200 ppm it becomes negligible), it is necessary to further reduce NO_x by injecting ozone to oxidize the remaining NO to NO_2 . The mixture then passes to the final denitrification section of the absorber and is passed countercurrent to a Mg(OH)₂ slurry. This section of the absorber is described by the following reactions.⁹⁷

$$2NO_2(g) \rightarrow N_2O_4(g)$$
 (2-8)

$$N_2O_4(g) \neq N_2O_4(aq)$$
 (2-9)

$$2N_2O_4(aq) + 2M_g(OH)_2(s) \rightarrow M_g(NO_3)_2(aq) + M_g(NO_2)_2(aq) + 2H_2O$$
 (2-48)

The clean flue gas leaves the top of this absorber section, is passed to a reheater for plume buoyancy and sent to the stack. Part of the liquid effluent from this section is recycled to the tops of the absorber sections while the rest drops to the holding tank. The slurry solution is pumped to a thickener which separates the soluble nitrite (NO_2) and nitrate (NO_3) salts from the solid magnesium sulfite. The overflow from the thickener passes to a NO_2 decomposition reactor where sulfuric acid is added.⁹⁸

$$3Mg(NO_2)_2(aq) + 2H_2SO_4(aq) \rightarrow 2MgSO_4(aq) + Mg(NO_3)_2(aq) + 4NO(g) \uparrow + 2H_2O$$

(2-49)

The NO off-gas passes through an oxidizer where it is oxidized by air to NO_2 and sent to the first denitrification section of the absorber. The effluent from the decomposition reactor is mixed with the thickener bottoms and pumped to a second oxidizer.⁹⁹

$$MgSO_{3} \cdot 6H_{2}O(s) + \frac{1}{2}O_{2}(g) \rightarrow MgSO_{4}(aq) + 6H_{2}O \qquad (2-50)$$

The magnesium sulfate formed in the oxidizer is treated with calcium nitrate in a gypsum production reactor.²⁵

$$Ca(NO_{3})_{2}(aq) + MgSO_{4}(aq) + 2H_{2}O \rightarrow CaSO_{4} \cdot 2H_{2}O(s) + Mg(NO_{3})_{2}(aq)$$
(2-51)

The products of this reaction are sent to a centrifuge to remove the solid gypsum by-product. The liquid from the centrifuge goes to another decomposition reactor where makeup lime slurry is added.¹⁰⁰

$$Mg(NO_{3})_{2}(aq) + Ca(OH)_{2}(s) \rightarrow Ca(NO_{3})_{2}(aq) + Mg(OH)_{2}(s) \qquad (2-52)$$

The magnesium hydroxide product is separated in a thickener and recycled to the absorbers. The thickener overflow stream is split and part is recycled to the gypsum production reactor and the rest is concentrated to form low-grade liquid fertilizer by-product, $Ca(NO_3)_2$.

Since the processes in this category are all very different, especially with respect to chemistry, generalization of typical ranges of operating variables is not meaningful and, therefore, not presented. No published economics for these processes were found.

Presently, the equimolar absorption-type oxidation-absorption processes are still in the pilot-unit stage of development. Table 2.3.10-1 presents a list of all oxidation-absorption process vendors and their project's status of development. These processes have not yet been applied to oil-fired boilers.

Vendor	Status of Development		
Kawasaki Heavy Industries	1975: 5000 Nm ³ /hr flue gas from coal- fired boiler		
Tokyo Electric-MHI (NO _X only)	1974: 100,000 Nm ³ /hr flue gas from natural gas-fired boiler		
Ube Industries	No information available		

TABLE 2.3.10-1. PROCESS VENDORS OF OXIDATION-ABSORPTION PROCESSES¹⁰⁰

2.3.10.2 System Performance--

No oil-fired tests have been performed. Very little information has been published on any of the tests conducted.

The production of nitrate salts poses a potential secondary pollution problem. The plan for reclaiming and concentrating the nitrates as $Ca(NO_3)_2(aq)$ for liquid fertilizer is questionable as the by-product is of low quality and may not be easily marketable in the U.S. Also, the gypsum by-product would be contaminated with various nitrate and sulfite salts, and therefore, would probably be useful only as landfill material. Much corrosion-resistant material is necessary due to the utilization of ozone and circulating magnesium slurries. The three absorber sections, with their respective operating conditions, and ozone generation present complex process control problems. The process steps of several absorber sections in series (large fan requirements), ozone generation (corona discharge), flue gas reheat (inline heater), and by-product and wastewater treatment are all energy intensive and present technical and economic disadvantages when compared to other simpler FGT processes.

2.4 CONTROLS FOR NATURAL GAS-FIRED BOILERS

In the previous two sections which discuss controls for coal and oilfired boilers, up to ten different process types are presented. Many of these process types are not considered here for application to natural gasfired boilers for two reasons. First, natural gas-fired boilers have no SO₂ emission problems, and therefore, the simultaneous systems are not considered. Second, these boilers have no particulate emissions, and as a result, the systems designed specifically for high particulate applications are not considered. This leaves two systems to be considered for application to natural gas-fired boilers and these are discussed in the following sections.

2.4.1 Selective Catalytic Reduction-Fixed Packed Bed Reactor

2.4.1.1 System Description--

Fixed packed bed systems are applicable only to flue gas streams containing less than 20 mg/Nm^3 of particulates. As such, they are applicable to natural gas-fired boilers.

The primary feature of these systems is the reactor which contains the catalyst. As the name implies, the granular catalyst is randomly packed in a stationary bed. An example of a typical fixed bed reactor is shown in Figure 2.4.1-1. The important features of the reactor are:

- the catalyst
- the catalyst support
- the gas distributor

The catalyst can be either spherical or cylindrical in shape. Spherical granules typically range in size from 4-10 mm in diameter.¹⁰³ The composition varies from process to process and most formulations are proprietary. The catalyst is supported either by inert packing (as shown in Figure 2.4.1-1) or by a perforated support plate (Figure 2.4.1-2). The catalyst supports

2-165

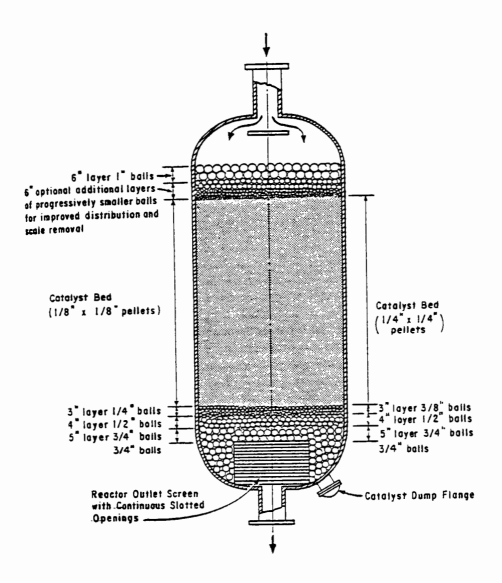


Figure 2.4.1-1. Example of typical fixed packed bed reactor.¹⁰¹

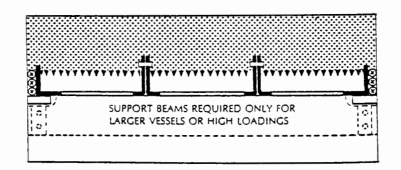


Figure 2.4.1-2. Example of catalyst support plate. 102

hold the catalyst fixed in place in order to prevent both mobilization of the particles by the gas stream and catalyst rearrangement which would allow channelling of the flue gas. The gas distributor can be a perforated plate or similar device which spreads the gas flow across the entire cross-section of the catalyst bed.

A typical fixed bed SCR process layout is presented in Figure 2.4.1-3. Several arrangements are possible, however, for application to new boilers this arrangement is the most desirable.

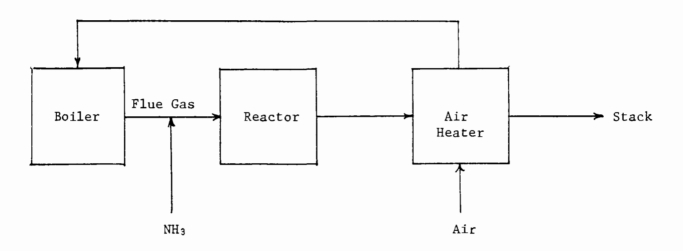


Figure 2.4.1-3. Process layout for fixed bed SCR process.

The principle of operation of these systems involves a gas phase reaction between ammonia (NH₃) and NO_{\times} (NO and NO₂). These reactions are presented most accurately by¹²

$$4NH_3 + 4NO + O_2 \neq 4N_2 + 6H_2O$$
 (2-53)

$$4NH_3 + 2NO_2 + O_2 \neq 3N_2 + 6H_2O$$
 (2-54)

The first reaction predominates since flue gas NO_{χ} is typically 90-95 percent NO. As shown, the NO_{χ} is reduced to molecular nitrogen (N₂) which exits with the flue gas stream.

The primary design equation used with these processes is the standard equation for reactor design,¹³ represented by

$$\frac{V}{F} = \int_{0}^{x} \frac{dx}{r}$$
(2-3)

The reaction rate, r, for each NO reduction reaction can be represented by

$$r = k[NH_3]^{a}[NO]^{b}[O_2]^{c}$$
(2-4)

where k is the reaction rate constant

[NH3], [NO], [O2] are the reactant concentrations a, b, c, are empirically determined exponents

The reaction rate is different for each catalyst formulation and, therefore, values for k, a, b, and c must be determined for the particular catalyst to be used before any design can be performed. The reaction rate constant is usually described by the Arrhenius equation

$$-\frac{E}{RT}$$

$$k = Ae$$
(2-5)

where A is the frequency factor E is the activation energy R is the universal gas constant, and T is the temperature Values for k, a, b and c for two catalyst formulations are shown in Table 2.4.1-1. Values for other catalyst formulations will be different.

TABLE 2.4.1-1.

REACTION RATE DATA FOR TWO CATALYST FORMULATIONS¹¹ Catalyst: V205 on Al203 $k = 2.05 \times 10^3 e^{-\frac{9650}{RT}}$ a = 0.30b = 0.22c = 0.05Catalyst: Fe-Cr on Al₂O₃ $k = 3.25 \times 10^3 e^{-\frac{10,860}{RT}}$ a = 0.45b = 0.10c = 0.15

The most important design and operating variables are similar to those for moving bed systems using granular catalysts. These are listed, along with typical ranges, in Table 2.4.1-2.

<u> </u>	Typical Range			
Variable	(For Oil)	(For Gas)		
Gas Velocity, m/s	0.5 - 1.5	0.5 - 1.5		
Bed Depth, m	0.2 - 0.6	0.2 - 0.4		
Space Velocity, hr^{-1}	6,000 - 10,000	8,000 - 15,000		
Pressure Drop, mmH ₂ O	40 - 80	40 - 70		
Temperature, °C	350 - 400	300 - 400		

TABLE 2.4.1-2. DESIGN AND OPERATING VARIABLES FOR FIXED PACKED BED SYSTEMS¹⁴

Other variables that affect the process are:

- flue gas flow rate
- NO_x control level
- NO_v concentration
- boiler load variation

The flue gas flow rate and NO_X control level determine the catalyst volume required (hence reactor size). Increases in either parameter also increase the reactor size. The NO_x concentration is primarily a function of fuel type used in the standard boilers. Higher concentrations require larger NH3 storage and vaporization equipment; reactor size is not affected. Boiler load can affect several things including flue gas temperature, flow rate and NOx concentration. It is usually necessary to maintain reaction temperatures of 350 to 400°C. Temperature control equipment may be necessary to accomodate large boiler load variations which cause lower flue gas temperatures. Where these variations are present, some equipment overdesign may be warranted to insure a constant control level. These variables are discussed in more detail in the section on moving bed SCR systems for coal-fired boilers, Section 2.2.2. Costs of fixed packed bed systems range from \$16-49/kW (capital) and 1.2-1.8 mills/kWh (operating). These costs are based on utility applications as well as a variety of process and operating conditions.

There are vendors of fixed packed bed SCR systems and all are Japanese. Vendors are listed in Table 2.4.1-3 and the scale of development is also noted. Fixed packed systems have been applied to industrial and utility boilers in Japan. Existing installations are shown in Tables 2.4.1-4 and 2.4.1-5. Currently, there are no installations in the U.S.

2.4.1.2 System Performance--

Typical performance data for fixed packed bed SCR systems are shown in Figures 2.4.1-4 through 2.4.1-8. These data indicate that NO_{χ} removals 90 percent and higher are achievable with these systems. This allows them to be considered for all control levels of interest in this study.

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TABLE 2.4.1-3.	VENDORS	OF SCR	FIXED	BED	SYSTEMS
	FOR GAS-	-FIRED	APPLICA	ATION	1S ²¹

Vendor	Notes		
Sumitomo	Tested on commercial scale equipment		
Hitachi Zosen	Tested on commercial scale equipment		
Hitachi, Ltd.	Tested on commercial scale equipment		
Mitsubishi Heavy Industries	Tested on commercial scale equipment		
Ishikawajima-Harima Heavy Industries	Tested on commercial scale equipment		
Mitsui Toatsu Chemical	Has not been to boilers		
Kawasaki Heavy Industries	Tested on pilot scale equipment		
Mitsubishi Kakoki Kaisha	Tested on commercial scale equipment		

TABLE 2.4.1-4. EXISTING FGT INSTALLATIONS OF SCR FIXED BED SYSTEMS GAS-FIRED INDUSTRIAL BOILERS²¹

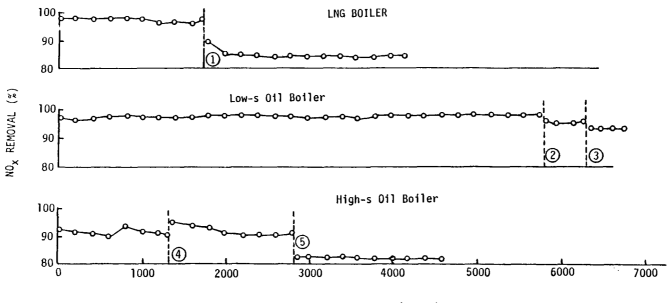
Location	User	Process Developer	Fuel	Capacity (Nm ³ /hr)	Completion Date
Takaishi	Osaka Gas	Mitsubishi H.I.	LNG	30,000	December 1976

TABLE 2.4.1-5. EXISTING FGT INSTALLATIONS OF SCR FIXED BED SYSTEMS GAS-FIRED UTILITY BOILERS²¹

Location	User	Process Developer	Fuel	Capacity (Nm ³ /hr)	Completion Date
Kokura	Kyushu Electric	Mitsubishi H.I.	LNG	3,380,000*	October 1978
Chita	Chubu Electric	Hitachi, Ltd.	LNG	4,000,000*	April 1978

*Flow rate is combined value from two boilers.

.





Circled figures show times when SV and $\mathrm{NH}_3/\mathrm{NO}$ mole ratio were changed.

1	. sv	10,000	20,000	hr-1	2.	. SV	10,000	15,000 hr ⁻¹	•
3	. sv	15,000	20,000	hr-1	4.	. sv	6,200	4,500 hr ⁻¹	L
5	. sv	4,500	6,200	and the	mole r	ratio	0.95	0.83	

Figure 2.4.1-4. Test results at gas- and oil-fired boilers.¹²⁴

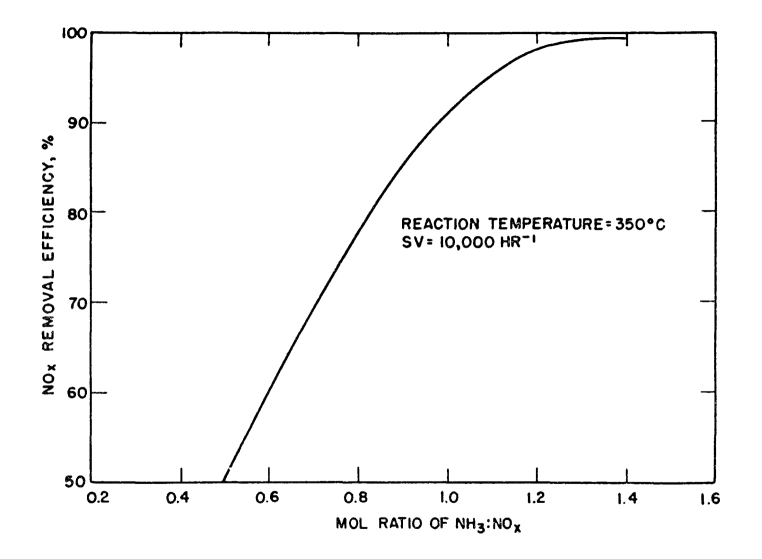


Figure 2.4.1-5. Characteristic curve of the effect of mol ratio of $NH_3:NO_{\times}$ on NO_{\times} removal efficiency for Hitachi, Ltd. Process.¹²⁵

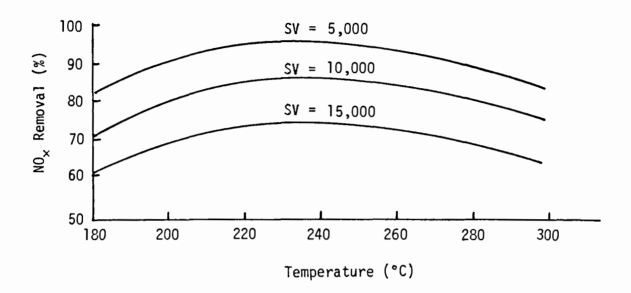


Figure 2.4.1-6. Performance of catalyst MTC-102 (flue gas by LPG burning).¹²⁶

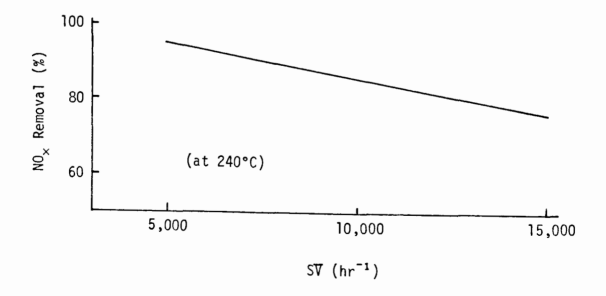


Figure 2.4.1-7. SV and NO_x removal (MTC-102) (flue gas by LPG burning).¹²⁷

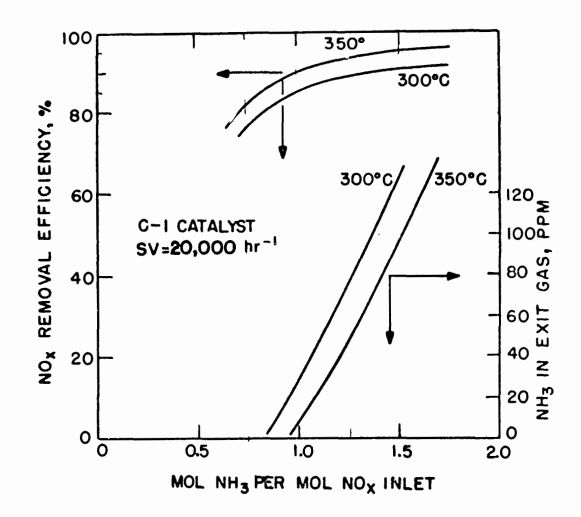


Figure 2.4.1-8. Relationship among inlet NH₃:NO_× mol ratio, NO_× removal efficiency, and exiting NH₃ concentration using the Sumitomo Chemical C-1 Catalyst.¹²⁸

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2.4.2 Absorption-Oxidation

2.4.2.1 System Description--

Absorption-oxidation processes remove NO_{\times} from flue gas by absorbing the NO or NO_{\times} into a solution containing an oxidant which converts the NO_{\times} to a nitrate salt. Two types of gas/liquid contactors can be used and examples of each type are shown in Figure 2.4.2-1. Both perforated plate and packed towers accomplish NO_{\times} absorption by generating high gas/liquid interfacial areas. The choice of one type of contactor is a design decision made to achieve a given removal for the least cost.

A generalized process flow diagram is shown in Figure 2.4.2-2. Flue gas is taken from the boiler after the air preheater. Before the gas can be sent to the NO_{χ} absorber, it must be SO_2 -free since SO_2 consumes prohibitive amounts of the costly liquid-phase oxidant. This is not a problem with natural gas fired-boilers since they have no SO_2 emissions. In most cases, the oxidant is permanganate (MnO_4). The flue gas enters the distributing space at the bottom of the NO_{χ} absorber, below the packing or plates. The gas passes upward through the column, countercurrent to the flow of the liquid absorbent/oxidant (usually a KOH solution containing KMnO₄). NO_{χ} is absorbed and then oxidized over the length of the column according to the following reactions.³¹

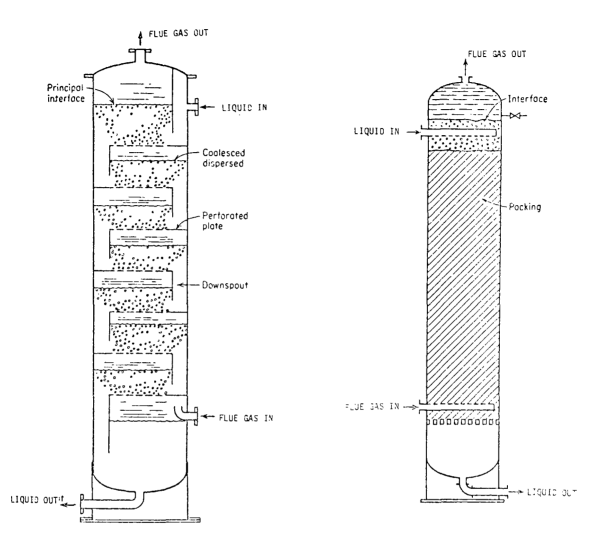
$$NO(g) \rightarrow NO(aq)$$
 (2-6)

$$NO(aq) + KMnO_4(aq) \rightarrow KNO_3(aq) + MnO_2(s) \downarrow$$
(2-7)

$$2NO_2(g) \rightarrow N_2O_4(g)$$
 (2-8)

$$N_2O_4(g) \rightarrow N_2O_4(aq)$$
 (2-9)

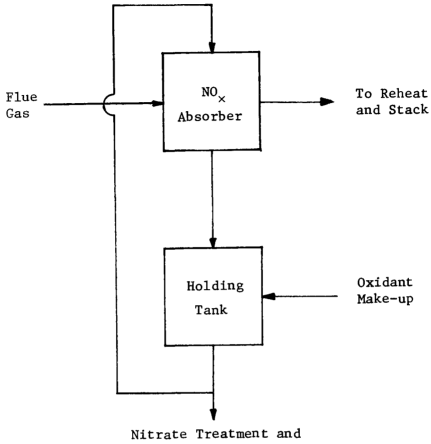
$$N_2O_4(aq) + 2K_2MnO_4(aq) + 2KMnO_4(aq) + 2KNO_2(aq)$$
 (2-10)



Perforated Plate Absorber

Packed Absorber

Figure 2.4.2-1. Gas/liquid contactor options for Absorption-Oxidation Processes.²⁹



Oxidant Regeneration

Figure 2.4.2-2. Process flow diagram for absorptionoxidation process.³⁰

Since most of the NO_{χ} from combustion processes occurs as NO_{χ}^{32} reactions 2-6 and 2-7 predominate. The clean gas passes out of the top of the absorber to a heater for plume buoyancy and is sent to the stack. The absorbing solution drops to a holding tank where makeup KOH and/or KMnO₄ are added. This solution flows to a centrifuge to separate the solid MnO₂ which is then electrolytically oxidized to MnO₄. The remaining solution is either concentrated in an evaporator to form a weak KNO₃ solution or is electrochemically treated to produce a weak HNO₃ solution and a mixed stream of KOH and KNO₃.

The fundamental design equation used for gas absorption column design is

$$\int_{Y_{a}}^{Y_{b}} \frac{dy}{(y-y^{\star})} = \left(\frac{K_{a}}{G_{y}}\right) Z \qquad (2-11)$$

- where y = bulk NO_x concentration (mole fraction) of gas phase at any
 given point in column
 - y-y* = overall driving force for absorption (y* being the NO_x concentration of a gas in equilibrium with a given liquid NO_x concentration)
 - $Y_{b} = inlet NO_{x}$ concentration
 - $Y_a = outlet NO_x$ concentration
 - Ky = overall gas-phase mass transfer coefficient, lb-moles NO_x/
 (ft²)(hr)(mole fraction)
 - a = area of gas-liquid interface per unit packed volume, ft^2/ft^3
 - G_y = molal gas mass velocity, lb-moles flue gas/(ft²)(hr)
 - Z = length of packed section of column, ft

In a column containing a given packing or plate configuration and being irrigated with a certain liquid flow, there is an upper limit to the gas flow rate. This limit's superficial gas velocity (volumetric gas flow rate/ cross-sectional area of column) is called the flooding velocity. At this point, the gas flow completely impedes the downward motion of the liquid and blows the liquid out of the top of the column. The gas velocity, obviously, must be lower than the flooding velocity. How much lower is a design decision. Often, it is an economic tradeoff between power costs and equipment costs. A low gas velocity will lower the pressure drop and, hence, the power costs but the absorber will have a larger diameter and cost more. High gas velocities have an opposite effect. Usually the optimum gas velocity is about one-half the flooding velocity.³⁴ The height of the column depends on the desired level of removal and on the rate of mass transfer. The latter is a major problem for these systems trying to achieve large NO_x reductions since NO is relatively insoluble in water. This can be seen in Table 2.4.2-1.

	Boiling Point, °C	Solubility in Cold Water (0°C), cm ³	Solubility in Hot Water (60°C), cm ³
NO	-151.8	7.34/100 cc H ₂ 0	2.37/100 cc H ₂ 0
NO ₂	21.2	soluble, decomposes	

TABLE 2.4.2-1. NITROGEN OXIDES CHARACTERISTICS³⁵

One can see that NO has a very limited solubility in water and, since most NO_{\times} is present as NO, the rate of mass transfer (absorption) is going to be relatively slow. This means that the absorber must be tall with a high liquid flow rate. Table 2.4.2-2 presents the effects of boiler/flue gas variables on the design of absorption-oxidation systems.

Variable	Design Effect		
Presence of particulates	Requires prescrubber		
Presence of SO ₂	Requires FGD pretreatment		
Increased gas flow	Requires larger column diameter; increased liquid flow rate		
Increased NO _x concentration	Requires larger column height; increased oxidant concentration		

TABLE 2.4.2-2. SYSTEM DESIGN CONSIDERATIONS

Both flue gas flow rate and NO_x concentration can be affected by boiler operating conditions. Therefore a change in load on an industrial boiler may alter these variables markedly. The absorber must be designed to accommodate any anticipated load changes. The column size and the liquid and oxidant flows must be designed for each application after examining the boiler operating history and establishing ranges of variation.

None of the sources consulted for this study could supply typical ranges for operating variables such as liquid/gas ratio, reagent concentrations or pressure drops and, as a result, none are presented here. Economic data were not presented either. One source did estimate the removal for absorptionoxidation processes to be 85 percent.³⁶

Presently, absorption-oxidation processes are still in the pilot unit stage of development. Table 2.4.2-3 presents a list of absorptionoxidation process vendors and the status of development of their projects. One can see from the table that no gas-fired flue gas tests have been performed.

Vendor	Status of Development		
Hodogaya	No information available; stopped development on process		
Kobe Steel	1974: 1000 Nm ³ /hr gas from iron-ore sintering furnace; stopped development on process		
MON (Mitsubishi Metal, MKK, Nikon Chemical)	1974: 4000 Nm ³ /hr flue gas from oil-fired boiler		
Nissan Engineering	1972: 4 pilot plants, 100-2000 Nm ³ /hr tail gas from HNO ₃ plant		

TABLE 2.4.2-3. PROCESS VENDORS OF ABSORPTION-OXIDATION PROCESSES 37,38

2.4.2.2 System Performance--

No gas-fired tests have been made. No information has been published on tests conducted with other fuels. The relative insolubility of NO in water may present a major obstacle to achieving the stringent level of control (90 percent NO_{χ} reduction) by absorption-oxidation processes. Another primary drawback of absorption-oxidation systems is the production of nitrate salts (see Equation 2-7), a secondary pollutant. These processes probably could not be applied on a large scale as wastewater treatment systems (chemical or biological) do not remove nitrogen compounds from the wastewater.³⁹ Trying to recover the nitrates as nitric acid for industrial use or potassium nitrate for fertilizer does not seem promising as the by-products are of low quality. Also, the use of an expensive, liquid-phase oxidant requires stainless steel and other corrosion resistant materials of construction. The process steps of oxidant regeneration (electrolysis) and flue gas reheat (inline heater) are all energy intensive and present technical and economic disadvantages.

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SECTION 3

CANDIDATES FOR BEST SYSTEMS OF EMISSION REDUCTION

The ten systems discussed in Section 2 are not applicable to all combinations of boiler types and fuels of interest in this study. However, several of these systems may be applicable to a specific boiler/fuel combination (*i.e.*, capable of removing sufficient NO_{\times} to meet proposed emission regulations). In this section, NO_{\times} control techniques which are applicable to the various boilers and fuels considered in this study are selected. The section is organized to compare NO_{\times} -only and simultaneous NO_{\times}/SO_{\times} reduction systems separately. The result is a set of candidate control techniques that will be evaluated in detail in subsequent sections to determine the "best" system for NO_{\times} control by FGT.

3.1 CRITERIA FOR SELECTION

Two sets of evaluation criteria are used to determine the set of candidate systems. One is the level of NO_{\times} control desired which determines the set of systems available for further evaluation. The other is a set of evaluation criteria that will allow comparison of the systems capable of meeting a particular level of control.

3.1.1 Factors Considered in Selection of Best Systems

A consistent set of rating criteria was used to evaluate and compare each of the FGT systems described in section 2 that are capable of achieving the proposed NO_{\times} removal levels. These criteria and the weighting factors are shown in Table 3.1.1-1. As can be seen, the criteria receiving most emphasis are status of development, economics, performance, and reliability.

3-1

Evaluation Category	Total Points
Performance	14
Operational/Maintenance Impacts on Performance	7
Preliminary Environmental Impacts	9
Preliminary Economic Impacts	15
Preliminary Energy/Material Impacts	10
Boiler Operation and Safety	4
Reliability	14
Status of Development	16
Adaptability to Existing Sources	6
Compatability with Other Control Systems	5
	100

TABLE 3.1.1-1. RATING CRITERIA AND WEIGHTING FACTORS

Emphasis is placed on the most developed FGT systems since they represent the most likely controls to be applied if a high degree of NO_{\times} control is required on industrial boilers. An FGT system must achieve the necessary NO_{\times} reduction and do so as economically as possible, hence the heavy emphasis on performance and economics. These are important considerations for any application. Reliability is heavily weighted because it is common for an industrial boiler to supply one or several continuous manufacturing processes. A high reliability is required to avoid frequent boiler shutdowns with subsequent loss of revenues due to dependency of the manufacturing process on the boiler.

It should be pointed out that only large differences in point values are significant while small differences are not. For example, ratings which differ by a factor of two are significant. However, two ratings 10 points apart do not necessarily indicate the superiority of one process. A more detailed breakdown of the evaluation criteria and the point values assigned is presented in Table 3.1.1-2. The basis for the detailed breakdown is discussed below. The analysis of each system using these criteria is discussed in Section 3.2.

	Item	Quality	Points
1.	Performance		
	a. Desired control level (stringent, intermediate, or moderate) as percent of system's maximum design capability $\left(\frac{\text{Desired Control Level}}{\text{Maximum Design Control Level}} \right)$ x 100	<70 70 - 80 80 - 90 90 -100 >100	8 6 4 2 No Go
	b. Particulate handling capability	Great Some None	4 1 0
	c. Load following ability	Good Fair Poor	2 1 0
2.	Operation and Maintenance İmpacts on Performance		
	a. Moving parts	Few Many	1 0
	b. Solids handling	No Yes	1 0
	c. Process separability	Once-through Regenerable	1 0
	d. Flue gas composition sensitivity	No Yes	1 0
	e. Prescrubbing necessary	No Yes	1 0
	f. Process stability	Simple process & insensitive control needs	2
		Complex process or sensitive control needs	1
		Complex process & sensitive control needs	0
3.	Preliminary Environmental Impacts		
	a. Secondary pollutants - Air	None Potential Some Major	3 2 1 0
	- Liquid	None Some Major	3 1 0

TABLE 3.1.1-2. SPECIFIC POINT VALUES ASSOCIATED WITH SELECTION FACTORS

	Item	Quality	Points
3.	a. Secondary pollutants (Cont'd) – Solid	None Some Major	3 1 0
4.	Preliminary Economic Impacts		
	a. Capital investment	<50% mean 50% mean 75% mean Mean	7 6 5 4
	b. Operating costs	125% mean 150% mean >150% mean	3 2 1
	c. Marketable by-product	Potential None	1 0
5.	Preliminary Energy/Material Impacts		
	a. Electrical demand	<1% output 1 - 2% 2 - 3% 3 - 4% 4 - 5% >5%	5 4 3 2 1 0
	b. Auxiliary fuel use	No Yes	1 0
	c. Energy intensive regeneration or by-product treatment	None Some Heavy	2 1 0
	d. Raw material demand	Light Moderate Heavy	2 1 0
6.	Boiler Operation and/or Safety		
	Boiler impacts or safety hazards	None Potential Yes	4 2 0
7.	Reliability		
	a. Plugging and scaling	None Some Much	5 2 0

TABLE 3.1.1-2. (Continued)

	Item	Quality	Points
7.	b. Simplicity - Number process steps	<3	6
		3	5
		4	4
		5	3
		6	2
		7	1
		>7	0
	c. Material of construction	Carbon steel	3
		Some corrosion resistant	
		material	1
		Much corrosion resistant	
		material	0
8.	Development Status		
	a. Scale demonstrated	Commercial	10
		Prototype	8
		Pilot	5
		Bench	2
		Conceptual	0
	b. Length of operation	>5000 hours	3
	.	3000 - 5000	2
		1000 - 3000	1
		<1000	0
	c. Uncertainties in technology	No	3
		Yes	0
9.	Adaptability to Existing Sources		
	a. Retrofit	Easy	3
		Difficult	0
	b. Land required	Small	3
		Large	0
.0.	Compatability with Other Control Systems		
	a. FGD	Yes	3
		No	0
	b. ESP, other	Yes	2
	-	No	0

TABLE 3.1.1-2. (Continued)

3.1.1.1 Performance--

A primary concern in the selection of an NO_{\times} flue gas treatment system is the system's performance. The first aspect to consider here is the NO_{\times} removal capability. This study is organized by different levels of NO_{\times} control (stringent, intermediate, moderate). The processes' maximum removal capability is compared to these various control levels to show the ease with which the system can meet the removal requirement. Another measurement of a system's performance is its load following capability--how well the system responds to a sudden change in boiler load. Generally, large, complex systems do not respond to load changes as quickly as small, simple systems. Slow response is a disadvantage since it may result in increased emissions during load changes.

3.1.1.2 Operational and Maintenance Impacts--

This category is important for several reasons. A system with difficult operational steps or high maintenance requirements is not as desirable since it will require more manpower and increase operating costs. Reliability may also be adversely affected. For most FGT systems, this type of data is not available. In this study these impacts are inferred by examining each system and applying engineering judgment. The more mechanically complex a system is, the more likely it is to have operation and maintenance problems.

3.1.1.3 Preliminary Environmental Impacts--

This category, along with the economic and energy categories, relies on published information for data. Detailed analyses of the candidate systems in these areas will be conducted in a subsequent section. The data presented in this section are used for comparison purposes only. Obviously it is undesirable for an FGT system to remove NO_{\times} at the expense of emitting a secondary pollutant. For this reason secondary pollutants (air, liquid, and solid) emitted by the process, or potentially so, are identified. Systems with no secondary pollutants receive the highest ratings.

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3.1.1.4 Preliminary Economic Impacts--

With an industrial boiler it is probable that application of FGT will affect the price of products from a new or modified facility and thereby affect the salability of these products. For this reason, the lowest cost system that will adequately control NO_X is desirable. The areas considered are capital investment (\$/kW), operating costs (mills/kWh), and credits for marketable by-products. Cost data that are available are primarily for utility installations. While there is some economy of scale in the investment cost due to the large size of the facilities, the values are adequate for preliminary cost comparisons. Sample economy of scale calculations showing how the preliminary economic figures were generated are contained in Appendix II.

3.1.1.5 Preliminary Energy/Material Impacts--

It is desired to minimize energy and raw material consumption by an FGT process since this also minimizes operating costs. In addition, dependence on outside factors such as raw material supplies is reduced. The main system parameters considered are the electrical demand of the system, use of auxiliary fuels and energy, and intensive regeneration or by-product treatment processes. Also, heavy raw material demands are noted. Again, utility data are used for comparative purposes since very little industrial boiler data are available.

3.1.1.6 Boiler Operation and/or Safety--

It is desirable to minimize impacts of the FGT system on the boiler. The main areas of potential impacts are air heater fouling, duct scaling and stack corrosion. These impacts as well as safety aspects of the process are determined by inspection of the process equipment and chemistry.

3.1.1.7 Reliability--

Reliability data are not generally available for all of the process types considered. Many have not been applied on commercial scale equipment. Some reliability data are available for SCR systems, but data from other

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systems are necessary before the reliability of SCR systems can be compared on a relative basis. For most systems it can be said that simplicity is concommitant with reliability and this concept is used in the evaluation.

3.1.1.8 Development Status--

A crucial consideration in the selection of the best NO_{\times} control techniques by flue gas treating is the status of development of the processes. Presently, there are but a few commercial-size NO_{\times} FGT units in operation on industrial boilers--all in Japan. Because most of the flue gas treatment development work has been conducted fairly recently, it is vital that those systems which have been demonstrated most fully be given primary consideration for implementation to industrial boilers. For this study, availability by the year 1981 was estimated using the current status of development and reported on-going development. The size of the unit, length of operation, and any uncertainties in technology were all taken into account.

3.1.1.9 Adaptability to Existing Sources--

Since applying FGT to modified existing sources is generally more difficult than with new sources, the ease of retrofit was examined. Structural and equipment modifications necessary for retrofit are considered since existing boilers are not constructed to accommodate FGT systems. Land requirements of the FGT system are also considered, since existing industrial boilers are not necessar'ly located near large land areas. Quite frequently, they are located in the center of a plant and surrounded by equipment. Small systems requiring little boiler modification are desired.

3.1.1.10 Compatibility with Other Control Systems--

This category is related to retrofit and new installation. Where additional cont ol equipment is existing or planned for installation, an FGT system which loes not affect and is not affected by other control systems is desirable. This aspect of the processes is determined by inspection of the chemistry and equipment of the FGT system as well as other pollutant control systems.

3.1.2 Selection of Control Levels--Moderate, Stringent, and Intermediate

		Load		Uncon	trolled N	O _× Emissi	ions	
Fuel	Туре	(MW _t)		(1b/hr)		(11	0/10 ⁶ Btu	1)
Gas	Firetube	4.4		2.63			0.18	
Gas	Watertube	44		26.26			0.18	
0il- dist.	Firetube	4.4		2.38			0.16	
0il - dist.	Watertube	44		23.76			0.16	
0il- resid.	Watertube	44		60.00			0.40	
0il- resid.	Watertube	8.8		12.00			0.40	
			High S Eastern	Low S Eastern	Low S Western	High S 	Low S E	Low S
Coal	Underfeed	8.8	19.05	16.35	23.40	0.64	0.55	0.78

The control levels selected are applied to the following boilers:

			Eastern	Eastern	Western	E	E	W
Coal	Underfeed Stoker	8.8	19.05	16.35	23.40	0.64	0.55	0.78
Coal	Chaingrate	22	47.70	40.80	58.65	0.64	0.54	0.78
Coal	Spreader Stoker	44	95.40	81.45	117.15	0.64	0.54	0.78
Coal	Pulverized Coal	58.6	152.46	130.50	187.56	0.76	0.65	0.94

These NO_X emission levels are all lower than the following average State Implementation Plan (SIP) requirements except for one oil-fired boiler, one coal-fired boiler burning high sulfur eastern coal, and all coal-fired boilers burning low sulfur western coal.

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Fuel	$\frac{NO_{\times}\left(\frac{1b}{10^{6} Btu}\right)}{10^{6} Btu}$
Coal	0.7
Oil	0.3
Gas	0.2

The moderate level of control is defined as representing that level which is achievable applying techniques in current practice within industry. This is the least stringent emission reduction achievable applying accepted engineering practice. For FGT systems, this represents an NO_{\times} removal of approximately 70 percent. When considering NO_{\times} FGT, it is not reasonable to consider a removal level less than 70% since such levels can probably be achieved by combustion modification techniques at lower costs. Allowable NO_{\times} emissions at this control level are shown below:

Fuel	$\underline{\text{Emission Level}\left(\frac{1b \text{ NO}_X}{10^6 \text{ Btu}}\right)}$
Coal	0.24
Oil	0.09
Gas	0.06

Most of the control techniques are capable of controlling the standard boilers with the highest NO, emissions at this level.

The stringent level of control is defined as a technology-forcing level and represents the most rigorous control which might be considered. This represents an NO_{χ} removal of 90 percent. Allowable emissions at this control level are shown below:

Fuel	$\underline{\text{Emission Level}\left(\frac{1b \text{ NO}_{X}}{10^{6} \text{ Btu}}\right)}$
Coal	0.08
011	0.03
Gas	0.02

These systems are operating at their upper limit of practical NO_{\times} removal capability to achieve this level of control and are definitely technology-forcing.

Intermediate level of control is defined as a level between moderate and stringent and probably representing a technological or cost breakpoint. At this point in time, it is difficult to say if those logical breakpoints exist and, if so, where they are. Therefore, the intermediate level was chosen between moderate and stringent levels. The intermediate levels of control considered here represent about 80 percent NO_{\times} removal. Allowable emissions at this level are shown below:

Fuel	Emission Level	$\frac{1b \text{ NO}_{X}}{10^6 \text{ Btu}}$
Coal	0.16	
0il	0.06	
Gas	0.04	

The best FGT systems should be able to achieve steady-state control at this level. This control level provides an alternative choice between the leaststringent and technology-forcing options.

The allowable emission rates for each of the control levels are summarized in Table 3.1.2-1.

TABLE 3.1.2-1.	CONTROLLED	EMISSION	LEVELS	IN	THIS	STUDY	(1b/10°	Btu)	
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	Moderate	Intermediate	Stringent
Coal	0.24	0.16	0.08
0i1	0.09	0.06	0.03
Gas	0.06	0.04	0.02

3.2 BEST CONTROL SYSTEMS FOR COAL-FIRED BOILERS

A three phase selection process was used to determine the best NO_X control systems. The first phase involves comparing the maximum removal level obtainable by each process with the level of control desired---moderate, intermediate or stringent. Those process types which cannot achieve this level are eliminated from further consideration. The remaining process types are then evaluated using the criteria established in Section 3.1.1. The result is a set of process types that are most desirable for a particular consideration of special characteristics of the process types in the set in order to determine the best system candidates. For example, all SCR processes may rate high for application to gas-fired boilers. However, the SCR fixed packed bed process may be more applicable than the moving bed or parallel flow SCR processes since ability to tolerate particulates is not required for gas-fired boiler applications.

For use in the application of the selection factors, tables are compiled which list the process features pertinent to each selection factor. The data in these tables was derived from information presented in Section 2. For coal-fired boilers, this information is presented in Tables 3.2-1 and 3.2-2.

3.2.1 Moderate Reduction Controls

The first phase evaluation eliminated the adsorption process from consideration since it cannot achieve 70% NO_x reduction at high NO_x concentrations (400-600 ppm). Application of the selection factors resulted in numerical ratings for the remaining processes as shown in Table 3.2.1-1. As can be seen, the four SCR processes were superior. The fixed packed bed technique was eliminat, i since it would rapidly plug due to the high particulate levels encountered with coal-fired applications. Therefore, the candidate systems for moderate control of coal-fired boilers are SCR parallel flow and SCR moving bed for NO_x-only removal and SCR parallel flow for simultaneous NO_x/SO_x removal.

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	Performance	Operational and maintenance impacts	Preliminary environmental impacts	Preliminary economic impacts	Preliminary energy and material impacts
SCR Fixed Packed Bed	Capable of attaining >90% NO _x control level; cannot be used with high particulate levels; good load following capability.	Few moving parts; gas phase chemistry; simple process - good controllability; need high removal of particulates - ESP; large pressure drop.	Potential for some NH3 and NH4 HSO4 emissions.	20 MW estimates: Capital: \$130/kW ¹ Operating: 2.1 mills/kWh ¹ , ² Cost is higher than other SCR's due to ESP.	Electrical usage: 1.2% of total output; large NH3 demand (1:1 NH3:NO _x mole ratio); may require auxiliary heater.
SCR Moving Bed	Capable of attaining >90% NO _X control level; can be used with some particulates (up to 2.0 g/Nm ³); adequate load following capability.	Noving parts, solids handling - increased maintenance; gas phase chemistry; fairly simple - controllable; need particulate removal; low pressure drop.	Potential for some NH3 and NH4HSO4 emissions.	20 MW estimates: Capital: \$92/kW ³ Operating: 2.0 mills/kWh	Electrical usage: unknown - should be <1%; large NH ₃ demand (1:1 NH ₃ :NO, mole ratio); may require auxiliary heater; greater catalyst attrition due to moving bed.
SCR Parallel Flow	Capable of attaining >90% NO _X control level; can be used with full particulate loading (up to 20 g/Nm ³); good load following capability.	Few moving parts; gas phase chemistry; simple process - good controllabiltiy; moderate pressure drop; no particulate removal needed.	Potential for some NH3 and NH4HSO4 emissions.	20 MW estimates: Capital: \$44/kW" Operating: 1.5 mills/kWh	Electrical usage: 0.2% of total output; large NH ₃ demand (1:1 NH ₃ :NO _X mole ratio); may require auxiliary heater.
Absorption- Oxidation	No removal data are available - should be able to achieve moderate control level; can be used with full particulate loading; fair load following capability.	Complex process with sensitive control needs; sensitive to flue gas sulfur content – separate SO _x scrubber before NO _x absorber; prescrubber needed to remove particulates and C1 ; very large pressure drops.	NO3 ⁻ salts in wastewater.	20 MW estimates: none available, but since process contains extra scrubber train, Capltal: \$500/kW Operating: 8 mills/kWh	Electrical usage: unknown, estimate ~3%; uses large amounts of gas-phase oxidant and by-product treatment materials.

TABLE 3.2-1. COMPARISON INFORMATION OF NO -ONLY SYSTEMS FOR COAL-FIRED BOILERS

	Boiler operation and/or safety	Reliability	Status of development	Adaptability to existing sources	Compatibility with other control systems
SCR Fixed Packed Bed	No safety hazards.	Catalyst easily plugged; possible NH4HSO4 scaling; simple - few process steps; little corrosion resistant material.	Has only been tested on bench-scale (8 oil- and numerous gas-fired commercial operations).	Some difficulty; few pieces of process equipment; little land needed.	Excessive reheat required if after FGD; needs ESP.
SCR Moving Bed	No safety hazards.	Little catalyst plugging; possible NH4HSO4 scaling; fairly simple - few process steps; little corrosion resistant material.	Has only been tested on bench-scale (5 oil- and 3 coke oven gas-fired commercial operations).	Some difficulty; few pieces of process equipment; little land needed.	Excessive reheat required if after FGD; needs particulate removal.
SCR Parallel Flow	No safety hazards.	Little catalyst plugging (must be packed well); possible NH4HSO4 scaling; simple - few process steps; little corrosion resistant material.	Has only been tested on bench-scale; pilot plants due to start up in 1979 (some oil-fired operations); commercial operation by 1981.	Some difficulty; catalyst can be placed in duct between economizer and preheater without a separate reactor; few pleces of equipment; little land needed.	Completely compatible with FGD.
Absorption- Oxidation	Oxidant handling could be hazardous.	Numerous process steps and corrosion resistant material.	Has not been tested on coal-fired flue gas (a few pilot plants treating oil-fired and furnace gases).	Much land needed for numerous pieces of process equipment and wastewater treatment.	Existing FGD would be helpful as process cannot tolerate sulfur.

TABLE 3.2-1. (Continued)

	Performance	Operational and maintenance impacts	Preliminary environmental impacts	Preliminary economic impacts	Preliminary energy and material impacts
SRC Parallel Flow	Capable of attaining 90% control of both NO _X and SO _X ; can be applied to gases with high particu- late loadings; process can follow boiler load easily through use of gas bypass arrangement.	Process has several sections but all except NO _X /SO _X reactor are based on well established technology; average maintenance require- ments.	Potential NH₃ emissions.	20 MW estimates: Capital: \$475/kW Operating: 5 mills/kWh	Electrical usage: 1,5% of total output; also consumes NH3, naphtha, and steam.
Adsorption	Capable of attaining 60% NO _X control level; cannot be used with high particulate levels; poor load following capability; primarily SO _X removal.	Many moving parts, hot solids handling; complex process; need ESP for particulate removal; major mainte- nance requirements; high pressure drop.	Ash disposal.	20 MW estimates: Capital: \$215 kW⁵ Operating: 2.3 mills/kWh	Electrical usage: unknown - should be ≃2%; activated char usage high due to attrition.
Electron Beam Radiation	Capable of attaining 80% NO_x control level; cannot be used with particulates; fair load following capability; also removes SO_x .	Simple process but complex control; sensitive to flue gas composition (at least 1% O ₂ and H ₂ O>NO _X).	H ₂ SO ₄ mist and a powder containing ammonium nitrates and sulfates are generated.	20 MW estimates: Capital: \$202 kW ⁶ Operating: unknown Electricity is only major.	Electrical usage: 3.3% of total output (excluding ESP); treatment of by-produc is unknown.
Absorption- Reduction	Capable of attaining 85% NO _X control level; can be used with full particulate loading; good load following capability; removes SO _X .	Complex process with very sensitive control needs; sensitive to flue gas composition (low 0_2 and $SO_X:NO_X$ ratio >2.5); need prescrubber to remove particulates and Cl ⁻ .	Possibility of plume from absorbent (sulfate or NH3).	20 MW estimates: Capital: \$413/kW ⁷ Operating: 7.4 mills/kWh Gypsum by-product (landfill).	Electrical usage: 1.8% of total output; large amounts of chelating compound, absorbent, and regeneration chemical; are used.
Oxidation- Absorption- Reduction	Capable of attaining 90% NO _X control level; can be used with full particulate loading; poor load following capability - oxidant generation lagtime; removes SO _X .	Complex process with very sensitive control needs; prescrubber needed to remove particulates and Cl ⁻ ; large pressure drop.	NO3 ⁻ or N-S salts or NH3 ⁻ based compounds in wastewater.	Economic estimates: unknown for coal-fired plant; gypsum by-product (landfill).	Electrical usage: 9.0% of total output; uses large amounts of gas-phase oxidant and by-product treatment materials.
Oxidation~ Absorption	Capable of attaining 90% NO _X control level; can be used with full particulate loading; poor load following capability; removes SO _X .	Complex process with very sensitive control needs; prescrubber needed; large pressure drop.	NO3 ⁻ salts in wastewater.	Economic estimates: unknown; gypsum by-product and liquid fertilizer, or HNO3.	Electrical usage: unknown (will be ≃10% of total output): use: large amounts of gas- phase oxidant and by-product treatment materials.

TABLE 3.2-2. COMPARISON INFORMATION OF SIMULTANEOUS NO_/SO_ SYSTEMS FOR COAL-FIRED BOILERS

	Boiler operation and/or safety	Reliability	Status of development	Adaptability to existing sources	Compatibility with other control systems
SCR Parallel Flow	H, usage may present safety hazard.	Process steps well established; should be reliable.	SO ₂ system has been tested on coal-fired flue gas; NO _x /SO _x operation with coal- fired flue gas to begin late 1979; pilot unit tests; SO ₂ work up and H ₂ generation not tested, but are established technology.	Will need land for equipment.	Compatible with particulate systems.
Adsorption	Possible safety hazard due to poor char distribution in beds,	Char plugged by particulates; numerous process steps; some corrosion resistant material in high temperature zones.	One prototype unit treating coal-fired flue gas.	Need land for pieces of process equipment.	Suitable for placement after ESP; not useful with FGD system as NO _X removal is secondary.
Electron Beam Radiation	Radiation safety hazards are unknown as are those of byproduct.	Few process steps; stainless steel reactor.	Has not been tested on coal-fired flue gas (one pilot plant treat- ing gas from sintering machine); uncertain by-product treatment method.	Need land for pieces of process equipment.	Needs ESP; with or without existing FGD but capital cost will be the same.
Absorption- Reduction	No safety hazards.	Many process steps; much glass- and elastomer-lined equipment.	Has not been tested on coal-fired flue gas (several pilot plants treating oil-fired flue gas); NO _X absorption chemistry uncertain.	Much land needed for numerous pieces of process equipment.	Cannot be used in conjunction with FGD.
Dxidation- Absorption- Reduction	Gas-phase oxidant presents serious safety hazard.	Numerous process steps and corrosion resistant material; oxidant generation system subject to periodic failure.	Has not been tested on coal-fired flue gas (6 prototype units treating oil-fired flue gas in operation).	Much land needed for numerous pieces of process equipment, oxidant generation, and wastewater treatment.	Cannot be used in conjunction with FGD.
Dxidation- Absorption	Gas-phase oxidant presents serious safety hazard.	Numerous process steps and corrosion resistant material; oxidant generation system subject to periodic failure.	One pilot plant treating flue gas from coal-fired boiler.	Much land needed for numerous pieces of process equipment, oxidant generation, and wastewater treatment.	Compatible.

TABLE 3.2-2. (Continued)

Control technique	Total point rating	Candidate system	Comments
NO _X -Only			
SCR Fixed Packed Bed	69	no	Adversely affected by particulates
SCR Moving Bed	70	no	Adversely affected
SCR Parallel Flow	83	yes	by particulates
Absorption-Oxidation	43	no	Low rating
Simultaneous NO _X /SO _X			
SCR Parallel Flow	72	yes	
Adsorption	NA	no	
Electron Beam Radiation	41	no	Low rating
Absorption-Reduction	52	no	Low rating
Oxidation-Absorption-Reduction	51	no	Low rating
Oxidation-Absorption	51	no	Low rating

TABLE 3.2.1-1. CANDIDATE SYSTEMS SELECTION: COAL-FIRED BOILERS - MODERATE CONTROL

A detailed listing of how each process was evaluated on each selection factor is contained in Tables A3.1 and A3.2 in the Appendix.

3.2.2 Stringent Reduction Controls

In a similar manner, candidate systems for stringent control were selected. The results appear in Table 3.2.2-1. A detailed listing of the selection factors and point values for each system is contained in Tables A3.3 and A3.4. The candidate systems selected are SCR parallel flow and SCR moving bed for NO_{χ} -only removal and SCR parallel flow for simultaneous NO_{χ}/SO_{χ} removal.

3.2.3 Intermediate Reduction Controls

The selection results for this level are presented in Table 3.2.3-1. Detailed application of the selection factors is presented in Tables A3.5 and A3.6. The candidate systems selected are SCR parallel flow and SCR moving bed for NO_x -only removal and SCR parallel flow for simultaneous NO_x/SO_x removal.

3.3 BEST CONTROL SYSTEMS FOR OIL-FIRED BOILERS

The control systems for oil-fired boilers were evaluated using the same method described in the previous section on coal-fired boilers. Tables 3.3-1 and 3.3-2 present a side-by-side comparison of all potential systems with data categorized with respect to the selection factors. The information in this table is summarized from Section 2. The table is similar in many respects to the equivalent table for coal. This is due to the fact that, since FGT systems are applied after the boiler, they are relatively insensitive to the types of fuel burned. Two notable exceptions are particulate and sulfur emissions which are a function of the fuel type. Process characteristics that change with fuel type are noted in the table.

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Control technique	Total point rating	Candidate system	Comments
NO _× -Only			
SCR Fixed Packed Bed	62	no	Adversely affected by particulates
SCR Moving Bed	60	no	Adversely affected
SCR Parallel Flow	73	yes	by particulates
Absorption-Oxidation	NA	no	
Simultaneous NO _× /SO _×			
SCR Parallel Flow	68	yes	
Adsorption	NA	no	
Electron Beam Radiation	NA	no	
Absorption-Reduction	NA	no	
Oxidation-Absorption-Reduction	48	no	Low rating
Oxidation-Absorption	49	no	Low rating

TABLE 3.2.2-1. CANDIDATE SYSTEMS SELECTION: COAL-FIRED BOILERS - STRINGENT CONTROL

Control technique	Total point rating	Candidate system	Comments
10x-0nly			
SCR Fixed Packed Bed	67	no	Adversely affected by particulates
SCR Moving Bed	69	no	Adversely affected
SCR Parallel Flow	81	yes	by p articul ates
Absorption-Oxidation	43	no	Low rating
imultaneous NO _× /SO _×			
SCR Parallel Flow	70	yes	
Adsorption	NA	no	
Electron Beam Radiation	41	no	Low rating
Absorption-Reduction	50	no	Low ratin g
Oxidation-Absorption-Reduction	49	no	Low r ating
Oxidation-Absorption	46	no	Low rating

TABLE 3.2.3-1. CANDIDATE SYSTEMS SELECTION: COAL-FIRED BOILERS - INTERMEDIATE CONTROL

	Performance	Operational and maintenance impacts	Preliminary environmental impacts	Preliminary economic impacts	Preliminary energy and material impacts
SCR Fixed Packed Bed	Capable of achieving >90% NO _X reduction; cannot be used with high particulate levels; good load following capability.	Few moving parts; gas phase chemistry; sImple process - good controllability; need high removal of particulates - ESP; large pressure drop.	Potential for some NH, and NH,HSO, emissions.	20 MW estimates: Capital: \$70/kW ^{0,9} Operating: 1.9 mills/kWh ^{10,11}	Electrical usage: unknown for oil-fired plant; moderate NH3 demand {1:1 NH3:NOx mole ratio); may require auxiliary heater.
SCR Moving Bed	Capable of achieving >90% NO _X reduction; can tolerate particulate level of most oils (<1 g/Mm ³); adequate load following capability.	Some moving parts, solids handling – increased maintenance; gas phase chemistry; simple – controllable; low pressure drop.	Potential for some NH3 and NH4HSO4 emissions.	20 MW estimates: Capital: \$70/kW ^{11,12} Operating: 1.8 mills/kWh	Electrical usage: unknown for oil-fired plant; moderate NH3 demand (l:l NH3:NO _x mole ratio); may require auxiliary heater; greater catalyst attrition due to moving hed.
SCR Parallel Flow	Capable of achieving >90% NO _X reduction; can tolerate full particulate loading (up to 20 g/Nm ³); good load following capability.	Few moving parts; gas phase chemistry; simple process - good controllability; no particulate removal needed; moderate pressure drop.	Potential for some NH3 and NH4HSO4 emissions.	20 MW estimates: Capital: \$39/kW ¹³ Operating: unknown.	Electrical usage: unknown for oll-fired plant; moderate NH3 demand (1:1 NH3:NO _X mole ratio); may require auxiliary heater.
Absorption- Oxidation	No removal data are available; can tolerate particulates; fair load following capability; removes SO _x .	Complex process with sensitive control needs; sensitive to flue gas sulfur content - separate SO _X scrubber before NO _X absorber; prescrubber needed; very Jarge AP.	NO3 ⁻ salts in wastewater.	Economic estimates: unknown.	Electrical usage: unknown; uses large amounts of liquid phase oxidant and regeneration materials.

TABLE 3.3-1. COMPARISON INFORMATION OF NO_x-ONLY SYSTEMS FOR OIL-FIRED BOILERS

	Boiler operation and/or safety	Reliability	Status of development	Adaptability to existing sources	Compatibility with other control systems
SCR Fixed Packed Bed	No safety hazards.	Catalyst easily plugged; possible NH ₄ HSO4 scaling; simple - few process steps; little corrosion resistant material.	8 commercial operations in Japan.	Some difficulty; few pieces of process equipment; little land required.	Excessive reheat required if after FGD; needs ESP.
SCR Moving Bed	No safety hazards.	Some catalyst plugging; possible NH4HSO4 scaling; fairly simple - few process steps; little corrosion resistant material.	6 commercial οil-fired operations in Japan.	Some difficulty; few pieces of process equipment; little land required.	Ехсеввіve reheat required if after FGD.
SCR Parallel Flow	No safety hazards.	Little catalyst plugging; possible NHuHSO4 scaling; simple - few process steps; little corrosion resistant material.	Numerous commercial oil-fired operations in Japan.	Some difficulty; catalyst can be placed in duct between economizer and preheater without a separate reactor; few pieces of equipment; little land needed.	Completely compatible with FGD.
Absorption- Oxidation	Oxidant handling can be dangerous.	Numerous process steps and corrosion resistant material.	2 pilot plants treating oil-fired flue gas.	Much land needed for numerous pieces of process equipment and wastewater treatment.	Existing FGD would be helpful as process cannot tolerate sulfur.

TABLE 3.3-1. (Continued)

	Performance	Operational and maintenance impacts	Preliminary environmental impacts	Preliminary economic impacts	Preliminary energy and material impacts
SCR Parallel Flow	Capable of attaining 90% control of both NO _x and SO ₂ ; can be used with full particulate loading; good load following capability.	Most of the process steps are based on well established technology; average maintenance require- ments.	Potential NH3 emissions.	Economic estimates: unknown for oil-fired plant; assumed to be similar to those for coal. 20 MW estimates for coal: Capital: \$475/kW Operating: 5 mills/kWh	Electrical usage: unknown for oil-firec plant; assumed to be similar to those for coal, i.e. 1.5% of boiler output as electricity; also uses steam, naphtha and NH ₃ .
Adsorption	Capable of attaining 60% NO _X reduction; cannot be used with high particulate levels; poor load following capability; primarily SO _X removal.	Many moving parts, hot solids handling – high maintenance; complex process; may need particulate removal on residual oils; large pressure drop.	Ash disposal.	Ecomonic estimates: unknown for oil-fired plant; elemental S by- product.	Electrical usage: unknown for oil-fired plant; large activated char demand due to attrition.
Electron Beam Radiation	Capable of attaining 80% NO _X removal; cannot be used with particulates; fair load following capability; also removes SO _X .	Simple process with complex control; sensitive to flue gas composition (at least $1Z O_2$ and $H_2 O>NO_X$); may need particulate removal on residual oils.	H ₂ SO, mist and a powder containing ammonium-nitrates and sulfates are generated.	Economic estimates: unknown for oil-fired plant; electricity is primary operating expense.	Electrical usage: unknown for oil-fired plant; treatment of by-product is unknown
Absorption- Reduction	Capable of attaining 85% removal; can tolerate particulates; good load following capability; cannot be used on distillate oil; also removes SO _x .	Complex process with very sensitive control needs; sensitive to flue gas composition (low O_2 and $SO_X:NO_X$ ratio >2.5); need prescrubber; large ΔP .	Possibility of plume (sulfite or NH3) from absorbent.	20 MW estimates: Capital: \$187/kW ^{14,15} Operating: 5.4 mills/kWh Gypsum by-product (landfill).	Electrical usage: 1.8% of total output; extremely large amounts of chelating compound; absorbent and regeneration chemicals are used.
Oxidation- Absorption- Reduction	Capable of attaining 90% NO _x reduction; can tolerate particulates; poor load following capability - oxidant gereration lagtime; cannot be used on distil- late oils; removes SO _x .	Complex process with very sensitive control needs; prescrubber needed; large pressure drop.	NO3 ⁻ or N-S salts or NH-base compounds in wastewater.	20 MW estimates: Capital: \$231/kW ^{16,17} Operating: 6.4 mills/kWh Gypsum by-product (landfill).	Electrical usage: 5-10% of total output; large amount of gas phase oxidant and by-product treat- ment materials.
Oxidation- Absorption	Capable of attaining 90% NO_X reduction; can tolerate particulates; poor load following capability; removes SO_X .	Complex process with very sensitive control needs; prescrubber needed; large pressure drop.	NO3 ⁻ salts in wostewater.	Economic estimates: unknown; gypsum by-product and liquid fertilizer or HNO3.	Electrical usage: unknown (will be 5- 10% of total output); uses large amounts of gas phase oxidant and by-product treat- ment materials.

	Boiler operation and/or safety	Reliability	Status of development	Adaptability to existing sources	Compatibility with other control systems
SCR Parallel Flow	H ₂ is potential safety hazard.	Process steps well established. Should be reliable.	Both SO, and NO, removal systems have been tested on oil. SO2 workup and H ₂ generation steps not tested but are established technology.	Will need land for equipment	Compatible with par- ticulate control systems
Adsorption	Possible safety hazard due to poor char distri- bution in beds.	Particulate plugging; numerous process steps; some corrosion resistant material in high temp- erature areas.	No tests on oil-fired gas.	Need land for pieces of process equipment	Not useful with FGD systems NO _X removal is secondary.
Electron Beam Radiation	Radiation safety hazards are unknown as are those of by-product.	Few process steps; stainless steel reactor.	One oil-fired pilot plant; by-product treating method is uncertain.	Need land for pieces of process equipment.	Operate with or without FGD but capital cost is same.
Absorption- Reduction	No safety hazards.	Many process steps; much glass - and elastomer-lined equipment.	3 pilot plants treating oil-fired flue gas; NO _x absorption mechanism uncertain.	Much land needed for numerous pleces of process equipment.	Cannot be used in conjunction with FGD.
Oxidation- Absorption- Reduction	Gas phase oxidant presents serious safety hazard.	Numerous process steps and corrosion resistant material; oxidant gen- eration system subject to periodic failure.	6 prototype units treat- ing oil-fired flue gas in operation.	Much land needed for numerous pleces of process equipment, oxidant generation and wastewater treatment.	Cannot be used in conjunction with FGD.
Oxidation- Absorption	Gas phase oxidant presents serious safety hazards.	Numerous process steps and corrosion resistant material; oxidant gener- ation system subject to periodic failure.	One bench-scale test on oil-fired flue gas.	Much land needed for numerous pieces of process equipment, oxidant gener- ation, and wastewater treatment.	Compatible

TABLE 3.3-2. (Continued)

VICTOR AND AND ADDRESS
3.3.1 Moderate Reduction Controls

One system, adsorption, was eliminated because it was not capable of achieving sufficient emission reduction. The remaining systems were rated using the selection factors and the results are presented in Table 3.3.1-1. A detailed breakdown of this evaluation is contained in Tables A3.7 and A3.8. SCR fixed packed bed was selected as the NO_{χ} -only candidate system for distillate-oil-fired boilers since these have low particulate emissions. For resid-fired boilers, which have higher particulate emissions, the NO_{χ} -only candidate systems are SCR parallel flow and SCR moving bed and the simultaneous NO_{χ}/SO_{χ} candidate system is SCR parallel flow.

3.3.2 Stringent Reduction Controls

The results of system evaluations for stringent control levels are shown in Table 3.3.2-1. The detailed evaluation breakdown is contained in Tables A3-9 and A3-10. The candidate systems are the same as for moderate control.

3.3.3 Intermediate Reduction Controls

The results of system evaluations for intermediate control levels are shown in Table 3.3.3-1. A detailed breakdown of the selection factor ratings is presented in Tables A3.11 and A3.12. The candidate systems are the same as for the other two levels: NO_{χ} -only, SCR fixed packed bed for distillate oil plus SCR parallel flow and SCR moving bed for resid oil; simultaneous NO_{χ}/SO_{χ} , SCR parallel flow.

3.4 BEST CONTROL SYSTEMS FOR GAS-FIRED BOILERS

Table 3.4-1 compares all of the FGT systems as applied to gas-fired boilers for each of the selection factors. This table was used to arrive at the point values shown on the candidate selection tables.

Control technique	Total point rating	Candidate system	Comments
NO _× -Only			
SCR Fixed Packed Bed	81	yes	Distillate oil-fired boilers
SCR Moving Bed	88	yes	Residual oil-fired boilers
SCR Parallel Flow	90	yes	Residual oil-fired boilers
Absorption-Oxidation	53	no	Low rating
Simultaneous NO _× /SO _×			
SCR Parallel Flow	75	yes	Residual oil-fired boilers
Adsorption	NA	no	
Electron Beam Radiation	47	no	Low rating
Absorption-Reduction	58	no	Low rating
Oxidation-Absorption-Reduction	59	no	Low rating
Oxidation-Absorption	52	no	Low rating

TABLE 3.3.1-1. CANDIDATE SYSTEMS SELECTION: OIL-F	FIRED BOILERS - MODERATE CONTROL
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Control technique	Total point rating	Candidate system	Comments
NO _x -Only			
SCR Fixed Packed Bed	74	yes	Distillate oil-fired boilers
SCR Moving Bed	81	yes	Residual oil-fired boilers
SCR Parallel Flow	83	yes	Residual oil-fired boilers
Absorption-Oxidation	NA	no	
Simultaneous NO_X/SO_X			
SCR Parallel Flow	71	yes	
Adsorption	NA	no	
Electron Beam Radiation	NA	no	
Absorption-Reduction	NA	no	
Oxidation-Absorption-Reduction	54	no	Low rating
Oxidation-Absorption	50	no	Low rating

TABLE 3.3.2-1. CANDIDATE SYSTEMS SELECTION: OIL-FIRED BOILERS - STRINGENT CONTROLS

Control technique	Total point rating	Candidate system	Comments
NO×-Only			
SCR Fixed Packed Bed	79	yes	Distillate oil-fired boilers
SCR Moving Bed	86	yes	Residual oil-fired boilers
SCR Parallel Flow	88	yes	Residual oil-fired boilers
Absorption-Oxidation	54	no	Low rating
Simultaneous NO _X /SO _X			
SCR Parallel Flow	73	yes	Residual oil-fired boilers
Adsorption	NA	no	
Electron Beam Radiation	47	no	Low rating
Absorption-Reduction	58	ņo	Low rating
Oxidation-Absorption-Reduction	55	no	Low rating
Oxidation-Absorption	48	no	Low rating

TABLE 3.3.3-1. CANDIDATE SYSTEMS SELECTION: OIL-FIRED BOILERS - INTERMEDIATE CONTROL

	Performance	Operational and maintenance impacts	Preliminary environmental impacts	Preliminary economic impacts	Preliminary energy and material impacts
SCR Fixed Packed Bed	Capable of achieving >90% NO _X removal; good load following capability.	Few moving parts; gas phase chemistry; simple process - good controllability; large pressure drop.	Potential for some NH3 and NH4HSO4 emissions.	20 MW estimates for clean gas: Capital: \$27/kW ['] Operating: 1.2 mills/kWh	Electrical usage: unknown for gas-fired flue gas; light NH ₃ demand (l:l NH ₃ :NO, mole ratio).
SCR Moving Bed	Capable of achieving >90% NO _x removal; adequate load following capability.	Some moving parts, solids handling – increased maintenance; gas phase chemistry; simple – controllable; low pressure drop.	Potential for some NH3 and NH4HSO4 emissions.	Economic estimates: unknown for gas-fired plant.	Electrical usage: unknown for gas-fired flue gas; light NH ₃ demand (1:1 NH ₃ :NO _X mole ratio).
SCR Parallel Flow	Capable of achieving >90% NO _X removal; good load following capability.	Few moving parts; gas phase chemistry; simple process - good controllability; moderate pressure drop.	Potential for some NH3 and NH4HSO4 emissions.	Economic estimates: unknown for gas-fired plant,	Electrical usage: unknown for gas-fired flue gas; light NH3 demand (l:l NH3:NO _X mole ratio).
Absorption- Oxidation	No removal data are available; fair load following capability.	Complex process with sensitive control needs; very large pressure drop.	NO3 salts in waste- waters.	Economic estimates: unknown.	Electrical usage: unknown; uses large amounts of liquid phase oxidant and regeneration materials.

TABLE 3.4-1. COMPARISON INFORMATION OF NO_×-ONLY SYSTEMS FOR GAS-FIRED BOILERS

TABLE	3.4-1.	(Continued)

	Boiler operation and/or safety	Reliability	Status of development	Adaptability to existing sources	Compatibility with other control systems
SCR Fixed Packed Bed	No safety hazards.	Possible NH,HSO,scaling; simple - few process steps; little corrosion resistant material.	Numerous commercial operations in Japan.	Some retrofit difficulty; few pieces of process equipment - little land required.	Compatible
SCR Moving Bed	No safety hazards.	Possible NH,HSO, scaling; simple - few process steps; little corrosion resistant material.	Three commercial coke oven gas operations in Japan.	Some retrofit difficulty; few pieces of process equipment - little land required.	Compatible
SCR Parallel Flow	No safety hazards.	Possible NH ₄ HSO4 scaling; simple - few process steps; little corrosion resistant material.	No commercial operations (many oil-fired; not necessary for gas-fired - no particulates)	Some retrofit difficulty; few pieces of process equipment – little land required; if space exists, catalyst can fit in duct.	Compatible
Absorption - Oxidation	Oxidant handling could be dangerous.	Numerous process steps and corrosion resistant material.	Pilot plants treating off gases from HNO; and steel plants.	Much land needed for numerous pieces of process equipment and wastewater treatment.	Compatible

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3.4.1 Moderate Reduction Controls

The first cut in FGT systems applied to gas-fired boilers eliminated one process due to insufficient emission reduction and five processes due to their removal of SO_x which is not present in gas-fired flue gas. This can be seen in Table 3.4.1-1 which presents the results of the candidate selection. SCR fixed packed bed was chosen as the candidate system. SCR parallel flow and SCR moving bed were eliminated since their specialized ash handling characteristics are not required for this application. A detailed selection factor rating breakdown is contained in Table A3.13.

Control technique	Total point rating	Candidate system	Comments
NO _x -Only			
SCR Fixed Packed Bed	93	Yes	
SCR Moving Bed	91	No ∫	These specialized catalyst
SCR Parallel Flow	93	No 🔪	These specialized catalyst arrangements are not neces- sary for gas-fired sources.
Absorption - Oxidation	n 58	No	Low rating

TABLE 3.4.1-1. CANDIDATE SYSTEMS SELECTION: GAS-FIRED BOILERS - MODERATE CONTROL

3.4.2 Stringent Reduction Controls

The results of system evaluations for stringent control levels are shown in Table 3.4.2-1. The detailed evaluation breakdown is contained in Table A3.14. The candidate system is SCR fixed packed bed.

0110		DIRINOLI	
Control technique	Total point rating	Candidate system	Comments
NOOnly			
SCR Fixed Packed Bed	83	Yeş	
SCR Moving Bed	81	No	These specialized catalyst arrangements are not neces-
SCR Parallel Flow	83	No	sary for gas-fired sources.
Absorption - Oxidation	n NA	No	

TABLE 3.4.2-1. CANDIDATE SYSTEMS SELECTION: GAS-FIRED BOILERS - STRINGENT CONTROL

3.4.3 Intermediate Reduction Controls

The results of system evaluations for intermediate control levels are shown in Table 3.4.3-1. The detailed evaluation is presented in Table A3.15. The candidate system is SCR fixed packed bed.

Control technique	Total point rating	Candidate system	Comments
NO _x -Only		-	
SCR Fixed Packed Bed	87	Yes	
SCR Moving Bed	85	No	These specialized catalyst arrangements are not neces-
SCR Parallel Flow	87	No	sary for gas-fired sources.
Absorption - Oxidatio	n 58	No	Low rating

TABLE 3.4.3-1. CANDIDATE SYSTEMS SELECTION: GAS-FIRED BOILERS - INTERMEDIATE CONTROL

3.5 SUMMARY

A candidate or set of candidates has now been chosen for each of the standard boilers under consideration. These are shown in Table 3.5-1. These systems will be analyzed in detail in the subsequent sections in order to determine the best overall system for NO_{\times} reduction by FGT on industrial boilers. The major performance characteristics for the candidate processes are presented in Table 3.5-2.

TABLE 3.5-1. SUMMARY OF CANDIDATE SYSTEMS: ALL LEVELS OF CONTROL

Fuel	Candidate Systems
Coal	SCR Parallel Flow
Residual Oil	SCR Parallel Flow SCR Moving Bed
Distillate Oil	SCR Fixed Packed Bed
Natural Gas	SCR Fixed Packed Bed

	Collection Efficiency	Environmental Impacts	Energy Impacts	Reliability	Commercial Availability
SCR-fixed packed bed	>90% NO _X reduction	Possible NH ₃ and NH4HSO4 emissions	Coal-need ESP elec.= 1.2% of total power input	Simple, few process steps; catalyst easily plugged	Coal-bench scale; 8 oil- and numerous gas-fired commercial operations
SCR-parallel flow	>90% NO _X reduction	Possible NH ₃ and NH4HSO4 emissions	Coal-no ESP elec.=0.2% of total power output	Simple, few process steps; little plugging	Coal-bench scale; numerous oil-fired commercial operations
SCR-moving bed	>90% NO _X reduction	Possible NH3 and NH4HSO4 emissions	Coal-some particulate removal needed elec.<1% of total power output	Simple, few process steps; moving parts and solids handling; some plugging	Coal-bench scale; 5 oil-fired and 3 coke oven gas commercial operations

TABLE 3.5-2.	MAJOR	PERFORMANCE	CHARACTERISTICS	OF	CANDIDATE	SYSTEMS

REFERENCES

- Faucett, H.L., et αl. Technical Assessment of NO_× Removal Processes for Utility Applications. EPA-600/7-77/127. November 1977. pp. 210, 268, 301.
- Rosenberg, H.S., et αl. State-of-the-Art Review of Stack Gas Treatment Techniques for NO_x Control. EPRI-Batelle. April 1976. p. A-47.
- 3. Faucett, H.L., op. cit., pp. 233, 319.
- 4. Ibid., p. 218.
- 5. Ibid., p. 204.
- 6. Ibid., p. 166.
- 7. Ibid., p. 33
- 8. Ibid., p. 347.
- 9. Rosenberg, H.S., op. cit., p. A-14.
- 10. Faucett, H.L., op. cit., pp. 280, 327.
- 11. Rosenberg, H.S., op. cit., p. A-23.
- 12. Faucett, H.L., op. cit., p. 242.
- 13. Faucett, H.L., op. cit., p. 223.
- 14. Ibid., p. 41
- 15. Rosenberg, H.S., op. cit., p. A-139.
- 16. Ibid., pp. A-106, A-116.
- 17. Faucett, H.L., op. cit., pp. 48, 117, 127.

SECTION 4

COST ANALYSIS OF CANDIDATES FOR BEST EMISSION CONTROL SYSTEMS

4.1 NO_X-ONLY SYSTEMS

4.1.1 Introduction

This section considers the costs involved with applying the "best" NO_X FGT systems selected in Section 3 to the standard boilers. The costs presented are based on several factors. First, typical process layouts were determined to establish the equipment requirements. Material balances are established for each case and the equipment sized. Process layouts and material balances for all nineteen cases considered in detail are presented in Appendices 3, 4 and 5 for coal, oil, and gas sources, respectively. Purchased equipment lists for each process considered are shown in Table 4.1.1-1. The equipment is selected and sized by using standard engineering techniques. Example calculations for equipment size and energy usage are presented in the Appendix 8. Energy usage for all systems consists only of electricity and steam. Other costs were based on cost factors supplied by references 1 and 2 as well as other sources.

All of the equipment listed in Table 4.1.1-1 will require some maintenance. The items requiring the most maintenance are the pump, fan motor drive, vaporizer, screen, catalyst elevator, baghouse/blower, and all associated process control elements. The catalyst has a lifetime of about one year and its regeneration is presently uncertain. Therefore, in this analysis, it is replaced annually and represents both a capital and operating cost.

4-1

Parallel Flow SCR	Moving Bed SCR	Fixed Packed Bed SCR
Reactor	Reactor	Reactor
Catalyst	Catalyst	Catalyst
Fan Motor Drive	Catalyst Screen	Fan Motor Drive
NH ₃ Storage Tank	Catalyst Elevator	NH ₃ Storage Tank
NH ₃ Transfer Pump	Baghouse/Blower	NH ₃ Transfer Pump
NH ₃ Vaporizer	Fan Motor Drive	NH ₃ Vaporizer
	NH_3 Storage Tank	
	NH ₃ Transfer Pump	
	NH ₃ Vaporizer	

TABLE 4.1.1-1. PURCHASED EQUIPMENT FOR NO, FGT SYSTEMS

The cost bases can be separated into several areas. Costs of materials associated with all of the processes evaluated are presented in Table 4.1.1-2. Sources of the costs are also shown. Several costs were determined by multiplying a factor times another cost. This is common with this type of economic analysis and the cost factors used are shown in Table 4.1.1-3. Direct costs were determined on a full year basis and then multiplied by the boiler load factor to determine the annual direct costs. Load factors for the standard boilers are shown in Table 4.1.1-4. The capital recovery factor was calculated from the formula:

$$\frac{i(1+i)^{n}}{(1+i)^{n}-1}$$

For i = 0.10 (interest) and n = 15 (years) the capital recovery factor is 0.13147.

The costs of each equipment item was determined using a variety of cost references shown in Table 4.1.1-5. Installation costs were provided in the references. As with the annual costs, some of the capital costs were determined by multiplying a factor times another cost. A list of the factors used in the capital cost estimates is contained in Table 4.1.6.

4-2

Item	Cost Used	Reference
Direct Labor, \$/manhour	12.02	1
Maintenance Labor, \$/manhour	14.63	1
Electricity, mills/kWh	25.8	1
Ammonia, \$/ton delivered	130	1
Steam, \$/1000 1b	3.50	2
Catalyst, \$/ft ³ Parallel Flow Moving Bed Fixed Packed Bed	212 282 282	3 4 4

TABLE 4.1.1-2. ANNUAL COST PARAMETERS USED IN COST ANALYSIS

TABLE 4.1.1-3. ANNUAL COST FACTORS

Item	Amount	Reference
Maintenance Materials	3% of turnkey costs	5
Payroll Overhead	30% of direct labor	1
Plant Overhead	26% of labor, parts & maintenance	1
General and Administrative Expenses (G&A), Taxes & Insurance	4% of total turnkey costs	1
Capital Recovery Factor (10% interest rate)	13.147% of total turnkey costs	1

TABLE 4.1.1-4. LOAD FACTORS FOR THE STANDARD BOILERS

Fuel	Load Factor
Coal	0.60
Residual Oil	0.55
Distillate Oil and Natural Gas	0.45

Equipment Item	Reference
Reactor	6
Catalyst	3,4
Fan Motor Drive (Incremental)	7,8
NH ₃ Storage Tank	9
NH ₃ Transfer Pump	10
NH ₃ Vaporizer	11
Vibrating Screen	12
Catalyst Elevator	13,8
Baghouse	14

TABLE 4.1.1-5. SOURCES OF COSTS FOR SPECIFIC EQUIPMENT ITEMS

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TABLE 4.1.	1-6.	CAPTTAL.	COST	FACTORS	

TABLE 4.1.1-0.	CAFITAL COST FACTORS	
Item	Amount	Reference
Engineering	10% of installed cost of largest NO _× removal system considered (pulverized coal boiler; stringe level of control)	
Construction and Field Expense	10% of installed cost	1
Contractor Fee	10% of installed cost	1
Start-up	2% of installed cost	1
Performance Tests	\$2000	1
Contingency	Coal: 20% of total direct and indirect costs Oil and Gas: 15% of total direc and indirect costs	1 t
Working Capital	25% of total direct operating costs	1

Capital costs were escalated to June 1978 costs using standard cost indices. For example, costs in Guthrie⁶ are based on June 1970 costs. Cost indices for this year and June 1978 for various types of equipment are shown in Table 4.1.1-7. To obtain the mid 1978 costs the costs given in Guthrie are multiplied by the escalation index.

Item	June 1970 Index	June 1978 Index	Escalation Index (E.I.) (1978/1970)
Fabricated Equipment	124.0	237.4	1.91
Process Machinery	122.7	226.6	1.85
Pipes, Valves & Fittings	133.0	268.4	2.02
Process Instruments	132.0	214.8	1.63
Pumps & Compressors	124.1	258.2	2.08
Electrical Equipment	98.9	167.9	1.70
Miscellaneous	118.5	250.1	2.11
Construction Labor	134.8	184.3	1.37

TABLE 4.1.1-7. CHEMICAL ENGINEERING COST INDICES¹⁵

The labor requirements were determined from the basis for an economic analysis performed by a process vendor which indicated a requirement of one person/shift/day per reactor.¹⁶ Equipment life was estimated at 15 years based on the average lifetime of chemical processing equipment.¹⁷ Capital costs were annualized over a 15 year period to give constant annual costs for the life of the boiler.

The capital and operating costs were collected and presented in a consistent set of table, and an annualized cost was calculated. These comprehensive tables are contained in separate appendices and discussed in the subsequent subsections. Costs for modified or reconstructed facilities will most likely be slightly higher than those for new facilities. This is due to the fact the major cost items--*i.e.* the fan motor drive, reactor plus catalyst, and NH₃ storage tanks--are the same for both applications. There may be some increased costs where additional ductwork, boiler modification or flue gas heating is necessary and these factors are highly site specific. The cost of a retrofit will have to be determined for each application since it is dependent on site specific factors.

The cases considered include only one type of coal, low sulfur western. Other coal types are not considered since process costs do not vary significantly with coal type. Two of the most significant cost items for FGT systems are the reactor plus catalyst and the fan motor drive. These equipment items are sized and costed based on flue gas flow rate which does not vary significantly with coal type. Since including all three coal types would not provide additional information, only low sulfur western was considered. Since all catalysts considered in this study are assumed to be resistant to SO_X poisoning, low sulfur coal was chosen since it had the highest NO_X emissions and flue gas flow rate. Therefore, use of this coal provides a "worst case" analysis.

SIP control levels are not considered since in many cases no control is required. On cases that require some control the level can be easily attained through use of combustion modifications. In no instance is FGT required to meet the average SIP levels, except possibly in California. Los Angeles hourly maximum NO_× concentration occassionally exceeds the state standard by a factor of $3.^{18}$ Because of the topographic characteristics of the area and its high concentration of mobile sources (automobiles which also have mileage requirements to meet), strict legislation for stationary sources has been proposed that would require NO_× FGT on boilers.

4-6

4.1.2 Control Costs for Coal-Fired Boilers

Equipment costs are determined from equipment sizing calculations which are in turn determined from material balances. Material balances for coalfired boilers are contained in Appendix 3. These and the factors discussed in Section 4.1.1 were used to compute the various cost values. The cost components are broken down into individual capital and operating costs in Appendices 6 and 7, respectively.

The annualized costs for each of the standard boilers are summarized in Tables 4.1.2-1 through 4.1.2-4. The costs are also presented as a percent of cost of the uncontrolled boiler. These data are also plotted in Figures 4.1.2-1 through 4.1.2-4 to show the sensitivity of the process costs to control level. The slight nonlinearities are a result of the cost of catalyst which increases while several equipment costs and labor costs are constant for all control levels.

The cost effectiveness of the various applications can be assessed by dividing the annual cost by the annual NO_X removal. The results of this calculation are presented in Table 4.1.2-5. As can be seen, the effectiveness of the parallel flow system on the largest boiler indicates an optimum at 70% while the smallest boiler exhibits an optimum at 90%. There are three primary cost components that determine these results: equipment costs per unit size, catalyst costs, and labor costs. On the smaller boilers the equipment required is obviously smaller and its costs per unit size is greater due to the lack of economy of scale. This is shown directly by the improving cost effectiveness with boiler size (less \$ required per kg NO $_{ imes}$ removed). Now, on the smaller boilers, the catalyst costs are not as dominant as the labor costs. This is due to less catalyst required by the smaller boilers, yet the operating and maintenance labor requirements for the NO_{\times} systems on smaller boilers are comparable to those of larger boilers (at least within the size range of the standard boilers). What this means is that on a small NO_X system where maintenance and operation

4-7

		Syste	211		An	nual costs	Impacts* % increase
	tandard bo input (MBtu/hr)	ilers Type	Type and level of control [†]	Control efficiency (%)	\$/J/S	(\$/MBtu/hr)	in costs over uncontrolled boiler
8.8 Low Sulfu Weste Coal		Package Watertube Underfeed Stoker	PF SCR Intermediate	80	0.0134	(3920)	10.7

.

TABLE 4.1.2-1. COSTS OF NO_× FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS

*Based only on Annual Costs [†]PF = Parallel Flow SCR

		System			Annu	al costs	Impacts* % increase
Hea MW	at input (MBtu/hr)	Туре	Type and Contr	Control efficiency (%)	\$/J/S	(\$/MBtu/hr)	in costs over uncontrolled boiler
22	(75)	Package Watertube	PF SCR Stringent	90	0.00882	(2620)	9.1
Low Sulfu Weste Coal		Chaingrate	PF SCR Intermediate	80	0.00769	(2270)	7.9
			PF SCR Moderate	70	0.00687	(2030)	7.1

TABLE 4.1.2-2. COSTS OF NO $_{\times}$ FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS

*Based only on Annual Costs.

†PF = Parallel Flow SCR

		System			Annu	al costs	Impacts* % increase
Heat MW	t input (MBtu/hr)	Туре	Type and level of control [†]	Control efficiency (%)	\$/J/S	(\$/MBtu/hr)	in costs over uncontrolled boiler
44	(150)	Field-	PF SCR	80	0.00567	(1680)	7.2
		Erected	Intermediate				
.ow Sulfur		Watertube					
		Spreader Stoker					
lestern Coal		SLOKEL					

TABLE 4.1.2-3. COSTS OF NO $_{\times}$ FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS

*Based only on Annual Costs [†]PF = Parallel Flow SCR

TABLE 4.1.2-4. COSTS OF NO $_{\times}$ FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS

		System			Annu	ual costs	Impacts* % increase
Hea MW	t input (MBtu/hr)	Туре	Type and level of control [†]	Control efficiency (%)	\$/J/S	(\$/MBtu/hr)	in costs over uncontrolled boiler
58.6 Low	(200)	Field- Erected Watertube	PF SCR Stringent	90	0.00599	(1760)	7.9
Sulfur Vester Coal		Pulverized Coal	PF SCR Moderate	70	0.00433	(1270)	5.7

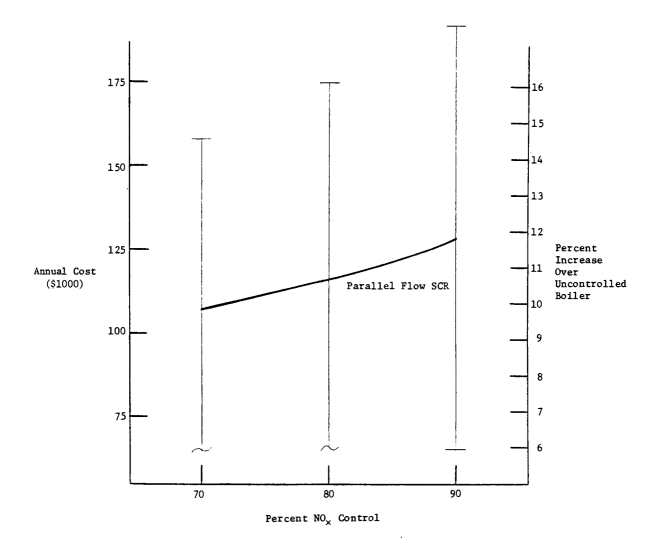


Figure 4.1.2-1. Annual cost of $\rm NO_{\times}$ control systems applied to underfeed stoker standard boiler.

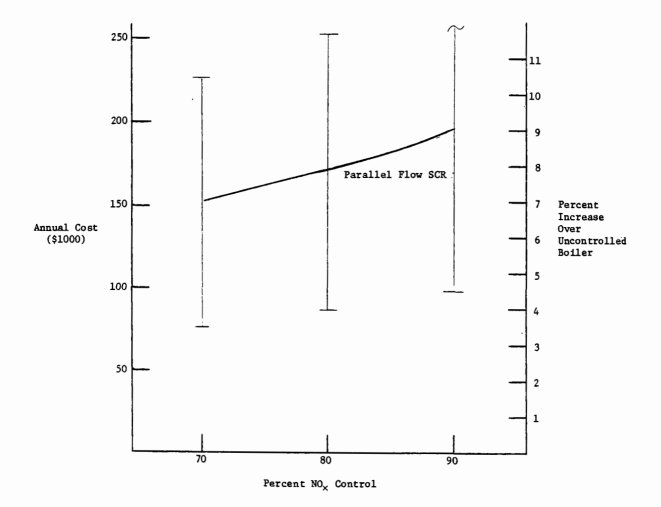


Figure 4.1.2-2. Annual cost of $\rm NO_X$ control systems applied to chaingrate standard boiler.

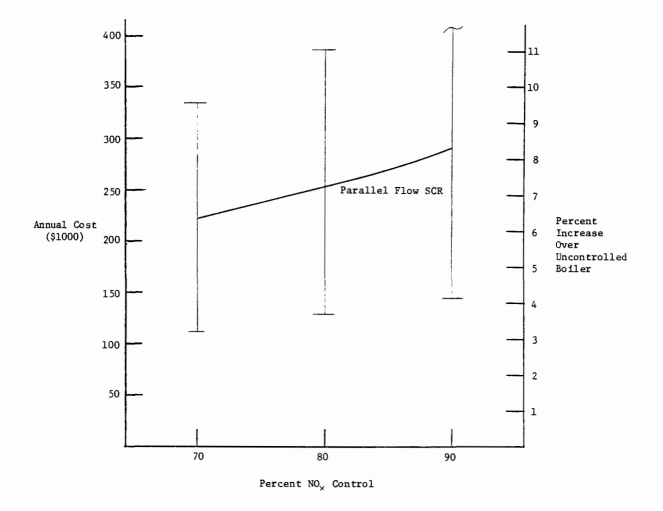


Figure 4.1.2-3. Annual cost of NO_{X} control systems applied to spreader stoker standard boiler.

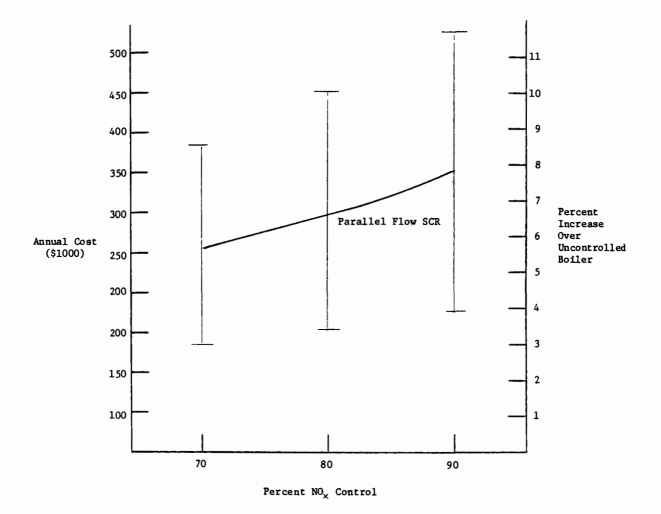


Figure 4.1.2-4. Annual cost of NO_{\times} control systems applied to pulverized coal standard boiler.

		Per	cent NO _× cont	rol
Boiler type	Control type	90	80	70
Underfeed	PF	2.57	2.64	2.75
Chaingrate	PF	1.56	1.53	1.56
Spreader Stoker	PF	1.16	1.13	1.14
Pulverized Coal	PF	0.874	0.836	0.813

TABLE 4.1.2-5. COST EFFECTIVENESS OF NO FGT (\$/kg NO, removed)

personnel are going to be needed regardless, the NO_{\times} system might as well be a little larger to remove additional NO_{\times} . This trend is just the opposite for larger boilers where catalyst costs become dominant. It requires larger amounts of expensive catalyst to remove the additional NO_{\times} and thus increases the cost substantially and decreases the cost effectiveness of the system at higher removal levels. The data presented in Table 4.1.2-5 is plotted in Figure 4.1.2-5.

The cost of applying NO_× FGT to modified or reconstructed facilities is likely to be higher than the cost for applications to new facilities. All of the equipment for new installations will be necessary for retrofit installations, however, additional equipment may also be necessary. Specific costs for retrofit applications were not calculated here, but can be estimated. In a study for the Japanese Environment Agency, five process vendors prepared economic analyses for three applications:¹⁹

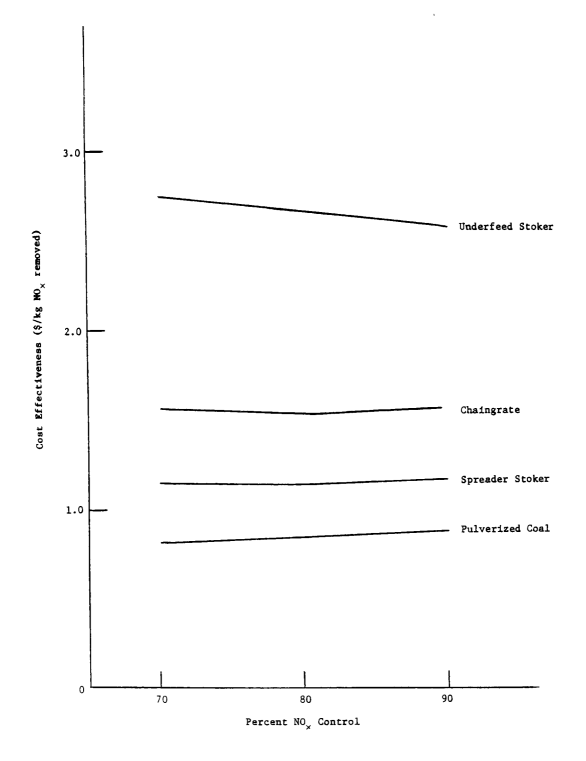


Figure 4.1.2-5. Cost effectiveness of parallel flow SCR NO_X control systems applied to the coal-fired standard boilers.

- 1) new boiler,
- 2) retrofit for gas taken upstream of the air preheater requiring additional ductwork and a fan, and
- 3) retrofit for gas taken downstream of the ESP including gas/gas heat exchanger, heater, fan.

The relative costs for each system treating $40,000 \text{ Nm}^3/\text{hr}$ of flue gas is shown in Table 4.1.2-6.

System	Relative annualized cost
1	1.00
2	1.23
3	2.20

TABLE 4.1.2-6. RELATIVE COSTS OF RETROFIT SCR SYSTEMS

These results indicate that SCR applications to modified or reconstructed facilities can cost from 25 to 120 percent more than applications to new boilers.

4.1.3 Costs To Control Oil-Fired Boilers

The cost calculations presented in this section are based on material balances performed for each case considered. The material balances are presented in Appendix 4. These are used to size the equipment which is subsequently costed. The costing techniques are described in principle in Section 4.1.1. The cost components are broken down into individual capital and operating costs in Appendices 6 and 7. The annualized costs are summarized in Tables 4.1.3-1 and 4.1.3-2. These tables show the annual cost as a percentage of the uncontrolled boiler cost. These values are plotted as a function of control level in Figures 4.1.3-1 and 4.1.3-4.

The parallel flow system shows slightly more sensitivity to control level. This is most likely due to the catalyst which is the most significant cost component. The parallel flow catalyst is about as expensive as the moving bed catalyst per cubic meter, but has a lower space velocity. This causes the parallel flow systems to have a higher catalyst cost component. The nonlinearity is due to this fact combined with the fact that the cost/ unit of equipment increases as size (*i.e.*, control level) decreases.

The cost effectiveness is also determined in Table 4.1.3-3 where the cost per kg of NO_{χ} removed is presented. The cost for the distillate oil-fired boiler is very high due primarily to poor economy of scale since the boiler is small. These costs are plotted in Figures 4.1.3-5 and 4.1.3-6.

The cost differences between the two systems applied to the residual oil-fired boilers are not significant within the accuracy of this cost estimate (±50 percent). The table indicates that the cost effectiveness of the moving bed system increases as removal level increases. This seems to be due to the effect of a greater economy of scale with the larger systems. The reactor is smaller than the parallel flow so the catalyst cost is not as dominant a cost component as the labor cost component. There are several different types of parallel flow type reactors. Some of them consume more energy and cost more than the moving bed reactors, as described here. However, reactors using thin-wall honeycomb or plate catalysts developed recently in Japan are reported to require less energy and cost less than moving bed reactors, and have been used for virtually all of the new SCR plants for dirty or semi-dirty gases.

	System					ual Costs	Impacts* % increase
	tandard boi	llers	Type and	Control			in costs over
Hea ^{MW} t	t input (MBtu/hr)	Туре	level of control [†]	efficiency (%)	\$/J/S	(\$/MBtu/hr)	uncontrolled boiler
4.4	(15)	Package Firetube	FPB SCR Stringent	90	0.0154	(4500)	12.1
Disti Oil	llate	Scotch	FPB SCR Moderate	70	0.0145	(4240)	11.4
44	(150)	Package Watertube	FPB SCR Stringent	90	0.0040	.(1170)	7.5
			FPB SCR Moderate	70	0.0031	(915)	5.6

TABLE 4.1.3–1. COSTS OF NO_{\times} FGT CONTROL TECHNIQUES FOR OIL-FIRED BOILERS

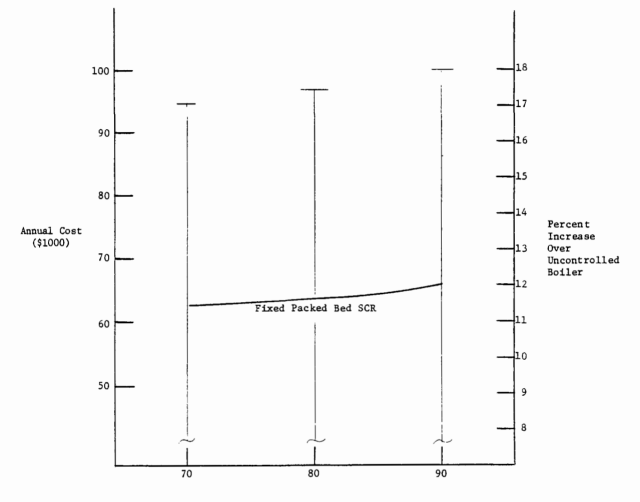
*Based only on Annual Costs †FPB = Fixed Packed Bed

		Sys	tem		Annu	al costs	Impacts* % increase
	tandard bot t input	ilers	Type and level	Control efficiency			in costs over uncontrolled
MWt	(MBtu/hr)	Туре	of control [†]	(%)	\$/J/S	(\$/MBtu/hr)	boiler
8.8	30	Package Watertube	PF SCR Stringent	90	0.0123	(3600)	14
Residu Fuel (FF SCR Moderate	70	0.0110	(3200)	12
			MB SCR Stringent	90	0.0148	(4330)	16
			MB SCR Moderate	70	0.0137	(4010)	15
44 Residu	(150)	Package Watertube	PF SCR Stringent	90	0.00502	(1490)	7.0
Fuel (PF SCR Moderate	70	0.00408	(1210)	5.7
			MB SCR Stringent	90	0.00457	(1360)	6.4
			MB SCR Moderate	70	0.00377	(1120)	5.3

TABLE 4.1.3-2. COSTS OF NO $_{\times}$ FGT CONTROL TECHNIQUES FOR OIL-FIRED BOILERS

* Based only on Annual Costs

#PF = Parallel Flow
MB = Moving Bed



Percent NO_{\times} Control

Figure 4.1.3-1. Annual cost of NO_X control system applied to 4.4 MW distillate oil-fired standard boiler.

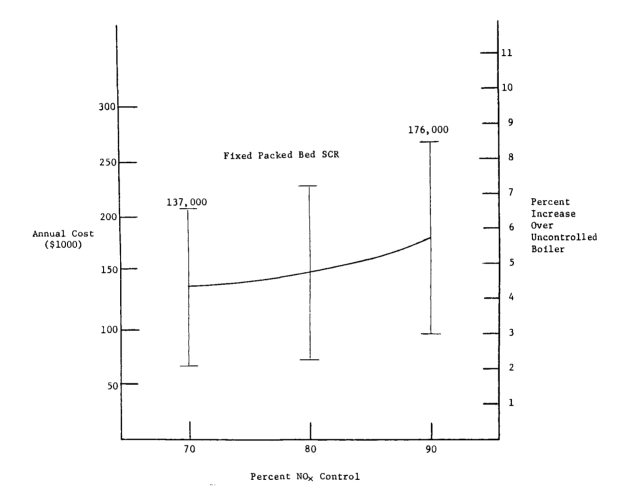
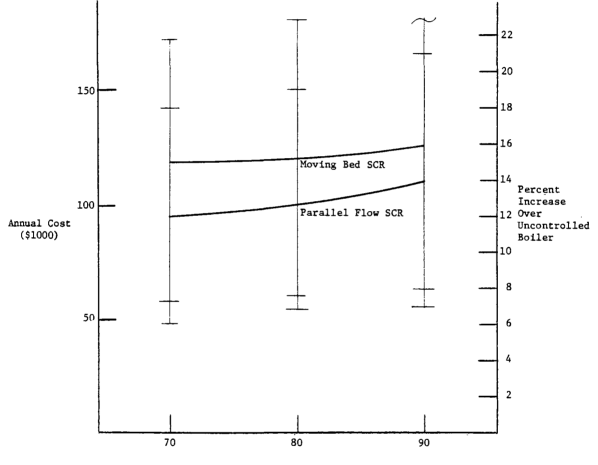
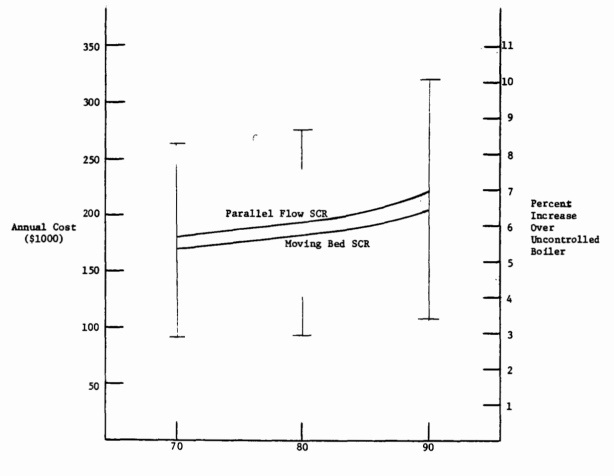


Figure 4.1.3-2. Annual cost of $\rm NO_{\times}$ control system applied to 44 MW distillate oil-fired standard boiler.



Percent NO_X Control

Figure 4.1.3-3. Annual cost of $\rm NO_{\times}$ control systems applied to 8.8 MW residual oil-fired standard boiler.



Percent NO_× Control

Figure 4.1.3-4. Annual cost of $\rm NO_X$ control systems applied to 44 MW residual oil-fired standard boiler.

	Boiler size,	_	Percent NO _X o	control (\$/kg NC)x removed)
Fuel	MWt	Control type*	90	80	70
Distillate Oil	4.4	FPB	17.6	19.0	21.4
Distillate Oil	44	FPB	.3.8	3.8	3,8
Residual Oil	8.8	PF MB	5.7 6.9	6.0 7.3	6.6 8.2
Residual Oil	44	PF MB	1.89 1.72	1.85 1.75	1.97 1.84

TABLE 4.1.3-3. COST EFFECTIVENESS OF NO_X FGT

* FPB = Fixed Packed Bed SCR

PF = Parallel Flow SCR

MB = Moving Bed SCR

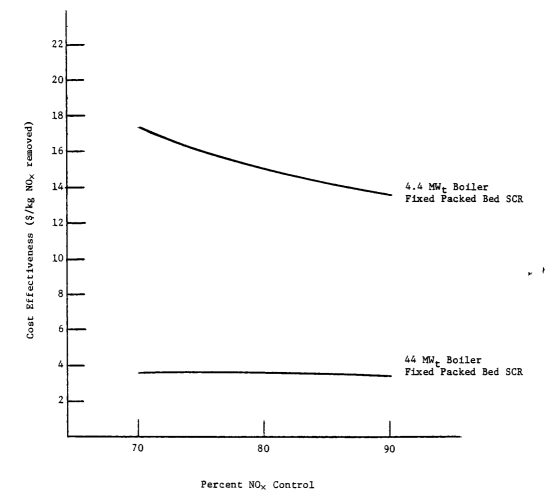
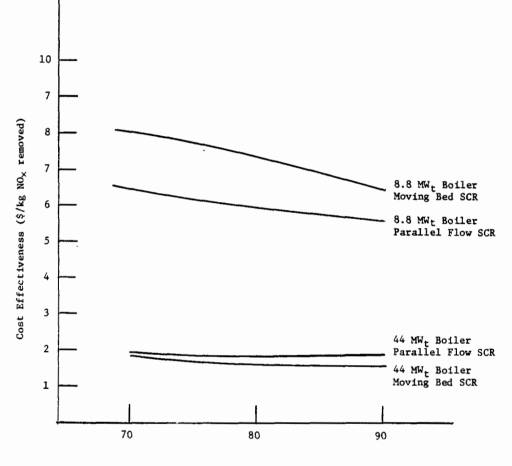


Figure 4.1.3-5. Cost effectiveness of FGT systems applied to distillate oil-fired boilers.



Percent NO_× Control

Figure 4.1.3-6. Cost effectiveness of FGT systems applied to residual oil-fired boilers.

The overall conclusion is the systems applied to the residual oil-fired boilers to not appear to be cost sensitive with respect to control level. The small distillate oil-fired boiler appears to be very sensitive since higher control can be achieved with only slightly higher annualized costs.

The cost of applying NO_× FGT to modified or reconstructed facilities is likely to be higher than the cost for applications to new facilities. All of the equipment for new installations will be necessary for retrofit installations, however, additional equipment may also be necessary. Specific costs for retrofit applications were not calculated here, but can be estimated. In a study for the Japanese Environment Agency, five process vendors prepared economic analyses for three applications:¹⁹

- 1) new boiler,
- 2) retrofit for gas taken upstream of the air preheater requiring additional ductwork and a fan, and
- retrofit for gas taken downstream of the ESP including gas/gas heat exchanger, heater, fan.

The relative costs for each system treating 40,000 Nm /hr of flue gas are shown in Table 4.1.3-4.

System	Relative annualized cost			
1	1.00			
2	1.23			
3	2.20			

TABLE 4.1.3-4. RELATIVE COSTS OF RETROFIT SCR SYSTEMS

These results indicate that SCR applications to modified or reconstructed facilities can cost from 25 to 120 percent more than applications to new boilers.

4.1.4 Control Costs for Natural Gas-Fired Boilers

This section presents cost calculations for a FGT system applied to the natural gas-fired standard boiler. The calculations are based on material balances contained in Appendix 5. The costing techniques have been described in Section 4.1.2.

The cost components are broken down into individual capital and operating costs in Appendices 6 and 7. Both total annualized costs and costs as a percentage of the uncontrolled boiler cost are shown in Table 4.1.4-1. The data presented are also plotted in Figures 4.1.4-1 and 4.1.4-2 to show the sensitivity to control level. The costs are fairly linear with control level indicating only a slight sensitivity.

The cost effectiveness of NO_{\times} control on natural gas-fired boilers is determined in Table 4.1.4-2 and these values are plotted in Figure 4.1.4-3. The smaller systems are not as cost effective as the larger systems since the cost per unit size of equipment is less for larger systems. The cost effectiveness of the small system is more sensitive to control level due to the influence of labor costs, which are constant for all control levels.

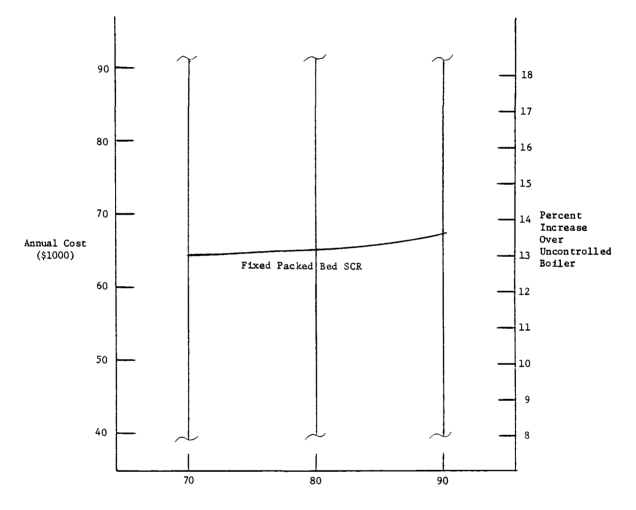
The cost of applying NO_{\times} FGT to modified or reconstructed facilities is likely to be higher than the cost for applications to new facilities. All of the equipment for new installations will be necessary for retrofit installations, however, additional equipment may also be necessary. Specific costs for retrofit applications were not calculated here, but can be estimated. In a study for the Japanese Environment Agency, five process vendors prepared economic analyses for three spplications:¹⁹

- 1) new boiler,
- retrofit for gas taken upstream of the air preheater requiring additional ductwork and a fan, and

System					Annı	ual costs	Impacts* % increase
Heat	ndard boi Input 18tu/hr)	llers Type	level		\$/J/S	(\$/MBtu/hr)	in costs over uncontrolled boiler
4.4 Natu r al Gas	(15)	Package Firetube Scotch	FPB SCR Stringent	90	0.0154	(4510)	13.6
			FPB SCR Moderate	70	0.0146	(4290)	13.0
44 Natural Gas	(150)	Package Watertube	FPP SCR Stringent	90	0.0040	(1160)	7.5
			FPB SCR Moderate	70	0.0029	(863)	5.6

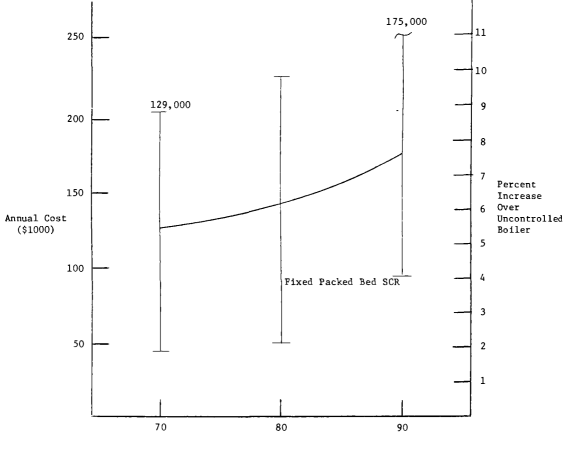
TABLE 4.1.4-1. COSTS OF NO $_{\times}$ FGT CONTROL TECHNIQUES FOR NATURAL GAS-FIRED BOILERS

*Based only on Actual Costs †FPB = Fixed Packed Bed



Percent NO_{χ} Control

Figure 4.1.4-1. Annual cost of $\rm NO_X$ control system applied to 4.4 MW natural gas-fired standard boiler.



Percent NO_X Control

Figure 4.1.4-2. Annual cost of $\rm NO_X$ control system applied to 44 MW natural gas-fired standard boiler.

		\$/kg NO _X removed			
	Boiler size,		Pe	rcent NO _X cont	rol
Fuel	^{MW} t	Control type	90	80	70
atural Gas	4.4	Fixed Packed Bed	16.0	17.5	19.7
atural Gas	44	Fixed Packed Bed	3.4	3.3	3.2

TABLE 4.1.4-2. COST EFFECTIVENESS OF NO_X FGT

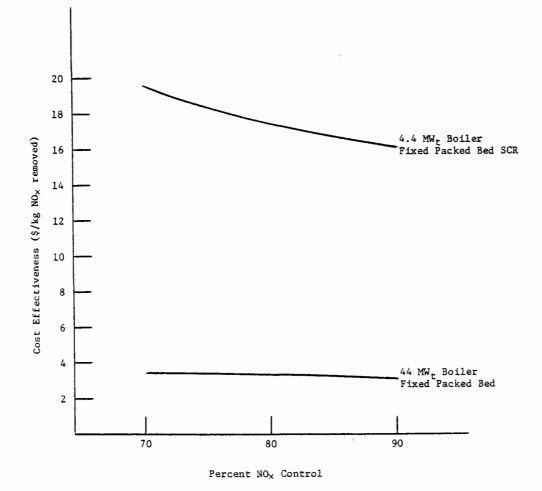


Figure 4.1.4-3. Cost effectiveness of FGT systems applied to natural gas-fired boilers.

 retrofit for gas taken downstream of the air preheater including gas/gas heat exchanger, heater, fan.

The relative costs for each system treating 40,000 Nm³/hr of flue gas is shown in Table 4.1.4-3.

System	Relative annualized cost
1	1.00
2	1.23
3	2.20

TABLE 4.1.4-3. RELATIVE COSTS OF RETROFIT SCR SYSTEMS

These results indicate that SCR applications to modified or reconstructed facilities can cost from 25 to 120 percent more than applications to new boilers.

4.1.5 Summary

In all cases the catalyst cost is a significant capital cost. Other significant capital cost components are labor, fan motor drive, and NH₃ storage tanks. The most significant operating cost component in all cases was labor. The smaller systems are more significantly affected by this fact than are the larger systems. As a result, the costs for small systems are high, not only because they lack economy of scale, but due to labor considerations as well. As a result the size of the unit has a greater effect on costs than does control level. This dramatic effect is most readily observed in the cost effectiveness numbers. The systems exhibited an order of magnitude larger cost/kg of NO_X when applied to the smallest systems.

4.2 NO_X/SO_X SYSTEM

4.2.1 Introduction

This section considers the costs of applying the NO_X/SO_X system selected in Section 3 to two coal-fired and one oil-fired boilers. The costing techniques are the same as used with the NO_X -only processes and will not be repeated here. The equipment items are more numerous due to the higher number of process operations associated with the process. These items are listed in Table 4.2.1-1.

 NO_X/SO_X Parallel Passage	
Reactors (2)	
Catalyst	
Fan Motor Drive	
NH₃ Storage Tank	
NH₃ Transfer Pump	
NH ₃ Vaporizer	
Naphtha Reformer	
H ₂ SO ₄ Plant	
Compressor/Gasholder	

TABLE 4.2.1-1. PURCHASED EQUIPMENT FOR NO_{\times} FGT SYSTEMS

With the coal-fired boilers, both high and low sulfur coals were analyzed. However, only one set of control levels are considered (80 percent NO_X , 85 percent SO_X) and therefore, it is not possible to present costs as a function of control level as is done in the NO_X -only section. Instead, the costs are plotted against flue gas flow rate to show the effect of unit size on cost. The results for the residual oil-fired boiler are presented in tabular form, but not plotted since only one boiler and control level are considered.

4.2.2 Control Costs for Coal-Fired Boilers

The equipment listed in the previous table is sized based on material balances performed for each case. These balances are presented in Appendix 3. Detailed breakdowns of both capital and operating costs are presented in Appendices 6 and 7, respectively.

The annualized costs for the standard boilers considered are presented in Table 4.2.2-1 and plotted in Figure 4.2.2-1. The costs are significantly higher than those for the NO_X -only processes because the additional requirement of SO₂ removal necessitates the use of small processing units for H₂ production and SO₂ workup. In a real world situation where several industrial boilers operate at a single location, it will be possible to reduce costs by having large, central units for H₂ production and SO₂ workup. This option is not considered here since the cost impact is a function of the total number of boilers serviced by the central facilities and this is entirely site specific.

The cost to retrofit such a process can be calculated from the data presented in Section 4.1.2. Depending on the modifications required by the retrofit, the additional cost will be increased by an amount equivalent to 25 to 120 percent of the cost of an average NO_X -only system. The cost of special equipment necessary for SO_2 processing is not affected by a retrofit application.

4.2.3 Control Costs for the Oil-Fired Boiler

The equipment items necessary to treat flue gas from the residual oilfired boiler are the same as for the coal-fired boilers. The annualized cost of the dry NO_X/SO_X process applied to the residual oil-fired standard boiler

	System				Annu	ual Costs	Impacts* % increase	
Hea MW	ut input (MBtu/hr)	Туре	Control efficiency (%)	\mathtt{Coal}^\dagger	\$/J/S	(\$/MBtu/hr)	in costs over uncontrolled boiler	
8.8	(30)	Package Watertube	90% NO _× 85% SO _×	HSE	0.0811	(23,690)	75	
		Underfeed Stoker		LSW	0.0527	(15,400)	49	
58.6	(200)	Field Erected	90% NO _X 85% SO _X	HSE	0.0309	(9,025)	42	
		Watertube Pulverized Coal	Â	LSW	0.0153	(4,468)	21	

TABLE 4.2.2-1. COSTS OF NO_X/SO_X FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS

*Based only on Annual Costs †HSE = High sulfur eastern coal (3.5% S) LSW = Low sulfur western coal (0.6% S)

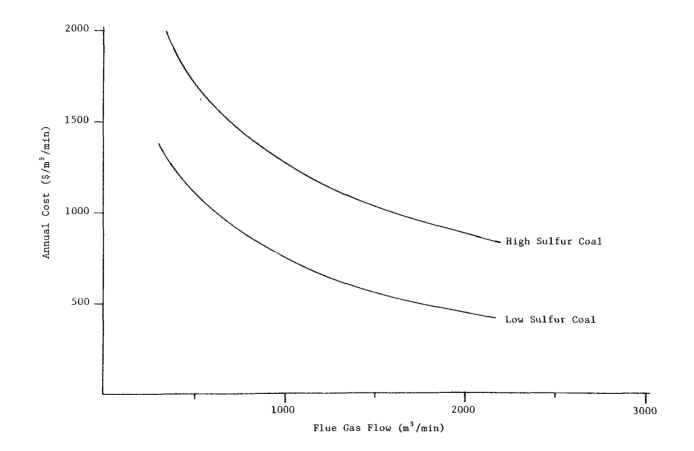


Figure 4.2.2-1. Annual cost of parallel flow SCR NO_X/SO_X FGT for coal-fired boilers

is shown in Table 4.2.3-1. Detailed capital and annual costs are presented in detail for this case in the appendices. Retrofit considerations for this case are similar to those for the coal cases. The increased cost of a retrofit will be increased by the same dollar amount as the cost increase incurred when a similarly sized NO_X -only unit is retrofitted.

Standard boiler Heat input			Annual costs		Impact* % increase in costs over uncontrolled	
MW	(MBtu/hr)	Туре	Control Level	\$/J/S	(\$/MBtu/hr)	boiler
44	(150)	Package Watertube	90% NO _× 85% SO _×	0.0249	(7280)	43

TABLE 4.2.3-1. COSTS OF THE DRY NO_X/SO_X CONTROL TECHNIQUE FOR THE RESIDUAL OIL-FIRED BOILER

*Based only on Annual Costs

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SECTION 5

ENERGY IMPACT

5.1 NO_×-ONLY SYSTEMS

5.1.1 Introduction

The three types of control systems selected in Section III for further comparison are analyzed with respect to energy requirements. All of the control systems are basically similar in principle and differ mainly in the design parameters. There are also a few differences in equipment requirements. Energy consumption steps considered in this analysis are listed in Table 5.1.1-1 for each of the control systems considered. It was assumed that flue gas could be taken from the boiler between the economizer and air heater at a temperature of 375°C. This eliminates the need for flue gas heating and heat exchange equipment. Since the hot flue gas is returned to the boiler upstream of the air heater, there is no loss in boiler efficiency.

Energy consumption was calculated using the design information and standard engineering principles. Design information from a variety of process developers was compared and used to generate a range of values or specific values. A range of values was determined for design parameters which changed with control level. Specific values for analysis were chosen from this range based on the level of control being considered, *e.g.* for 70% control a value at the lower end of the range was used. Design data used in this analysis is presented in Table 5.1.1-2.

NO _× FGT system	Energy consumption step (equipment)	Type of energy consumed
Parallel Flow SCR	Reactor Draft Loss (Fan) Liquid NH ₃ Transfer (Pump) NH ₃ Vaporization (Vaporizer) NH ₃ Dilution	Electrical Electrical Steam Steam
Moving Bed SCR	Reactor Draft Loss (Fan) Liquid NH ₃ Transfer (Pump) Catalyst Screening & Transfer (Elevator) Baghouse Draft Loss (Blower) NH ₃ Vaporization (Vaporizer) NH ₃ Dilution	Electrical Electrical Electrical Electrical Steam Steam
Fixed Packed Bed SCR	Reactor Draft (Fan) Liquid NH3 Transfer (Pump) NH3 Vaporization (Vaporizer) NH3 Dilution Soot Blowing-Distillate Oil Boiler Only	Electrical Electrical Steam Steam Steam

TABLE 5.1.1-1. AREAS OF ENERGY CONSUMPTION IN NO $_{\times}$ FGT SYSTEMS^{1,2}

TABLE 5.1.1-2.		IGN PARAMETERS USED FO)R
	ENERGY IMPACT	F CALCULATIONS ^{1,2,3}	

	Range or specific values used		
Parameter	Parallel flow	Moving bed	Fixed packed bed
Space velocity	3000-5000	6000-10000	6000-10000
$\rm NH_3:NO_{x}$ mole ratio	0.7-1.0	0.7-1.0	0.7-1.0
Dilution ratio (moles steam/mole M	₩ ₃) 5:1	5:1	5:1
Dilution steam pressure	30 psig	30 psig	30 psig
Flue gas temperature	375°C	375°C	375°C
Pressure drop	80-160 mmH ₂ 0	40-80 mmH ₂ 0	$\sim 125 \text{ mmH}_2\text{O}$
Catalyst type	Square honeycomb	Ring	Spherical pellet
Void fraction of packed catalyst particles	0.67-0.7	0.52	0.26

Steam was chosen as the NH₃ dilution gas because of its ease of application and safety considerations. Air, at 20:1 air:NH₃ mole ratio, can also be used as an NH₃ diluent. Its use requires a compressor or blower and a motor which are high maintenance items. Also, at dilution ratios less than 20:1 there is an explosion hazard. The optimum choice would ordinarily be made by comparing the operating costs of steam use versus the capital charges of the air handling equipment plus the operating costs of electricity. This optimization is beyond the scope of this study and is site specific.

The analyses conducted in this study assumed that the boilers were operated constantly at full load and, therefore, had constant flue gas temperatures. However, it is possible that the boiler may experience large and frequent load swings which result in a variable flue gas temperature. FGT systems in this service will require flue gas heating in order to maintain sufficiently high temperatures. Temperature control can be accomplished by either a heater or a slipstream around the economizer. The heater will effectively decouple the FGT system from the boiler and does not require flow control of a flue gas slipstream. The economizer bypass will not derate the boiler since it will only be required during low load situations. Energy usage calculations were not made for either of these approaches since the amount of heating necessary is likely to be different for each boiler application.

5.1.2 Energy Impact of Controls for Coal-Fired Boilers

This subsection presents the results of calculations on the energy requirements of the candidate control systems applied to the standard boilers. One simplification was made in order to reduce the number of cases necessary for consideration and that is that only one coal was analyzed for each boiler. The justification for this simplification is presented below.

The result of the energy impact analyses indicate that the most significant energy consumption occurs in the fan required to overcome the reactor pressure drop and NH₃ dilution by steam. Coal sulfur content does not

significantly affect the fan requirements which are a function of flue gas flow rate and control level. NH_3 dilution steam is affected; however, energy consumption of this step is approximately a third of the fan requirement. This is illustrated in an example calculation in Table 5.1.2-1. As can be seen the sulfur content of the coal does not significantly affect the total energy requirements especially when compared to the effect of control level.

The low sulfur coal was used for the analyses since the NO_{\times} emissions were somewhat higher and, therefore, energy usage for the other coals will not exceed those presented here.

Also, SIP control levels were not considered since in cases where control is required, it can be achieved through use of combustion modifications. The typical SIP control levels are shown in Table 5.1.2-2.

Fuel	NO _x emissions, $\frac{1b}{10^5}$ Btu	SIP level, $\frac{1b}{10^6}$ Btu	Required control efficiency
3.5% S Coal	0.64	0.7	0
0.9% S Coal	0.55	0.7	0
0.6% S Coal	0.78	0.7	10%

TABLE 5.1.2-2. SIP CONTROL LEVELS⁴

Material balances were performed for each of the 7 cases considered for the coal-fired standard boilers. The results of these calculations appear in Appendices 3, 4, and 5. These results were used to calculate energy requirements of the control systems and an example calculation is presented in Appendix 8.

The results of the energy requirement calculations are presented in Tables 5.1.2-3 through 5.1.2-6. Each table represents one standard boiler and all control types and levels are included. It should be noted that the megawatt values shown for electrical usage are thermal megawatts and not

a) Effect of sulfur content	Energy usage (MW thermal)		
	0.6% S coal	0.9% S coal	3.5% S coal
	(187.56 1b NO _× /hr)	(130.50 1b NO _× /hr)	(152.46 1b NO _× /hr)
Energy Consumer			
Fan	0.91	0.88	0.88
Liquid NH Pump	0.00373	0.00373	0.00373
NH Vaporizer	0.0383	0.0275	0.0325
NH Dilution Steam	0.325	0.234	0.275
Total	1.275	1.145	1.191
	10% Differ	1	
	7 Differ	% ence	

TABLE 5.1.2-1.RELATIVE SIGNIFICANCE OF PARAMETERS CONSIDERED IN ENERGY ANALYSIS
Example:Example:Pulverized Coal Boiler, 90% Control, Parallel Flow SCR

b) Effect of removal level	90% removal	70% removal	
Total Energy Consumed	1.28	0.821	
(MW thermal)			
	3 Diffe	6% rence	

		S	ystem				Energy consu	mption	
	Standard boiler			Control		Energy consumed		% increase	
Hea MW	t input (MBtu/hr)	Туре	Type and level of control	efficiency %	Energy types	by cont MW t	rol device (MBtu/hr)	in energy use over uncontrolled boiler	
			SCR Parallel Flow		Electrical	0.161	(0.549)		
58.6	(200)	Field Erected, Watertube, Pulverized Coal	Moderate	70	Steam	0.0797	(0.272)	0.41	
			SCR Parallel Flow		Electrical	0.268	(0.912)		
			Stringent	90	Steam	0.108	(0.364)	0.64	

TABLE 5.1.2-3. ENERGY CONSUMPTION FOR NO_× FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS

TABLE 5.1.2-4. ENERGY CONSUMPTION FOR NO_{\times} FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS

_		Sy:	stem	Energy consumption				
	Standard bo		Type and level	Control	_	-	y consumed	% increase
Hea MW	t input (MBtu/hr)	Туре	of control	efficiency %	Energy types	<u>by con</u> MW t	(MBtu/hr)	in energy use over uncontrolled boiler
44	(150)	Field Erected,	SCR Parallel Flow		Electrical	0.126	(0.428)	
		Watertube, Spreader Stoker	Intermediate	80	Steam	0.0568	(0.194)	0.41

		S	ystem		Energy Consumption				
	Standard Boi Input (MBtu/hr)	<u>ler</u> Туре	Туре & Level of Control	Control Efficiency %	Energy Types	Energy C <u>by Contro</u> MW _t (% Increase in Energy Use Over Uncontrolled Boile	
			SCR Parallel Flow		Electrical	0.0408	(0.139)		
			Moderate	70	Steam	0.0253	(0.0862)	0.30	
22	(75)	Package Watertube	SCR Parallel Flow		Electrical	0.0505	(0.172)		
		Chaingrate	Intermediate	80	Steam	0.0289	(0.0988)	0.36	
			SCR Parallel Flow		Electrical	0.0669	(0.228)		
			Stringent	90	Steam	0.0337	(0.115)	0.46	

TABLE 5.1.2-5. ENERGY CONSUMPTION FOR NO× FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS

TABLE 5.1.2-6. ENERGY CONSUMPTION FOR NO_× FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS

			System				Energy consu	mption
Standard boiler Heat input Type		Type and level	Control efficiency	Energy	Energy consumed by control device		% increase in energy use over	
MW	(MBtu/hr)	<i>.</i>	of control	7	types	Mwt	(MBtu/hr)	uncontrolled boiler
8.8	(30)	Package	SCR Parallel Flow		Electrical	0.0185	(0.0630)	
		Watertube Underfeed Stoker	Intermediate	80	Steam	0.0113	(0.0387)	0.34

electrical megawatts. The data appearing in these tables is summarized in Table 5.1.2-7. This data is plotted in Figures 5.1.2-1 through 5.1.2-4. Each figure represents one standard boiler and shows the effect of removal level on energy usage for both types of FGT candidate systems: parallel flow SCR and moving bed SCR. Figure 5.1.2-5 presents energy usage for all boilers and FGT systems as a percent of the boiler heat input.

In general, energy usage seems to increase with control level in a nonlinear manner. This is also true with regard to energy usage as a percent of boiler input. Also, it appears that more energy is used per mole of NO_X controlled with the larger boilers. This increase with boiler size is not a physical phenomena of SCR systems but rather an idiosyncrasy of the reactor design scheme. In keeping reactor geometry consistent from boiler to boiler, the linear gas velocity (m/s) decreased slightly with boiler size resulting in a corresponding slight decrease in pressure drop. However, the pressure drops of all the standard boilers are in the range of commercial operations and the differences in energy usage as a percentage of boiler heat input of the standard boilers are not large.

The effect of this energy usage on economics is examined in Section IV.

Very little work has been done with SCR systems to affect reductions in energy consumption. Problem solving efforts thus far have been directed toward improving the reliability of operating units and applying the control techniques to coal-fired flue gas. It is likely that only after the processes have been applied and demonstrated on coal-fired units will the overall energy consumption be examined in detail. It should be noted that the SCR processes are the least energy intensive of all of the FGT control systems mentioned ir Section II.^{1,5}

There are two areas in which there is a potential for energy savings. These are control of excess air and NH₃ dilution. By using only as much excess air as necessary, the energy required for pressure drop will be reduced. This has a twofold effect. Not only is the flow through the

	Pulveri	zed coal	Sprea	der stoker	Chai	ngrate	Underfeed stoker		
	Total thermal kW	% of boiler heat input							
Parallel Flow SCR									
90% Removal	376	0.64	-	-	101	0.46	-	-	
80% Removal	-	-	183	0.41	80	0.36	30	0.34	
70% Removal	241	0.41	-	-	66	0.30	-	_	

TABLE 5.1.2-7. SUMMARY OF ENERGY REQUIREMENTS FOR COAL-FIRED INDUSTRIAL BOILERS

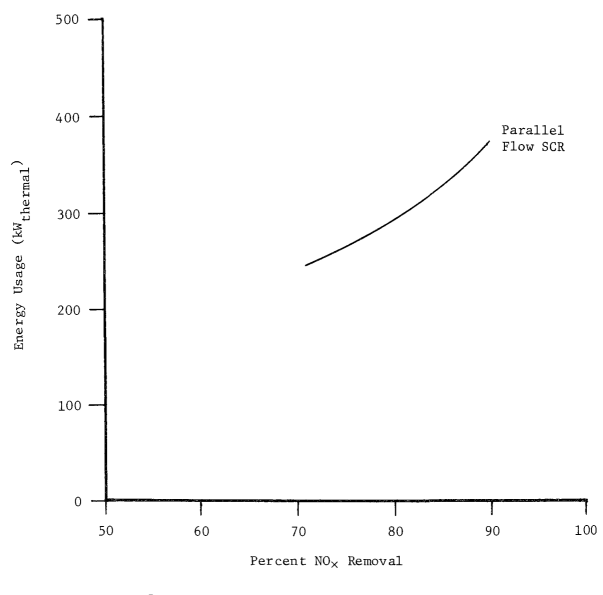


Figure 5.1.2-1. Energy usage of NO_{\times} control systems for pulverized coal standard boiler.

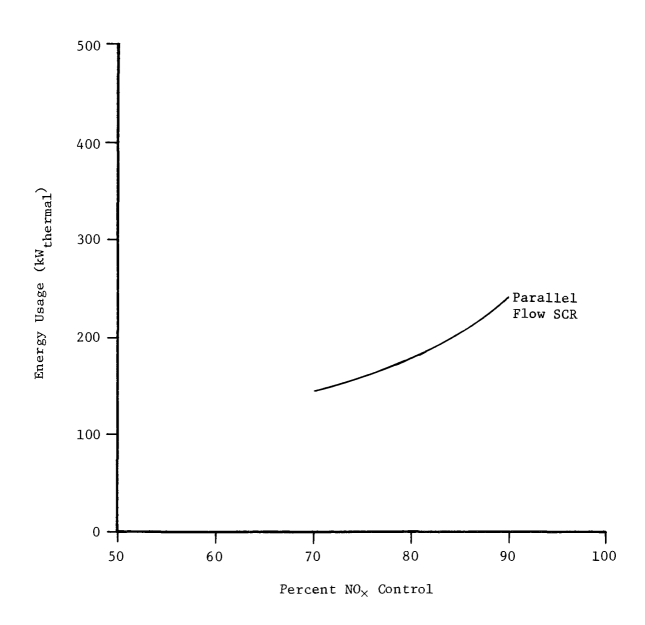


Figure 5.1.2-2. Energy usage of NO_X control systems for spreader stoker standard boiler.

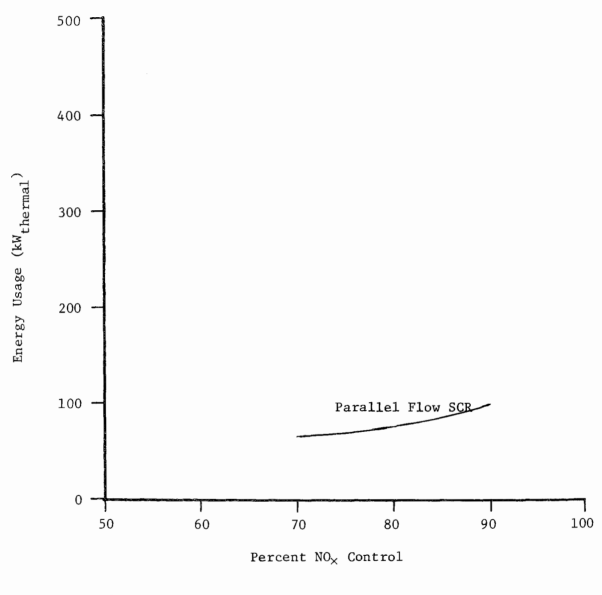


Figure 5.1.2-3. Energy usage of $\rm NO_{\times}$ control systems for chaingrate standard boiler.

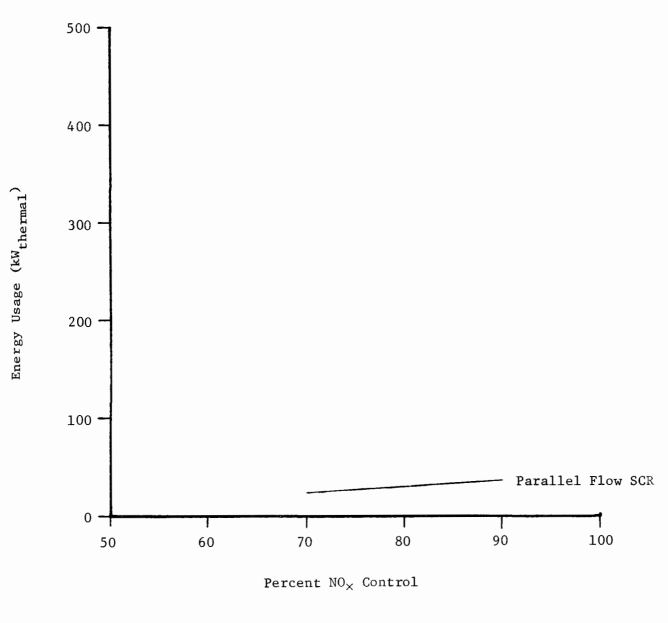


Figure 5.1.2-4. Energy usage of $\rm NO_{\times}$ control systems for underfeed stoker standard boiler.

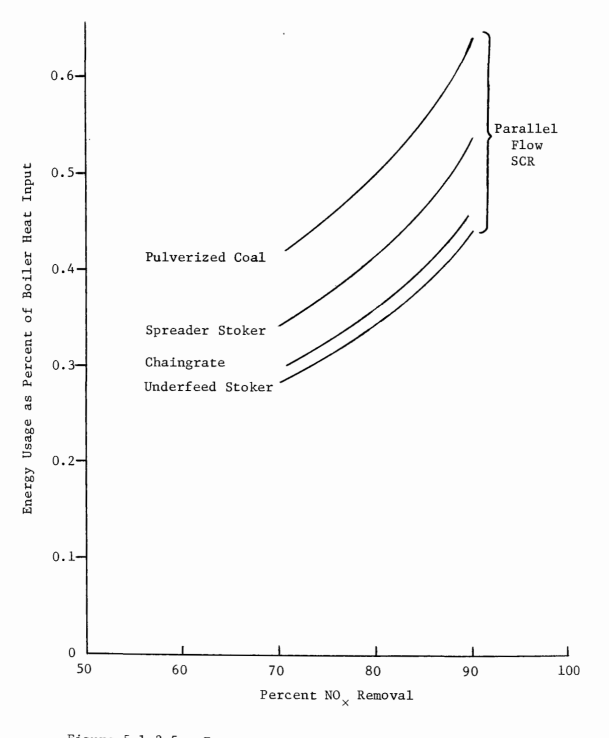


Figure 5.1.2-5. Energy usage of NO_{\times} control systems as percent of boiler heat input.

reactor reduced, but the required reactor volume itself is reduced by lower flue gas flow rates. It is likely that a boiler equipped with combustion modifications will utilize low excess air for NO_{χ} control. Energy consumption by NH₃ dilution might be reduced by using air instead of steam at a specific site. Use of air is less safe since some air:NH₃ mixtures can be explosive.

The energy impact of FGT controls applied to modified or reconstructed facilities (retrofit application) will be the same or greater than that for new facilities. If flue gas can be taken from the economizer of the existing boiler at 350-400°C and returned upstream of any existing heat exchange equipment, then there will be no additional energy impact.

If the flue gas is only available at a lower temperature (<350-400°C) then a heater will be required. The energy impact of the heater will depend on the temperature of the flue gas. If the temperature is that of the outlet gas of the standard boilers (approximately 180°C), calculations indicate that energy requirement would be more than tripled even if heat exchange equipment is used to recover 85% of the energy supplied by the heater. The heater will probably be oil-fired for ease of control.

These results indicate that, on retrofit applications, there is a considerable energy incentive to obtain the flue gas at the necessary reaction temperature in order to avoid flue gas heating. Other energy impacts would be the same as those for new facilities.

5.1.3 Energy Impact of Controls for Oil-Fired Boilers

In this subsection, the results of energy impact calculations for the candidate FGT systems as applied to the standard oil-fired boilers are presented. The combinations considered are

Boiler Size, MW _t	Fuel	FGT System
8.8, 44	Residual Oil	Parallel Flow SCR
8.8, 44	Residual Oil	Moving Bed SCR
4.4, 44	Distillate Oil	Fixed Packed Bed SCR

Also, two levels of control are considered for each combination.

The first step in performing this energy impact analysis was to calculate general material balances. The result of these balances were used to determine energy requirements for each process step. Energy consuming steps and the types of energy used were presented earlier in Table 5.1.1-1. All calculations are similar to the example case presented in Appendix 8.

The results of these calculations are presented in Tables 5.1.3-1 and 5.1.3-2. The data in Table 5.1.3-1 represents energy consumption for residual oil-fired boilers. Two candidate systems and two levels of control are considered. Table 5.1.3-2 shows energy consumption for application of a fixed packed bed SCR process to the standard boiler firing distillate oil.

All energy values presented are on a thermal basis. Actual electrical usages have been converted to a heat input basis by assuming 10,000 Btu/hr per kW.

The data appearing in Tables 5.1.3-1 and 5.1.3-2 is summarized in Table 5.1.3-3 and is plotted in Figures 5.1.3-1 through 5.1.3-4. The first two figures show thermal energy usage as a function of NO_X control for all fuels and control systems. The next two figures illustrate energy usage as a percent of boiler heat input for all cases.

Energy usage increases in a nonlinear manner with control level. The energy usage as a percent of boiler input is also nonlinear. On this basis the fixed packed bed SCR appears to be the most energy intensive and the moving bed SCR the least. It is difficult to draw any definite conclusions

		Sys	tem		Energy consumption				
Heat	Standard bo t input	iler Type	Type and level	Control efficiency	Energy		consumed	% increase in energy use over	
MWt	(MBtu/hr)		of control	%	types	MWt	(MBtu/hr)	uncontrolled boiler	
3.8	(30)	Package Watertube	SCR Parallel Flow Moderate	70	Electrical Steam	0.011 0.0071	(0.0367) (0.0242)	0.20	
			SCR Parallel Flow Stringent	90	Electrical Steam	0.018 0.0095	(0.0597) (0.0323)	0.31	
			SCR Moving Bed Moderate	70	Electrical Steam	0.0094 0.0071	(0.0322) (0.0242)	0.19	
			SCR Moving Bed Stringent	90	Electrical Steam	0.014 0.0095	(0.0462) (0.0323)	0.26	
4	(150)	Package Watertube	SCR Parallel Flow Moderate	70	Electrical Steam	0.0813 0.0253	(0.277) (0.0864)	0.24	
			SCR Parallel Flow Stringent	90	Electrical Steam	0.134 0.0337	(0.458) (0.115)	0.38	
			SCR Moving Bed Moderate	70	Electrical Steam	0.0570 0.0253	(0.195) (0.0864)	0.19	
			SCR Moving Bed Stringent	90	Electrical Steam	0.0918 0.0337	(0.314) (0.115)	0.29	

TABLE 5.1.3-1. ENERGY CONSUMPTION FOR NO $_{\times}$ FGT CONTROL TECHNIQUES FOR RESIDUAL OIL-FIRED BOILERS

			ystem	Energy consumption				
	Standard bo	iler		Energy consumed			% increase	
Heat	t_input Type Type and level efficiency E		Energy	by conti	col device	in energy use over		
MWt	(MBtu/yr)		of control	%	types	MWt	(MBtu/yr)	uncontrolled boile
4.4	(15)	Package Firctube Scotch	SCR Fixed Packed Bed Moderate	70	Electrical Steam	0.00994 0.00697	(0.0339) (0.0238)	0.38
			SCR Fixed Packed Bed Stringent	90	Electrical Steam	0.0158 0.00888	(0.0539) (0.0302)	0.56
44	(150)	Package Watertube	SCR Fixed Packed Bed Moderate	70	Electrical Steam	0.200 0.0734	(0.682) (0.251)	0.62
			SCR Fixed Packed Bed Stringent	90	Electrical Steam	0.121	(0.414) (0.241)	0.44

TABLE 5.1.3-2. ENERGY CONSUMPTION FOR NO_{\times} FGT CONTROL TECHNIQUES FOR DISTILLATE OIL-FIRED BOILERS

	8.8 MW _t	Residual oil	44 MW _t F	Residual oil	4.4 MW _t I)istillate oil	44 MW _t I	Distillate oil
	Total thermal kW	% of boiler heat input	Total thermal kW	% of boiler heat input	Total thermal kW /	% of boiler heat input	Total thermal kW	% of boiler heat input
Parallel Flow SCR								
90% Removal	27	0.31	168	0.38		*		*
70% Removal	18	0.20	107	0.24		*		*
Moving Bed SCR								
90% Removal	23	0.26	126	0.29		*		*
70% Removal	17	0.19	82	0.19		*		*
Fixed Packed Bed SCR								
90% Removal		*		*	25	0.56	274	0.62
70% Removal		*		*	17	0.38	192	0.44

TABLE 5.1.3-3. SUMMARY OF ENERGY REQUIREMENTS FOR OIL-FIRED INDUSTRIAL BOILERS

*Not considered as a candidate system.

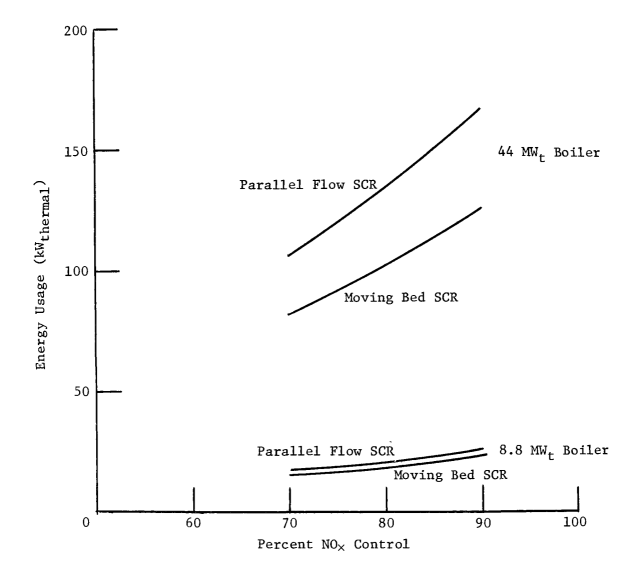


Figure 5.1.3-1. Energy usage of $\rm NO_X$ control systems for residual oil-fired standard boilers.

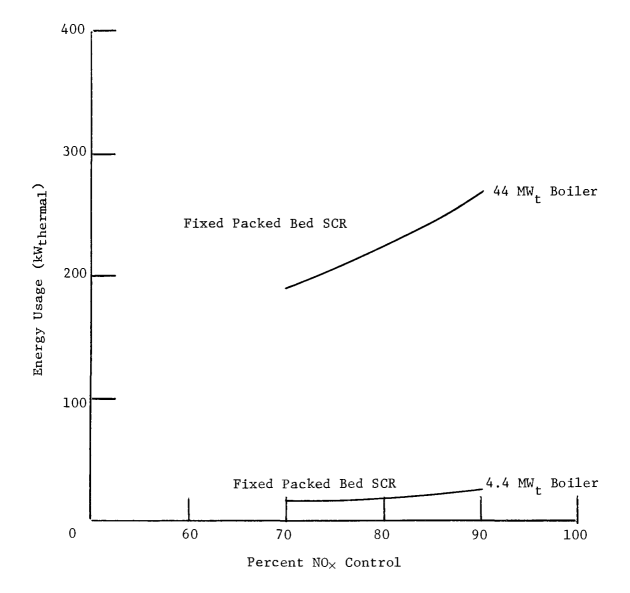


Figure 5.1.3-2. Energy usage of NO_X control systems for distillate oil-fired boilers.

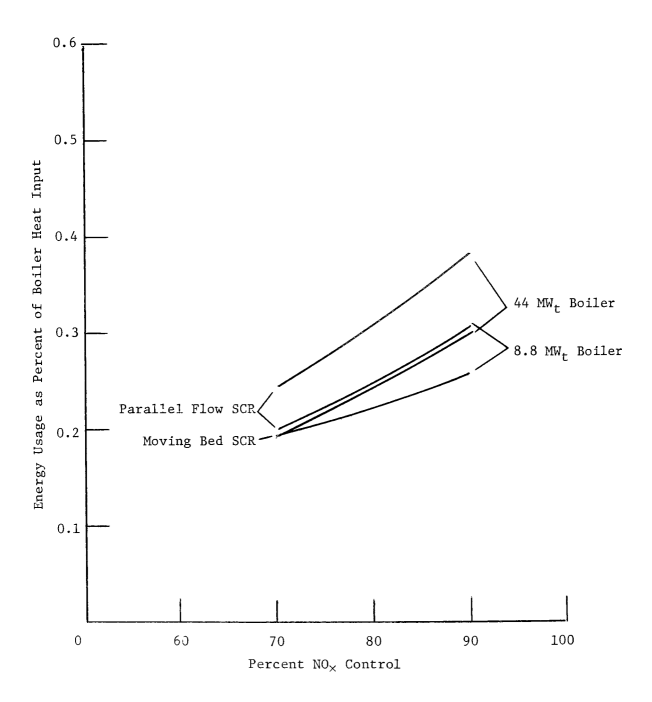
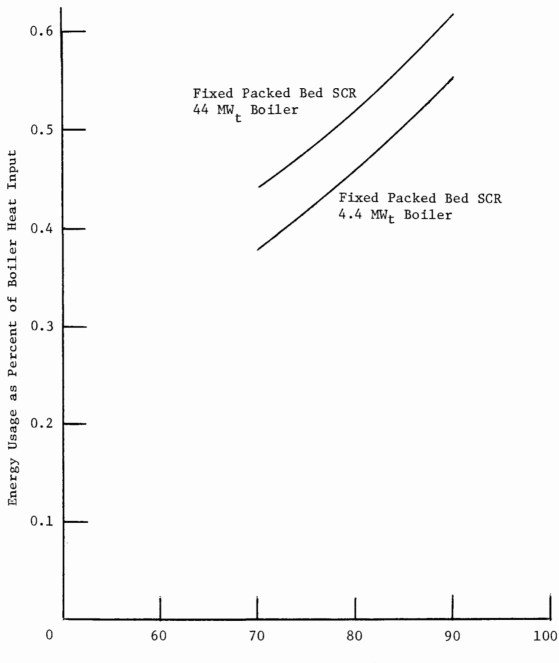


Figure 5.1.3-3. Energy usage of NO_X control systems applied to residual oil-fired boilers as percent of boiler heat input.



Percent NO_X Control

Figure 5.1.3-4. Energy usage of $\rm NO_{\times}$ control systems applied to distillate oil-fired boilers as percent of boiler heat input.

when comparing the two fuels since the size of the standard boilers is different by an order of magnitude. It can be said then, for the residual oil case, the moving bed systems are less energy intensive than parallel flow systems due to the moving beds¹ lower pressure drop across the length of the reactor.

It is not clear as to whether or not these systems have been optimized with respect to energy usage. The technology is relatively new and problem solving efforts are probably directed toward improving reliability and operability. It does seem possible that there is an optimum catalyst size and reactor volume that would minimize the pressure drop. Another potential method of lowering the pressure drop is to minimize the excess air. This reduces both the required reactor volume and the ΔP . It is likely that a boiler equipped with low NO_x burners will utilize low excess air for NO_x control.

 $\rm NH_3$ dilution by air instead of steam might possibly use less energy. There is, however, a safety aspect to consider since some air/ $\rm NH_3$ mixtures (<20:1) are explosive.

The energy impact of FGT controls applied to modified or reconstructed facilities (retrofit application) will be the same or greater than that for new facilities. If flue gas can be taken from the economizer of the existing boiler at 350-400°C and returned upstream of any existing heat exchange equipment, then there will be no additional energy impact.

If the flue gas is only available at a lower temperature (<350-400°C) then a heater will be required. The energy impact of the heater will depend on the temperature of the flue gas. If the temperature is that of the outlet gas of the standard boilers (approximately 180°C) calculations indicate that energy requirement would be more than tripled even if heat exchange equipment is used to recover 85% of the energy supplied by the heater. The heater will probably be oil-fired for ease of control.

These results indicate that, on retrofit applications, there is a considerable energy incentive to obtain the flue gas at the necessary reaction temperature in order to avoid flue gas heating. Other energy impacts would be the same as those for new facilities.

5.1.4 Energy Impact of Controls for Natural Gas-Fired Boilers

This subsection presents the results of energy and material balance for natural gas-fired industrial boilers. For new facilities two standard boilers and one NO_{χ} FGT system is considered. Results are presented for two levels of control.

The data presented is the result of several calculations. First, material balances were performed and the necessary equipment sized. Then, knowing the equipment size and flow rates, it was possible to calculate energy usage for each process step.

The results of these calculations are presented in Table 5.1.4-1. Both thermal energy requirements and energy requirements as a percentage of boiler heat input are shown. The candidate system for natural gas-fired boilers is fixed packed bed SCR. For the calculations, it is assumed that flue gas is available from the boiler economizer at 375°C and can be returned upstream of the air heater. Therefore, no energy is necessary for flue gas heating.

The data appearing in Table 5.1.4-1 is summarized in Table 5.1.4-2 and plotted in Figures 5.1.4-1 and 5.1.4-2. Figure 5.1.4-1 presents total energy usage and Figure 5.1.4-2 shows the energy usage as a percent of boiler heat input.

There are some areas of potential energy usage reduction. The catalyst particle size and reactor volume may be optimized to minimize reactor pressure drop. Reduction of excess air may also reduce the pressure drop and this may be standard practice on boilers with low NO_x burners. It may be more

TT IT TET		System						
Hea	Standard boild input	er	Type and level	Control efficiency	Energy			% increase in energy use over
MWt	(MBtu/hr)		of control	%	types	MWt	(MBtu/yr)	uncontrolled boiler
4.4	(15)	Package Firetube Scotch	SCR Fixed Packed Bed Moderate	70	Electrical Steam	0.0108 0.00106	(0.0369) (0.00363)	0.27
			SCR Fixed Packed Bed Stringent	90	Electrical Steam	0.0173 0.00133	(0.0590) (0.00455)	0.42
44	(150)	Package Watertube	SCR Fixed Packed Bed Moderate	70	Electrical Steam	0.123 0.0110	(0.421) (0.0345)	0.30
			SCR Fixed Packed Bed Stringent	90	Electrical Steam	0.203 0.0142	(0.692) (0.0483)	0.49

TABLE 5.1.4-1. ENERGY CONSUMPTION FOR NO_× FGT CONTROL TECHNIQUES FOR NATURAL GAS-FIRED BOILERS

TABLE 5.1.4-2.	SUMMARY	OF	ENERGY	REQUIREMENTS	FOR	NATURAL
	GAS-FIRE	DE	BOILERS			

	Nat Total thermal kW	ural gas % of boiler heat input
4.4 MW _t Boiler Fixed Packed Bed <u>SCR</u>		
90% Removal	19	0.42
70% Removal	12	0.27
44 MW _t Boiler Fixed Packed Bed SCR		
90% Removal	217	0.49
70% Removal	134	0.30

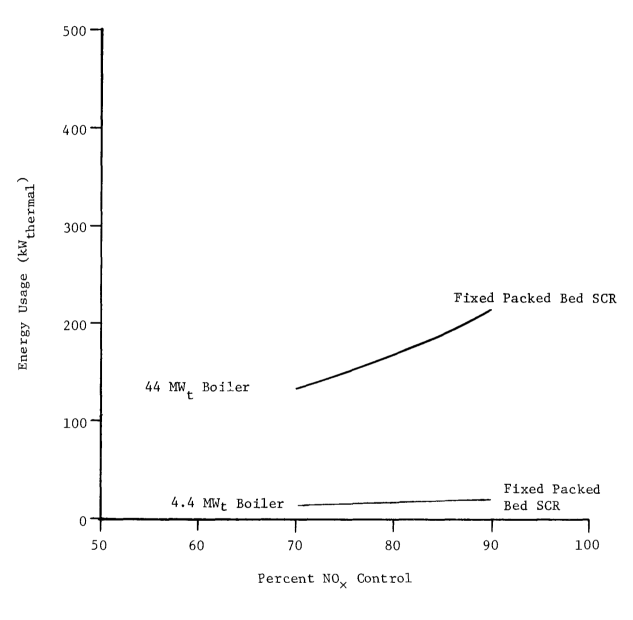


Figure 5.1.4-1. Energy usage of $\rm NO_X$ control systems for natural gas-fired standard boiler.

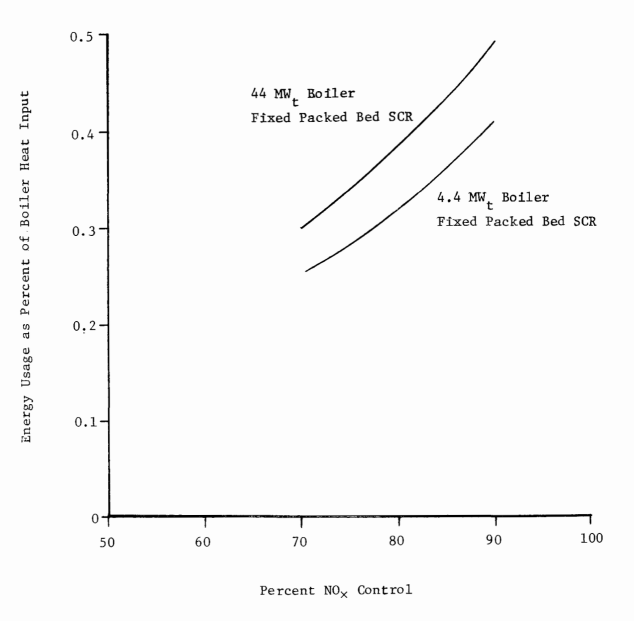


Figure 5.1.4-2. Energy usage of $\rm NO_X$ control systems as percent of boiler heat input.

energy efficient to use air instead of steam for NH_3 dilution, however, there is an explosion hazard with some air: NH_3 mixtures (<20:1).

The energy impact of FGT controls applied to modified or reconstructed facilities (retrofit application) will be the same or greater than that for new facilities. If flue gas can be taken from the economizer of the existing boiler at 350-400°C and returned upstream of any existing heat exchange equipment, then there will be no additional energy impact.

If the flue gas is only available at a lower temperature (<350-400°C) then a heater will be required. The energy impact of the heater will depend on the temperature of the flue gas. If the temperature is that of the outlet gas of the standard boilers (approximately 180°C) calculations indicate that energy requirement would be more than tripled even if heat exchange equipment is used to recover 85% of the energy supplied by the heater. The heater will probably be oil-fired for ease of control.

These results indicate that, on retrofit applications, there is a considerable energy incentive to obtain the flue gas at the necessary reaction temperature in order to avoid flue gas heating. Other energy impacts would be the same as those for new facilities.

5.2 NO_X/SO_X SYSTEMS

5.2.1 Introduction

This section considers the energy impacts associated with applying the UOP NO_X/SO_X FGT system to three industrial boilers. The combinations analyzed are presented in Table 5.2.1-1.

The NO_{χ}/SO_{χ} system has several more energy inputs than the NO_{χ} only systems; however, much of this energy is recovered by the air preheater resulting in an energy credit. The areas of energy utilization are shown in Table 5.2.1-2.

NO _× /SO _×			Control Level		
System	Boiler	Fuel*	NO×	S0 _×	
UOP	Pulverized Coal	LSW HSE	80	85	
UOP	Underfeed Stoker	LSW HSE	80	85	
UOP	Oil-Fired	Residual Oil	80	85	

TABLE 5.2.1-1. NOX/SOX FGT/BOILER COMBINATIONS ANALYZED FOR ENERGY IMPACT

* LSW = Low sulfur western coal (0.6% S)
HSE = High sulfur eastern coal (3.5% S)

TABLE 5.2.1–2. AREAS OF ENERGY UTILIZATION IN THE $\rm NO_X/SO_X$ FGT SYSTEM

Process Step	Type of Energy Consumed				
Reactor Draft Loss (Fan)	Electrical				
Liquid NH_3 Transfer (Pump)	Electrical				
NH_3 Vaporization (Vaporizer)	Steam				
NH ₃ Dilution	Steam				
Naphtha Reformer	Electrical, Steam, Fuel				
Compressor/Gasholder	Electrical				
H ₂ SO ₄ Plant	Electrical, Steam				

For each case, a heat and material balance is performed and these are contained in Appendices 4 and 5. These are used to size the equipment and determine the energy requirements. These requirements are listed in tabular form in each section and summarized. Since only one removal level is considered, the energy usage is not plotted against removal level as in the NO_v -only section.

5.2.2 Energy Impact of NO_X/SO_X Controls for Coal-Fired Boilers

Energy usage by these NO_X/SO_X applications is fairly evenly divided among three energy types: electrical, steam and fuel. These data are presented in Tables 5.2.2-1 and 5.2.2-2. Also shown in the tables are the heat credits for energy recovered by the air preheater.

The net energy usage by the NO_X/SO_X system is higher than that of the NO_X -only systems. When put on the basis of percent increase in energy over that of the uncontrolled boiler, the energy usage appears to be a function of the coal sulfur content, but not unit size. Removal level will also impact the energy usage; however, the magnitude of this impact is not known. Energy usage is summarized in Table 5.2.2-3 and plotted in Figure 5.2.2-1.

	Pulver	rized coal	Underfeed stoker			
Fuel	The rmal kW	% of boiler heat input	The r mal kW	% of boiler heat input		
Low Sulfur Western Coal	1,240	2.1	200	2.3		
High Sulfur Eastern Coal	11,200	7.7	680	7.7		

TABLE 5.2.2-3. SUMMARY OF ENERGY USAGE OF $\rm NO_{\times}/\rm SO_{\times}$ SYSTEMS APPLIED TO COAL-FIRED BOILERS

System						Knergy Consumption			
Standard Boiler Heat Input Type		Boiler Type	Coal Type	Type and Level	Control Bfficiency (% NO _x /SO _x)	Energy Types	Energy Consumed By Control Device		% Increase in Energy Use Over
MWt	(MBtu/hr)			of Control			MWt	(MBtu/hr)	Uncontrolled Boiler
58.6	200	Field Erected Watertube	High Sulfur Eastern Coal	SCR Parallel Flow Intermediate		Electrical	9.45	(32.25)	
		Pulverized Coal				Steam	2.7 9	(9.52)	7.7
						Fuel	5.18	(17.69)	
						Heat Credit	-(6.24)	-(21.3)	
58.6	200		Low Sulfur Western Coal	SCR Parallel Flow Intermediate	80/85	Electrical Steam Fuel Heat Credit	0.941 0.703 1.09 -(1.49)	(3.21) (2.40) (3.72) -(5.1)	2.1

TABLE 5.2.2-1. ENERGY CONSUMPTION FOR NO_X/SO_X FGT CONTROL TECHNIQUES FOR COAL FIRED BOILERS

System					Energy Consumption					
Hea ^{MW} t	Standard at Input (MBtu/hr)	Boiler Type	Coal Type	Type and Level of Control	Control Efficiency (% NO _X /SO _X)	Energy Types		y Consumed trol Device (MBtu/hr)	% Increase in Energy Use Over Uncontrolled Boiler	
8.8	8.8 30	Package Watertube Underfeed	High Sulfur Eastern Coal	SCR Parallel Flow Intermediate	80/85	Electrical Steam	0.428 0.416	(1.46)	7.7	
					Fuel	0.774	(2.64)			
						Heat Credit	-(0.935)	-(3.19)		
8.8	30			Low Sulfur Western Coal	SCR Parallel Flow Intermediate	80/85	Electrical	0.151	(0.52)	
			western ood		Steam	0.102	(0.348)	2.3		
						Fuel	0.163	(0.558)		
-						Heat Credit	-(0.217)	-(0.74)		

TABLE 5.2.2-2. ENERGY CONSUMPTION FOR NO_{\times}/SO_{\times} FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS

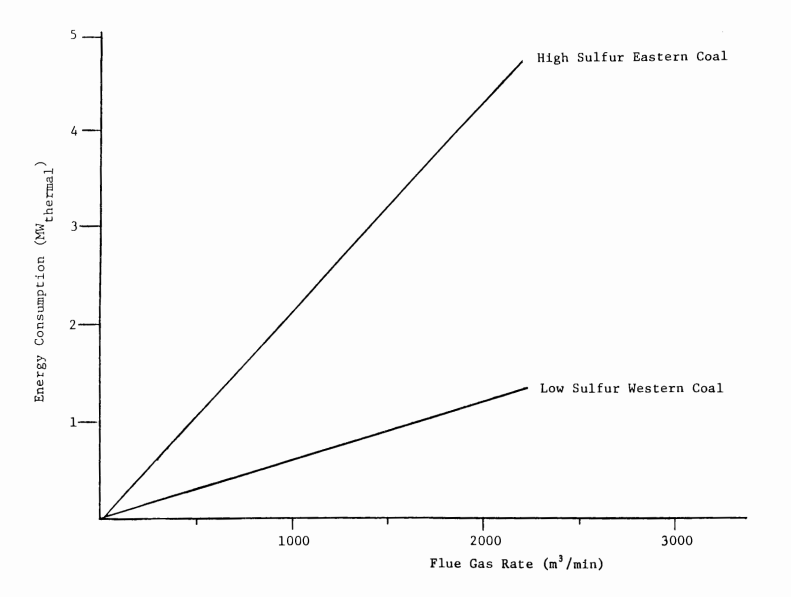


Figure 5.2.2-1. Energy consumption of parallel flow SCR NO_X/SO_X FGT systems for coal-fired boilers.

The energy impact of FGT controls applied to modified or reconstructed facilities (retrofit application) will be the same or greater than that for new facilities. If flue gas can be taken from the economizer of the existing boiler at 350-400°C and returned upstream of any existing heat exchange equipment, then there will be no additional energy impact.

If the flue gas is only available at a lower temperature (350-400°C) then a heater will be required. The energy impact of the heater will depend on the temperature of the flue gas. The energy used in heating the gas is not completely lost since the air preheater can recover about 85% of the energy supplied by the gas heater. The heater will probably be oil-fired for ease of control.

These results indicate that, on retrofit applications, there is an energy incentive to obtain the flue gas at the necessary reaction temperature in order to avoid flue gas heating. Other energy impacts would be the same as those for new facilities.

5.2.3 Energy Impact of NO_X/SO_X Controls for Oil-Fired Boilers

Three types of energy are utilized by these systems--electrical, steam and fuel. The amounts of each type are shown in Table 5.2.3-1. Energy consumption of each energy type is of the same order of magnitude. Also shown is the heat credit that is obtained by partially recovering heat from the energy inputs with the air preheater.

Since only one case is considered, the variables that affect the energy impact cannot be quantified. It can be stated qualitatively, however, that the primary variables that affect energy usage are oil sulfur content and control level (both NO_X and SO_X). The effect of fuel sulfur content was examined in the section on coal-fired applications.

System						Energy Consumption				
Hea Wt	Standard at Input (MBtu/hr)	Boiler Type	Oil Type	Type and Level of Control	Control Efficiency (Z NO _X /SO _X)	Energy Types		y Consumed trol Device (MBtu/hr)	% Increase in Energy Use Over Uncontrolled Boiler	
\$4	150	Package Watertube	Residual	SCR Parallel Flow Intermediate	80/85	Electrical	1.26	(4.29)		
						Steam	1.16	(3.96)	4.4	
						Fuel	2.16	(7.37)		
						Heat Credit	-(2.63)	-(8.98)		

TABLE 5.2.3-1. ENERGY CONSUMPTION FOR NO_{\times}/SO_{\times} FGT CONTROL TECHNIQUES FOR OIL FIRED BOILERS

The energy impact of FGT controls applied to modified or reconstructed facilities (retrofit application) will be the same or greater than that for new facilities. If flue gas can be taken from the economizer of the existing boiler at 350-400°C and returned upstream of any existing heat exchange equipment, then there will be no additional energy impact.

If the flue gas is only available at a lower temperature (<350-400°C) then a heater will be required. The energy impact of the heater will depend on the temperature of the flue gas. The energy used in heating the gas is not completely lost since the air preheater can recover about 85% of the energy supplied by the heater. The heater will probably be oil-fired for ease of control.

These results indicate that, on retrofit applications, there is an energy incentive to obtain the flue gas at the necessary reaction temperature in order to avoid flue gas heating. Other energy impacts would be the same as those for new facilities.

5.3 SUMMARY

All of the NO_X -only systems and cases required <1% of the total heat input to the boiler. By far, the item contributing the most to energy consumption was the flue gas fan which supplied the draft loss caused by the catalyst bed.

The parallel flow systems appear to use more energy than the moving bed systems; however, both are of the same order of magnitude. Within the accuracy of the calculations, the systems examined should be considered to have approximately similar energy impacts.

The NO_X/SO_X systems require 2-8% of the total heat input to the boiler. This is primarily due to the requirement for SO_2 workup. Although this requirement is higher than that for NO_X -only processes, it may be less than that for the combination of NO_X -only and FGD.

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- 3. Perry, Robert H. <u>Chemical Engineers Handbook</u>. 5th Edition. 1973. McGraw-Hill. pp. 5-52, 53.
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SECTION 6

ENVIRONMENTAL IMPACT OF CANDIDATES FOR BEST EMISSION CONTROL SYSTEMS

6.1 INTRODUCTION

The three best candidate control techniques identified in Section III are as follows:

SCR - Fixed Packed Bed SCR - Parallel Flow SCR - Moving Bed

These techniques have similar environmental concerns as they all utilize NH3 to reduce NO, to N₂. All are capable of achieving the levels of control considered in this study, although the applicability of a particular system is fuel dependent. While NO, reduction is the primary beneficial environmental impact of these systems, particulate removal is a secondary beneficial impact of the moving bed systems. The moving bed system will reduce the level of particulates in the flue gas by 70-80%.1,2 The particulates are embedded on the catalyst (rings or granules) as the catalyst moves downward through the reactor. The dirty catalyst is removed to a vibrating screen which separates the dust and the clean catalyst is then recycled to the top of the reactor. Pilot plant tests on the moving bed reactor have shown it capable of handling <1 g/Nm³ of particulates.³ The uncontrolled particulate levels in the flue gas from the pulverized coal (5-9 g/Nm^3), spreader stoker (3.5-6.3 g/nM^3), chaingrate and underfeed (both 1.4-2.4 g/Nm³) standard boilers are all greater than this figure. As a result, moving bed systems are not considered for application to the coal-fired standard boilers. The fixed packed bed system

cannot tolerate particulates so it is applied only to natural gas- and distillate oil-fired flue gas which have low particulate loadings (13 and 19 mg/Nm³, respectively, for the standard boilers). Conversely, the parallel flow system can tolerate full particulate loadings (up to 20 g/nM³)⁴ as the open passageways present unobstructed paths for particulates to pass through with little impaction on the catalyst surface.

There are some potential adverse environmental impacts of the selective catalytic reduction processes. First, the use of NH₃ as the gaseous reducing agent introduces the possibility of ammonia emissions. Commercial operations of the three reactor types on industrial and utility boilers have demonstrated emissions of <10 ppm NH₃ at the NH₃:NO_x mole ratio required for stringent level of control. These levels are shown graphically in Figures 6.1-1 through 6.1-3 on the following pages. (These plots are composites of the available commercial data.) The only data available on NH₃ emissions are from Japanese process vendors and these data indicate NH₃ emissions to be <10 ppm. This number may be optimistic, especially considering that currently there is no continuous monitoring technique for measuring NH₃ in the presence of SO_x. The data, therefore, represent spot measurements and not continuous data. It seems reasonable to assume that 10 ppm represents a minimum level of NH₃ emissions.

A potential environmental problem in commercial SCR operations is the formation of ammonium bisulfate, NH_4HSO_4 , or ammonium sulfate, $(NH_4)_2SO_4$. The presence of NH_3 , SO_3 , and H_2O in the hot flue gas leads to the formation of liquid NH_4HSO_4 upon cooling to approximately $180-220^{\circ}C$ by the following reaction.

$$\mathrm{NH}_{3}(g) + \mathrm{SO}_{3}(g) + \mathrm{H}_{2}\mathrm{O}(g) \stackrel{\neq}{\leftarrow} \mathrm{NH}_{4}\mathrm{HSO}_{4}(\ell) \tag{6-1}$$

This can create a plugging and corrosion problem in heat exchange equipment, particularly for boilers burning medium- or high-sulfur fuels. Further cooling to about 190°C precipitates the formation of solid ammonium sulfate $[(NH_4)_2SO_4]$ by the following reaction.

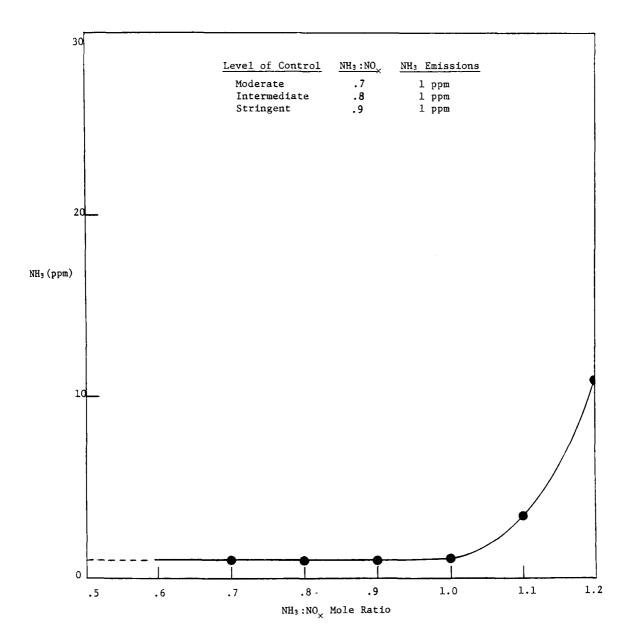


Figure 6.1-1. NH₃ Emissions - Fixed Packed Bed Reactor. 5,6,7

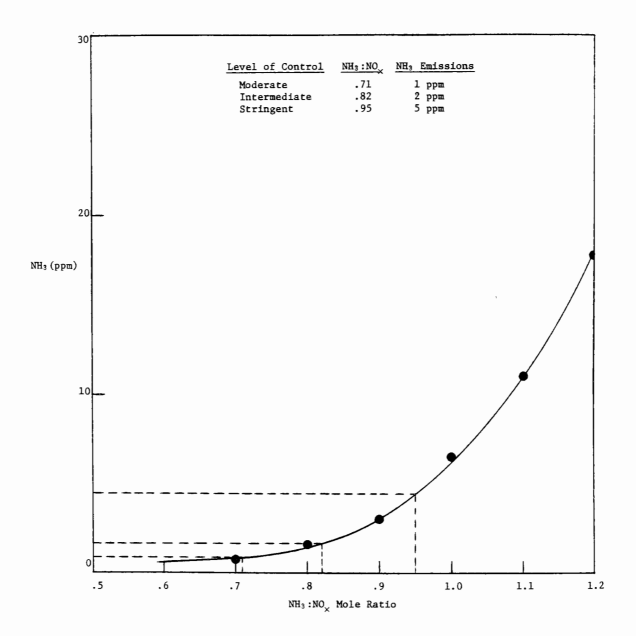


Figure 6.1-2. NH₃ Emissions - Parallel Flow Reactor.^{8,9,10,11,12}

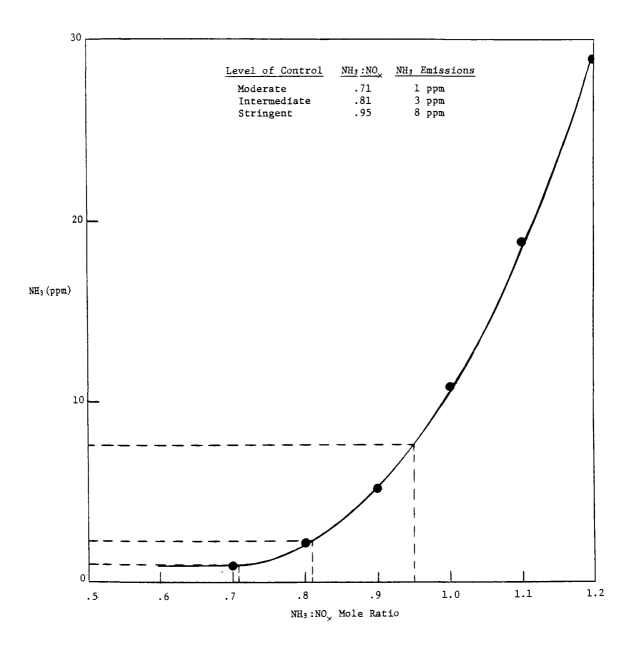


Figure 6.1-3. NH₃ Emissions - Moving Bed Reactor. 13,14,15,16,17

$$\mathrm{NH}_{4}\mathrm{HSO}_{4}(l) + \mathrm{NH}_{3}(g) \stackrel{\tau}{\leftarrow} (\mathrm{NH}_{4})_{2}\mathrm{SO}_{4}(s) \tag{6-2}$$

The impact of the solid sulfate and liquid bisulfate on downstream particulate collection equipment and FGD systems is unknown at present and is currently being investigated by the EPA and others. It is speculated that minor, if any, amounts of these sulfates will be emitted to the atmosphere in situations where particulate control equipment exists downstream of the NO_X control system.

The final environmental concern of the SCR processes is disposed of spent catalyst. Catalysts such as titanium dioxide (TiO_2) and vanadium pentoxide (V_2O_5) are probably recycled due to their high cost. To date, virtually no catalyst has been used commercially yet for over 10,000 hours, and, as a result, there has been no commercial experience on the treatment of spent catalyst. Reprocessing or disposal of spent catalyst will most likely be carried out by the catalyst vendor.²⁷ This question is not currently addressed in literature. Another potential problem related to catalysts is that of catalyst erosion, especially with the moving bed systems. Catalyst erosion may generate small particulates which may present a stack fume problem if particulate control devices are not present or not effective at removing the catalyst particles. No problems of this nature have been reported at this time.

6.2 ENVIRONMENTAL IMPACTS OF CONTROLS FOR COAL-FIRED BOILERS

6.2.1 Air Pollution

The emission rates for primary and secondary pollutants are presented in Tables 6.2.1-1 through 6.2.1-12 on the following pages. There are three tables for each of the 4 coal-fired standard boilers. Each table is broken down according to coal type (high sulfur eastern and low sulfur western) and control level (uncontrolled, moderate, intermediate, and stringent). For each entry the impact on the primary pollutant, NO_x , is shown. Then, the adverse impact of the secondary pollutant, NH_3 , is given for each case.

6-б

		Standard Hea	Boiler: at Rate: Coal:	Pulveriz 200 MBtu High Sul				
			NOv	Particu	lates	N	Н з	
Control level	Type of control	g/s (1b/hr)	ng/J (1b/MBtu)	g/s (1b/hr)	ng/J (1b/MBtu)	g/s (1b/hr)	ng/J (1b/MBtu)	Bisulfate
Uncontrolled	-	19.2 (152.46)	328.0 (.762)	181.2 (1436.5)	3090 (7.18)	0	0	0
Moderate-70%	SCR-Parallel Flow	5.77 (45.7)	98.2 (.229)	Negligi	ole Effect	.0154 (.122)	.261 (.000608)	See Text
Stringent-90%	SCR-Parallel Flow	1.92 (15.2)	32.8 (.0762)	Negligi	ole Effect	.0767 (.608)	1.31 (.00304)	See Text
	. AIR POLLUTION IMPA	Standard		Pulveriz 200 MBtu	zed Coal			
			NOX	Parti	culates		NH 3	
Control level		g/s	ng/J	g/s	ng/J	g/s	ng/J	
	Type of control	(1b/hr)	(1b/MBtu)	(1b/hr)	(1b/MBtu)	(1b/hr)	(1b/MBtu)	Bisulfate
Uncontrolled	Type of control	(1b/hr) 23.7 (187.56)				(1b/hr) 0	0.	Bisulfate 0
Uncontrolled Moderate-70%	Type of control - SCR-Parallel Flow	23.7	(1b/MBtu) 403.0	(1b/hr) 113.5 (900.3)	(1b/MBtu) 1936		(1b/MBtu)	

(.0938)

(.632)

(.00316)

(18.8)

7

TABLE 6.2.1-1. AIR POLLUTION IMPACTS FROM BEST NO_× FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS^{1,2,8-17}

				150 MBtu	r Stoker 1/hr lfur Easter	n		
Control level	Type of control	g/s (1b/hr)	NO _X ng/J (1b/MBtu)	Partia g/s (1b/hr)	culates ng/J (lb/MBtu)	N (1b/hr)	H ₃ ng/J (1b/MBtu)	Bisulfate
Uncontrolled	_	12.0 (95.4)	273.0 (.636)	111.0 (876.4)	2512 (5.84)	0	0	0
Intermediate-80%	SCR-Parallel Flow	2.41 (19.1)	54.7 (.127)	Negligi	ble Effect	.0266 (.211)	.604 (.00140)	See Text

TABLE 6.2.1-3. AIR POLLUTION IMPACTS FROM BEST NO FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS^{1,2,8-17}

TABLE 6.2.1-4. AIR POLLUTION IMPACTS FROM BEST NO X FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS 1,2,8-17

			Standard B Heat		Spreader St 150 MBtu/hn Low Sulfur	r		
Control level	Type of control	g/s (1b/hr)	NO _x ng/J (1b/MBtu)	Part g/s (lb/hr)	iculates ng/J (lb/MBtu)	g/s (1b/hr)	NH ₃ ng/J (1b/MBtu)	Bisulfate
Uncontrolled	-	14.8 (117.15)	336.0 (.781)	69.1 (548.3)	1572 (3.66)	0	0	0
Intermediate-80%	SCR-Parallel Flow	2.95 (23.4)	67.2 (.156)	Neglig	ible Effect	.0273 (.217)	.622 (.00145)	See Text

		Sta	ndard Boil Heat Ra Co	te: 75	ingrate MBtu/hr h Sulfur E	astern		
		N	0 _x	Partic	ulates	NH3	L	
Control level	Type of control	g/s (1b/hr)	ng/J (1b/MBtu)	g/s (1b/hr)	ng/J (1b/MBtu)	g/s (1b/hr)	ng/J (1b/MBtu)	Bisulfate
Uncontrolled	-	6.02 (47.7)	273.0 (.636)	21.2 (168.5)	966.0 (2.25)	0	0	0
Moderate-70%	SCR-Parallel Flow	1.80 (14.3)	82.0 (.191)	Negligibl	e Effect	.00662 (.0525)	.301 (.000700)	See Text
Intermediate-80%	SCR-Parallel Flow	1.20 (9.54)	54.7 (.127)	Negligib1	e Effect	.0132 (.105)	.602 (.00140)	See Text
Stringent-90%	SCR-Parallel Flow	.602 (4.77)	27.3 (.0636)	Negligibl	e Effect	.0331 (.262)	1.50 (.00350)	See Text

TABLE 6.2.1-5. AIR POLLUTION IMPACTS FROM BEST NO_× FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS^{1,2,8-17}

TABLE 6.2.1-6. AIR POLLUTION IMPACTS FROM BEST NO_× FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS^{1,2,8-17}

		1	rd Boiler: Heat Rate: Coal:	75 MBt		rn		
Control level	Type of control	N g/s (1b/hr)	O _X ng/J (1b/MBtu)	Partic g/s (lb/hr)	culates ng/J (1b/MBtu)	NI 	H ₃ ng/J (1b/MBtu)	Bisulfate
Uncontrolled	-	7.40 (58.65)	336.0 (.782)	13.3 (105.6)	605.0 (1.41)	0	0	0
Moderate-70%	SCR-Parallel Flow	2.22 (17.6)	101.0 (.235)	Negligib]	le Effect	.00683 (.0542)	.311 (.000723)	See Text
Intermediate-80%	SCR-Parallel Flow	1.48 (11.7)	67.2 (.156)	Negligib]	le Effect	.0137 (.108)	.621 (.00145)	See Text
Stringent-90%	SCR-Parallel Flow	.740 (5.87)	33.6 (.0782)	Negligib]	le Effect	.0342 (.271)	1.55 (.00361)	See Text

		-	Boiler: at Rate: Coal:	Underfee 30 MBtu/ High Sul		n		
Control level	Type of control	g/s (1b/hr)	NO _x ng/J (1b/MBtu)	Partic g/s (1b/hr)	ulates ng/J (1b/MBtu)	<u>NH</u> g/s (1b/hr)	ng/J (1b/MBtu)	Bisulfate
Uncontrolled	-	2.40 (19.05)	273.0 (.635)	8.49 (67.31)	965.0 (2.24)	0	0	0
Intermediate-80%	SCR-Parallel Flow	.480 (3.81)	54.6 (.127)	Negligit	le Effect	.00529 (.0419)	.601 (.00140)	See Text

TABLE 6.2.1-7. AIR POLLUTION IMPACTS FROM BEST NO_× FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS^{1,2,8-17}

TABLE 6.2.1-8. AIR POLLUTION IMPACTS FROM BEST NO FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS 1,2,8-17

		Standard Hea	t Rate:	Underfeed 30 MBtu/h Low Sulfu	4			
Control level	Type of control	l g/s (1b/hr)	NO _X ng/J (1b/MBtu)	Partic g/s (lb/hr)	ulates ng/J (1b/MBtu)	l g/s (1b/hr)	<u>NH₃</u> ng/J (1b/MBtu)	Bisulfate
Uncontrolled	-	2.95 (23.40)	335.0 (.780)	5.31 (42.12)	604.0 (1.40)	0	0	0
Intermediate~80%	SCR-Parallel Flow	.590 (4.68)	67.1 (.156)	Negligib	le Effect	.00544 (.0431)	.618 (.00144)	See Text

		Standa	ard Boilen Heat Rate Coal	e: 200 M	erized Coa 1Btu/hr Sulfur Ea					
Control level	Type of control	g/s (1b/hr)	NO _x ng/J (1b/MBtu)	g/s (1b/hr)	50 ₂ ng/J (1b/MBtu)	Parti g/s (lb/hr)	culates ng/J (1b/MBtu)	g/s (lb/hr)	NH ₃ ng/J (1b/Mßtu)	Bisulfate
Uncontrolled	-	19.2 (152.5)	328.0 (0.762)	142.0 (1127.0)	2423.0 (5.64)	181.0 (1437.0)	3090.0 (7.18)	0	0	0
Intermediate (80% NO _×) (85% SO ₂)	SRC-Parallel Flow	3.85 (30.5)	65.6 (0.153)	21.3 (169.0)	363.0 (0.865)	Negligible	Effect	0.307 (2.43)	5.22 (0.0122)	See Text

TABLE 6.2.1-9. AIR POLLUTION IMPACTS FROM BEST NO_×/SO_× FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS

TABLE 6.2.1-10. AIR POLLUTION IMPACTS FROM BEST NO_{\times}/SO_{\times} FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS

		Stan		te: 200	verized Co MBtu/hr Sulfur We	1				
Control level	Type of control	g/s (1b/hr)	NO _x ng/J (1b/MBtu)	g/s (1b/hr)	50 ₂ ng/J (1b/MBtu)	Parti g/s (15/hr)	culates ng/J (1b/MBtu)	g/s (1b/hr)	NH ₃ ng/J (1b/MBtu)	Bisulfate
Uncontrolled	_	23.7 (187.6)	403.0 (0.938)	30.0 (237.6)	511.0 (1.19)	113.5 (900.3)	1936.0 (4.50)	0	0	0
Intermediate (80% NO _×) (85% SO ₂)	SRC-Parallel Flow	4.73 (37.5)	80.6 (0.188)	4.49 (35.6)	7.65 (0.178)	Negligible	e Effect	0.318 (2.52)	5.42 (0.0126)	See Text

		Star		te: 30	-					
Control level	Type of control	g/s (1b/hr)	NO _x ng/J (1b/MBtu)	g/s (1b/hr)	502 ng/J (1b/MBtu)	Partic g/s (lb/hr)	culates ng/J (1b/MBtu)	g/s (1b/hr)	NH ₃ ng/J (1b/MBtu)	Bisulfate
Uncontrolled	-	2.40 (19.05)	273.0 (0.635)	21.3 (168.9)	2421.0 (5.63)	8.49 (67.31)	965.0 (2.24)	0	0	0
Intermediate (80% NO _×) (85% SO ₂)	SRC-Parallel Flow	0.481 (3.81)	54.6 (0.127)	3.20 (25.3)	363.0 (0.845)	Negligibl	e Effect	0.0529 (0.419)	6.01 (0.0140)	See Text

TABLE 6.2.1-11. AIR POLLUTION IMPACTS FROM BEST NO_×/SO_× FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS

TABLE 6.2.1-12. AIR POLLUTION IMPACTS FROM BEST NO /SO FGT CONTROL TECHNIQUES FOR COAL-FIRED BOILERS

		Stan	dard Boil. Heat Ra Co	te: 30	erfeed Sto MBtu/hr Sulfur Wo					
Control level	Type of control	g/s (1b/hr)	NO _x ng/J (1b/MBtu)	g/s (1b/hr)	50 ₂ ng/J (1b/MBtu)	<u>Parti</u> g/s (1b/hr)	culates ng/J (1b/MBtu)] g/s (1b/hr)	NH ₁ ng/J (1b/MBtu)	Bisulfate
Uncontrolled	-	2.95 (23.40)	335.0 (0.780)	4.49 (35.6)	510.0 (1.19)	5.31 (42.12)	604.0 (1.40)	0	0	0
Intermediate (80% NO _x) (85% SO ₂)	SRC-Parallel Flow	0.590 (4.58)	67.1 (0.156)	0.674 (5.34)	76.8 (0.179)	Negligibl	e Effect	0.0544 (0.431)	6.18 (0.0144)	See Text

Table 6.2.1-13 shows the uncontrolled NO_{\times} emissions for each standard boiler and the SIP level for each fuel. One can see that a majority of the uncontrolled emissions are less than SIP allowable levels. For coal, the worst case is the pulverized coal boiler burning low sulfur western coal. To meet the SIP control level the degree of removal required is as follows.

% reduction =
$$\frac{.938 - .7}{.938} \times 100\% = 25\%$$

This level of control is easily obtained via combustion modifications,¹⁸ therefore, this study does not address the SIP control level.

Also, in Tables 6.2.1-1 through 6.2.1-12 the secondary pollutant bisulfate is not quantified, for several reasons. First, kinetic and thermodynamic data for the reaction

$$NH_3(g) + SO_3(g) + H_2O(g) \rightarrow NH_4HSO_4(l)$$

have not yet been evaluated. Therefore, the extent of reaction cannot be determined for the residence time of the flue gas in the duct. Second, bisulfate emissions are not constant since they are at a high level during soot blowing and at a lower level during other periods. Finally, it is not known how much of the bisulfate is caught by downstream particulate removal equipment (assuming that the equipment is present) and how much is emitted. A hypothetical calculation can be made for the case of the pulverized coal standard boiler burning high sulfur eastern coal (Table 6.2.1-1) equipped with an SCR parallel flow control system operating at the stringent level of control. The amount of bisulfate formed is as follows.

	Uncontrolled standard boilers									
SIP control	Natural gas	Distillate oil	Residual oil	Coal type	Underfeed stoker	Chaingrate	Spreader stoker	Pulverized coal		
.2	.175				····					
.3		.159	.400							
				High S Eastern	.635	.636	.636	.762		
.7				Low S Eastern	.545	. 544	.543	.653		
				Low S Western	.780	.782	.781	.938		
	.2 .3	control gas .2 .175 .3	control gas o11 .2 .175 .3 .159	control gas oil oil .2 .175 .3 .159 .400	SIP Natural Distillate Residual Coal control gas oil oil type .2 .175 .3 .159 .400 .3 .159 .400 High S Eastern Low S Eastern .7 Low S Eastern	SIP Natural Distillate Residual Coal Underfeed control gas oil oil type stoker .2 .175 .3 .159 .400 .3 .159 .400 High S Eastern .635 .7 Eastern .545 Low S Low S	SIP Natural Distillate Residual Coal Underfeed Chaingrate .2 .175 .3 .159 .400 High S Eastern .635 .636 .7 Eastern .545 .544 Low S Low S .545 .544	SIP Natural Distillate Residual Coal Underfeed Chaingrate Spreader .2 .175 .3 .159 .400 High S Eastern .635 .636 .636 .7 Eastern .545 .544 .543 Low S Low S Low S .545 .544 .543		

.

TABLE 6.2.1-13. NO $_{\rm X}$ EMISSION LEVELS AND SIP CONTROL LEVELS (All values in 1b/10⁶ Btu)

Assuming all $NH_3 \rightarrow NH_4HSO_4$:

wt. bisulfate =
$$\left(.0767 \frac{\text{g NH}_3}{\text{s}}\right) \left(\frac{\text{g-mole NH}_3}{17.0 \text{g NH}_3}\right) \left(\frac{\text{g-mole NH}_4 \text{HSO}_4}{\text{g-mole NH}_3}\right)$$

= $\left(\frac{115 \text{g NH}_4 \text{HSO}_4}{\text{g-mole NH}_4 \text{HSO}_4}\right)$
= $\left(.519 \frac{\text{g NH}_4 \text{HSO}_4}{\text{s}}\right) = \left(19 \frac{\text{tons}}{\text{year}}\right)$

There are beneficial aspects of this reaction. The bisulfate formation ties up SO_3 which is more hazardous than SO_2 and is difficult to catch with FGD.¹⁹ If the bisulfate can be collected adequately and disposed of safely, an overall improvement could be achieved.

The fate of bisulfate formed in this manner has not been completely resolved and is currently an aspect of NO, control receiving much attention. In Japan there have been problems at some installations of precipitation of the bisulfate or sulfate on elements of regenerative air preheaters and tubes of tubular air preheaters. This deposit can be removed by periodically water-washing the air preheater.²⁰ Bisulfate or sulfate particulates that pass through the air preheater may be collected by downstream particulate control or FGD equipment if such equipment exists. The collection efficiency of particulate control equipment cannot be determined quantitatively without knowing the particle size and resistivity (for ESP's) or the K-factor and concentration (for baghouses). These data are not currently known for these compounds, however, it is reasonable to expect that some fraction of the particles will be collected.²¹ Similarly, the collection efficiency for an FGD unit has not been examined. Where neither particulate control nor FGD equipment exist, there may be stack emissions of sulfates. An FGD system may also absorb NH3 emitted by an SCR system, however, the removal cannot be determined from the information currently available.

To reduce the adverse environmental effects, improved combustion control utilizing less 0_2 minimizes the formation of NO_{χ} . It also minimizes formation of SO_3 which is necessary for ammonium bisulfate formation. Careful operation of the FGT system should keep the NH₃ injection ratio as low as possible to minimize NH₃ emissions and bisulfate formation. Also, careful operation of downstream heat exchange equipment to keep the flue gas above the acid (SO_3) dew point is required. The use of corrosion-resistant material in any heat exchanger is advisable where NH₄HSO₄ deposits are probable.²² A multitube type heat exchanger with the tubes placed vertically is a possible configuration to prevent bisulfate deposits from causing problems. Any bisulfate liquid formed in the tube will drip downward as long as the temperature of the tube is kept above the melting point of bisulfate.²³ It will be necessary to design the exchanger out of corrosion resistant materials.

6.2.2 Water Pollution

There are no water streams that are associated with NO_{χ} -only SCR systems, however, there is one potential source of water pollution. In some Japanese installations NH_4HSO_4 deposits are removed from the air preheater by water washing. The blowdown from this operation will contain both ammonium and sulfate ions which, if not treated, present a water pollution source. Since the amounts of NH_4HSO_4 and water are not known it is possible to estimate the concentration or flow additional of this potential source.

6.2.3 Solid Waste

The only potential solid waste associated with this system is spent catalyst. Presently, the life of SCR catalysts is 1-2 years and the topic of recycling is not addressed in the literature. Since, to date, virtually no commercial SCR units have operated long enough to require catalyst replacement, there is no commercial experience on the treatment of spent catalyst.²⁷ With the high cost of some of the catalysts, recycling seems to be warranted. Recycling is feasible where the spent catalyst support is composed of valuable

materials such as titanium, stainless steel, or possibly a ceramic material. Alumina catalyst supports probably do not warrant recycling unless required for environmental reasons. These catalysts can probably be disposed of in the same manner as other industrial cataylsts.

The amount of catalyst that must be recycled or disposed of is one reactor volume since replacement involves total catalyst replacement. The actual frequency of catalyst replacement is unique to each specific process, however, catalyst lifetimes are typically one or more years.

Few process vendors have published their catalyst formulations since the field is very competitive at the present time. Base metal oxides are typically used.²⁴ The environmental impact of catalyst disposal will depend on what materials and compounds are involved.

6.2.4 Other Environmental Impacts

The only other environmental impact is a secondary impact. NH_3 is commonly made from natural gas and its consumption is considered a secondary environmental impact. This impact is quantified in a report on the impact of NO_{χ} regulations on the NH_3 industry. The report was prepared by TVA under contract to EPA-IERL. Other adverse environmental impacts (noise, thermal pollution, electrical discharges, etc.) are not present with SCR systems.

6.2.5 Environmental Impact on Modified and Reconstructed Facilities

The environmental impacts of a new facility and a retrofitted facility should be similar. There is not enough difference between new and retrofit systems to indicate that environmental impacts would be significantly different with retrofitted systems.

6.3 ENVIRONMENTAL IMPACTS OF CONTROLS FOR OIL-FIRED BOILERS

6.3.1 Air Pollution

Emission rates for primary and secondary pollutants are presented in Tables 6.3.1-1 and 6.3.1-2 on the next pages. The tables are organized by fuel type (residual and distillate oil); control level (uncontrolled, moderate, and stringent); and type of control (residual - SCR moving bed and SCR parallel flow; distillate - SCR fixed packed bed). The impact on the primary pollutant, NO_x , is given for each case. Also, the moving bed's impact on particulates of residual oil is shown. The adverse impact of the secondary pollutant, NH_3 , is given for each entry.

Table 6.2.1-13 shows that only the residual oil-fired flue gas has uncontrolled NO_{\times} emissions greater than the SIP control level. To achieve the SIP level of control the removal required is as follows.

% reduction =
$$\frac{0.4-0.3}{0.4} \times 100\%$$

= 25%

This control level is readily achieved by combustion modifications; hence, FGT to achieve the SIP control level is not examined for oil-fired boilers.

In Table 6.3.1-1 the secondary pollutant bisulfate is not quantified. This is due to a lack of developed kinetic and thermodynamic data to predict the extent of reaction. Also, removal levels are not constant since the degree of downstream particulate removal is uncertain. However, one can see that the bisulfate problem is worse for residual oil than for distillate oil because ther ' is more SO₃ available for reaction. Bisulfate is formed by a one-to-one reaction between NH₃, SO₃ and H₂O.

$$\mathrm{NH}_3(g) + \mathrm{SO}_3(g) + \mathrm{H}_2\mathrm{O}(g) \xrightarrow{} \mathrm{NH}_4\mathrm{HSO}_4(\ell)$$

Standard boiler				NO		Particulates		NH 3		
eat rate MBtu/hr)	Туре	Control level	Type of control	g/s (lb/hr)	ng/J (1b/MBtu)	g/s (1b/hr)	ng/J (1b/MBtu)	g/s (lb/hr)	ng/J (1b/MBtu)	Bisulfate
150 R	Residual	Uncontrolled	-	7.57 (60.0)	172.0 (.400)	4.16 (33.0)	94.6 (.220)	0	0	0
		Moderate - 70%	SCR Moving Bed	2.27 (18.0)	51.6 (.120)	1.25 (9.90)	28.4 (.0660)	.00957 (.0759)	.218 (.000506)	See Text
		Stringent - 90%	SCR Moving Bed	.757 (6.00)	17.2 (.0400)	1.25 (9.90)	28.4 (.0660)	.0766 (.607)	1.74 (.00405)	See Text
		Moderate - 70%	SCR Parallel Flow	2.27 (18.0)	51.6 (.120)	Neglig	ible Effect	.00957 (.0759)	.218 (.000506)	See Text
		Stringent - 90%	SCR Parallel Flow	.757 (6.00)	17.2 (.0400)	Neglig	ible Effect	.0479 (.380)	1.09 (.00253)	See Text
30 R.	Residual	Uncontrolled	-	2.02 (16.0)	229.0 (0.533)	.580 (4.60)	65.9 (.153)	0	0	0
		Moderate - 70%	SCR Moving Bed	0.606 (4.80)	68.7 (0.160)	.174 (1.38)	20.1 (.0459)	.00201 (.0160)	.229 (.000533)	See Text
		Stringent - 90%	SCR Moving Bed	0.202 (1.60)	22.9 (0.0533)	.174 (1.38)	20.1 (.0459)	.0161 (.128)	1.83 (.00426)	See Text
		Moderate - 70%	SCR Parallel Flow	0.606 (4.80)	68.7 (0.160)	Neglig	ible Effect	.00201 (.0160)	.229 (.000533)	See Text
		Stringent - 90%	SCR Parallel Flow	0.202 (1.60)	22.9 (0.0533)	Negligible Effect		.0101 (.0798)	1.14 (.00266)	See Text
75	Distillat	te Uncontrolled	-	2.99 (23.76)	68.0 (0.158)	1.02 (8.10)	46.4 (.108)	0	0	0
		Moderate - 70%	SCR Fixed Packed Bed	0.898 (7.13)	20.4 (0.047)	Neglig	ible Effect	.00502 (.0398)	.228 (.000531)	Less than Residual
		Stringent - 90%	SCR Fixed Packed Bed	0.299 (2.38)	6.80 (0.0158)	Neglig	ible Effect	.00502 (.0398)	.228 (.000531)	Less than Residual
15	Distilla	te Uncontrolled	-	.300 (2.38)	68.2 (.159)	Neglig	ible Amount	0	0	()
		Moderate - 70%	SCR Fixed Packed Bed	.0900 (.714)	20.5 (.0476)			.00109 (.00864)	.248 (.000576)	Less than Residual
		Stringent - 90%	SCR Fixed Packed Bed	.0300 (.238)	6.82 (.0159)			.00109 (.00864)	.248 (.000576)	Less than Residu al

TABLE 6.3.1-1. AIR POLLUTION IMPACTS FROM BEST NO_× FGT CONTROL TECHNIQUES FOR OIL-FIRED BOILERS^{1,2,5-17}

TABLE 6.3.1-2. AIR POLLUTION IMPACTS FROM BEST NO_×/SO_× FGT CONTROL TECHNIQUES FOR OIL-FIRED BOILERS

Boiler	Type:	
Heat	Rate:	150 MBtu/hr
	0il:	Residual

		NOX		\$0 ₂		Particulates		NH ₃		
Control level	Type of control	g/s (1b/hr)	ng/J (1b/MBtu)	g/s (1b/hr)	ng/J (1b/MBtu)	g/s (1b/hr)	ng/J (1b/MBtu)	g/s (1b/hr)	ng/J (1b/MBtu)	Bisulfate
Uncontrolled	-	7.57 (60.0)	172.0 (0.400)	59.4 (471.0)	1350.0 (3.14)	4.16 (33.0)	94.6 (0.220)	0	0	0
Tntermediate (80% NO _x) (85% SO ₂)	SRC-Parallel Flow	1.51 (12.0)	34.4 (0.0800)	8.91 (70.7)	203.0 (0.471)	Negligib:	le Effect	0.191 (1.52)	4.36 (0.0101)	See Text

The flue gas SO3 concentration can be calculated as follows:

= 3.0%

 $\frac{\text{Residual 0il}}{\text{Flue gas}} = 46,700 \frac{\text{ft}^3}{\text{min}} \left(\frac{32 + 460 \ ^\circ\text{R}}{400 + 460 \ ^\circ\text{R}}\right) \left(\frac{1\text{b-mole}}{359 \ \text{scf}}\right) \left(\frac{60 \ \text{min}}{\text{hr}}\right)$ $= 4465 \ \text{lb-moles/hr}$ $\text{SO}_2 = 471.0 \ \frac{1\text{bs}}{\text{hr}} \left(\frac{1\text{b-mole}}{64.0 \ 1\text{b}}\right)$ $= 7.359 \ \text{lb-moles/hr}$

Fuel S

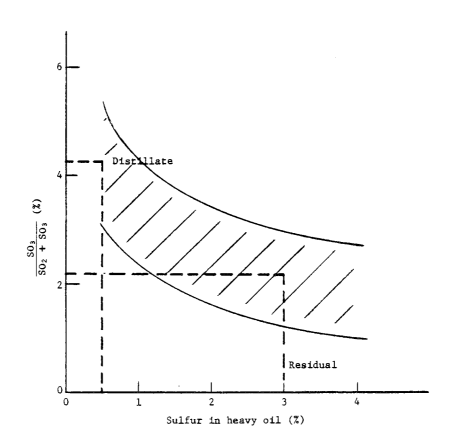


Figure 6.3.1-1. Formation ratio of SO_3 ²⁵

from Figure 6.3.1-1 % SO₃ = 2.3%

The SO_3 concentration can be determined by calculation to be

$$[SO_3] = 39 \text{ ppm}$$

Distillate Oil

Flue gas
$$= \left(5000 \frac{\text{ft}^3}{\text{min}}\right) \left(\frac{32 + 460 \text{ }^\circ \text{R}}{350 + 460 \text{ }^\circ \text{R}}\right) \left(\frac{1\text{b-mole}}{359 \text{ } \text{scf}}\right) \left(\frac{60 \text{ } \text{min}}{\text{hr}}\right)$$
$$= 507.6 \text{ } 1\text{b-moles/hr}$$
$$= 7.67 \frac{1\text{bs}}{\text{hr}} \left(\frac{1\text{b-mole}}{64.0 \text{ } 1\text{b}}\right)$$
$$= .1198 \text{ } 1\text{b-moles/hr}$$
Fuel S
$$= 0.5\%$$

from Figure 6.3.1-1 % SO₃ = 4.3%

The SO_3 concentration can be determined by calculation to be

$$[SO_3] = 11 \text{ ppm}$$

One can see in Figure 6.3.1-2 below that the residual oil-fired flue gas will form bisulfate at a higher temperature (earlier in the exchanger). Also, if the NH₃ concentration does not become limiting, the greater SO₃ concentration will drive the equilibrium of the reaction further to the right, creating more bisulfate and eventually sulfate, $(NH_4)_2SO_4$.

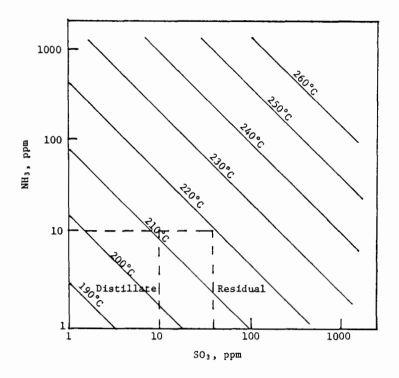


Figure 6.3.1-2. Temperatures below which NH4HSO4 forms.²⁶

Removing the SO3 as bisulfate using particulate control equipment may be a more effective method of removing SO3 from the environment than FGD. This would be a beneficial impact of bisulfate formation. A downstream FGD system could potentially absorb the small NH3 emissions and, therefore, NH3 emissions may be negligible if FGD is used on conjunction with FGT. The level of removal that can be achieved by an FGD scrubber has not yet been examined. Also, the effect of absorbed NH3 on the FGD chemistry has not been resolved, although this question is being studied by the EPA. This is due primarily to the fact that there is only one installation where FGD is applied downstream of an SCR unit and data from this Japanese installation has not been published in the U.S. Several things can be done to reduce the adverse environmental impacts. Combustion control with less 02 minimizes formation of NO $_{\times}$ and SO3. This would be the case for a boiler equipped with low NO $_{\times}$ burners. A minimum NH injection ratio is needed for low NH $_3$ emissions and bisulfate formation. Heat exchanger temperatures must be kept above bisulfate formation and acid condensation points. Use of corrosionresistant material is warranted where bisulfate deposits are probable. Verticle tube heat exchangers are preferable since they are less prone to plugging.

6.3.2 Water Pollution

There are no water streams that are associated with NO_{χ} -only SCR systems, however, there is one potential source of water pollution. In some Japanese installations NH_4HSO_4 deposits are removed from the air preheater by water washing. The blowdown from this operation will contain both ammonium and sulfate ions which, if not treated, present a water pollution source. Since the amounts of NH_4HSO_4 and water are not known, it is impossible to estimate the concentration or flow rate of this potential source.

6.3.3 Solid Waste

The only potential solid waste associated with this system is spent catalyst. Presently, the life of SCR catalysts is 1-2 years and the topic of recycling is not addressed in the literature. Since, to date, virtually no commercial SCR units have operated long enough to require catalyst replacement, there is no commercial experience on the treatment of spent catalyst.²⁷ With the high cost of some of the catalysts, recycling seems to be warranted. Recycling is feasible where the spent catalyst support is composed of valuable materials such as titanium, stainless steel or possibly a ceramic material. Alumina catalyst supports probably do not warrant recycling unless required for environmental reasons. These catalysts can probably be disposed of in the same manner as other industrial catalysts.

The amount of catalyst that must be recycled or disposed of is one reactor volume since replacement involves total catalyst replacement. The actual frequency of catalyst replacement is unique to each specific process, however, catalyst lifetimes are typically one or more years.

Few process vendors have published their catalyst formulations since the field is very competitive at the present time. Base metal oxides are typically used.²⁴ The environmental impact of catalyst disposal will depend on what materials and compounds are involved.

6.3.4 Other Environmental Impacts

The only other environmental impact is a secondary impact. NH_3 is commonly made from natural gas and its consumption is considered a secondary environmental impact. This impact will be quantified in a forthcoming report on the impact of NO_{χ} regulations on the NH_3 industry. The report is being prepared by TVA under contract to EPA-IERL.

Other adverse environmental impacts (noise, thermal pollution, electrical discharges, etc.) are not present with SCR systems.

6.3.5 Environmental Impacts on Modified and Reconstructed Facilities

The environmental impacts of new and retrofitted facilities should be similar. There is not enough difference between new and retrofit systems to indicate that environmental impacts would be significantly different with retrofitted systems.

6.4 ENVIRONMENTAL IMPACTS OF CONTROLS FOR GAS-FIRED BOILERS

6.4.1 Air Pollution

Emission rates for primary and secondary pollutants are listed in Table 6.4.1-1. The table is organized according to control level (uncontrolled, moderate, and stringent). The impact on the primary pollutant, NO_{χ} , is given for each case, as is that of the secondary pollutant, NH_3 . There is an insignificant amount of particulates in the flue gas and, therefore, these are not considered to be a pollutant. There is also no problem with bisulfate formation since the fuel has only a trace of sulfur.

Table 6.2.1-13 shows the uncontrolled NO_{\times} emission for the natural gasfired standard boiler to be less than the SIP control level.

Standard	l boiler			N	10 _×	NH 3		
Heat rate (MBtu/hr)	е Туре	Control level	Type of control	g/s (1b/hr)	ng/J (1b/MBtu)	g/s (1b/hr)	ng/J (1b/MBtu)	
15	Firetube	Uncontrolled	_	0.332 (2.63)	75.4 (.175)	0.	0	
		Moderate-70%	SCR Fixed Packed Bed	0.0995 (.789)	22.6 (.0526)	0.00113 (.00898)	0.257 (.000598)	
		Stringent-90%	SCR Fixed Packed Bed	.0332 (.263)	7.54 (.0175)	0.00113 (.00898)	0.257 (.000598)	
150	Watertube	Uncontrolled	-	3.31 (26.26)	75.3 (.175)	0	0	
		Moderate-70%	SCR Fixed Packed Bed	0.993 (7.88)	22.6 .0525)	0.00511 (.0405)	0.232 (.000540)	
		Stringent-90%	SCR Fixed Packed Bed	0.331 (2.63)	7.53 (.0175)	0.00511 (.0405)	0.232 (.000540)	

TABLE 6.4.1-1. AIR POLLUTION IMPACTS FROM BEST NO_× FGT CONTROL TECHNIQUES FOR GAS-FIRED BOILERS^{5,6,7}

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The only environmental impacts are NO_{\times} and NH_3 emissions. The uncontrolled NO_{\times} emissions, $0.332 \frac{g}{s}$, are the lowest for all standard boilers except distillate oil. Moderate - stringent controls reduce this figure to $0.0995-0.0332 \frac{g}{s}$. NH_3 emissions for all control levels are 1 ppm (Figure 6.1-1). This corresponds to a mass rate of $0.00113 \frac{g}{s}$.

To reduce the adverse environmental impacts, combustion controls utilizing less O_2 minimizes NO_{\times} formation could be implemented. NH_3 emissions are presently quite low. Care needs to be taken to see that an excessive $NH_3:NO_{\times}$ injection ratio is not used thus increasing the low emission level.

6.4.2 Water Pollution

There are no water streams in SCR fixed packed bed systems.

6.4.3 Solid Waste

The only potential solid waste associated with this system is spent catalyst. Presently, the life of SCR catalysts is 1-2 years and the topic of recycling is not addressed in the literature. Since, to date, virtually no commercial SCR units have operated long enough to require catalyst replacement, there is no commercial experience on the treatment of spent catalyst.²⁷ With the high cost of some of the catalysts, recycling seems to be warranted. Recycling is feasible where the spent catalyst support is composed of valuable materials such as titanium, stainless steel, or possibly a ceramic material. Alumina catalyst supports probably do not warrant recycling unless required for environmental reasons. These catalysts can probably be disposed of in the same manner as other industrial catalysts.

The amount of catalyst that must be recycled or disposed of is one reactor volume since replacement involves total catalyst replacement. The actual frequency of catalyst replacement is unique to each specific process, however, catalyst lifetimes are typically one or more years.

Few process vendors have published their catalyst formulations since the field is very competitive at the present time. Base metal oxides are typically used.²⁴ The environmental impact of catalyst disposal will depend on what materials and compounds are involved.

6.4.4 Other Environmental Impacts

The only other environmental impact is a secondary impact. NH_3 is commonly made from natural gas and its consumption is considered a secondary environmental impact. This impact will be quantified in a forthcoming report on the impact of NO_{\times} regulations on the NH_3 industry. The report is being prepared by TVA under contract to EPA-IERL.

Other adverse environmental impacts (noise, thermal pollution, electrical discharge, etc.) are not present with SCR systems.

6.4.5 Environmental Impacts on Modified and Reconstructed Facilities

The environmental impacts of new and retrofitted systems should be similar. Retrofitted systems are not so different as to create a greater adverse environmental impact for these systems.

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- 24. Ibid., p. 3-2.
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- 27. Ando, J., Letter of April 4, 1979 to J. David Mobley.

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SECTION 7

EMISSION SOURCE TEST DATA

7.1 INTRODUCTION

Test data from operating units are necessary to demonstrate that the control technology will perform as claimed. For this purpose the most meaningful test data are those that represent 24 hour averages over 30 days of continuous operation. At the present time, very little of this type of information is published. However, some continuous data have been presented at recent seminars and obtained from the process operators.

The EPA approved test methods are the same for all fuel types and are discussed here to avoid unnecessary repetition. There are two methods for measuring NO_{\times} (expressed as NO_2) in gas streams. First, the EPA Reference Method 7 is for the determination of nitrogen oxides emissions from stationary sources. Presently, this method is the only one approved by the EPA for measuring NO_{\times} levels in flue gas from industrial boilers for emission source test data.

Method 7 is based on grab sampling for wet chemical analysis and is used for spot-checking SCR systems' performance and in calibrating an instrument analyzer. Continuous monitoring by Method 7 for process control purposes is impractical as the method requires a collected sample to set a minimum of 16 hours. However, continuous data can be developed using Method 7 by taking samples at several intervals during a 24 hour period and computing a 24 hour average. Daily values computed in this manner can represent continuous data when computed for a period of 30 days or more.

The second method, EPA Reference Method 20, is for the determination of nitrogen oxide emissions from stationary gas turbines. While this method is not approved for industrial boilers, it is applicable to continuous monitoring due to its utilization of an instrumental analyzer based on chemiluminescence. This instrument provides a sound basis for process control, which is most important when hourly ambient NO_v standards are in effect.

The methodology and test procedures for each method are described in the following paragraphs. With Method 7, grab samples are collected in an evacuated flask containing a dilute sulfuric acid-hydrogen peroxide $(H_2SO_4-H_2O_2)$ absorbing solution. The nitrogen oxides, except nitrous oxide (N_2O) , are measured colorimetrically using the phenoldisulfonic acid (PDS) procedure. The apparatus for this system is shown in Figure 7.1-1.

A 25 ml aliquot of absorbing solution is added to the flask. The flask is stoppered and then evacuated by use of the pump. After checking for leaks, the probe and vacuum tube are purged using the squeeze bulb. Then the flask valve is turned to the "sample" position allowing the gas to enter the flask. When the pressures in the flask and sample line (i.e., duct) are equalized the flask is isolated, disconnected from the sampling train, and shaken for 5 minutes. The sample flask is allowed to set for a minimum of 16 hours. After transferring the sample and then washing out the sample flask into a volumetric flask, 25 ml aliquot is pipetted into a porcelain evaporating dish. This aliquot is evaporated to dryness on a steam bath and allowed to cool. Two ml of phenoldisulforic acid solution is added to the dried residue and the residue is ground to a powder with a polyethylene policeman. After adding deionized, distillated water and concentrated sulfuric acid, concentrated ammonium hydroxide is added dropwise until the pH is 10. The contents of the flask are mixed thoroughly and the absorbance of a sample is measured by a spectrophotometer. The total mass of NO_{X} per sample is expressed by the following equation:

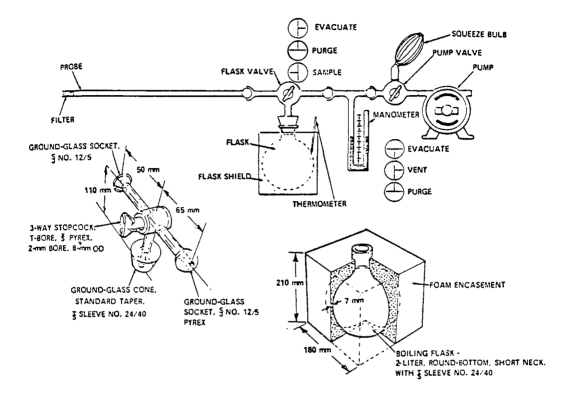


Figure 7.1-1. Sampling train, flask valve, and flask.¹

$$m = 2 K A$$

where $m = mass of NO_{\times} as NO_{2}$ in gas sample, μg $K_{c} = spectrophotometer calibration factor$ A = absorbance of sample

The sample volume, dry basis, corrected to standard conditions is found by the equation

$$V_{s} = 0.3858 \frac{^{\circ}K}{\text{mmHg}} \left(V_{f} - 25\text{ml} \right) \left(\frac{P_{f}}{T_{f}} - \frac{P_{i}}{T_{i}} \right)$$

where V_s = sample volume, ml V_f = volume of flask and valve, ml P_f = final absolute flask pressure, mmHg T_f = final absolute flask temperature, °K P_i = initial absolute flask pressure, mmHg T_i = initial absolute flask temperature, °K

Finally, the NO_{\times} concentration in the gas sample is determined by

$$C = 10^3 \frac{mg/m^3}{\mu g/m1} \left(\frac{m}{V_s} \right)$$

Method 20, for determining nitrogen oxides emissions from stationary gas turbines, utilizes an instrumental analyzer to which a continuous gas sample from the exhaust stream is conveyed. The apparatus for this system is shown in Figure 7.1-2. Particulate matter and water vapor are the primary interferring species for most instrumental analyzers, but these are removed by the filter and condenser, respectively, present in the sampling train. In application to SCR systems on boilers, the presence of NH₃ may interfere with the instruments performance. This problem can be circumvented via the use of an ammonia decomposition catalyst before the probe measuring the reactor outlet NO_{γ} concentration.

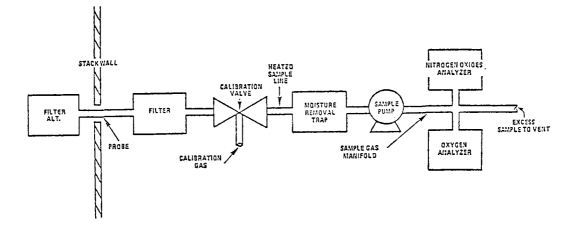


Figure 7.1-2. Measurement system design for stationary gas turbine tests.² 7.2 EMISSION SOURCE TEST DATA FOR COAL-FIRED BOILERS

At this writing, there have been only a few pilot plant tests performed in Japan utilizing SCR systems to treat coal-fired flue gas. Little data from these tests have been released. Most all SCR work has been done recently in Japan (the oldest SCR system, on an oil-fired boiler, has been operating for 5 years). Because of Japan's lack of large coal reserves, there are few coal-fired utility or industrial boilers in the country. However, more coalfired boilers are planned for the near future which will utilize imported coal. This is a result of the scarcity and high cost of cleaner fuels.

There are two coal-fired utility boilers equipped with SCR NO_x removal systems due to start-up in 1980.³ Hokkaido Electric Company plans a 90 MW coal-fired boiler with an SCR unit to be started up at Tomato in March 1980. The Electric Power Development Corporation has a 250 MW coal-fired SCR unit due for completion at Takehara in November, 1980. Also, in the United States,

there are 2 SCR pilot plants presently under construction at coal-fired utilities located in Tampa, Florida and Albany, Georgia. These are scheduled for completion in 1979. Once these units are in operation it will be possible to obtain more test data.

The available coal-fired source test data is summarized below. Figure 7.2-1 shows the performance of a cylindrical catalyst treating coal-fired flue gas after particulate removal. Figures 7.2-2 and 7.2-3 show the performance of a parallel flow and a moving bed reactor, respectively, treating coal-fired flue gas after an ESP. Figure 7.2-4 is for a parallel flow system. One can see from the plots that the SCR systems are capable of achieving the stringent level of control.

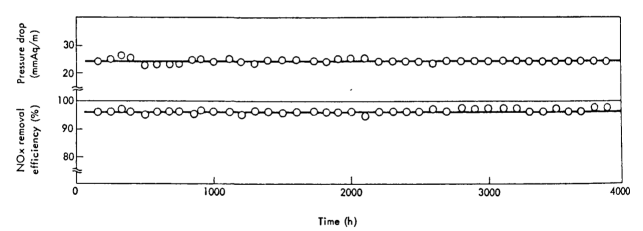


Figure 7.2-1. Change of NO_× removal efficiency and pressure drop (Kawasaki Heavy Industries process, Takehara power station, Hiroshima, Japan).⁴

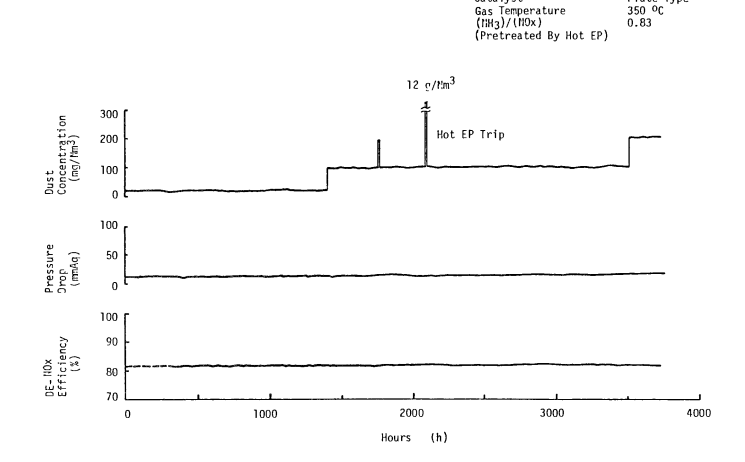


Plate Type 350 ^OC 0.83

Catalyst

Figure 7.2-2. Pilot plant test of a parallel flow reactor treating a flue gas from a coal-fired utility boiler (Hitachi, Ltd. process, unknown location, Japan).⁵

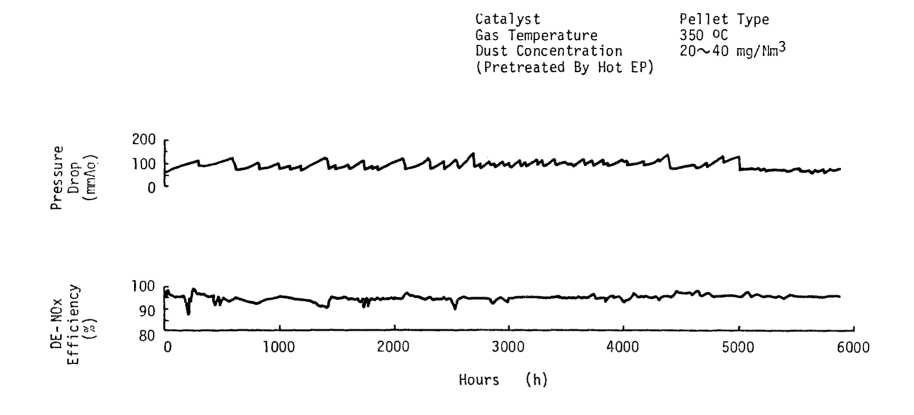


Figure 7.2-3. Pilot plant test of an intermittent-moving bed reactor treating a flue gas from a coal-fired utility boiler (Hitachi, Ltd. process, unknown location, Japan).⁶

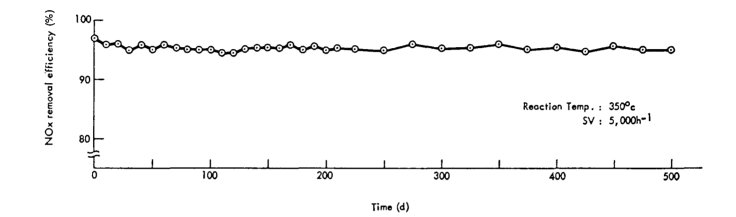


Figure 7.2-4. Durability test of NO_× removal catalyst (Kawasaki Heavy Ind. process, Takehara power station, Hiroshima, Japan).

7.3 EMISSION SOURCE TEST DATA FOR OIL-FIRED BOILERS

While there are a number of commercial SCR systems presently treating oil-fired flue gas in Japan, the data on these units are limited mostly to a single reported removal level. Catalyst life tests on heavy oil-fired pilot unit equipment do provide an indication as to how commercial units will perform. These continuous test results are shown in Figures 7.3-1 through 7.3-3. Most of the data available are presented as summaries of pilot test results and are usually expressed in plots of NO_x removal (%) vs. NH₃:NO_x mole ratio or reactor space velocity (hr⁻¹). In a few cases, tables of operating parameters of commercial SCR plants are given. These results are given in Tables 7.3-1 through 7.3-3 on the following pages.

Most data give only point values of removal and not a set of continuous data. In addition, the test method and boiler operating conditions are not given. Included in these figures are data recently obtained from commercial Japanese installations on industrial boilers and, as such, they represent the most complete set of continuous data currently available.

Summaries of the oil-fired industrial (larger than 3 MW) and utility SCR plants in Japan are shown in Tables 7.3-4 and 7.3-5. These tables are presented since they represent locations where operating data on SCR units can be obtained. The data can be obtained by either contacting the boiler owners and requesting available data or by arranging for independent on-site testing.

The data shown in Figures 7.3-4 and 7.3-5 represent the most recently available continuous daily average data available from SCR systems applied to industrial boilers. EPA Method 20 was used to obtain the data in both of the figures. The average removal level represents the level of control necessary to meet the local Japanese emission regulations. Continuous daily averages are not available from most SCR units; however, Table 7.3-6 shows maximum and minimum NO_{\times} removal values for several industrial units.

7-10

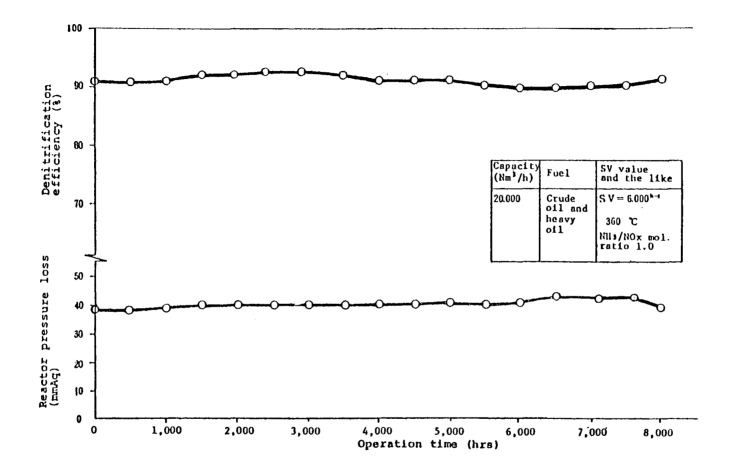


Figure 7.3-1. Catalyst life test results (IHI process, Taketoyo power station, Japan).⁷

7-11

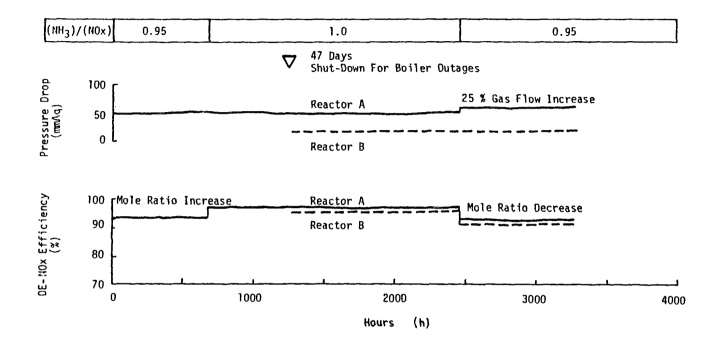
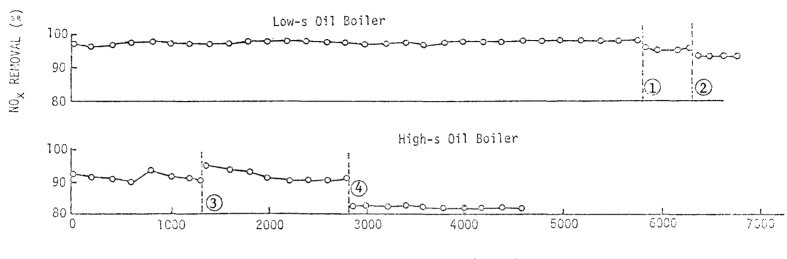


Figure 7.3-2. Pilot plant test of a parallel flow reactor treating a flue gas from a high sulfur heavy oil-fired utility boiler (Hitachi, Ltd. process, unknown location, Japan).⁹

Ν.



OPERATION PERIOD (Hours)

Circled figures show times when SV and NH_3/NO mole ratio were changed.

1.	SV	$10,000 - 15,000 \text{ hr}^{-1}$	3.	SV $6,200 - 4,500 \text{ hr}^{-1}$
2.	SV	$15,000 - 20,000 \text{ hr}^{-1}$	4.	SV $4,500 - 6,200$ and the mole
				ratio 0.95 – 0.83.

Figure 7.3-3. Test results of oil-fired boilers (Hitachi, Ltd. process, unknown location, Japan).¹⁶

	Idemitsu Kosan	Shindaikyowa Petrochemical	Kawasaki Steel
Completed	Oct. 1975	Nov. 1975	Nov. 1976
Plant site	Chiba	Yokkaichi	Chiba
Gas source	FCC-CO Boiler and furnace	Oil-fired Boiler	Iron-ore Sintering machine
Capacity (Nm ³ /hr)	350,000	440,000	762,000
Load factor (%)	50-100	50-100	70-100
Pretreatment of gas	Heating	EP*, FGD, Heating	EP, FGD WEP [†] , Heating
Reactor inlet			
NO _× (ppm)	230	150	200-300
SO _× (ppm)	50-80	80-130	5-20
Dust (mg/Nm ³)	20-50	30-100	3-10
0 ₂ (%)	2.3	3.2	11.2
Reactor type	Fixed bed	Fixed bed	Fixed bed
Reaction temp.	400	420	
$\mathrm{NO}_{\mathrm{x}}/\mathrm{NH}_{3}$ ratio	1.0	1.0	1.0
Catalyst No.	204	304	304
$SV (hr^{-1})$	5,000	10,000	4,000
NO _x removal (%)	93	80‡	95
Pressure drop by SCR reactor (mmH ₂ 0)	170	160	50
Catalyst life	l year	l year	l year

TABLE 7.3-1. OPERATION PARAMETERS OF MAJOR PLANTS CONSTRUCTED BY HITACHI ZOSEN¹⁰

*Electrostatic precipitator

 $^{\rm t}{\rm Wet}$ electrostatic precipitator

 $\dot{\dagger}$ Including leakage in heat exchanger

	Mitsui Petro- Chemical Co.	
Capacity (Nm ³ /hr)	200,000	220,000
Gas composition		
NO _× (ppm)	190	150
SO _× (ppm)	None	300
Dust (mg/Nm ³)	20-50	100-150
Catalyst and reactor		
Catalyst carrier	A1203	TiO ₂
Catalyst shape	Granule	Tube
SV (hr ⁻¹)	2,600	4,000
Temperature (°C)	350	350-400
$\rm NH_3/NO_{\times}$ mole ratio	1.0	1.0
NO _x removal (%)	Above 90	Above 90
Total pressure drop (mmH ₂ 0)		180
Leak NH ₃ (ppm)		Below 10
Operation start	Oct. 1975	July 1977
Plant cost (10 ⁶ yen)		260
Denitrification cost		
(yen/kWhr)*		

TABLE 7.3-2. SCR PLANTS BY MITSUI ENGINEERING & SHIPBUILDING CO.¹¹

*Including 7 years depreciation.

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		Co	Commercial			
	Pilot	No. 1	No. 2			
Gas for SCR (NM ³ /hr)	30,000	240,000	300,000			
Fuel	0i1(S=0.7%)					
Load fluctuation	60-100%	60-100%	60-100%			
Stack height (m)	70	140	140			
Inlet gas composition						
02 (%)	6	6	6			
SO _x (ppm)	400	400	400			
NO _x (ppm)	200	200	200			
Particulates after EP (mg/Nm ³)	5-20	5-10	10-20			
FGD unit	None	Scheduled	Scheduled			
$SV (hr^{-1})$	5,000	5,000	5,000			
Temperature (°C)	320	320	320			
NO, removal (%)	Over 90	Over 90	Over 90			
^ NH₃/NO mole ratio	1.0	1.0	1.0			
Leak ammonia (ppm)	10-20	10-20	10-20			
Type of reactor	Fixed bed	Fixed bed	Moving bed			
Pressure drop (mmH ₂ O)						
Reactor		200				
Total system		500				
Plant completed	July 1973	Mar. 1976	Oct. 1976			

TABLE 7.3-3. OPERATION DATA OF SCR PLANTS FOR DIRTY GAS¹²

Company	Site	Capacity (Nm ³ /hr)	Reactor type*	Completion date
Sumitano Chemical	Sodegaura	30,000	FPB	July 1973
Kurabo	Hirakata	30,000	MB	August 1975
Mitsui Petrochemical	Chiba	200,000	FPB	October 1975
Idemitsu Kosan	Chiba	350,000	FPB	October 1975
Shindaikyowa P.C.	Yokkaichi	440,000	FPB .	November 1975
Sumitomo Chemical	Sodegaura	240,000	FPB	March 1976
Fuji Oil	Chiba	70,000	PF	July 1976
Sumitomo Chemical	Sodegaura	300,000	MB	September 1976
Sumitomo Chemical	Sodegaura	300,000	MB	October 1976
Nisshin Steel	Amagasaki	19,000	FPB	July 1977
Nisshin Steel	Amagasaki	20,000	FPB	August 1977
Chiyoda Kenzai	Kaizuka	15,000	MB	October 1977
Fuji Oil	Sodegaura	200,000	PF	January 1978
Ajinomoto	Kawasaki	180,000	PF	January 1978
Nippon Oils & Fats	Amagasaki	20,000	MB	April 1978
Nippon Yakin	Kawasaki	14,000	FPB	July 1978

TABLE 7.3-4. OIL-FIRED INDUSTRIAL SCR PLANTS $^{1\,3}$

*FPB = Fixed Packed Bed
MB = Moving Bed
PF = Parallel Flow

Power company	Site	Capacity (Nm ³ /hr)	Reactor type*	Completion date
Kansai Electric	Kainan	300,000	FPB	June 1977
Company C	-	1,010,000	PF	February 1978
Kansai Electric	Amagasaki	410,000	-	June 1978
Company A	-	490,000	PF	June 1978
Kansai Electric	Osaka		PF	July 1978
Company D	-	490,000	PF	July 1978
Company G	-	1,000,000	PF	April 1979
Chugoku Electric	Kudamatsu	1,900,000	PF	July 1979
Chubu Electric	Chita	1,920,000	PF	February 1980
Tohoku Electric	Niigata	1,660,000	PF	August 1981

TABLE 7.3-5. OIL-FIRED UTILITY SCR PLANTS^{1 4}

*FPB = Fixed Packed Bed

PF = Parallel Flow

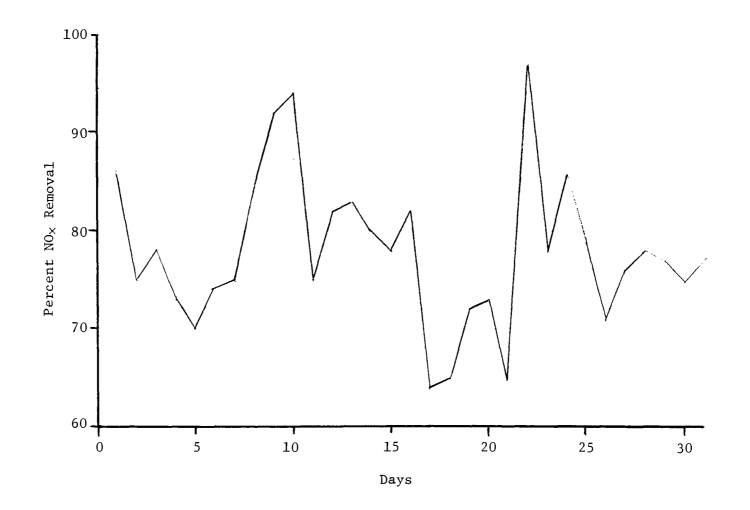


Figure 7.3-4. NO_X removal for the month of May 1977 (Hitachi Zosen fixed bed process, Shindaikyowa Petrochemical, Yokkaichi, Japan, Chemiluminescence Method).²⁰

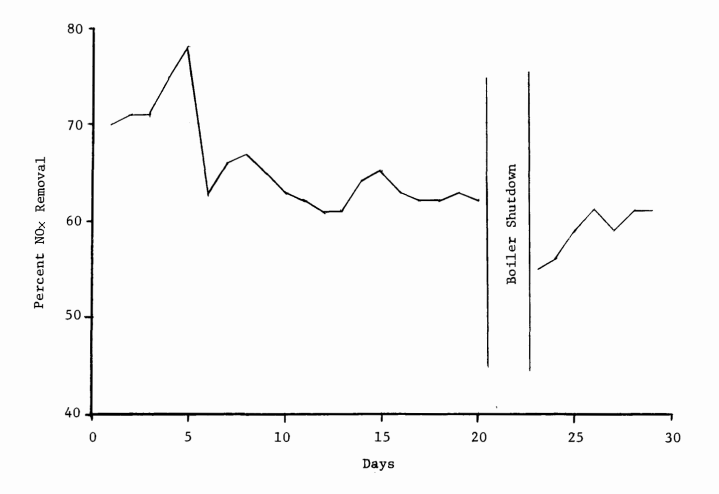


Figure 7.3-5. NO_X removal for August, 1978 (MHI process, Fuji Oil, Sodegaura, Japan, PDS/chemiluminescence method).²⁰

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Plant owner	Plant site	Capacity* (MW)	Process	Percent NO removal range
Nippon Yakin	Kawasaki	5	Fixed bed + sodium scrubbing	86-98
Shindaikyowa	Yokkaichi	135	Sodium scrubbing + fixed bed	53-62
Fuji Oil	Sodegaura	23	Fixed (parallel passage) bed	93-97
Fuji Oil	Sodegaura	53	Fixed (honeycomb) bed	60-80
Kurabo	Hirakata	10	Moving bed (continuous)	90-94
Nippon Oil & Fats	Amagasaki	7	Moving bed (intermittent)	94–97
Kansai Paint	Amagasaki	5.3	Fixed bed	90-92
Nisshin Steel	Amagasaki	6.5	Moving bed	94–96

TABLE 7.3-6. NO $_{\times}$ REMOVAL LEVELS AT SEVERAL JAPANESE INDUSTRIAL BOILERS WITH NO $_{\times}$ CONTROL BY SCR²⁰

*Assumed to be \mathtt{MW}_{e} equivalent.

7.4 EMISSION SOURCE TEST DATA FOR GAS-FIRED BOILERS

Although gas-fired boilers, both industrial and utility, are numerous in Japan, few have been equipped with SCR units so far. This is due to the fact that combustion modifications on the boilers have been installed because of their lower cost and the lack of fuel-bound nitrogen to contend with. The data available on gas-fired SCR systems in Japan are presented in Figures 7.4-1 through 7.4-5 on the following pages. Figure 7.4-1 is a plot of a long-term performance test while the rest are summaries of pilot tests.

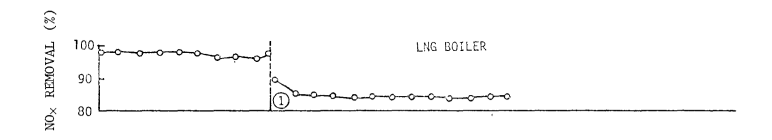
Summaries of the gas-fired industrial and utility SCR plants in Japan are shown in the table below.

Company	Site	Capacity (Nm³/hr)	Reactor type*	Completion date
Osaka Gas	Takaishi	15,000x2	FPB	December 1976
Chubu Electric	Chita	1,910,000	FPB	April 1978
Kyushu Electric	Kokura	1,610,000	FPB	July 1978
Chubu Electric	Chita	1,910,000	FPB	September 1978
Hyushu Electric	Kokura	1,610,000	FPB	December 1978

TABLE 7.4-1. GAS-FIRED SCR PLANTS^{1 5}

*FPB = Fixed Packed Bed

As with oil-fired installations, the type of test data desired may exist, but has not yet been published. To obtain this data it will be necessary to contact the boiler owners to possibly get available data or conduct on-site testing.



OPERATION PERIOD (Hours)

Circled figure shows time when SV and NH3/NO mole ratio were changed.

1. SV $10,000 - 20,000 \text{ hr}^{-1}$

Figure 7.4-1. Test results of gas-fired boilers (Hitachi, Ltd. process, unknown location, Japan).¹⁶

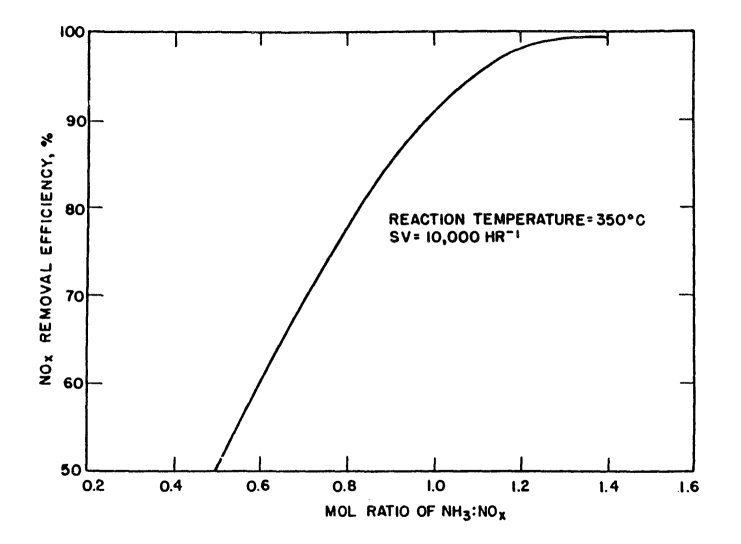


Figure 7.4-2. Characteristic curve of the effect of mole ratio of $NH_3:NO_X$ on NO_X removal efficiency for Hitachí, Ltd. process.¹⁷

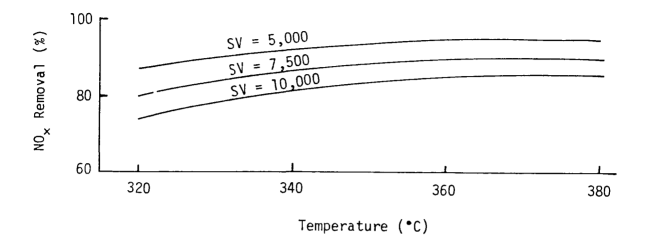


Figure 7.4-3. Performance of catalyst MTC-102 (Mitsui Toatsu process, unknown location, Japan).¹⁸

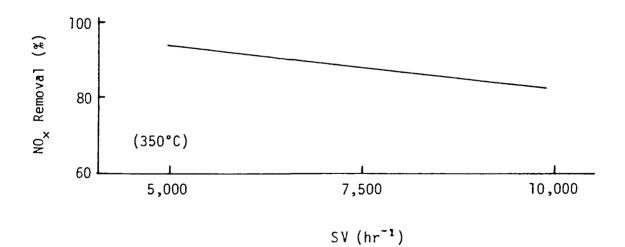


Figure 7.4-4. SV and NO_{\times} removal (MTC-102) (Mitsui Toatsu process, unknown location, Japan).¹⁸

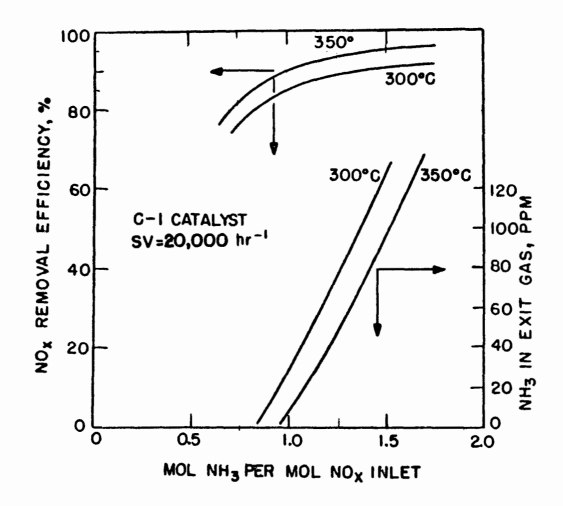


Figure 7.4-5. Relationship among inlet $NH_3:NO_x$ mol ratio, NO_x removal efficiency, and exiting NH_3 concentration using the Sumitomo Chemical C-1 Catalyst.¹⁹

REFERENCES

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1.	Environmental Reporter, Appendix A, Oct. 21, 1977, p. 92.
2.	Federal Register, Volume 42, Number 191, Oct. 3, 1977, p. 53790.
3.	Ando, J. NO _x Abatement from Stationary Sources in Japan. EPA draft report in preparation. October 1978, pp. 1-35.
4.	Niwa, Senji. Characteristic of Cylindrical DeNO _X Catalyst for a Coal-Fired Boiler. Paper presented at EPRI NO _X Control Technology Seminar II. Denver, Colorado. November 8-9, 1978. p. 21-8.
5.	Kuroda, H., Nakajima, F. Experiences of NO Removal in Pilot Plants and Utility Boilers. Paper presented at EPRI NO Control Technology Seminar II. Denver, Colorado. November 8-9, 1978. p. 20-13.
6.	<i>Ibid.</i> , p. 20-10.
7, .	Ando, J., op. cit., p. 4-96.
8.	Niwa, S., op. cit., p. 21-5.
9.	Kuroda, H., et al., op. cit., p. 20-12.
10.	Ando, J., op. cit., p. 4-21.
11.	<i>Ibid.</i> , p. 4-71.
12.	<i>Ibid.</i> , p. 4-5.
13.	Ibid., p. 3-4, 3-5.
14.	<i>Ibid.</i> , p. 1-35
15.	<i>Ibid.</i> , p. 3-4, 3-5.
16.	Ibid., p. 4-43.
17.	Faucett, H. L., <i>et al.</i> Technical Assessment of NO _x Removal Processes for Utility Applications. EPA $600/7-77-127$. November 1977. p. 214.
18.	Ando, J., op cit., p. 4-121.
19.	Faucett, H. L., op. cit., p. 298.

7-27

APPENDIX 1

DETAILED SYSTEM EVALUATIONS

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	Performance	Operation and maintenance	Environ- mental impact	Economic impact	Energy/ material impact	Boiler operation and safety	Relia- bility	Status of development	Adapt~ ability	Compati- bility	Total
SCR Fixed Packed Bed	8	6	7	11	7	3	9	10	3	5	69
SCR Moving Bed	8	4	7	14	7	3	9	10	3	5	70
SCR Parallel Flow	12	7	7	14	8	3	14	10	3	5	83
Absorption - Oxidation	7	2	6	2	3	2	7	6	3	5	43

TABLE A1.1. POINT VALUE NO_x-ONLY PROCESS RATINGS: COAL-FIRED BOILERS - MODERATE CONTROL

TABLE A1.2. POINT VALUE SIMULTANEOUS NO_{\times}/SO_{\times} PROCESS RATINGS: COAL-FIRED BOILERS - MODERATE CONTROL

	Performance	Operation and maintenance	Environ- mental impact	Economic impact	Energy/ material impact	Boiler operation and safety	Relia- bility	Status of development	Adapt- ability	Compati- bility	Total
SCR Parallel Flow	12	3	9	11	5	3	12	13	2	2	72
Adsorption	*										
Electron Beam Radiation	4	5	3	6	4	2	6	3	3	5	41
Absorption - Reduction	10	2	7	10	4	4	7	3	3	2	52
Oxidation - Absorption/Reductio	n 10	3	6	6	1	2	8	10	3	2	51
Oxidation - Absorption	8	4	6	5	1	2	8	11	3	5	51

*Not Applicable - Does not meet removal requirements.

		Operation and	Environ- mental	Economic	Energy/ material	Boiler operation	Relia-	Status of	Adapt-	Compati	_
	Performance	maintenance	impact	impact	impact	and safety	bility	development	ability	bility	Total
SCR Fixed	· · · · · · · · · · · · · · · · · · ·										
Packed Bed	4	6	7	8	7	3	9	10	3	5	62
SCR Moving											
Bed	4	4	7	8	8	3	9	10	3	5	60
SCR Parallel											
Flow	8	7	7	8	8	3	14	10	3	5	73
Absorption -											
Oxidation	*										

TABLE A1.3. POINT VALUE NO_x-ONLY PROCESS RATINGS: COAL-FIRED BOILERS - STRINGENT CONTROL

*Not Applicable - Does not meet removal requirements

TABLE A1.4.	POINT VALUE SIMULTANEOUS NO_X/SO_X PROCESS RATINGS:
	COAL-FIRED BOILERS - STRINGENT CONTROL

	Performance	Operation and maintenance	Environ- mental impace	Economic impact	Energy/ material impact	Boiler operation and safety	Relia- bility	Status of development	Adapt- ability	Compati bility	- Total
SCR Parallel Flow	8	3	9	11	5	3	12	13	2	2	68
Adsorption	*										
Electron Beam Radiation	*										
Absorption - Reduction	*										
Oxidation - Absorption/Reductio	n 6	3	6	6	1	2	8	10	3	2	48
Oxidation - Absorption	6	4	6	5	1	2	8	11	3	5	49

*Not Applicable - Does not meet removal requirements.

	Performance	Operation and maintenance	Environ- mental impact	Economic impact	Energy/ material impact	Boiler operation and safety	Relia- bility	Status of development	Adapt- ability	Compati- bility	Total
SCR Fixed Packed Bed	6	6	7	11	7	3	9	10	3	5	67
SCR Mo ving Bed	6	4	7	14	8	3	9	10	3	5	69
SCR Parallel Flow	10	7	7	14	8	3	14	10	3	5	81
Absorption - Oxidation	7	2	6	2	3	2	7	6	3	5	43

TABLE A 1.5. POINT VALUE NO_×-ONLY PROCESS RATINGS: COAL-FIRED BOILERS - INTERMEDIATE CONTROL

TABLE A1.6. POINT VALUE SIMULTANEOUS NO×/SO× PROCESS RATINGS: COAL-FIRED BOILERS - INTERMEDIATE CONTROL

	Performance	Operation and maintenance	Environ- mental impact	Economic impact	Energy/ material impact	Boiler operation and safety	Relia- bility	Status of development	Adapt- ability	Compati- bility	Total
SCR Parallel Flow	10	3	.9	11	5	3	12	13	2	2	70
Adsorption	*										
Electron Beam Radiation	4	5	3	6	4	2	6	3	3	5	41
Absorption - Reduction	8	2	7	10	4	4	7	3	3	2	50
Oxidation - Absorption/Reduction	n 8	3	6	6	1	2	8	10	3	2	49
Oxidation - Absorption	6	4	6	5	1	2	8	11	3	5	46

*Not Applicable - Does not meet removal requirements.

	Performance	Operation and maintenance	Environ- mental impact	Economic impact	Energy/ material impact	Boiler operation and safety	Relia- bility	Status of development	Adapt- ability	Compati- bility	Total
SCR Fixed Packed Bed	8	6	7	14	8	3	11	16	3	5	81
SCR Moving Bed	12	5	7	14	9	3	14	16	3	5	88
SCR Parallel Flow	12	7	7	14	9	3	14	16	3	5	90
Absorption - Oxidation	7	3	6	2	4	2	8	13	3	5	53

TABLE A1.7. POINT VALUE NO_x-ONLY PROCESS RATINGS: OIL-FIRED BOILERS - MODERATE CONTROL

TABLE A1.8. POINT VALUE SIMULTANEOUS NO_{\times}/SO_{\times} PROCESS RATINGS: OIL-FIRED BOILERS - MODERATE CONTROL

	Performance	Operation and maintenance	Environ- mental impact	Economic impact	Energy/ material impact	Boiler operation and safety	Relia- bility	Status of development	Adapt- ability	Compati- bility	Total
SCR Parallel Flow	12	4	9	11	7	3	12	13	2	2	75
Adsorption	*										
Electron Beam Radiation	4	5	3	6	4	2	8	7	3	5	47
Absorption - Reduction	8	3	7	9	5	4	7	10	3	2	58
Oxidation - Absorption/Reductio	n 10	4	6	6	2	2	8	16	3	2	59
Oxidation - Absorption	8	5	6	5	2	2	9	7	3	5	52

*Not Applicable - Does not meet removal requirements.

TABLE A1.9. POINT VALUE NO_×-ONLY PROCESS RATINGS: OIL-FIRED BOILERS - STRINGENT CONTROL

	Performance	Operation and maintenance	Environ- mental impact	Economic impact	Energy/ material impact	Boiler operation and safety	Relia- bility	Status of development	Adapt- ability	Compati- bility	Total
SCR Fixed Packed Bed	4	6	7	8	8	3	11	16	3	5	74
SCR Moving Bed	8	5	7	8	9	3	14	16	3	5	81
SCR Parallel Flow	8	7	7	8	9	3	14	16	3	5	83
Absorption - Oxidation	*										

*Not Applicable - Does not meet removal requirements

TABLE A1.10.	POINT VALUE SIMULTANEOUS NO_{\times}/SO_{\times} PROCESS RATINGS:	
	OIL-FIRED BOILERS - STRINGENT CONTROL	

	Performance	Operation and maintenance	Environ- mental impact	Economic impact	Energy/ material impact	Boiler operation and safety	Relia- bility	Status of development	Adapt- ability	Compati- bility	Total
SCR Parallel Flow	8	4	9	11	7	3	12	13	2	2	71
Adsorption	*										
Electron Beam Radiation	*										
Absorption - Reduction	*										
Oxidation - Absorption/Reduction	n 6	4	6	6	2	2	8	16	3	2	54
Oxidation - Absorption	6	5	6	5	2	2	9	7	3	5	50

*Not Applicable - Does not meet removal requirements

	Performance	Operation and maintenance	Environ- mental impact	Economic impact	Energy/ material impact	Boiler operation and safety	Relia- bility	Status of development	Adapt- ability	Compati- bility	Total
SCR Fixed Packed Bed	6	6	7	14	8	3	11	16	3	5	79
SCR Moving Bed	10	5	7	14	9	3	14	16	3	5	86
SCR Parallel Flow	10	7	7	14	9	3	14	16	3	5	88
Absorption - Oxidation	7	3	6	3	4	2	8	13	3	5	54

TABLE A1.11. POINT VALUE NO - ONLY PROCESS RATINGS: OIL-FIRED BOILERS - INTERMEDIATE CONTROL

TABLE A1.12. POINT VALUE SIMULTANEOUS NO_X/SO_X PROCESS RATINGS: OIL-FIRED BOILERS - INTERMEDIATE CONTROL

	Performance	Operation and maintenance	Environ- mental impact	Economic impact	Energy/ material impact	Boiler operation and safety	Relia- bility	Status of development	Adapt- ability	Compati- bility	Total
SCR Parallel Flow	10	4	9	11	7	3	12	13	2	2	73
Adsorption	*										
Electron Beam Radiation	4	5	3	6	4	2	8	7	3	5	47
Absorption - Reduction	8	3	7	9	5	4	7	10	3	2	58
Oxidation - Absorption/Reductio	n 8	4	6	4	2	2	8	16	3	2	55
Oxidation - Absorption	6	5	6	3	2	2	9	7	3	5	48

*Not Applicable - Does not meet removal requirements.

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	Performance	Operation and maintenance	Environ- mental impact	Economic impact	Energy/ material impact	Boiler operation and safety	Relia- bility	Status of development	Adapt ability	Compati- bility	Total
SCR Fixed Packed Bed	12	7	8	14	10	4	14	16	3	5	93
SCR Moving Bed	12	5	8	14	10	4	14	16	3	5	91
SCR Parallel Flow	12	7	8	14	10	4	14	16	3	5	93
Absorption - Oxidation	8	4	7	2	6	2	8	9	3	5	58

TABLE A1.13. POINT VALUE NO - ONLY PROCESS RATINGS: GAS-FIRED BOILERS - MODERATE CONTROL

TABLE A1.14.	POINT	VALUE NO×-ON	LY PROCESS	RATINGS:	GAS-FIRED	BOILERS	- STRINGENT	CONTROL

	Performance	Operation and maintenance	Environ- mental impact	Economic impact	Energy/ material impact	^	Relia- bility	Status of development	Adapt- ability	Compati- bility	Total
SCR Fixed Packed Bed	8	7	8	8	10	4	14	16	3	5	83
SCR Moving Bed	8	5	8	8	10	4	14	16	3	5	81
SCR Pa rallel Flow	8	7	8	8	10	4	14	16	3	5	83
Absorption - Oxidation	*										

*Not Applicable - Does not meet removal requirements.

	Performance	Operation and maintenance	Environ- mental impact	Economic impact	Energy/ material impact	Boiler operation and safety	Relia- bility	Status of development	Adapt- ability	Compati- bility	Total
SCR Fixed Packed Bed	10	7	8	14	10	4	14	16	3	5	87
SCR Moving Bed	10	5	8	14	10	4	14	16	3	5	85
SCR Parallel Flow	10	7	8	14	10	4	14	16	3	5	87
Absor pti on - Oxi dati on	8	4	7	2	6	2	8	9	3	5	58

TABLE A1.15. POINT VALUE NO_X-ONLY PROCESS RATINGS: GAS-FIRED BOILERS - INTERMEDIATE CONTROL

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APPENDIX 2

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EXAMPLE OF TECHNIQUE FOR ECONOMIC SCALING

SAMPLE CALCULATION

Most of the available economic data are for utility boilers. Various base capacities are utilized in the process developers' economic calculations. For the preliminary economic screening of alternative processes a base capacity of 20 MW was selected to represent industrial boilers. The capital cost figures were adjusted to a 20 MW cost by using the six-tenths rule.¹

$$I = I_{B} \left(\frac{Q}{Q_{B}} \right)^{-6}$$
 (Reference 1)

where I = estimated 20 MW investment I_B = known base investment Q = 20 MW Q_B = known base capacity

A sample calculation for the SCR-parallel passage (coal) process is shown below.

Size = 250 MW
Investment = \$4,000,000
I = (\$4,000,000)
$$\left(\frac{20 \text{ MW}}{250 \text{ MW}}\right)^{.6}$$

= (\$4,000,000) (.220)
= \$879,000
Capital Cost (\$/kW) = $\frac{$879,000}{(20 \text{ MW})\left(\frac{1000 \text{ kW}}{\text{MW}}\right)}$
= \$44/kW

A2-2

The results of these calculations for all systems considered are contained in Tables A2.1 and A2.2.

Reference: 1. Rudd, D. F., C. C. Watson, Strategy of Process Engineering, 1968, p.121.

Note: The six-tenths factor was used only during this preliminary phase to put reported costs for the various processes on a consistent basis. The six-tenths rule was not used in the development of process economics for this report.

	Coal	0i1	Gas
SCR fixed packed bed	130*	70	27
SCR parallel flow	44	39	NA
SCR moving bed	92*	70	NA
SCR parallel flow, NO_{x}/SO_{x}	475	NA	NA
Adsorption	215	NA	NA
Electron Beam Radiation	302	NA	NA
Absorption - Reduction	413	187	NA
Absorption - Oxidation	NA	NA	NA
Oxidation - Absorption - Reduction	NA	231	NA
Oxidation - Absorption	NA	NA	254

TABLE A2.1. ECONOMICS - CAPITAL COST, \$/kW

*Includes particulate removal

NA = Not Available

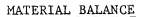
	Coal	0i1	Gas
SCR fixed packed bed	2.1	1.9	1.2
SCR parallel flow	1.5	NA	NA
SCR moving bed	2.0	1.8	NA
SCR parallel flow, NO $_{\rm x}/$ SO $_{\rm x}$	5.0	NA	NA
Adsorption	2.3	NA	NA.
Electron Beam Radiation	NA	NA	NA
Absorption - Reduction	7.4	5.4	NA
Absorption - Oxidation	NA	NA	NA
Oxidation - Absorption - Reduction	NA	6.4	NA
Oxidation - Absorption	NA	NA	NA

TABLE A2.2. ECONOMICS - OPERATING COST, MILLS/kWh

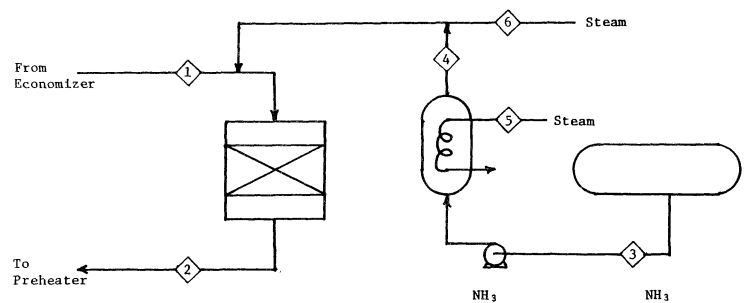
NA = Not Available

APPENDIX 3

MATERIAL BALANCES FOR COAL-FIRED BOILERS

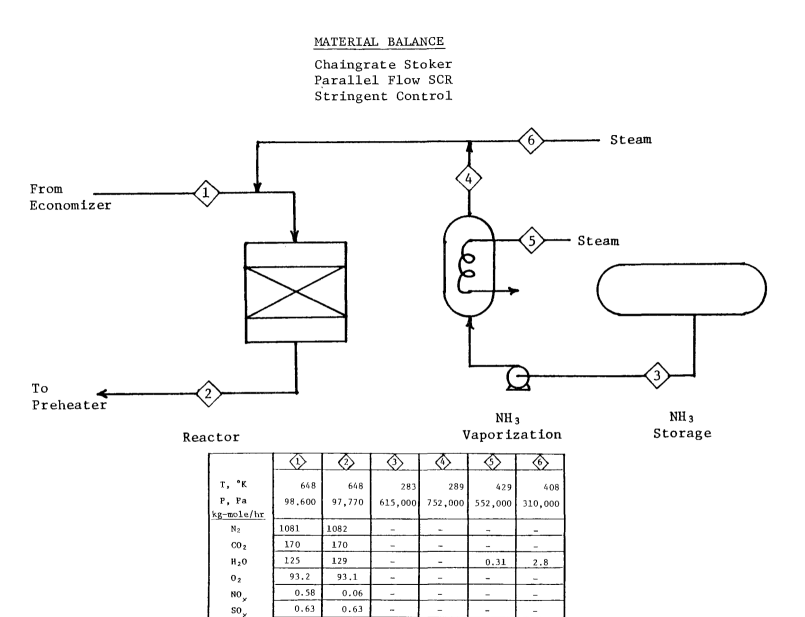


Underfeed Stoker Parallel Flow SCR Intermediate Control



ctor		Vaporization						
			\Diamond	$\langle \bullet \rangle$	$\langle \rangle$	6		
т, °К	648	648	283	289	429	408		
P, Pa <u>kg-mole/hr</u>	98,600	98,040	615,000	752,000	552,000	310,000		
N ₂	431	431	-	~	-	-		
CO 2	67.9	67.9	_	-	-	_		
H ₂ O	49.9	51.2	-	-	0.10	0.94		
02	37.2	37.2	~	_	~	-		
NO×	0.23	0.05	-	-	-	-		
so	0.25	0.25	-	-	-	-		
NH 3	-	<.01	0.19	0.19	-	-		

Storage



0.01

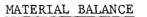
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NH₃

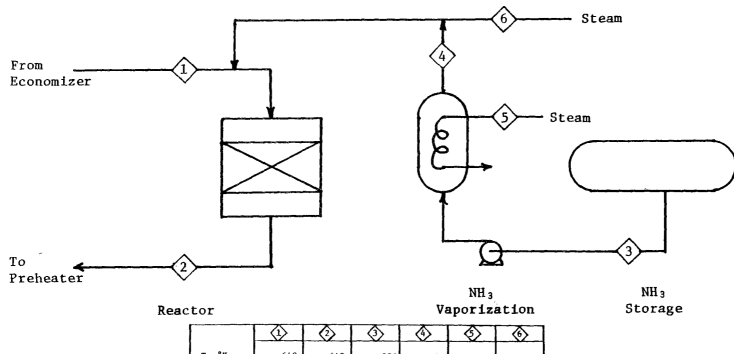
0.55

0.55

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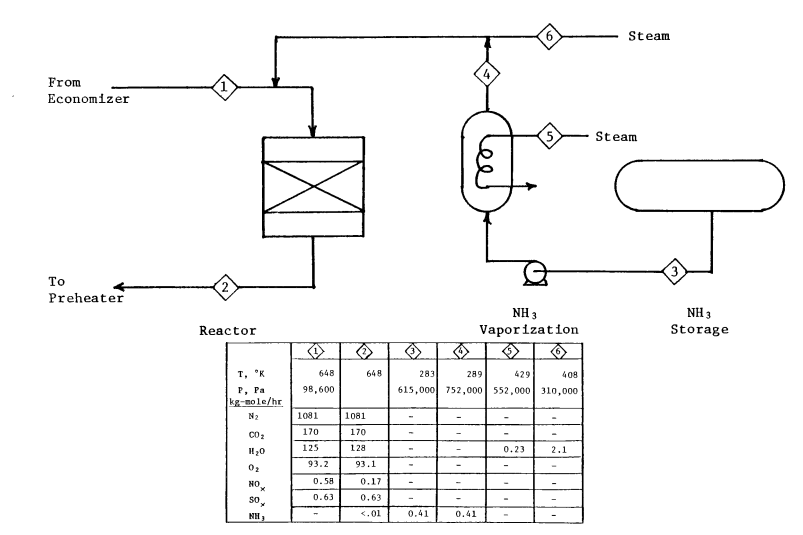


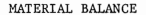
Chaingrate Stoker Parallel Flow SCR Intermediate Control



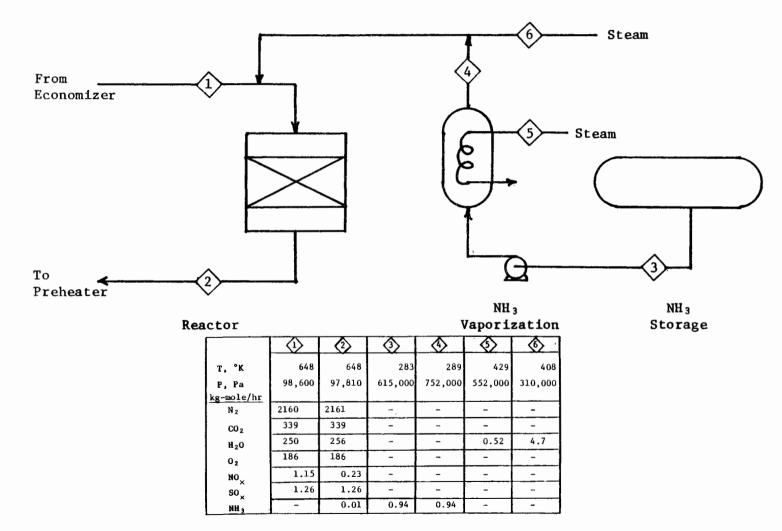
			\bigcirc		\odot	\odot
т, °К	648	648	283	289	429	408
P, Pa kg-mole/hr	98,600		615,000	752,000	552,000	310,000
N2	1081	1082	~		-	-
C0 2	170	170	-	~	-	~
HzO	125	128	-	-	0.26	2.4
0 z	93.2	93.1	-	-	-	-
NO _×	0.58	0.12	-	-	-	-
sox	0.63	0.63	-	-	-	-
NH	-	<.01	0.48	0.48	-	-

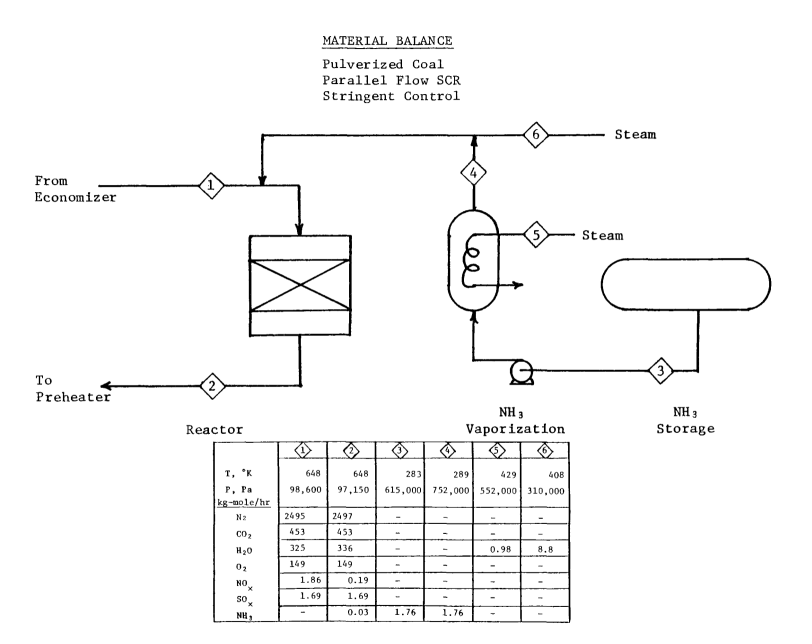
Chaingrate Stoker Parallel Flow SCR Moderate Control

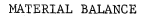




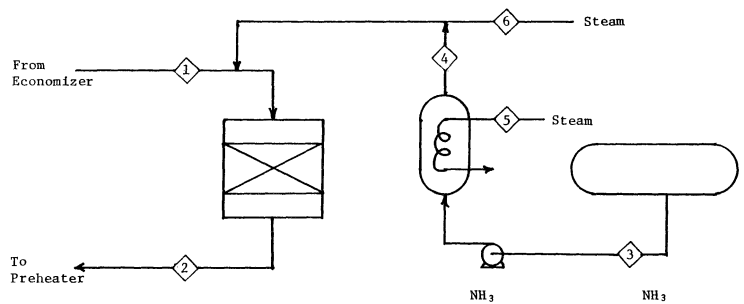
Spreader Stoker Parallel Flow SCR Intermediate Control







Pulverized Coal Parallel Flow SCR Moderate Control

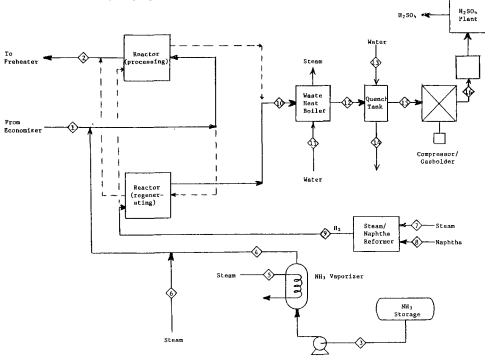


ctor				v	aporiz	
			3	\bigcirc	$\langle \rangle$	\diamond
т, °к	648	648	283	288	429	408
P, Pa	98,600	97,730	615,000	752,000	552,000	310,000
kg-mole/hr						
N ₂	2495	2496 -	-	-	_	-
CO 2	453	453	1	-	-	-
H₂O	325	334	1	-	0.73	6.6
02	149	149	-	-	-	-
NOX	1.86	0.56	-	_	-	-
sox	1.69	1.69	-	-	-	-
NH 3		<.01	1.32	1.32	-	-

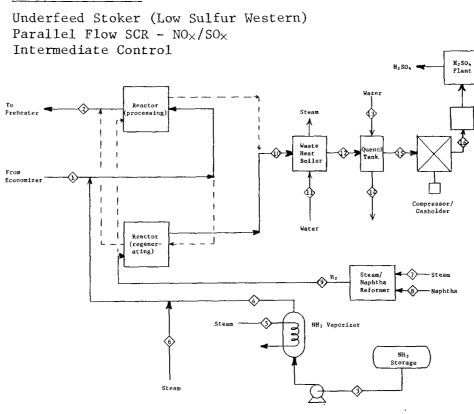
Storage

A3-8

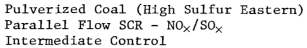
Underfeed Stoker (High Sulfur Eastern) Parallel Flow SCR - NO_X/SO_X Intermediate Control

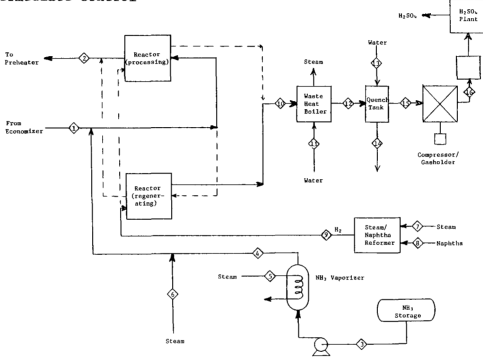


			3	4	5	6	\Diamond	8	(9)			12	(13)	14	15	16
т, °к	648	648	283	289	429	405	789	666	704	700	411	450	300	346	389	389
P, Pa	98,600	96,150	615,000	752,000	552,000	310,000	665,000	860,000	274,000	177,000	345,000	170,000	276,000	276,000	165,000	∿i.3x107
kg-mole/hr														ļ		
N ₂	428	428	-	-	-	-	-	-		-	-	-	-	-		-
CO2	62.2	62.2	-	-	-	-	-	-	-	-	~	-	-	-	-	-
HzO	42.2	43.6	-	-	0.14	1.18	7.20	-	2.52	9.71	2.93	9.71	7.54	7.54	9.71	-
Oz	36.8	36.8	-	-	-	-	-	-	-	-	-	-	-	-	-	-
NO×	0.20	0.04	-	-	-	-	-	-	-	-	-	•	-	-	-	-
so _×	1,20	0.18	-	-	-	-	-	-	-	1.02	-	1.02	-	<0.001	1.02	1.02
NH 3	-	0.01	0.24	0.24	-	-	-	-	-	-	-	-	-	-	-	-
H2	-	-	-	-	-	-	-	-	7.19	-	-	-	-	-	-	-
Naphthe	-	-	-	-	-	-	-	0.34	-	-	-	-	-	-	-	-

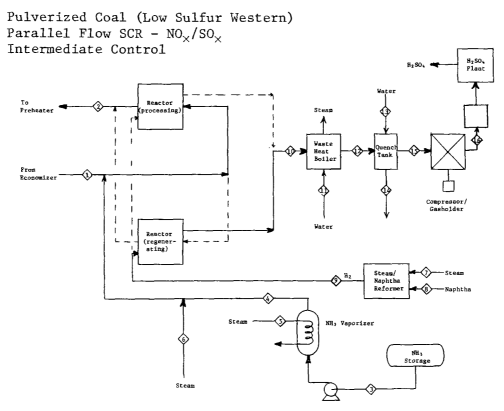


			$\langle \rangle$	$\langle 4 \rangle$	$\langle S \rangle$	6	$\langle i \rangle$	8	9	10			(13)	14	15	16
т, °к	648	648	283	289	429	408	789	666	704	700	411	450	300	346	389	389
P, Pa	98,600	96,150	615,000	752,000	552,000	310,000	665,000	860,000	274,000	177,000	345,000	170,000	276,000	276,000	165,000	~1.3x107
kg~mole/hr		L														
N ₂	431	431	- 1	-	-	- 1	-	-	-	-	-	-	-]	-	-
CO 2	67.9	67.9	-	-	-	- 1	-	-	-	-	~	-	-	-	-	-
H ₂ O	49.9	51.6	-	-	0.16	1.38	1.51	-	0.53	2.03	0.63	2.03	1.59	1.59	2.03	-
0z	37.2	37.2	-	-	-	-	-	-	-	-	-	-	-	-	-	-
NOx	0.23	0.05	-	-	-	-	-	-	-	-	-	-	-	-	-	-
so,	0.25	0.04	-	-	-	-	-	-	-	0.21	-	0.21	-	<0.0001	0.21	0.21
NH 3	-	0.01	0.28	0.28	-	-	-	-	-	-	-	-	-	-	-	-
H2	-	-	-	-	-	-	-	-	1.50		-	-	-	-	-	-
Naphtha	-	-	L	-	-	_	-	0.07	-	-	-	-	-	-	-	-





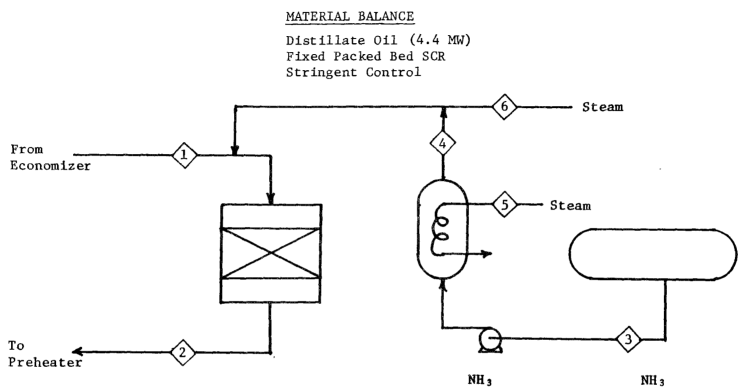
		2	\odot	4	5	\diamond	$\langle \rangle$	8	\diamond					14	15	(16)
т, °к	648	648	283	289	429	408	789	666	704	700	411	450	300	346	389	389
P, Pa	98,600	96,150	615,000	752,000	552,000	310,000	665,000	860,000	274,000	177,000	345,000	170,000	276,000	276,000	165,000	∿1.3×10 ⁷
kg-mole/hr		L			<u> </u>											
N 2	2470	2470	-	-	-	-	-	-	-	-	~	-	-	-	-	-
CO2	415	415	-	-	-	-	-	· _	-	-	-	-	-	-	-	-
H20	271	282	-	-	1.11	9.40	48.0	-	16.8	64.7	19.6	64.7	50.4	50.4	64.7	-
02	148	148	-	-	-	-	-	-	-	-	-	-	-	-	-	-
NO _x	1.57	0.31	-	-	-	-	-	-	-	-	-	-	-	-	~	-
so _×	7.99	1.20	-	-	-	-	-	-	-	6.79	-	6.79	-	<0.01	6.79	6.79
NH 3	-	0.07	1.88	1.88	-	-	-	-	-	-	-	-	-	-	-	-
H 2	-	-	-	-	-	-	-	-	47.9	-	-	-	-	-	-	-
Naphtha	-	-	-	-	-	-	-	2.23	-	-	-	-	-	-	-	-



			$\langle \rangle$	4	5	6	\Diamond		$\langle \mathfrak{I} \rangle$				(13)	4	<u>(15)</u>	16
т, °к	648	648	283	289	429	408	789	666	704	700	411	450	300	346	389	389
2, Pa	98,600	96,150	615,000	752,000	552,000	310,000	665,000	860,000	274,000	177,000	345,000	170,000	276,000	276,000	165,000	∿1.3x107
kg-mole/hr																
Nz	2490	2490	-	-	-	-	-	-	-	-	-	-	-	-	-	-
CO2	453	453	-	-	-	-	-	-	~	-	-		-	-	-	-
H2O	325	340	-	-	1.32	11.2	10.2	-	3.55	13.7	4.14	13.7	10.6	10.6	13.7	-
02	149	149	-	-	-	-	-	-	-	-	-		-	-	-	-
NO _×	1.86	0.37	-	-	-	-	-	-	-	-	-	-	-	-	-	-
SO _x	1.69	0.25	-	-	-	-	-	-	-	1.44	-	1.44	-	<0.001	1.44	1.44
NH 3	-	0.07	2.23	2.23	-	-	-	-	-	-	-	-	-	-	-	-
H2	-	-	-	-	-	-	-	-	10,1	-	-	-	-	-	-	-
Naphtha	-	- 1	-	-	-	-	-	0.47	-	-	-	-	-	-	~	

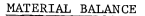
APPENDIX 4

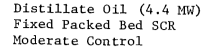
MATERIAL BALANCES FOR OIL-FIRED BOILERS

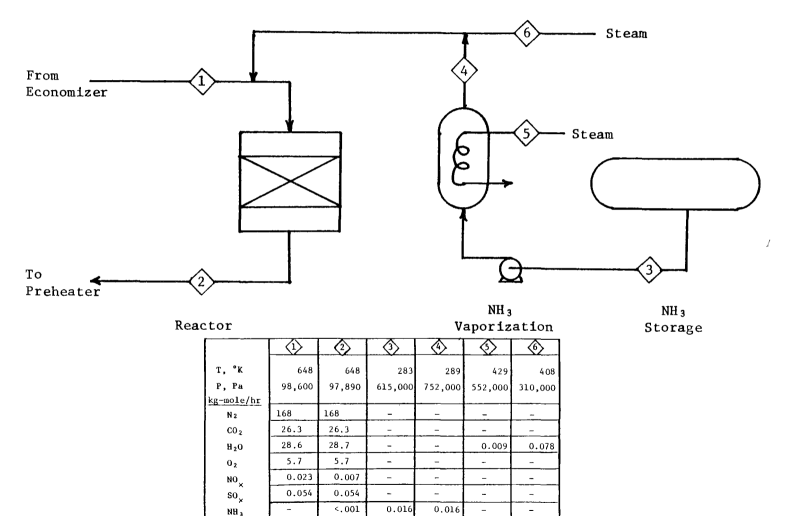


leactor				V	aporia	-
	$\langle i \rangle$	$\langle \rangle$	\odot	\diamond	\odot	<u>ن</u>
τ, °κ	648	648	283	289	429	408
P, Pa	98,600	97,420	615,000	752,000	552,000	310,000
kg-mole/hr						
N 2	168	168	~	-	-	-
CO₂	26.3	26.3	-	-	-	-
H₂O	28.6	28.7	-	-	0.011	0.101
02	5.7	5.7	-	-	~	-
NO _x	0.023	0.002	-	-	-	-
sox	0.054	0.054	-	-	-	-
NH ₃	-	<.001	0.020	0.020	~	-

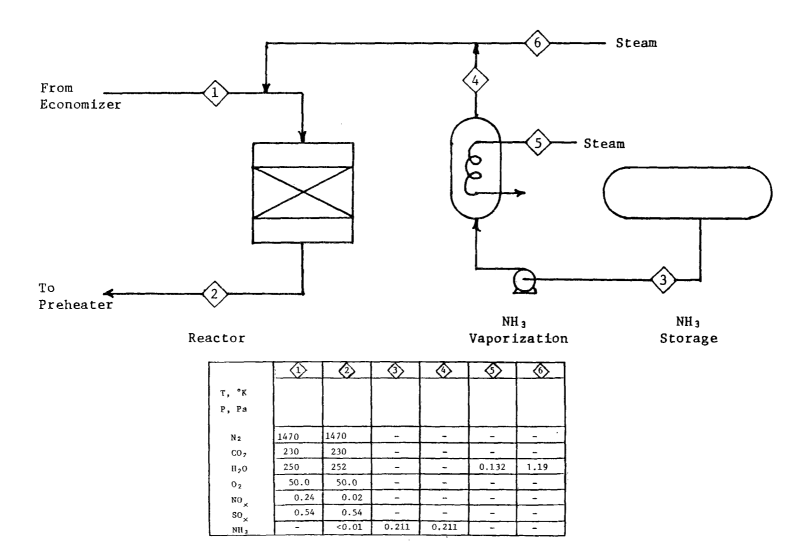
Storage



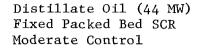


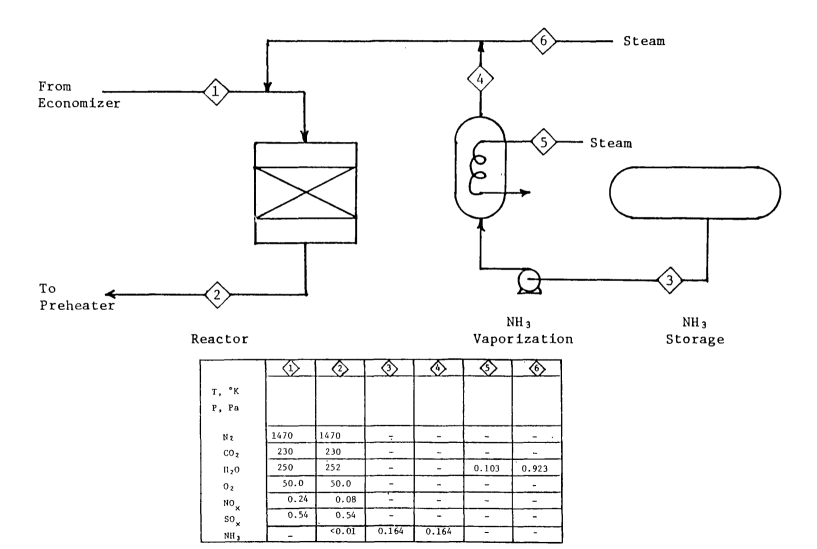


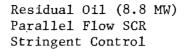
Distillate Oil (44 MW) Fixed Packed Bed SCR Stringent Control

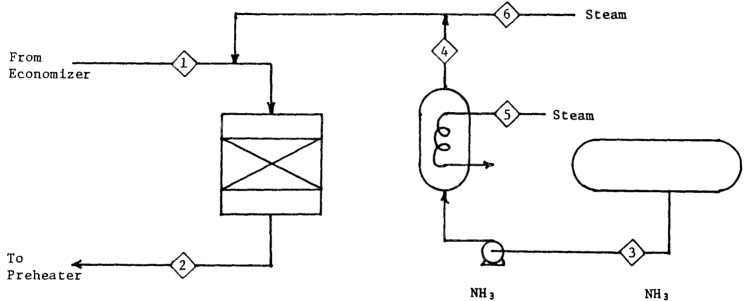










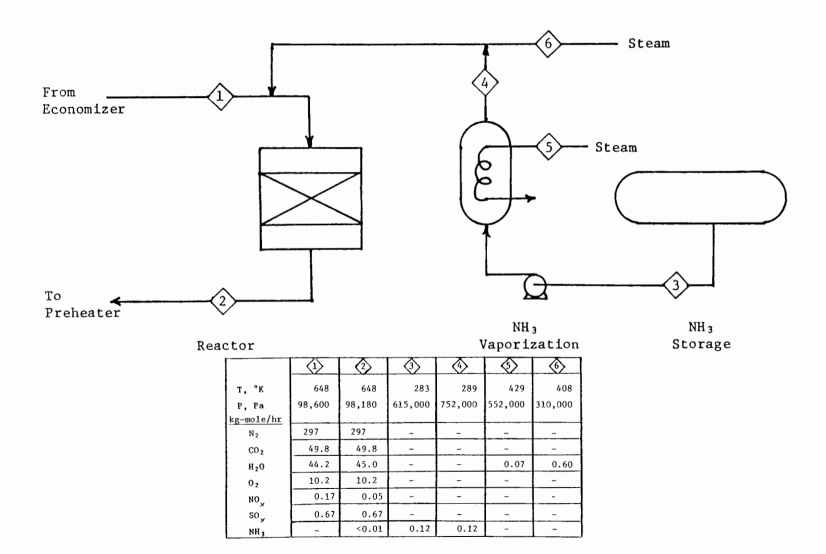


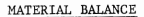
actor				Ve	aporiz	ation
•	$\langle \rangle$	2	\Diamond	٨	\Diamond	\diamond
Т, °К	648	648	283	289	429	408
P, Pa	98,600	97,890	615,000	752,000	552,000	310,000
kg-mole/hr						
N 2	297	297	-	-	-	
CO ₂	49.8	49.8			-	-
H ₂ O	44.2	45.2	-		0.09	0.78
02	10.2	10.2	-	-		
NO _x	0.17	0.02	-	-	~	-
sox	0.67	0.67	-	-	-	-
NH 3	-	<0.01	0.16	0.16	-	-

Storage

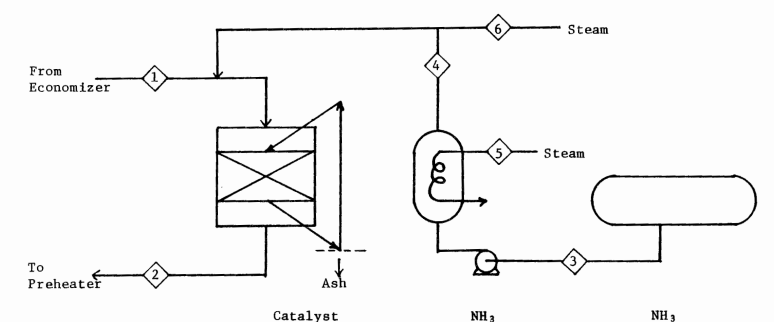


Residual Oil (8.8 MW) Parallel Flow SCR Moderate Control

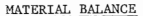




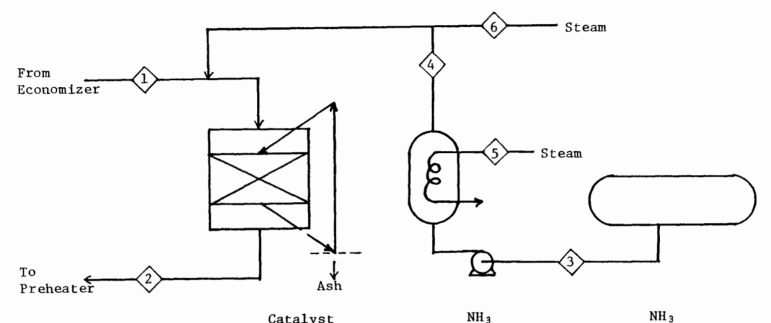
Residual Oil (8.8 MW) Moving Bed SCR Stringent Control



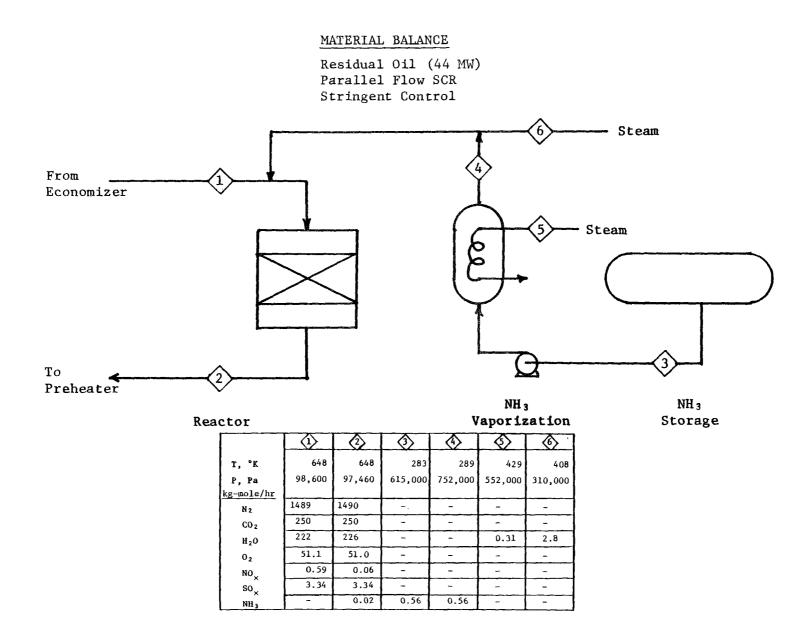
Reactor		Catal Handl	•		Vap	NH3 orizat	ion	NH ₃ Storage
		$\langle \rangle$	$\langle \rangle$	\Diamond	$\langle \! \! \ \ \ \ \ \ \ \ \ \ \ \ $	$\langle \rangle$	$\langle \diamond \rangle$	
	т, °к	648	648	283	289	429	408	
	P, Pa	98,600	98,160	615,000	752,000	552,000	310,000	
	kg-mole/hr							
	N ₂	297	297	-	-	-	-	
	CO2	49.8	49.8	-	-	-	-	
	H ₂ O	44.2	45.2	-	-	0.09	0.78	
	02	10.2	10.2	-	-	-	-	
	NO,	0.17	0.02	-	-	-	-	
	sox	0.67	0.67	-	-	-	-	
	NH 3	-	<0.01	0.16	0.16	-	-	



Residual Oil (8.8 MW) Moving Bed SCR Moderate Control

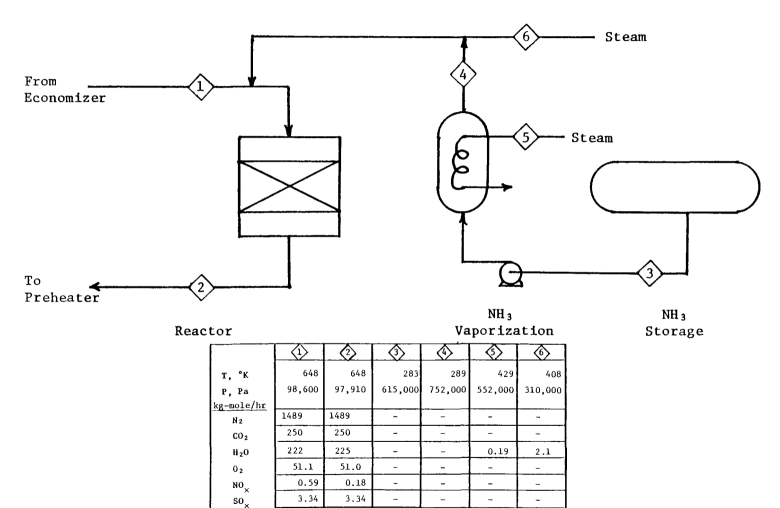


Reactor		alyst dling		Va	NH: aporiza	-		NH ₃ Storage
				\bigcirc	$\langle 4 \rangle$	$\langle S \rangle$	$\langle \delta \rangle$	
	т, °К	648	648	283	289	429	408	
	P, Pa	98,600	98,340	615,000	752,000	552,000	310,000	
	N 2	297	297		_	-	-	
	COz	49.8	49.8	-	-	-	-	
	H ₂ O	44.2	45.0	-	-	0.07	0.60	
	02	10.2	10.2		-	-	-	
	NO _×	0.17	0.05				_	
	sox	0.67	0.67	-	-		-	
	NR 3	-	<0.01	0.12	0.12			





Residual Oil (44 MW) Parallel Flow SCR Moderate Control



<.01

-

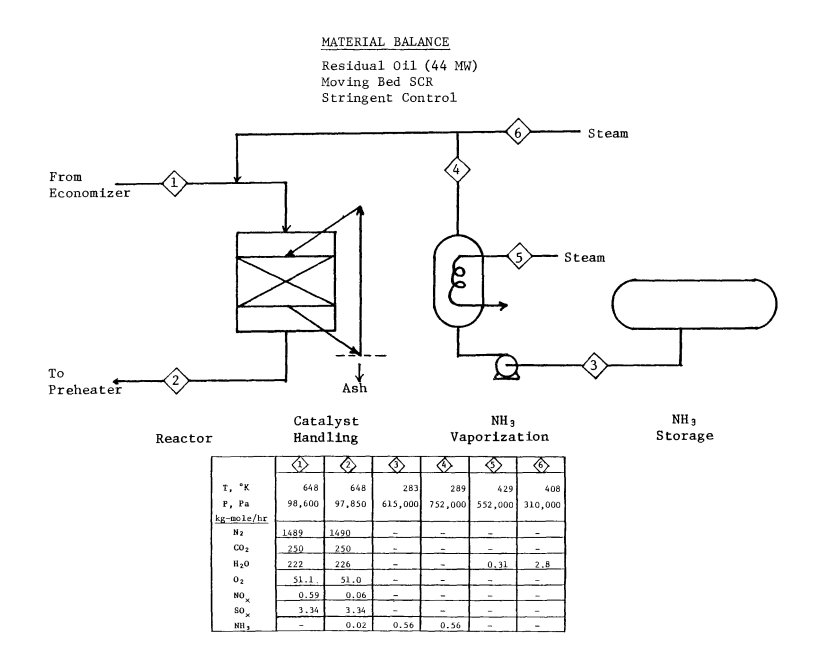
NH 3

0.42

0.42

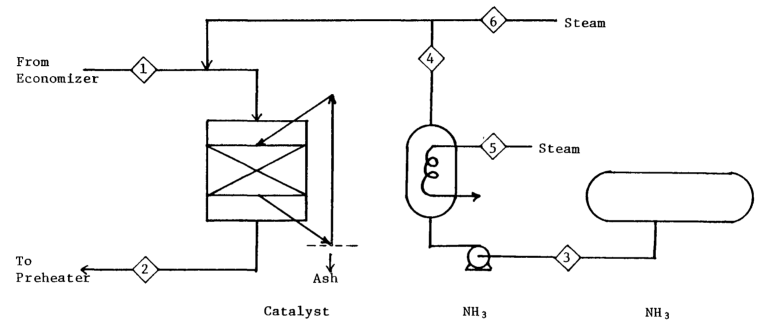
-

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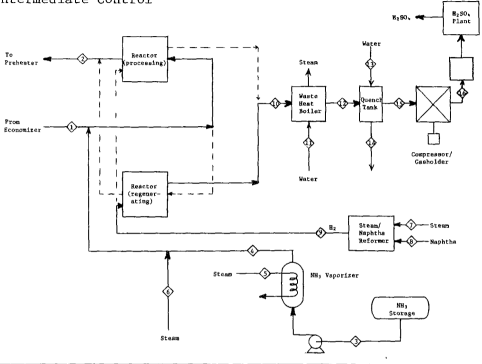
Residual Oil (44 MW) Moving Bed SCR Moderate Control



Reactor		Cata] Hand]	-		Vap	NH ₃ orizat	ion
		$\langle \hat{\mathbf{x}} \rangle$		\Diamond	$\langle \! \rangle$	$\langle \rangle$	6
	т, °к	648	648	283	289	429	408
	P, Pa kg-mole/hr	98,600	98,150	615,000	752,000	552,000	310,000
	N 2	1489	1489	-	-	-	-
	CO ₂	250	250	-	-	-	-
	H20	222	225	-	-	0.19	2.1
	02	51.1	51.0	-	-	-	-
	NOX	0.59	0.18	-	-	-	_
	so	3.34	3.34	-	-	-	-
	NH 3	-	<.01	0.42	0.42	-	-

Storage

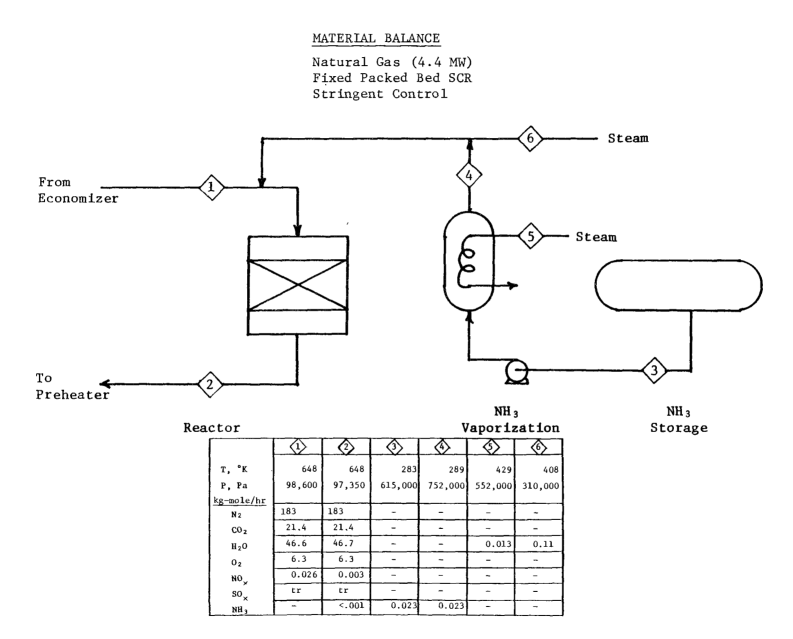
Residual Oil (44 MW) Parallel Flow SCR - NO_X/SO_X Intermediate Control



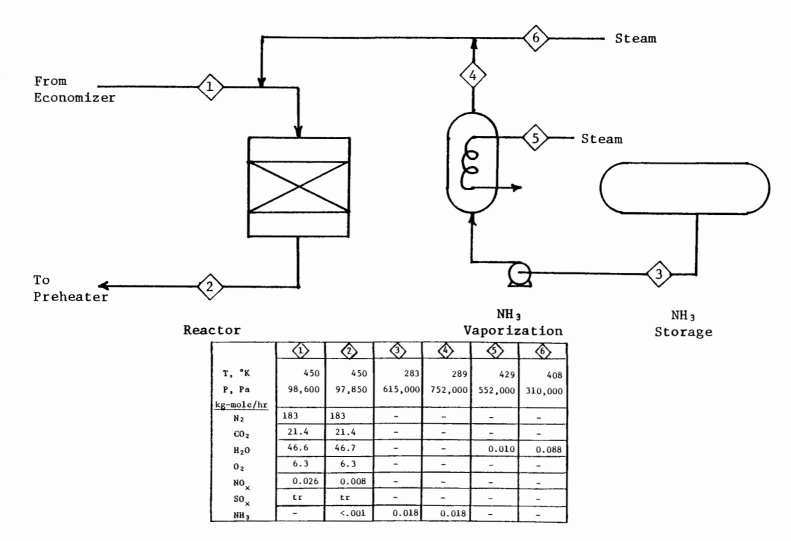
[$\langle \mathbf{j} \rangle$	$\langle 4 \rangle$	$\langle \rangle$	6	$\langle i \rangle$	<u></u>	<i>(</i>)				(1)	14		16
г, °х	648	648	283	289	429	408	789	666	704	700	411	450	300	346	389	389
2, Pa	98,600	96,150	615,000	752,000	552,000	310,000	665,000	860,000	274,000	177,000	345,000	170,000	276,000	276,000	165,000	∿1.3x10 ⁷
kg-mole/hr		 i														
N ₂	1490	1490	-	-	-	-	-	-	-	-	-	-	-	-	-	-
CO ₂	250	250	-	-	-	-	-	-	-	-	-	-	~	-	-	-
H20	222	226	-	-	0.42	3.54	20.1	-	7.01	27.0	8.17	27.0	21.0	21.0	27.0	-
0z	51.1	51.1	-	-	-	-	-	-	-	-	-	-	-	-	-	-
NOx	0.59	0.12	-	-	-	-	-	-	-	-	-	-	-	-	-	-
s0 _×	3.34	0.50	-	-	-	-	-	-	-	2.84	-	2.84	-	<0.01	2.84	2.84
NH 3	-	0.04	0.71	0.71	-	-	-	-	-	-	-	-	-	-	-	-
H 2	-	-	-	-	-	~	-	-	20.0	-	-	-	-	-	-	-
Naphtha	-] -] -] -	- 1	-	-	0.93	-	-	-	-	-	~	-	-

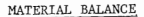
APPENDIX 5

MATERIAL BALANCES FOR NATURAL GAS-FIRED BOILERS

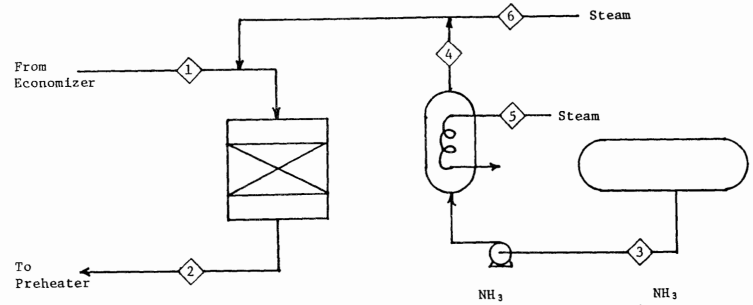


Natural Gas (4.4 MW) Fixed Packed Bed SCR Moderate Control





Natural Gas (44 MW) Fixed Packed Bed SCR Stringent Control



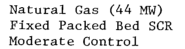
Reactor

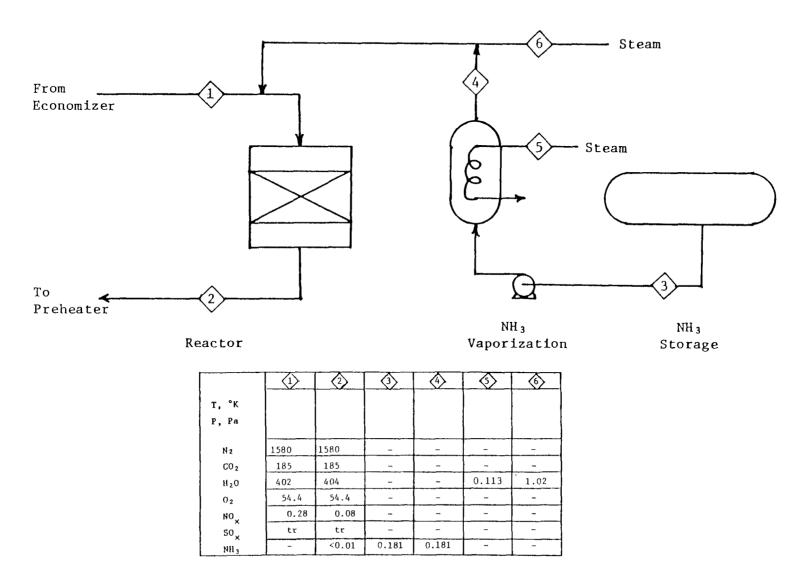
Vaporization

Storage

		$\langle 2 \rangle$	$\langle \mathfrak{T} \rangle$	$\langle \! \rangle$	$\langle S \rangle$	$\langle \bullet \rangle$
т, ⁰К						
P, Pa						
Nz	1580	1580		-	_	
CO ₂	185	185	-	-	-	-
H ₂ O	402	404	-	-	0.146	1.32
02	54.4	54.4	~	-	-	-
NO _×	0.28	0.02	-	-	-	-
sox	tr	tr	-	-	-	-
NH 3	-	<0.01	0.233	0.233	-	-

MATERIAL BALANCE





APPENDIX 6

CAPITAL COST BREAKDOWNS

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Table A6-1. CAPITAL COSTS

Boiler type: Underfeed Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR Control level: Intermediate

Equipment cost

Basic equipment (includes freight) <u>29.</u> Required auxiliaries <u>24</u> .	270	
Total equipment cost _53.	,470	
Installation costs, direct		
Piping14Insulation1Painting1Electrical1Instruments2	,290 ,080 ,760 330 ,350 ,430 ,500	
Total installation cost 42	,740	
Total Direct Costs (equipment + installat	ion) <u>96,210</u>	
Installation costs, indirect		
Construction and field expense 9 Construction fees 9 Start-up 1	,892 ,621 ,621 ,924 ,000	
Total Indirect Costs	67,058	
Contingencies	32,654	
Total lurnkey Costs (direct+indirect+cont	ingencies) <u>195,922</u>	
Land	490	
Working capital	16,675	
GRAND TOTAL (turnkey + land + working capital)		

\$213,090

Table A6-2. CAPITAL COSTS

Boiler type: Chaingrate Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR Control level: Stringent

Equipment cost Basic equipment (includes freight) 61,510 Required auxiliaries 81,000 Total equipment cost 142,510 Installation costs, direct Foundations and supports 9,190 29,010 Piping 3,900 Insulation Painting 700 Electrical 2,570 4,820 Instruments Installation labor 41,310 82,310 Total installation cost Total Direct Costs (equipment + installation) 224,820 Installation costs, indirect Engineering 43,892 Construction and field expense 22,482 Construction fees 22,482 Start-up 4,496 Performance tests 2.000 95,352 Total Indirect Costs 64,034 Contingencies Total Turnkey Costs (direct+indirect+contingencies) 384,206 961 Land 28,129 Working capital \$413,300 GRAND TOTAL (turnkey + land + working capital)

A6-3

Table A6-3. CAPITAL COSTS

Boiler type: Chaingrate Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR

Control level: Intermediate Equipment cost Basic equipment (includes freight) 49,880 Required auxiliaries 60,600 Total equipment cost 110,480 Installation costs, direct Foundations and supports 7,430 Piping 23,470 Insulation 3,090 Painting 570 Electrical 2,100 Instruments 3,930 Installation labor 32,920 73,510 Total installation cost Total Direct Costs (equipment + installation) Installation costs, indirect 43,892 Engineering 18,399 Construction and field expense Construction fees 18,399 Start-up 3,680 Performance tests 2,000 Total Indirect Costs Contingencies Total Turnkey Costs (direct+indirect+contingencies)324,432 Land

Working capital 24,294 GRAND TOTAL (turnkey + land + working capital)

\$349,540

183,990

86,370

54,072

811

Table A6-4. CAPITAL COSTS

Boiler type: Chaingrate Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR Control level: Moderate

Equipment cost

Basic equipment (includes freight) <u>42,000</u> Required auxiliaries <u>48,500</u>		
Total equipment cost90,500		
Installation costs, direct		
Foundations and supports6,150Piping19,270Insulation2,490Painting480Electrical1,780Instruments3,260Installation labor27,170		
Total installation cost <u>60,600</u>		
Total Direct Costs (equipment + installation)		
Installation costs, indirect		
Engineering43,892Construction and field expense15,110Construction fees15,110Start-up3,022Performance tests2,000		
Total Indirect Costs	79,134	
Contingencies	46,047	
Total Turnkey Costs (direct+indirect+contingencies) 276,281		
Land	691	
Working capital	21,894	
GRAND TOTAL (turnkey + land + working capital)		

\$298,870

Table A6-5. CAPITAL COSTS

Boiler type: Spreader Stoker Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR Control level: Intermediate

Equipment cost Basic equipment (includes freight) 74,330 Required auxiliaries 121,000 Total equipment cost 195.330 Installation costs, direct 10,260 Foundations and supports 32,610 Piping Insulation 4.130 Painting 800 Electrical 2,950 Instruments 5.600 Installation labor 49,790 Total installation cost 106,140 Total Direct Costs (equipment + installation) 301,470 Installation costs, indirect Engineering 43,892 Construction and field expense 30,147 Construction fees 30,147 Start-up 6,029 Performance tests 2,000 Total Indirect Costs 112,215 Contingencies 82,737 Total lurnkey Costs (direct+indirect+contingencies) 496,422 Land 1,241 Working capital 36,958 GRAND TOTAL (turnkey + land + working capital)

Table A6-6. CAPITAL COSTS

Boiler type: Pulverized Coal Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR Control level: Stringent

Equipment cost

Basic equipment (includes freight Required auxiliaries	104,470 188,000	
Total equipment cost	292,470	
Installation costs, direct		
Foundations and supports Piping Insulation Painting Electrical Instruments Installation labor	$ \begin{array}{r} 14,120 \\ 44,030 \\ 5,560 \\ 1,090 \\ 4,020 \\ 7,770 \\ 69,860 \\ \end{array} $	
Total installation cost		
Total Direct Costs (equipment + insta	11ation) <u>438,920</u>	
Installation costs, indirect		
Engineering Construction and field expense Construction fees Start-up Performance tests	<u>43,892</u> <u>43,892</u> <u>43,092</u> <u>8,778</u> <u>2,000</u>	
Total Indirect Costs	142,454	
Contingencies		
Total Turnkey Costs (direct+indirect+contingencies) <u>697,64</u> 9		
Land	<u> 1,744</u>	
Working capital	53,013	
GRAND TOTAL (turnkey + land + working capital)		

Table A6-7. CAPITAL COSTS

Boiler type: Pulverized Coal Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR Control level: Moderate

Equipment cost

Equipment cost		
Basic equipment (includes freight Required auxiliaries		
Total equipment cost	191,470	
Installation costs, direct		
Foundations and supports Piping Insulation Painting Electrical Instruments Installation labor	$ \begin{array}{r} 10,540 \\ 33,160 \\ 4,130 \\ 830 \\ 3,050 \\ 6,110 \\ 49,780 \\ \end{array} $	
Total installation cost	107,600	
Total Direct Costs (èquipment + insta	allation) <u>299,070</u>	
Installation costs, indirect		
Engineering Construction and field expense Construction fees Start-up Performance tests	<u>43,892</u> <u>29,907</u> <u>29,907</u> <u>5,981</u> <u>2,000</u>	
Total Indirect Costs	111,687	
Contingencies	82,151	
Total Turnkey Costs (direct+indirect+contingencies) 492,908		
Land	<u> 1,23</u> 2	
Working capital	<u> </u>	
GRAND TOTAL (turnkey + land + workin	g capital)	\$531 ,9 00

Table A6-8. CAPITAL COSTS

Boiler type: Underfeed Stoker Fuel: High Sulfur Eastern Coal Control technique: Parallel Flow SCR (NO_×/SO_×) Control level: Intermediate

Total Direct Costs (equipment + installation)	1,284,000		
Installation costs, indirect			
Engineering373,400Construction and field expense128,400Construction fees128,400Start-up25,700Performance tests4,000			
Total Indirect Costs	660,000		
Contingencies	389,000		
Total Turnkey Costs (direct+indirect+contingencies) 2,333,000			
Land	6,000		
Working capital	64,000		
GRAND TOTAL (turnkey + land + working capital)	\$2,403,000		

Table A6-9. CAPITAL COSTS

Boiler type: Underfeed Stoker Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR (NO_×/SO_×) Control level: Intermediate

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Total Direct Costs (equipment + insta	llation)	. 648,800	
Installation costs, indirect			
Engineering Construction and field expense Construction fees Start-up Performance tests	373,400 64,900 64,900 13,000 4,000		
Total Indirect Costs		520,200	
Contingencies		233,800	
Total Turnkey Costs (direct+indirect+contingencies) 1,403,000			
Land		4,000	
Working capital		43,000	
GRAND TOTAL (turnkey + land + working	capital)		<u>\$1,450,000</u>

TABLE A6-10. CAPITAL COSTS

Boiler type: Pulverized Coal Fuel: High Sulfur Eastern Coal Control technique: Parallel Flow SCR (NO_×/SO_×) Control level: Intermediate

Total Direct Costs (equipment + installation)	3,734,000	
Installation costs, indirect		
Engineering373,400Construction and field expense373,400Construction fees373,400Start-up74,700Performance tests4,000		
Total Indirect Costs	1,199,000	
Contingencies	987,000	
Total Turnkey Costs (direct+indirect+contingencies	s) <u>5,920,000</u>	
Land	15,000	
Working capital	180,000	
GRAND TOTAL (turnkey + land + working capital)	\$6,115,00	0

TABLE A6-11. CAPITAL COSTS

Boiler type: Pulverized Coal Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR (NO_X/SO_X) Control level: Intermediate

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Total Direct Costs (equipment + insta	allation)	1,793,000	
Installation costs, indirect			
Engineering Construction and field expense Construction fees Start-up Performance tests	373,400 179,300 179,300 35,900 4,000		
Total Indirect Costs		772,000	
Contingencies		513,000	
Total Turnkey Costs (direct+indirect-	-contingencies)	3,078,000	
Land	_	8,000	
Working capital		78,000	
GRAND TOTAL (turnkey + land + working	g capital)		\$3,164,000

Table A6-12. CAPITAL COSTS

Boiler type: Firetube (4.4 MW_t) Fuel: Distillate Oil Control technique: Fixed Packed Bed SCR Control level: Stringent

Equipment cost

Iquipment cost		
Basic equipment (includes freight Required auxiliaries) <u>8,230</u> <u>8,580</u>	
Total equipment cost	16,810	
Installation costs, direct		
Foundations and supports Piping Insulation Painting Electrical Instruments Installation labor	850 2,990 360 60 580 470 3,910	
Total installation cost	9,220	
Total Direct Costs (equipment + insta	11ation) <u>26,030</u>	
Installation costs, indirect		
Engineering Construction and field expense Construction fees Start-up Performance tests	<u>43,892</u> <u>2,603</u> <u>2,603</u> <u>521</u> <u>2,000</u>	
Total Indirect Costs	51,619	
Contingencies		
Total Turnkey Costs (direct+indirect+	contingencies) 89,296	
Land	223	
Working capital	9,892	
GRAND TOTAL (turnkey + land + working	; capital)	\$99,410

Table A6-13. CAPITAL COSTS

Boiler type: Firetube (4.4 MW_t) Fuel: Distillate Oil Control technique: Fixed Packed Bed SCR Control level: Moderate

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Equipment cost

Basic equipment (includes freight) 6,230 Required auxiliaries5,120	
Total equipment cost <u>11,350</u>	
Installation costs, direct	
Foundations and supports620Piping2,270Insulation280Painting40Electrical500Instruments350Installation labor2,980	
Total installation cost7,040	<u> </u>
Total Direct Costs (equipment + installation)	18,390
Installation costs, indirect	
Engineering43,892Construction and field expense1,839Construction fees1,839Start-up368Performance tests2,000	<u> </u>
Total Indirect Costs	49,938
Contingencies	10,248
Total Turnkey Costs (direct+indirect+continger	ncies) <u>78,576</u>
Land	196
Working capital	9,402
GRAND TOTAL (turnkey + land + working capital))

Table A6-14. CAPITAL COSTS

Boiler type: Watertube (44 MW_t) Fuel: Distillate Oil Control technique: Fixed Packed Bed SCR Control level: Stringent

Equipment cost Basic equipment (includes freight) 34,030 Required auxiliaries 86,140 Total equipment cost 120,170 Installation costs, direct Foundations and supports 3,280 Piping 10,420 Insulation 1,150 Painting 220 Electrical 1,660 Instruments 1,690 Installation labor 21.300 39,720 Total installation cost Total Direct Costs (equipment + installation) 159,890 Installation costs, indirect 43,890 Engineering Construction and field expense 15,990 Construction fees 15,990 Start-up 3,200 Performance tests 2,000 Total Indirect Costs 81,070 Contingencies 36.140 Total Turnkey Costs (direct+indirect+contingencies) 277.100 690 Land 27,810 Working capital GRAND TOTAL (turnkey + land + working capital)

305,600

Table A6-15. CAPITAL COSTS

Boiler type: Watertube (44 MW_t) Fuel: Distillate Oil Control technique: Fixed Packed Bed SCR Control level: Moderate

Control level: Moderate Equipment cost Basic equipment (includes freight) 25,400 51,680 Required auxiliaries 77,080 Total equipment cost Installation costs, direct Foundations and supports 2,500 Piping 8,000 Insulation 900 Painting 180 Electrical .280 Instruments 1,320 14,930 Installation labor 29,110 Total installation cost Total Direct Costs (equipment + installation) 106,190 Installation costs, indirect Engineering 43,890 Construction and field expense 10,620 Construction fees 10,620 Start-up 2,120 Performance tests 2,000 Total Indirect Costs 69,250 Contingencies 26,320 Total Turnkey Costs (direct+indirect+contingencies)201,760 Land 500 Working capital 21,510 GRAND TOTAL (turnkey + land + working capital)

223,800

Table A6-16. CAPITAL COSTS

Boiler type: Watertube (8.8 MWt) Fuel: Residual Oil Control technique: Parallel Flow SCR Control level: Stringent

Equipment cost

Basic equipment (includes freight) Required auxiliaries	<u>29,100</u> 23,800
Total equipment cost	52,900
Installation costs, direct	
Foundations and supports Piping Insulation Painting Electrical Instruments Installation labor	4,380 14,080 1,870 460 1,240 2,350 18,580
Total installation cost	43,000
Total Direct Costs (equipment + instal	lation) 95,900
Installation costs, indirect	
Engineering Construction and field expense Construction fees Start-up Performance tests	43,890 9,590 9,590 1,920 2,000
Total Indirect Costs	67,000
Contingencies	24,400
Total Turnkey Costs (direct+indirect+c	ontingencies) <u>187,300</u>
Land	500
Working capital	15,100
GRAND TOTAL (turnkey + land + working	capital)

\$202**,**900

Table A6-17. CAPITAL COSTS

Boiler type: Watertube (8.8 MW_t) Fuel: Residual Oil Control technique: Parallel Flow SCR Control level: Moderate

Equipment cost

Basic equipment (includes freight) 20,910 Required auxiliaries 14,300	
Total equipment cost35,210	
Installation costs, direct	
Foundations and supports2,830Piping9,850Insulation1,300Painting240Electrical1,010Instruments1,670Installation labor12,820	
Total installation cost29,720	<i>(</i>) , , , , , , , , , , , , , , , , , ,
Total Direct Costs (equipment + installation)	64,930
Installation costs, indirect	
Engineering43,890Construction and field expense6,490Construction fees6,490Start-up1,300Performance tests2,000	
Total Indirect Costs	60,170
Contingencies	18,770
Total lurnkey Costs (direct+indirect+contingend	ies) <u>143,900</u>
Land	400
Working capital	13,800
GRAND TOTAL (turnkey + land + working capital)	

\$158,100

Table A6-18. CAPITAL COSTS

Boiler type: Watertube (8.8 MW_t) Fuel: Residual Oil Control technique: Moving Bed SCR Control level: Stringent

Equipment cost

Basic equipment (includes freigh Required auxiliaries	t) <u>21,340</u> <u>15,740</u>	
Total equipment cost	37,080	
Installation costs, direct		
Foundations and supports Piping Insulation Painting Electrical Instruments Installation labor	3,070 $9,870$ $1,290$ 250 $1,010$ $1,710$ $12,100$	
Total installation cost	29,300	
Total Direct Costs (equipment + insta	allation) <u>66,380</u>	
Installation costs, indirect		
Engineering Construction and field expense Construction fees Start-up Performance tests	43,890 6,640 6,640 1,330 2,000	
Total Indirect Costs	60,500	
Contingencies	19,030	
Total Turnkey Costs (direct+indirect	+contingencies)145,910	
Land	400	
Working capital	20,350	
GRAND TOTAL (turnkey + land + workin	g capital)	\$ <u>166</u> ,700

Boiler type: Watertube (8.8 MW_t) Fuel: Residual Oil Control technique: Moving Bed SCR Control level: Moderate

Equipment cost

Basic equipment (includes freight Required auxiliaries) 17,440 9,440
Total equipment cost	26,880
Installation costs, direct	-
Foundations and supports Piping Insulation Painting Electrical Instruments Installation labor	2,540 8,090 1,060 200 870 1,390 9,810
Total installation cost	23,960

Total Direct Costs (equipment + installation) 50,840 Installation costs, indirect

Engineering43,890Construction and field expense5,080Construction fees5,080Start-up1,020Performance tests2,000	
Total Indirect Costs	57,070
Contingencies	16,190
Total Turnkey Costs (direct+indirect+contingencie	s)124,100
Land	300
Working capital	19,190
GRAND TOTAL (turnkey + land + working capital)	

\$144,600

Table A6-20. CAPITAL COSTS

Boiler type: Watertube (44 MW_t) Fuel: Residual Oil Control technique: Parallel Flow SCR Control level: Stringent

Equipment cost

Basic equipment (includes freight Required auxiliaries Total equipment cost	$\begin{array}{c} \underline{)} & 74,670 \\ \underline{113,000} \\ 187,670 \end{array}$
Installation costs, direct	
Foundations and supports Piping Insulation Painting Electrical Instruments Installation labor	$ \begin{array}{r} 10,770 \\ 34,420 \\ 4,620 \\ \underline{820} \\ 3,180 \\ 5,660 \\ 50,720 \\ \end{array} $
Total installation cost	110,190
Total Direct Costs (equipment + insta	allation) <u>297,860</u>
Installation costs, indirect	
Engineering Construction and field expense Construction fees Start-up Performance tests	43,892 29,786 29,786 5,957 2,000
Total Indirect Costs	<u> 111,421</u>
Contingencies	61,392
Total Turnkey Costs (direct+indirect+	contingencies) 470,673
Land	<u> 1,17</u> 7
Working capital	31,290
GRAND TOTAL (turnkey + land + working	g capital)

\$503,140

Table A6-21. CAPITAL COSTS

Boiler type: Watertube (44 MW_t) Fuel: Residual Oil Control technique: Parallel Flow SCR Control level: Moderate

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Equipment cost

Basic equipment (includes freight Required auxiliaries) 54,980 68,100
Total equipment cost	123,080
Installation costs, direct	
Foundations and supports Piping Insulation Painting Electrical Instruments Installation labor	$ \begin{array}{r} 7,900 \\ 25,320 \\ 3,310 \\ 600 \\ 2,360 \\ 4,310 \\ 35,890 \\ 70,600 \end{array} $
Total installation cost	<u>79,690</u>
Total Direct Costs (equipment + insta	11ation) <u>202,770</u>
Installation costs, indirect	
Engineering Construction and field expense Construction fees Start-up Performance tests	<u>43,892</u> <u>20,277</u> <u>20,277</u> <u>4,055</u> <u>2,000</u>
Total Indirect Costs	90,501
Contingencies	43,991
Total T rnkey Costs (direct+indirect+	contingencies) <u>337,262</u>
Land	843
Working capital	23,456
GRAND TOTAL (turnkey + land + working	capital)

\$378,190

Table A6-22. CAPITAL COSTS

Boiler type: Watertube (44 MW_t) Fuel: Residual Oil Control technique: Moving Bed SCR Control level: Stringent

Equipment cost

Equipment cost
Basic equipment (includes freight) 34,650 Required auxiliaries 89,550
Total equipment cost <u>124,200</u>
Installation costs, direct
Foundations and supports6,620Piping20,870Insulation2,650Painting510Electrical2,050Instruments3,620Installation labor26,110
Total installation cost <u>62,430</u>
Total Direct Costs (equipment + installation) <u>186,63</u> 0
Installation costs, indirect
Engineering43,892Construction and field expense18,663Construction fees18,663Start-up3,733Performance tests2,000
Total Indirect Costs86,951
Contingencies41.037
Total Turnkey Costs (direct+indirect+contingencies) <u>314,61</u> 7
Land787
Working capital31,621
GRAND TOTAL (turnkey + land + working capital)

Table A6-23. CAPITAL COSTS

Boiler type: Watertube (44 MW_t) Fuel: Residual Oil Control technique: Moving Bed SCR Control level: Moderate

Equipment cost

Basic equipment (includes freight Required auxiliaries) 26,170 57,550	
Total equipment cost	83,720	
Installation costs, direct		
Foundations and supports Piping Insulation Painting Electrical Instruments Installation labor	4,740 14,930 1,880 370 1,510 2,650 18,570	
Total installation cost	44,650	
Total Direct Costs (equipment + insta	11ation) <u>128,370</u>	
Installation costs, indirect		
Engineering Construction and field expense Construction fees Start-up Performance tests	43,892 12,837 12,837 2,567 2,000	
Total Indirect Costs	74,133	
Contingencies	30,376	
Total Turnkey Costs (direct+indirect+contingencies) <u>232,883</u>		
Land	582	
Working capital	26,223	
GRAND TOTAL (turnkey + land + working	; capital)	

<u>\$259,690</u>

Table A6-24. CAPITAL COSTS

Boiler type: Watertube (44 MW_t) Fuel: Residual Oil Control technique: Parallel Flow SCR (NO_×/SO_×) Control level: Intermediate

Total Direct Costs (equipment + installation)	2,323,000
Installation costs, indirect	
Engineering373,400Construction and field expense232,300Construction fees232,300Start-up46,500Performance tests4,000	
Total Indirect Costs	889,000
Contingencies	482,000
Total Turnkey Costs (direct+indirect+contingencie	s) <u>3,693,000</u>
Land	9,000
Working capital	99,000
GRAND TOTAL (turnkey + land + working capital)	\$ <u>3,801,00</u> 0

Table A6-25. CAPITAL COSTS

Boiler type: Firetube (4.4 MW_t) Fuel: Natural Gas Control technique: Fixed Packed Bed SCR Control level: Stringent

.

Equipment cost

Basic equipment (includes freight) Required auxiliaries	8,900
Total equipment cost	17,430
Installation costs, direct	
Foundations and supports Piping Insulation Painting Electrical Instruments Installation labor	$ \begin{array}{r} $
Total installation cost	9.390
Total Direct Costs (equipment + instal	lation) <u>26,820</u>
Installation costs, indirect	
Engineering Construction and field expense Construction fees Start-up Performance tests	43,892 2,682 2,682 536 2,000
Total Indirect Costs	51,792
Contingencies	11,792
Total 'urnkey Costs (direct+indirect+c	contingencies) 90,404
Land	226
Working capital	9,893
GRAND TOTAL (turnkey + land + working	capital)

\$100,520

Table A6-26. CAPITAL COSTS

Boiler type: Firetube (4.4 MW_t) Fuel: Natural Gas Control technique: Fixed Packed Bed SCR

Control level: Moderate Equipment cost Basic equipment (includes freight) 6,580 Required auxiliaries 5.320 Total equipment cost 11,900 Installation costs, direct Foundations and supports 650 Piping 2,330 Insulation 280 Painting 40 Electrical 510 Instruments 360 Installation labor 5,590 9,760 Total installation cost Total Direct Costs (equipment + installation) 21,660 Installation costs, indirect Engineering 43,892 Construction and field expense 2,166 Construction fees 2,166 Start-up 433 Performance tests 2,000 Total Indirect Costs 50,657 10,848 Contingencies Total Turnkey Costs (direct+indirect+contingencies) 83,165 Land

9,391 Working capital

GRAND TOTAL (turnkey + land + working capital)

\$92,760

208

Table A6-27. CAPITAL COSTS

Boiler type: Watertube (44 MW_t) Fuel: Natural Gas Control technique: Fixed Packed Bed SCR Control level: Stringent

Equipment cost

Basic equipment (includes freight Required auxiliaries) 33,410 87,700
Total equipment cost	121,110
Installation costs, direct	
Foundations and supports Piping Insulation Painting Electrical Instruments Installation labor	$ \begin{array}{r} 3,310\\ 10.480\\ -1,160\\ 220\\ 1,670\\ 1,710\\ 21,380 \end{array} $
Total installation cost	39,930
Total Direct Costs (equipment + instal	llation) <u>161,040</u>
Installation costs, indirect	
Engineering Construction and field expense Construction fees Start-up Performance tests	<u>43,890</u> <u>16,100</u> <u>16,100</u> <u>3,220</u> <u>2,000</u>
Total Indirect Costs	81,310
Contingencies	36,350
Total Turnkey Costs (direct+indirect+	contingencies) <u>278,700</u>
Land	700
Working capital	27,440
GRAND TOTAL (turnkey + land + working	capital)

306,800

Table A6-28. CAPITAL COSTS

Boiler type: Watertube (44 MW_t) Fuel: Natural Gas Control technique: Fixed Packed Bed SCR Control level: Moderate

Control level: Moderate Equipment cost Basic equipment (includes freight) 25,760 Required auxiliaries 52,580 Total equipment cost 78,340 Installation costs, direct 2,520 Foundations and supports Piping 8,050 Insulation 900 Painting 180 Electrical 1.280 Instruments 1.340 Installation labor 15,000 Total installation cost 29,270 Total Direct Costs (equipment + installation) 107,610 Installation costs, indirect 43,890 Engineering Construction and field expense 10,760 Construction fees 10,760 2,150 Start-up Performance tests 2,000 Total Indirect Costs 69.560 Contingencies 26,580 Total Turnkey Costs (direct+indirect+contingencies)203,750 510 Land Working capital 19,470

223,700

GRAND TOTAL (turnkey + land + working capital)

APPENDIX 7

ANNUAL COST BREAKDOWNS

Table A7-1. ANNUAL COSTS

Boiler type: Underfeed Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR Control level: Intermediate

Direct costs

Direct labor	\$15,780
Maintenance labor	28,830
Materials	3,527
Catalyst	14,501
Electricity	
Steam	776
Ammonia	2,431

Total direct cost

66,699

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Overhead

Payroll	4,734
Plant	12,516

17,250

Capital Charges

G&A, taxes & ins.	7,837
Capital recovery	25,750

Total capital charges 33,595

TOTAL ANNUALIZED COSTS

\$117,540

Table A7-2. ANNUAL COSTS

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Boiler type: Chaingrate Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR Control level: Stringent

Direct costs

Direct labor	\$15,780
Maintenance labor	28,830
Materials	6,916
Catalyst	48,590
Electricity	3.101
Steam	2.272
Ammonia	7,030

Total direct cost

112,519

18,130

Overhead

Payroll	4,734
Plant	13,396

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Capital Charges

G&A, ta:	xes & ins.	15,368
Capital	recovery	50,512

Total capital charges 65,880

TOTAL ANNUALIZED COSTS

\$196,530

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Table A7-3. ANNUAL COSTS

Boiler type: Chaingrate Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR Control level: Intermediate

Direct costs

Direct labor	<u>\$15,780</u>
Maintenance labor	28,830
Materials	5.840
Catalyst	36,329
Electricity	2,345
Steam	
Ammonia	6,110

Total direct cost

Overhead

Payroll	4,734
Plant	13,117

Total overhead cost

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97,177

Capital Charges

G&A,	taxes & ins.	12,977
Capit	tal recovery	42,653

Total capital charges 55,630

TOTAL ANNUALIZED COSTS

\$170,660

Table A7-4. ANNUAL COSTS

Boiler type: Chaingrate Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR Control level: Moderate

Direct costs

Direct labor	\$15,780
Maintenance labor	28,830
Materials	4,973
Catalyst	29,129
Electricity	1,892
Steam	1.713
Ammonia	5,256

Total direct cost

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87,573
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17,626

Overhead

Payroll	4,734
Plant	12,892

Total overhead co	st
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Capital Charges

G&A, tax	xes & ins.	11,051
Capital	recovery	36,323

Total capital charges 47,374

TOTAL ANNUALIZED COSTS

\$152,570

Table A7-5. ANNUAL COSTS

Boiler type: Spreader Stoker Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR Control level: Intermediate

Direct costs

Direct labor	\$15,780
Maintenance labor	28,830
Materials	8,936
Catalyst	72,600
Electricity	5,847
Steam	3,814
Ammonia	12,023

Total direct cost

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147,830
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Overhead

Payroll	4,734
Plant	13,922

Total overhead	cost	_18,656
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Capital Charges

G&A, taxes & ins.	19,857
Capital recovery	65,265

Total capital charges 85,122

TOTAL ANNUALIZED COSTS

\$251,610

Table A7-6. ANNUAL COSTS

Boiler type: Pulverized Coal Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR Control level: Stringent

Direct costs

\$ 15,780
28,830
12,558
112,800
12,417
7,133

Total direct cost

<u>212,053</u>

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Overhead

Payroll	4,734
Plant	14,864

Total overhead	cost	19,598

Capital Charges

G&A, taxes & ins.	27,906
Capital recovery	91,720

Total capital charges 119,626

TOTAL ANNUALIZED COSTS

\$351,280

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Table A7-7. ANNUAL COSTS

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Boiler type: Pulverized Coal Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR Control level: Moderate

Direct costs

Direct labor	\$15,780
Maintenance labor	28,830
Materials	8,872
Catalyst	67,800
Electricity	7,490
Steam	5,371
Ammonia	16,885

Total direct cost

151,028

Overhead

Payroll	4,734
Plant	13,905

Total overhead	cost	18,639

Capital Charges

G&A, taxes & ins.	19,716
Capital recovery	64,803

Total capital charges 84,519

TOTAL ANNUALIZED COSTS

\$254,190

Boiler type: Underfeed Stoker Fuel: High Sulfur Eastern Coal Control technique: Parallel Flow SCR $(NO_{\times}/SO_{\times})$ Control level: Intermediate

Direct costs

Direct labor	31,590
Maintenance labor	76,900
Materials	70,000
Catalyst	12,450
Electricity	19,730
Steam	26,100
Fuel	33,790
Boiler feed water	39,520
Ammonia	3,030
Heat credit	(-32, 270)
By-product credit	(-26,050)

Total direct cost

254,800

Overhead

Payroll	9,480	
Plant	46,410	
Total o	warhead cost	55

Capital Charges

G&A, taxes & ins.	93,320	
Capital recovery	306,700	
Total capital cha	arges	400,000

TOTAL ANNUALIZED COSTS

\$710,700

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Boiler type: Underfeed Stoker Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR (NO_X/SO_X) Control level: Intermediate

Direct costs

Direct labor	31,590
Maintenance labor	76,900
Materials	42,090
Catalyst	2,620
Electricity	6,990
Steam	6,400
Fuel	7,140
Boiler feed water	8,350
Ammonia	3,710
Heat credit	(-7,780)
By-product credit	(-5,490)

Total direct cost

172,500

Overhead

Payroll	9,480	
Plant	39,150	

Total overhead cost _____48,600

Capital Charges

G&A, taxes & ins.	_56,120_
Capital recovery	184,450

Total capi	tal charges.	240,600
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TOTAL ANNUALIZED COSTS

\$462,000

Table A7-10. ANNUAL COSTS

Boiler type: Pulverized Coal Fuel: High Sulfur Eastern Coal Control technique: Parallel Flow SCR (NO_X/SO_X) Control level: Intermediate

Direct costs

Direct labor	31,590
Maintenance labor	76,900
Materials	106,600
Catalyst	83,100
Electricity	128,100
Steam	175,200
Fuel	226,500
Boiler feed water	269,500
Ammonia	24,100
Heat credit	(-223,000)
By-product credit	(-173,900)

Total direct cost

724,700

Overhead

Payroll	9,480
Plant	55,920

Total overhead	cost	<u>_65,400</u>
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Capital Charges

G&A, taxes & ins. Capital recovery	236,800 778,300	
Total capital cha	arges	1,010,000

TOTAL ANNUALIZED COSTS

\$1,805,000

Boiler type: Pulverized Coal Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR (NO_X/SO_X) Control level: Intermediate

Direct costs

Direct labor	31,590
Maintenance labor	76,900
Materials	55,400
Catalyst	17,520
Electricity	43,560
Steam	44,140
Fuel	47,620
Boiler feed water	55,820
Ammonia	29,780
Heat credit	(-51,930)
By-product credit	(-36, 680)

Total direct cost

313,700

.

Overhead

Payroll	9,480	
Plant	42,610	

Total overhead cost 52,100

Capital Charges

23,100
04,700
(

Total	capital	charges	527,800
	-	0	

TOTAL ANNUALIZED COSTS

<u>\$893,600</u>

Boiler type: Firetube (4.4 MW_t) Fuel: Distillate Oil Control technique: Fixed Packed Bed SCR Control level: Stringent

Direct costs

Direct labor	11,835
Maintenance labor	21,623
Materials	1,205
Catalyst	3,861
Electricity	547
Steam	
Ammonia	197

Total direct cost

39,569

Overhead

Payroll	3,551
Plant	9,012

Total overhead	l cost	12,563
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Capital Charges

G&A, taxes & ins.	3,571
Capital recovery	11,740

Total capital charges <u>15,311</u>

TOTAL ANNUALIZED COSTS

\$67,440

Table A7-13. ANNUAL COSTS

Boiler type: Firetube (4.4 MW_t) Fuel: Distillate Oil Control technique: Fixed Packed Bed SCR Control level: Moderate

Direct costs

Direct labor	\$11,835
Maintenance labor	21,623
Materials	1.061
Catalyst	2,304
Electricity	345
Steam	291
Ammonia	148

Total d	irect	cost
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37,607
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Overhead

Payroll	3,551
Plant	8,975

Total overhead cost	12,526
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Capital Charges

G&A,	taxes	&	ins.	3,143
Capit	tal rec	201	very	10,331

Total capital charges 13,474

TOTAL ANNUALIZED COSTS

\$63,610

Boiler type: Watertube (44 MW_t) Fuel: Distillate Oil Control technique: Fixed Packed Bed SCR Control level: Stringent

Direct costs

Direct labor	14,480
Maintainence labor	26,440
Materials	8,310
Catalyst	47,380
Electricity	8,500
Steam	3.740
Ammonia	2,470

Total direct cost

111,270

Overhead

Payroll	4,340	
Plant	12,800	

Total overhead	lcost	17,140
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Capital Charges

G&A, taxes & ins.	<u>11,080</u>
Capital recovery	<u>36,430</u>

Total capital charges 47,510

TOTAL ANNUALIZED COSTS

175,900

.

Boiler type: Watertube (44 MW_t) Fuel: Distillate Oil Control technique: Fixed Packed Bed SCR Control level: Moderate

Direct costs

Direct labor Maintainence labor Materials Catalyst	$\frac{14,480}{26,440}$ 6,050 28,420
Electricity	<u>5,140</u>
Steam	<u>3,570</u>
Ammonia	<u>1,920</u>

Total direct cost

86,020

16,550

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Overhead

Payroll	4,340
Plant	12,210

|--|

Capital Charges

G&A, taxes & ins.	8,070
Capital recovery	26,530

Total capital charges <u>34,600</u>

TOTAL ANNUALIZED COSTS

137,200

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Boiler type: Watertube (8.8 MW_t) Fuel: Residual Oil Control technique: Parallel Flow SCR Control level: Stringent

60,260

15,780

Direct costs

Direct labor	14,480
Maintenance labor	26,430
Materials	3,090
Catalyst	13,090
Electricity	740
Steam	590
Ammonia	1,840

Total direct cost

Overhead

Payroll	4,340
Plant	11,440

Total overhead	cost
----------------	------

Capital Charges

G&A, taxes & ins.	$\frac{7,490}{24,620}$
Capital recovery	24,620

Total capital charges ______32,110

TOTAL ANNUALIZED COSTS

\$108,200

Boiler type: Watertube (8.8 MW_t) Fuel: Residual Oil Control technique: Parallel Flow SCR Control level: Moderate

55,370

16,100

Direct costs

Direct labor	14,480
Maintenance labor	26,430
Materials	4,320
Catalyst	7,870
Electricity	460
Steam	440
Ammonia	1,370

Total	direct	cost

Overhead

Payroll	4,340
Plant	11,760

Total	overhead	cost	

Capital Charges

G&A,	taxes & ins.	5,750
Capit	tal recovery	18,910

Total capital charges 24,660

TOTAL ANNUALIZED COSTS

\$96,100

Boiler type: Watertube (8.8 MW_t) Fuel: Residual Oil Control technique: Moving Bed SCR Control level: Stringent

Direct costs

Direct labor	14,465
Maintenance labor	52,855
Materials	2,410
Catalyst	8,660
Electricity	570
Steam	590
Ammonia	1,840

Total direct cost

```
<u>81,390</u>
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22,470

0verhead

Payroll	4,340
Plant	18,130

Capital Charges

G&A, ta:	xes & ins.	5,840
Capital	recovery	19,180

Total capital charges _____25,020

TOTAL ANNUALIZED COSTS

\$129,900

.

76,770

Boiler type: Watertube (8.8 MW_t) Fuel: Residual Oil Control technique: Moving Bed SCR Control level: Moderate

Direct costs

Direct labor	14,465
Maintenance labor	52,855
Materials	2,050
Catalyst	5,190
Electricity	400
Steam	440
Ammonia	1,370

Total	direct	cost
-------	--------	------

Overhead

Payroll	4,340
Plant	18,040

Total overh	nead cost	22,380
10102 01011		

Capital Charges

G&A, taxes & ins.	4,960
Capital recovery	16,320

Total capital charges 21,280

TOTAL ANNUALIZED COSTS

\$120,400

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Boiler type: Watertube (44 MW_t) Fuel: Residual Oil Control technique: Parallel Flow SCR Control level: Stringent

Direct costs

Direct labor	\$14,465
Maintenance labor	26,428
Materials	7,766
Catalyst	62,150
Electricity	5,697
Steam	2,089
Ammonia	6,565

Total direct cost

125,160

Overhead

Payroll	4.340
Plant	12,651

Total overhead cost

16,991

Capital Charges

G&A,	taxes & ins	•	18,827
Capit	al recovery	_	61,879

Total capital charges <u>80,706</u>

TOTAL ANNUALIZED COSTS

\$222,860

Table A7-21. ANNUAL COSTS

Boiler type: Watertube (44 MW_t) Fuel: Residual Oil Control technique: Parallel Flow SCR Control level: Moderate

Direct costs

Direct labor Maintenance labor Materials Catalyst	\$14,465 26,428 5,565 37,455
Electricity	3,457
Steam	1,516
Ammonia	4,938

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Total direct cost

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93,824
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Overhead

Payroll	4,340
Plant	12,079

Total	overhead	cost	16,419
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Capital Charges

G&A, ta:	xes & ins.	13,490
Capital	recovery	44,340

•

Total capital charges 57,830

TOTAL ANNUALIZED COSTS

\$181,180

Boiler type: Watertube (44 MW_t) Fuel: Residual Oil Control technique: Moving Bed SCR Control level: Stringent

Direct costs

,465
,855
,191
.510
820
2.080
560

Total direct cost

<u>126,481</u>

Overhead

Payroll	4,340
Plant	18,853

Total	overhead	cost	23,193
Total	overhead	cost	

Capital Charges

G&A, taxes & ins.	12,585
Capital recovery	41,363

Total capital charges _____53,948

TOTAL ANNUALIZED COSTS

\$203,620

Boiler type: Watertube (44 MW_t) Fuel: Residual Oil Control technique: Moving Bed SCR Control level: Moderate

Direct costs

<u>\$14,465</u>
52,855
3,842
24,900
2,350
1.540
4,940

Total direct cost

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104,892
```

Overhead

Payroll	4,340
Plant	18,502

Total overhead	cost	22,842
lotal overnead	COST	22,042

Capital Charges

G&A, taxes & ins.	9,315
Capital recovery	30,617

Total capital charges 39,932

TOTAL ANNUALIZED COSTS

\$167,670

Table A7-24. ANNUAL COSTS

Boiler type: Watertube (44 MW_t) Fuel: Residual Oil Control technique: Parallel Flow SCR (NO_×/SO_×) Control level: Intermediate

Direct costs

Direct labor	28,960
Maintenance labor	70,490
Materials	110,790
Catalyst	31,900
Electricity	53,330
Steam	66,780
Fuel	86,490
Boiler feed water	102,980
Ammonia	8,710
Heat credit	(-86, 530)
By-product credit	(-78,290)

Total direct cost

395,600

Overhead

Payroll		8,690	
Plant		_54,660_	
Total	overhead	cost	63,350

Capital Charges

G&A, taxes & ins. Capital recovery	147,720 485,520	
Total capital cha	arges	633,240

TOTAL ANNUALIZED COSTS

\$1,092,000

Table A7-25. ANNUAL COSTS

Boiler type: Firetube (4.4 MW_t) Fuel: Natural Gas Control technique: Fixed Packed Bed SCR Control level: Stringent

39,573

15,501

Direct costs

Direct labor	\$11,835
Maintenance labor	21,623
Materials	1,220
Catalyst	4,005
Electricity	600
Steam	70_
Ammonia	220

Total direct cost

Overhead

Payroll	3,551
Plant	9,016

Total overhe	ead cost	12,567
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Capital Charges

G&A, taxes & ins.	3,616
Capital recovery	11,885

Total capital charges

TOTAL ANNUALIZED COSTS

\$67**,**640

Boiler type: Firetube (4.4 MW_t) Fuel: Natural Gas Control technique: Fixed Packed Bed SCR Control level: Moderate

37,565

Direct costs

Direct labor	\$11,835
Maintenance labor	21,623
Materials	1,123
Catalyst	2,394
Electricity	370
Steam	50
Ammonia	170

Total direct cost	Total	direct	cost
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Overhead

Payroll	3,551
Plant	8,991

Total overhead	cost	12,542
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Capital Charges

G&A, taxes & ins.	3,327
Capital recovery	19,934

Total capital charges 14,261

TOTAL ANNUALIZED COSTS

\$64,370

Boiler type: Watertube (44 MW_t) Fuel: Natural Gas Control technique: Fixed Packed Bed SCR Control level: Stringent

Direct costs

Direct labor Maintainence labor	14,480
Materials	<u>26,440</u> 8,360
Catalyst	48,240
Electricity	8,630
Steam	880
Ammonia	2,730

Total direct cost

109,760

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Overhead

Payroll	4,340
Plant	12,810

Total overhead cost	17,510
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Capital Charges

G&A, taxes & ins.	11,150
Capital recovery	36,640

Total capital charges 47,790

TOTAL ANNUALIZED COSTS

8

174,700

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Boiler type: Watertube (44 MW_t) Fuel: Natural Gas Control technique: Fixed Packed Bed SCR Control level: Moderate

Direct costs

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Direct labor	14,480
Maintainence labor	r 26,440
Materials	6,110
Catalyst	28,920
Electricity	5,230
Steam	680
Ammonia	2,120

Total direct cost

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77,870

Overhead

Payroll	4,340
Plant	12,230

Total overhead cost

1	6.	5	70	

Capital Charges

G&A, tax	kes & ins.	8,150
Capital	recovery	26,790

Total capital charges <u>34.940</u>

TOTAL ANNUALIZED COSTS

129,400

APPENDIX 8

SAMPLE CALCULATIONS

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An example calculation is shown below to illustrate how the energy vaues were arrived at. The example illustrates the case of a Pulverized Coal standard boiler with a Parallel Flow reactor and stringent control. Calculations for the other standard boilers were performed in a similar manner.

Sample Calculation--

First, it is necessary to perform a combustion calculation to characterize the flue gas.

> 130% excess air Basis: Coal analysis: 1b/1b fuel fired C = 0.5760 H₂ = 0.320 O₂ = 0.1120 N₂ = 0.0120 S = 0.0060 H₂O = 0.2080 Ash = 0.05401.0000

The calculation is based on a method presented in <u>Steam</u>¹ and the values shown here are documented in the reference.

02	and	Air	required	for	combustion
				101	compuscion

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			0_2 , $\frac{1b}{1b \text{ fuel fired}}$		Air, $\frac{1b}{1b \text{ fuel fired}}$
C H ₂	0.5760 0.0320		x 2.66 = 1.532	,	x 11.53 = 6.641
0_2	0.1120		x 7.94 = 0.254	,	x 34.34 = 1.099
N ₂ S H ₂ O	0.120 0.0060 0.2080		x 1.00 = 0.006	,	x 4.29 = 0.026
Ash	0.0540				<u> </u>
Total	1.00		1.792		7.766
Less 0_2	in the fuel	-	0.112	-	0.482
Require	ment		1.680		7.284
Excess	(30%)		0.504		2.185
Total			2.184		9.469

Products of combustion

		lb/lb fuel
CO ₂ 0.5760 x 3.66	=	2.108
H_2O (0.032 x 8.94) + (0.2080) + (0.013 x 9.469)	=	0.617
0 ₂ excess	=	0.504
N ₂ 9.469 x (0.7685 + 0.0120)	=	7.391
NO_{\times} (specified by Acurex)	=	0.0090
SO_{x} (specified by Acurex)	=	0.0114
CO (specified by Acurex)	=	0.0005
HC as CH ₄ (specified by Acurex)	=	0.0001
Fly ash (specified by Acurex)	=	0.0432
		10.68
	c 1	

Fuel feed rate = $10.42 \frac{\text{tons}}{\text{hr}} \times \frac{2000 \text{ lb}}{\text{ton}} = 20,840 \frac{\text{lb fuel}}{\text{hr}}$

Flue Gas Composition:

	1b/hr	moles/hr	mole %
N ₂	154,000	5,496	72.8
CO2	43,900	998	13.2
H ₂ O	12,900	716	9.5
02	10,500	328	4.3
S0 ₂	238	4	0.1
NO_{\times}	188	4	0.1
со	10	-	-
HC (as CH4)	3	-	-
Fly ash	900	-	
	222,639	7,546	100.0

Average molecular weight =
$$\frac{222,639}{7,546}$$
 = 29.5 $\frac{1b}{1b}$ mole
= 29.5 $\frac{g}{g}$ mole

Flue gas flow rate

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$$G_{v} = \left(73,200 \ \frac{\text{ft}^{3}}{\text{min}}\right) \left(\frac{492^{\circ}\text{R}}{810^{\circ}\text{R}}\right) \left(\frac{1.698 \ \text{Nm}^{3}/\text{hr}}{\text{scfm}}\right) = 75,500 \ \frac{\text{Nm}^{3}}{\text{hr}}$$
Reactor Sizing

Next, it is necessary to size the reactor so the pressure drop across the reactor may be calculated. For the stringent level of control, a large reactor size and bed depth are used to ensure 90% NO_{\times} reduction.

Basis: ² Space velocity =
$$3000 \text{ hr}^{-1}$$
 (based on catalyst volume: $3000 \text{ catalyst volumes of flue gas per hour}$)
Bed depth = 4.5 m
Catalyst volume= $\frac{75,500 \text{ Nm}^3/\text{hr}}{3,000 \text{ hr}^{-1}}$ = 25.2 m³

To calculate the reactor volume, the specific surface areas of the pure catalyst and of the catalyst packed in a reactor are needed.

601 m²/m³ catalyst (20mm parallel plate)
194 m²/m³ packed reactor volume
Reactor volume =
$$(25.2 \text{ m}^3 \text{ catalyst}) \left(\frac{601 \text{ m}^2/\text{m}^3 \text{ catalyst volume}}{194 \text{ m}^2/\text{m}^3 \text{ packed reactor volume}} \right)$$

= 78 m³
= width² x depth (square reactor)
Therefore, width = 4.16 m

Pressure Drop

Now that the reactor geometry has been defined, the pressure drop across the reactor can be determined. For this calculation the following equation is used.

$$\Delta P = \frac{2 f_m G^2 L (1-\varepsilon)^{3-n}}{D_p g_c \rho \phi_s^{3-n} \varepsilon^3}$$
(A8-1)

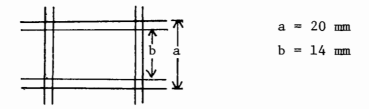
where ΔP = pressure drop across bed of granular solids, lb_f/ft^2

- f_{m} = friction factor: a function of modified Reynold's number (N_{Re}), dimensionless
- G = gas superficial mass velocity, lb_m/ft^2 sec
- L = bed depth, ft
- ε = void fraction, dimensionless
- n = exponent: a function of modified Reynold's number (N^{*}_{Re}), dimensionless
- D = average particle diameter: diameter of a sphere of the p same volume as the non-spherical particle, ft

- $g_c = dimensional constant, 32.2 \frac{lb_m ft}{sec^2 lb_f}$
- ρ = gas density, $1b_m/ft^3$

(The modified Reynolds number, N_{Re} , is defined as D_pG/μ , where μ = gas viscosity, lb_m/ft sec)

Parallel Flow Catalyst (a square passage was assumed for ease of calculation)



Cell length = 1 m (assumed, a common commercial cell length²)

In order to calculate a shape factor it is necessary to calculate the diameter of a sphere that has a volume of catalyst equivalent to a single passage of the square honeycomb catalyst.

Catalyst volume per passage = $[(20 \text{ mm})^2 - (14 \text{ mm})^2] 1000 \text{ mm}$ = 204,000 mm³ or 7.21 x 10⁻³ ft³ V sphere = $\frac{4}{3} \prod r_p^3 = 7.21 \times 10^{-3}$ $r_p = 0.120 \text{ ft}$ $D_p = 0.240 \text{ ft}$

The shape factor can now be calculated

$$\phi_{s} = \text{shape factor} = \frac{\text{surface area of sphere of equivalent volume}}{\text{actual surface area per passage}}$$

$$\phi_{s} = \frac{\Pi(0.240)\text{ft})^{2}}{(4)(14 \text{ mm})(1000 \text{ mm})} \left(\frac{10^{6} \text{ mm}^{2}}{\text{m}^{2}}\right) \left(\frac{.0929 \text{ m}^{2}}{\text{ft}^{2}}\right)$$

$$= 0.301$$

Void fraction =
$$\varepsilon = 1 - \frac{194 \text{ m}^2/\text{m}^3 \text{ packed reactor volume}}{601 \text{ m}^2/\text{m}^3 \text{ catalyst volume}}$$

 $\varepsilon = 1 - 0.323 \frac{\text{catalyst volume}}{\text{reactor volume}}$
 $\varepsilon = 0.677 \frac{\text{void volume}}{\text{reactor volume}}$

Gas Viscosity, μ^{0}

This viscosity of a gas mixture can be calculated from^4

$$\mu_{\text{mixture}}^{O} = \frac{\Sigma_{y_{i}} \mu_{i} (M_{i})^{\frac{1}{2}}}{\Sigma_{y_{i}} (M_{i})^{\frac{1}{2}}}$$
(A8-2)

where $\mu_{\text{mixture}}^{\circ}$ = viscosity of gas mixture, micropoise y_i = mole fraction of component i in mixture, dimensionless μ_i = gas viscosity of pure component i, micropoise M_i = molecular weight of component i, g/g mole

 μ_{i} can be calculated from[§]

$$\mu_{i} = \frac{33.3(M_{i}T_{c})^{\frac{1}{2}}}{V_{c}^{2/3}} \times f (1.33T_{r})$$
(A8-3)

where T_c = critical temperature of component i, °K

$$V_{c}$$
 = critical volume of component i, cm³/g-mole

$$T_r$$
 = reduced temperature = ratio of gas temperature to critical temperature (T/T_c) , dimensionless

 $f(T_r)$ = gas viscosity temperature function, dimensionless

Values for $\mu_{\mbox{i}}$ were calculated using data from Smith & VanNess 6

	T _c ,°k	T _r ,°k	$v_{c}, \frac{cm^{3}}{gmole}$	$f(1.33T_r)$	M	$\mu_{i,ft-sec}$
N ₂	126.2	5.33	89.5	3.07	28.02	2.04×10^{-5}
CO ₂	304.2	2.21	94.0	1.65	44.01	2.07×10^{-5}
H20	647.1	1.04	56.0	0.862	18.02	1.42×10^{-5}
02	154.6	4.35	73.4	2.68	32.00	2.41×10^{-5}

The following data were used with equation (A8-2) to calculate the gas viscosity

	y _i	$\mu_{i, ft-sec}$	M
N ₂	0.728	2.04×10^{-5}	28.02
CO ₂	0.132	2.07×10^{-5}	44.01
H ₂ O	0.095	1.42×10^{-5}	18.02
02	0.043	2.41×10^{-5}	32.00

with the following result

$$\mu_{\text{mixture}}^{\text{o}} = 2.00 \times 10^{-5} \frac{1b}{\text{ft-sec}}$$

Gas Superficial Mass Velocity, G

From the results of the combustion calculation (total mass flow of flue gas) and the reactor sizing calculation (reactor width), the superficial mass velocity can be found.

$$G = \frac{\left(222,600 \ \frac{1b}{hr}\right) \left(\frac{1 \ hr}{3600 \ sec}\right)}{(4.16m)^2 \left(10.76 \ \frac{ft^2}{m^2}\right)} = 0.332 \ \frac{1b}{ft^2 sec}$$

Modified Reynolds Number, N_{Re}^{-}

Using the results of the catalyst characterization, gas mixture viscosity, and mass velocity calculations the modified Reynolds number can be found.

$$N_{Re} = \frac{D_{p}G}{\mu^{o}} = \frac{(0.240 \text{ ft}) (0.332 \frac{1b}{\text{ft}^{2}-\text{sec}})}{2.00 \times 10^{-5} \frac{1b}{\text{ft}-\text{sec}}} = 3980$$

Knowing the modified Reynolds number the friction factor and exponent can then be determined.³

$$f_{m} = 0.7$$

n = 1.97

Flue Gas Density, ρ

Reaction T = $750^{\circ}F = 1210^{\circ}R$ Volumetric flue gas flow = 73,200 acfm @ $350^{\circ}F$ (PedCo) From combustion calculation: mass flow = $222,600 \frac{1b}{hr}$

Density =
$$\frac{\text{mass}}{\text{volume}}$$

 $\rho = \frac{222,600 \frac{1b}{hr}}{(73,200 \frac{ft^3}{min})(\frac{1210^{\circ}R}{350+460^{\circ}R})(\frac{60 \text{ min}}{hr})} = 0.0339 \frac{1b}{ft^3}$

All of these terms are then substituted into equation (A8-1) to determine the pressure drop.

$$\Delta P = \frac{2f_{m}G^{2}L (1-\varepsilon)^{3-n}}{D_{p}g_{c}\rho\phi_{s}^{3-n}\varepsilon^{3}}$$

$$= \frac{2 (0.7) \left(0.332 \frac{1b_{m}}{ft^{2}sec}\right) \left[(4.5m) \left(\frac{ft}{.3048m}\right)\right] (1-0.677)^{3-1.97}}{(0.240 \text{ ft}) \left(32.2 \frac{1b_{ft}}{sec^{2}1b_{f}}\right) \left(0.0339 \frac{1b_{m}}{ft^{3}}\right) (0.301)^{3-1.97} (0.677)^{3}}$$

$$= 30.2 \frac{1b_{f}}{ft^{2}} \left(\frac{ft^{2}}{144 \text{ in}^{2}}\right)$$

 $\Delta P = 0.210 \text{ psi} = 148 \text{ mm } H_2O$

Now the energy consumption of the various process steps can be calculated. The energy consuming items considered in this case are

> flue gas fan, liquid NH_3 pump, NH_3 vaporization, and NH_3 dilution.

Flue Gas Fan

The gas side horsepower (hp) can be calculated from⁷

$$(hp)_{gas} = 1.57 \times 10^{-4} \ Q \ \Delta P$$

$$(A8-4)$$

$$Q = ft^{3}/min$$

$$\Delta P = in \ H_{2}O$$

$$(hp)_{air} = (1.57 \times 10^{-4}) \ \left(73,200 \ \frac{ft^{3}}{min}\right) \ (148mm \ H_{2}O) \ \left(\frac{1 \ in}{25.4mm}\right)$$

$$= 67.0 \ hp$$

Fan efficiencies typically range from 40 to 70%.⁷ If an efficiency of 55% is assumed the shaft hp can be calculated

$$(hp)_{shaft} = \frac{67.0}{0.55} = 122 hp$$

In terms of electrical usage, this is

$$(122 \text{ hp})\left(\frac{0.7457 \text{ kW}}{\text{hp}}\right) = 90.8 \text{ kW}$$

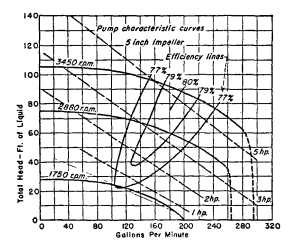
Liquid NH₃ Pump

For 90% removal an NH₃:NO mole ratio of 0.95 is typical²

$$NH_{3} \text{ requirement} = \left(188 \frac{1 \text{b } NO_{\times}}{\text{hr}}\right) \left(\frac{1 \text{ mole } NO_{\times}}{46 \text{ 1b } NO_{\times}}\right) \left(\frac{.95 \text{ mole } NH_{3}}{\text{mole } NO_{\times}}\right) \left(\frac{17 \text{ 1b } NH_{3}}{1 \text{ b mole}}\right)$$
$$= 66 \frac{1 \text{b } NH_{3}}{\text{hr}}$$

or 0.21 gpm

The following pump curve ⁸ indicates that a 0.5 hp centrifugal pump operated at 1750 rpm can supply 28 ft of NH₃ head (7.6 psi). This is adequate to transfer sufficient NH₃ to the vaporizer.



In terms of electrical usage this amounts to $(0.5 \text{ hp})\left(\frac{0.7457 \text{ kW}}{\text{hp}}\right) = 0.373 \text{ kW}$

NH₃ Vaporization

Looking at the worst case for NH_3 vaporization, a cold winter day at a Midwest location, the ambient air temperature might be, say, -10°F.

The pressure in the vaporizer is

Saturated vapor pressure $(-10^{\circ}F) = 23.7$ psia + Pump head $= \frac{7.6}{31.3}$ psia

The normal boiling point of NH_3 is -28.0°F.⁹ In order to determine the actual boiling point, and thus the heat load on the vaporizer, it is necessary to evaluate it at the higher pressure. This can be accomplished by use of the Clausius-Clapeyron equation.¹⁰

$$\Delta H_{vap} = -R \frac{d(\ln P^{sat})}{d(1/T)}$$
(A8-5)

At -28.0°F and 1 atm, $\Delta H_{vap} = 589.3 \frac{Btu}{1b}$

The Clausius-Clapeyron equation is used to determine the boiling temperature at the higher pressure of 31.3 psia. Use of the equation assumes a constant ΔH_{vap} , however, this is not strictly true. For this reason, two iterations are calculated. The first uses ΔH_{vap} at 14.7 psia and calculates an elevated boiling temperature at P = 31.3 psia. The ΔH_{vap} at this temperature is found from thermodynamic tables. Then the two ΔH_{vap} values are averaged to determine a pseudo constant ΔH_{vap} . This value is then used in the second iteration to determine a new and more accurate elevated boiling temperature.

Solving equation (A8-5) gives

$$\ln \frac{P_2}{P_1} = -\frac{\Delta H_{vap}}{R} \left(\frac{1}{T_2} - \frac{1}{T_1} \right)$$
(A8-6)

The data used to calculate T_2 are

$$P_{1} = 14.7 \text{ psia}$$

$$P_{2} = 31.3 \text{ psia}$$

$$T_{1} = -28.0^{\circ}\text{F} + 460 = 432^{\circ}\text{R}$$

$$\Delta H_{\text{vap}} = 589.3 \frac{Btu}{1b}$$

$$R = 1.986 \frac{Btu}{1b \text{ mole } ^{\circ}\text{R}}$$

Solution of equation (A8-6) for T_2 gives

$$T_2 = 462^{\circ}R = 2^{\circ}F$$

$$\Delta H_{vap} \text{ at } 2^{\circ}F \text{ is } 567.3 \frac{Btu}{1b}.^9$$

The two ΔH values are averaged to obtain a pseudo constant ΔH vap

$$\overline{\Delta H}_{vap} = \frac{589.3 + 567.3}{2} = 578.3 \frac{Btu}{1b}$$

Solving equation (A8-6) a second time using this new ΔH indicates a boiling temperature of 3°F. At this temperature, ΔH_{vap} is 566.5 $\frac{Btu}{1b}$.

Now the energy requirement for NH_3 vaporization can be calculated. The heat capacity of NH_3 at this temperature is 11

$$C_{p} [NH_{3}(l)] = 1.10 \frac{Btu}{lb^{\circ}F}$$

$$Q_{vaporization} = \left(66 \frac{1b NH_{3}}{hr}\right) \left[\left(1.10 \frac{Btu}{lb^{\circ}F}\right) [3^{\circ}F - (-10^{\circ}F)] + 566.5 \frac{Btu}{lb} \right]$$

$$= 38,300 \frac{Btu}{hr}$$

NH₃ Dilution

The NH₃ is diluted with 30 psig steam prior to injection. A 5:1 mole ratio is used.¹² The heat of vaporization of 30 psig steam is 929.0 $\frac{Btu_{13}}{1b}$.

Mass of steam injected =
$$\left(66 \frac{1b \text{ NH}_3}{\text{hr}}\right) \left(\frac{\text{mole NH}_3}{17.0 \text{ lb NH}_3}\right) \left(\frac{5 \text{ moles steam}}{\text{mole NH}_3}\right)$$

 $\left(\frac{18 \text{ lb steam}}{\text{mole steam}}\right) = 350 \frac{1b \text{ steam}}{\text{hr}}$
Q = (350 lb) $\left(\frac{929.0 \text{ Btu}}{1b}\right) = 325,000 \frac{\text{Btu}}{\text{hr}}$

Summarizing the energy consuming steps,

Item	Energy usage	Btu/hr	MW thermal
Flue gas fan	90.8 kW elec	908,000	0.266
Liquid NH ₃ pump	0.373 kW elec	3,730	0.00109
NH_3 vaporization	38,300 Btu/hr	38,300	0.0112
$\rm NH_3$ dilution	325,000 Btu/hr	325,000	0.0953
Total		1,275,030	0.374

The example calculation is continued below to show the methods used to arrive at annual cost figures for NO_{χ} control systems. The case is the Parallel Flow SCR system applied to the Pulverized Coal standard boiler and operated at the stringent level of control. The material balance and process flow diagram, as it appears in the Appendix, are presented in Figure A8-1 on the following page. First, each of the pieces of equipment, in succession, will be sized (including any necessary design calculations) and the F.O.B. costs determined. These results will then be utilized to determine the installed costs. Then, the direct operating costs are calculated. These costs are combined via the recommended format using the appropriate load factor to arrive at the annual costs.

From Figure A8-1, the NH₃ flow is shown to be 1.76 $\frac{\text{kg-mole}}{\text{hr}}$. Assuming the plant maintains a 15-day storage supply (large enough to survive delay in deliveries due to bad weather, strikes, etc.), the required NH₃ storage tankage is determined.

Gal Storage =
$$\left(1.76 \frac{\text{kg-mole}}{\text{hr}}\right) \left(\frac{17.0 \text{ kg}}{\text{kg-mole}}\right) \left(\frac{1b}{.454 \text{ kg}}\right) \left(\frac{\text{ft}^3}{39.0 \text{ lb}}\right)^{17} \left(\frac{7.48 \text{ gal}}{\text{ft}^3}\right) \left(\frac{24 \text{ hr}}{\text{day}}\right)$$

x (15 day supply) (A8-7)

= 4600 gal F.O.B. Equipment Cost = Base Cost (Unit Cost)^{Exponential Factor} x Pressure Factor

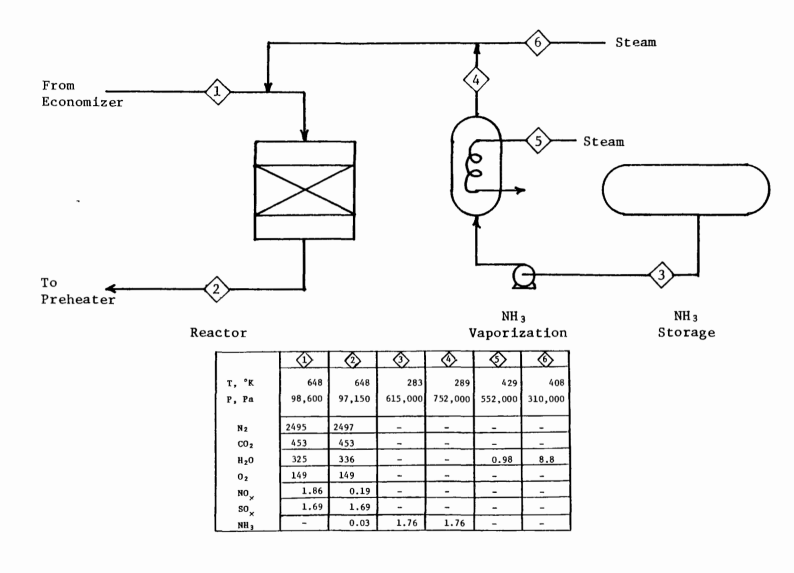


Figure A8-1. Material balance. Pulverized Coal Parallel Flow SCR Stringent Control

F.O.B. NH₃ Tank Cost (mid-1970)¹⁵ = 10,000
$$\left(\frac{\text{gal}}{10,000}\right)^{0.7}$$
 x 1.38 (A8-8)
= 10,000 $\left(\frac{4,600}{10,000}\right)^{0.7}$ x 1.38
= \$8,000

Next, the two liquid NH_3 pumps (one for a spare in case of failure) are examined. The volumetric liquid NH_3 flow rate can be determined from:

Liq. NH₃ Vol. Flow =
$$\left(1.76 \frac{\text{kg-mole}}{\text{hr}}\right) \left(\frac{17.0 \text{ kg}}{\text{kg-mole}}\right) \left(\frac{1b}{.454 \text{ kg}}\right)$$

 $x \left(\frac{\text{ft}^3}{39.0 \text{ lb}}\right) \left(\frac{7.48 \text{ gal}}{\text{ft}^3}\right) \left(\frac{\text{hr}}{60 \text{ min}}\right)$ (A3-9)
= 0.21 GPM

At this flow rate, a 0.5 hp centrifugal pump is adequate.⁸

F.O.B. Equipment Cost = H Pumps x Base Cost : F.O.B. Pump Cost (mid-1970)¹⁶ = 2 pumps x $\frac{$300}{pump}$

= \$600

To size the NH₃ vaporizer the sensible heat and the heat of vaporization for NH₃ is required. For the worst case at -10°F and 31 psia tank pressure, the heat required = $581 \frac{Btu}{1b \text{ NH}_3}$.

The heat load on the vaporizer is shown to be:

$$Q = \left(1.76 \frac{\text{kg-mole}}{\text{hr}}\right) \left(\frac{17.0 \text{ kg}}{\text{kg-mole}}\right) \left(\frac{1b}{.454 \text{ kg}}\right) \left(\frac{581 \text{ BTU required}}{1b \text{ NH}_3 \text{ vaporized}}\right)$$
$$= 38,300 \frac{\text{BTU}}{\text{hr}}$$

The heat transfer area required for the NH_3 vaporizer can be calculated from:

$$A = \frac{qF}{U_{c}(T_{n}-T)}^{17}$$
(A8-10)
where A = heat transfer surface area, ft²
q = heat transferred, BTU/hr
F = safety factor (assume 2.0)
U_{c} = heat transfer coefficient, BTU/hr ft² °F
T_{n} = temperature of heat medium, °F
(800 psia steam = 312°F)
T = fluid temperature, °F
(worst case: NH₃ = -10°F)
A = $\frac{(38,300 \frac{BTU}{hr})(2.0)}{(300 \frac{BTU}{hr-ft^{2}°F})^{18}(312 - (-10)°F)}$:

The smallest commercially available doube pipe heat exchanger is 1 ft^2 and the F.O.B. Vaporizer Cost (mid-1970) = \$300.¹⁹

Next, the reactor size is determined from the volumetric flue gas flow rate and the reactor space velocity.

Volume Flue Gas =
$$\left(73,200 \frac{\text{ft}^2}{\text{min}}\right) \left(\frac{492 \,^{\circ}\text{R}}{810 \,^{\circ}\text{R}}\right) \left(\frac{0.0283 \,\text{Nm}^3}{\text{scf}}\right) \left(\frac{60 \,\text{min}}{\text{hr}}\right)$$

= 75,500 $\frac{\text{Nm}^3}{\text{hr}}$ (A8-11)

Catalyst Volume²⁰ =
$$\frac{75,500 \text{ Nm}^3/\text{hr}}{3000 \text{ hr}^{-1} \text{ space velocity}} = 25.2 \text{ m}^3 = 889 \text{ ft}^3$$

(A8-12)

Reactor Volume²¹ = $(25.2 \text{ m}^3 \text{ catalyst}) \text{ x}$

$$\left(\frac{601 \text{ m}^2 \text{ surface area/m}^3 \text{ catalyst}}{194 \text{ m}^2 \text{ surface area/m}^3 \text{ packed volume}}\right)$$
= 78.0 m³ packed volume (A8-13)

Reactor Length²¹ = 4.5 m $\left(\frac{ft}{.3048m}\right)$ = 14.8 ft

Square Reactor Volume = $78.0 \text{ m}^3 = W^2 \text{L} = 4.5 W^2$

$$W^2 = \frac{78.0 \text{ m}^3}{4.5 \text{ m}^3} = 17.3 \text{ m}^2$$

$$W = 4.16 \text{ m} = 13.6 \text{ ft}$$

F.O.B. Reactor Material Cost (mid-1970) = $$16,000^{22}$

Finally, the draft fan motor drive must be determined.

Motor hp. =
$$\frac{0.000157 \text{ Q} \Delta \text{p}^{-7}}{55\% \text{ efficiency}}$$
 (A8-4)
= $\frac{0.000157 \left(73,200 \frac{\text{ft}}{\text{min}}\right) (148 \text{ mmH}_2\text{O}) \left(\frac{\text{in}}{25.4 \text{ mm}}\right)}{.55}$

= 122 hp

Motor Drive - F.O.B. Motor Cost
$$(\text{mid}-1970)^{23} = 5800 \left(\frac{\text{hp}}{70}\right)^{0.77}$$
 (A8-14)
= $5800 \left(\frac{122}{70}\right)^{0.77}$
= \$8,900

Each piece of equipment is factored by its respective escalation index to give a 1978 F.O.B. cost and an 8 percent freight charge for delivery is added to it.²⁴ The direct installation costs are determined by the appropriate factor multiplied times the 1970 F.O.B. equpment cost and that category's respective escalation index. These analyses for each equipment item are presented on the following pages.

NH₃ Storage Tank

F.O.B. Equipment Cost (mid-1970) = 10,000 $\left(\frac{\text{gal.}}{10,000}\right)^{0.7}$ x 1.38 = \$ 8,000

	June - 1978 Costs		
Equipment Cost (1978)			
Basic equipment =	F.O.B. 1970 x Escalation Index		
=	= \$8000 x 1.91	=	15,300
Freight =	= 0.08 x Basic Equipment	=	1,200
Required auxiliaries		=	N/A
1	Total Equipment Cost	=	16,500
Installation Costs, Direct	<pre>¹⁵ = F.O.B. 1970 x Installation Cost Fraction x Escalation Index</pre>		
Foundation and support	s = F.O.B. 1970 x 0.080 x 2.11	=	1,400
Ductwork		=	N/A
Stack		=	N/A
Piping	= F.O.B. 1970 x 0.153 x 2.02	=	2,500
Insulation		=	N/A
Painting	= F.O.B. 1970 x 0.012 x 2.11	=	200
Electrical	= F.O.B. 1970 x 0.007 x 1.70	=	100
Instruments	= F.O.B. 1970 x 0.118 x 1.63	=	1,500
Installation Labor	= F.O.B. 1970 x 0.352 x 1.37	=	3,900
	Total Installation Cost	=	9,600
	Total Direct Cost	=	\$2 6, 100

Liquid NH 3/Pumps

.

Pump and motor - F.O.B. Equipment Cost (mid-1970) = 300×2 = 600

June - 1978 Costs

Equipment Cost (1978)			
Basic equipment	= F.O.B. 1970 x Escalation Index		
	= \$600 x 2.08	=	1,250
Freight	= 0.08 x Basic equipment	=	100
Required auxiliaries		=	N/A
	Total Equipment Cost	=	<u>1,350</u>
Installation Costs, Direct ¹⁶	= F.O.B. 1970 x Installation Cost Fraction x Escalation Index		
Foundation and supports	= F.O.B. 1970 x 0.039 x 2.11	=	50
Ductwork		=	N/A
Stack		Ξ	N/A
Piping	= F.O.B. 1970 x 0.293 x 2.02	=	360
Insulation	= F.O.B. 1970 x 0.028 x 2.11	=	40
Painting	= F.O.B. 1970 x 0.008 x 2.11	=	10
Electrical	= F.O.B. 1970 x 0.303 x 1.70	=	310
Instruments	= F.O.B. 1970 x 0.029 x 1.63	=	30
Installation labor	= F.O.B. 1970 x 0.679 x 1.37	=	560
	Total Installation Cost		1,360
	Total Dimont Cost		60 710

Total Direct Cost \$2,710

NH₃ Vaporizer

```
1 ft<sup>2</sup> Double-Pipe Heat Exchanger
  (minimum size available)
```

Vaporizer - F.O.B. Equipment Cost (mid-1970) = \$300

June - 1978 Costs

Equipment Costs (1978)			
Equipment Costs (1978)			
Basic equipment	= F.O.B. 1970 x Escalation Index		
	= \$300 x 1.91	= 5	\$ 570
Freight	= 0.08 x Basic equipment	=	50
Required auxiliaries		=	N/A
	Total Equipment Cost	=	620
Installation Costs, Direct ¹⁹	= F.O.B. 1970 x Installation Cost Fraction x Escalation Index		
Foundation and supports	= F.O.B. 1970 x 0.038 x 2.11	=	50
Ductwork		=	N/A
Stack		=	N/A
Piping	= F.O.B. 1970 x 0.213 x 2.02	~	270
Insulation	= F.O.B. 1970 x 0.022 x 2.11	=	30
Painting	= F.O.B. 1970 x 0.002 x 2.11	=	
Electrical	= F.O.B. 1970 x 0.010 x 170	=	10
Instruments	= F.O.B. 1970 x 0.048 x 1.63	=	50
Installation labor	= F.O.B. 1970 x 0.467 x 1.37	=	400
	Total Installation Cost	=	810
	Total Direct Cost	=	\$1,430

Reactor

Reactor - F.O.B. Equipment Cost (mid-1970): Material : Fabricated (2x		= \$ 16,000
•		= 32,000
June - 1978 Co	osts	
Equipment Cost (1978)		
Basic equipment = F.O.B. 19	970 x Escalation Index	
= \$32,000 x	x 1.91	= 61,100
Required auxiliary:Catalyst = \$212 x ft ³ catalyst		= 188,000
Freight = 0.08 x Ba	asic equipment	= 4,890
Total Equip	pment Cost	= 254,000
Installation Costs, Direct ²² = F.O.B. 1970 p Fraction x Es	x Installation Cost scalation Index	
Foundation and supports = F.O.B. 1970	x 0.176 x 2.11	= 11,800
Ductwork		= N/A
Stack		= N/A
Piping = F.O.B. 1970 2	x 0.595 x 2.02	= 38,400
Insulation = F.O.B. 1970 2	x 0.080 x 2.11	= 5,400
Painting = F.O.B. 1970 2	x 0.013 x 2.11	= 880
Electrical = F.O.B. 1970 x	x 0.049 x 1.70 =	= 2,600
Instruments = F.O.B. 1970 x	x 0.114 x 1.63	= 6,000
Installation labor = [F.O.B. 1970	x 0.972 x 1.37	= 61,400
+ (Catalyst >	x 0.10)]	
Total Installat	tion Cost :	= 126,480
Total Direc	ct Cost	\$380,470

Draft Fan Motor Drive

June - 1978 Costs Equipment Cost (1978) Basic equipment = F.O.B. 1970 x Escalation Index = \$8900 x 2.08 = 18,500= 0.08 x Basic equipment 1,500 Freight N/A Required auxiliaries = 20,000 Total Equipment Cost Installation Costs, Direct²⁵ = F.O.B. 1970 x Installation Cost Fraction x Escalation Index 820 Foundation and supports = F.O.B. 1970 x 0.043×2.11 = N/A = Ductwork N/A = Stack = F.O.B. 1970 x 0.141 x 2.02 2,500 = Piping : = F.O.B. 1970 x 0.005 x 2.11 = 90 Insulation = N/A Painting = F.O.B. 1970 x 0.068 x 1.70 1,000 = Electrical = F.O.B. 1970 x 0.013 x 1.63 = 190 Instruments = F.O.B. 1970 x 0.295 x 1.37 = 3,600 Installation labor Total Installation Cost = 8,200

Total Direct Cost = \$28,200

The direct operating costs are shown below.

Ammonia

$$\left(1.76 \frac{\text{kg mole}}{\text{hr}}\right) \left(\frac{\$130}{\text{ton}}\right) \left(\frac{1 \text{ ton}}{2000 \text{ lb}}\right) \left(\frac{17 \text{ kg}}{\text{kg mole}}\right) \left(\frac{1b}{454 \text{ g}}\right) \left(\frac{1000 \text{ g}}{\text{kg}}\right) (8760 \text{ hr}) (0.6)$$
$$= \$22,535$$

Electricity

$$(0.268 \text{ MW}_{t}) \left(\frac{25.8 \text{ mills}}{\text{kwh}}\right) \left(\frac{\$1}{1000 \text{ mills}}\right) \left(\frac{342 \text{ kw}}{\text{MW}_{t}}\right) (8760 \text{ hr}) (0.6)$$

= \\$12,429

Steam

$$\left(\frac{9.78 \text{ kg mole}}{\text{hr}}\right) \left(\frac{18 \text{ kg}}{\text{kg mole}}\right) \left(\frac{1000 \text{ g}}{\text{kg}}\right) \left(\frac{1 \text{ 1b}}{454 \text{ g}}\right) \left(\frac{\$3.50}{1000 \text{ 1b}_{\text{m}}}\right) (8760 \text{ hr}) (0.6)$$
$$= \$7,133$$

:

The individual equipment costs and installation costs are summed and the totals entered in Table A8-1. The direct operating costs are entered in Table A8-2.

TABLE A8-1. CAPITAL COSTS

Boiler type: Pulverized Coal Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR Control level: Stringent

Equipment cost

Basic equipment (includes freight) Required auxiliaries	104,470 188,000		
Total equipment cost	292,470		
Installation costs, direct			
Foundations and supports Piping Insulation Painting Electrical Instruments Installation labor	14,120 44,030 5,560 1,090 4,020 7,770 69,860		
Total installation cost	146,450		
Total Direct Costs (equipment + instal	lation)	438,920	
Installation costs, indirect			
	43,892 43,892 43,892 8,778 2,000		
Total Indirect Costs		142,454	
Contingencies		116,275	
Total Turnkey Costs (direct+indirect+contingencies) <u>697,649</u>			
Land		1,744	
Working capital		54,288	
GRAND TOTAL (turnkey + land + working capital)			

\$753,620

Boiler type: Pulverized Coal Fuel: Low Sulfur Western Coal Control technique: Parallel Flow SCR Control level: Stringent

Direct costs

Direct labor	17,279
Maintainence 1	abor <u>31,569</u>
Materials	12,884
Catalyst	1 <u>13,081</u>
Electricity	12,417
Steam	_7,133
Ammonia	22,535

Total direct cost

216,900

Overhead

Payroll	<u>5,184</u>
Plant	16,050

Total overhe	ead cost	21_234
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Capital Charges

G&A, taxes & ins.	27,906
Capital recovery	91,720

Total capital charges 119,626

TOTAL ANNUALIZED COSTS

\$357,760

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⁷ AUTHOR(S) Gary D. Jones and K	evin L. Johnson	8. PEF	REORMING ORGAN	VIZATION REPORT NO.
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e e	Fossil Fuels	Flue Gas Treat		ÎB 21D
Assessments		Selective Cataly		1B
Industrial Processes		duction (SCR)	1	BH
Boilers			13	BA
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