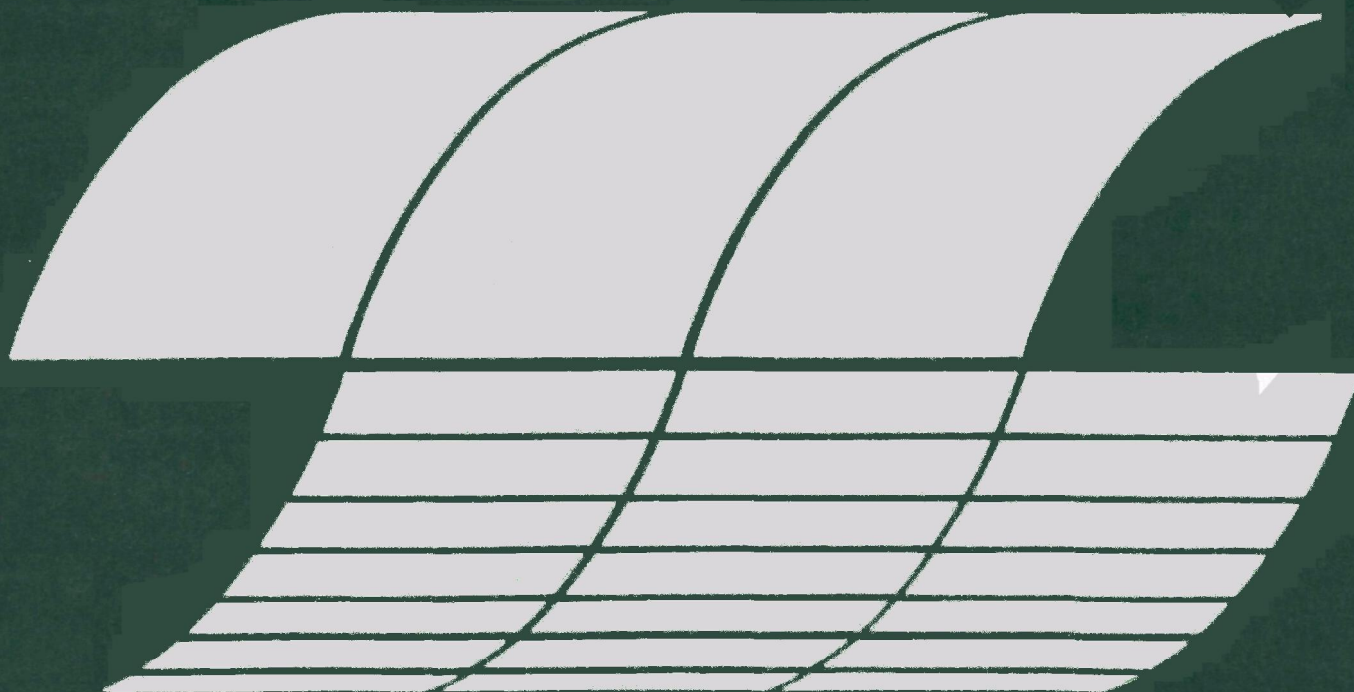


SO₂ ABATEMENT FOR STATIONARY SOURCES IN JAPAN

Interagency
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SO₂ ABATEMENT FOR STATIONARY SOURCES IN JAPAN

by

Jumpei Ando and B.A. Laseke

PEDCo. Environmental, Inc.
11499 Chester Road
Cincinnati, Ohio 45246

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EPA Project Officer: J. David Mobley

Industrial Environmental Research Laboratory
Office of Energy, Minerals, and Industry
Research Triangle Park, N.C. 27711

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FOREWORD

This report describes the status in Japan of technology for desulfurization of flue gases and for simultaneous removal of sulfur dioxide and nitrogen oxides from flue gas streams. The information is current through May 1976. The total capacity for flue gas desulfurization in Japan has reached 70 million normal cubic meters per hour (23,000 megawatt equivalent) and is expected to exceed 100 million cubic meters per hour (33,000 megawatt equivalent) by the end of 1977.

Ambient concentrations of sulfur dioxide in Japan have decreased markedly as a result of the desulfurization efforts by industry and the increasing use of low-sulfur fuels. The need for control of nitrogen oxides, however, has increased; efforts are being concentrated, therefore, on developing technologies for the simultaneous control of both classes of pollutants.

Section 1 of the report describes fuel use patterns in Japan, ambient concentrations of pollutants, and the current emission regulations.

Section 2 reviews the status of hydrodesulfurization of heavy oil, asphalt decomposition, and coal gasification

with emphasis on the new technology of asphalt decomposition to supply low-sulfur fuels.

Section 3 analyzes the status of flue gas desulfurization, including the major technical problems, trends, and economics.

Sections 4 and 5 describe in detail the applications of desulfurization technology by power companies and steel producers and the performance of new systems.

Sections 6 and 7 describe advanced processes for flue gas desulfurization and for simultaneous removal of the sulfur and nitrogen oxides.

Section 8 provides a comparative evaluation of advanced flue gas cleaning technologies in the United States and Japan.

REMARKS

The metric system is used in this report. Some of the conversion figures between the metric and English systems and abbreviations are shown below:

1 m (meter) = 3.3 feet

1 m³ (cubic meter) = 35.3 cubic feet

1 t (metric ton) = 1.1 short tons

1 kg (kilogram) = 2.2. pounds

1 liter = 0.26 gallon

1 kl (kiloliter) = 6.29 barrels

The capacity of SO₂ and NO_x removal plants is expressed in Nm³/hr (normal cubic meters per hour).

1 Nm³/hr = 0.59 standard cubic foot per minute

About 3,000 Nm³/hr is equivalent to 1 megawatt.

L/G ratio (liquid/gas ratio) is expressed in liters/Nm³.

1 liter/Nm³ = 7.4 gallons/1,000 standard cubic feet.

For monetary conversions, the exchange rate of 1 dollar = 300 yen is used.

SO₂ and NO_x removal costs are expressed in \$/kl oil. \$1/kl oil is equivalent to about 0.21 mil/kWh.

One liter heavy oil gives nearly 10,000 kcal (kilocalories).

1 kcal = 3.97 Btu.

Operability and availability, as referred to in the text, are defined as follows:

Operability (index): The number of hours the FGD system operated divided by the boiler operating hours in the period, expressed as a percentage.

Availability (index): The number of hours the FGD system was available for operation (whether operated or not) divided by hours in the period, expressed as a percentage.

ABBREVIATIONS

BPSD	Barrels per stream day
FGD	Flue gas desulfurization
HDS	Hydrodesulfurization
kW	Kilowatts
L/G	Liquid/gas ratio (see above)
MW ₃	Megawatts
Nm ³ /hr	Normal cubic meters per hour (see above)

TABLE OF CONTENTS

	<u>Page</u>
FORWARD	ii
CONVERSION FACTORS AND ABBREVIATIONS	iv
LIST OF FIGURES	ix
LIST OF TABLES	xii
LIST OF PHOTOS	xv
1. FUEL USAGE AND REGULATIONS ON SO _x AND NO _x	1-1
Supply and Usage of Energy	1-1
Emission and Regulation of SO _x	1-6
Total Mass Regulation of SO _x	1-11
Pollution-Related Health Damage Compensation Law	1-15
Emission and Regulation of NO _x	1-18
2. HYDRODESULFURIZATION AND DECOMPOSITION OF OIL AND GASIFICATION DESULFURIZATION	2-1
Status of Hydrodesulfurization (HDS)	2-1
Residual Oil Decomposition and Coal Gasification	2-4
3. GENERAL ASPECTS OF FLUE GAS DESULFURIZATION (FGD)	3-1
Trends	3-1
Major Processes and Systems	3-2
Wet Lime/Limestone Process	3-3

TABLE OF CONTENTS (continued)

		<u>Page</u>
	Indirect Lime/Limestone Process	3-12
	Other Processes (Recovery Processes)	3-16
	By-Products of FGD	3-21
	Wastewater and Gas Reheating	3-26
	Economic Aspects of FGD Systems	3-30
4.	MAJOR NEW FGD SYSTEMS FOR UTILITY BOILERS	4-1
	Status of FGD by Power Companies	4-1
	Plants Using the MHI Lime-Gypsum Process (Mitsubishi-JECCO Process)	4-6
	Mitsui-Chemico Limestone Gypsum Process at the Takasago Plant, Electric Power Development Co.	4-13
	Babcock-Hitachi Process at the Tamashima Plant, Chugoku Electric	4-17
	Kureha-Kawasaki Sodium-Limestone Process at the Sakaide Plant, Shikoku Electric	4-23
	Chiyoda Process at the Fukui Plant, Hokuriku Electric	4-29
5.	FLUE GAS DESULFURIZATION IN THE STEEL INDUSTRY	5-1
	Introduction	5-1
	MHI Process at the Mizushima Plant, Kawasaki Steel	5-3
	FGD Systems of Sumitomo Metal (Moretana Process)	5-7
	Kobe Steel Calcium Chloride Process	5-13
	Nippon Steel Slag Process (SSD Process)	5-17
6.	NEW FGD PROCESSES	6-1
	Status of New Developments	6-1
	Kawasaki Magnesium-Gypsum Process	6-2

TABLE OF CONTENTS (continued)

		<u>Page</u>
	MKK Lime-Gypsum Process Using Jet Scrubber	6-5
	Dowa Aluminum Sulfate Process	6-6
	Kurabo Ammonium Sulfate-Lime Process	6-15
	Kureha Sodium Acetate Process	6-18
	Mitsui-Chemico Magnesium Process	6-20
	Hitachi-Unitika Activated Carbon Process	6-23
7.	SIMULTANEOUS REMOVAL OF SO ₂ AND NO _x	7-1
	Outline	7-1
	Chemistry and Problems of Wet Processes	7-4
	Oxidation Reduction Processes	7-7
	Reduction Processes	7-13
	Dry Processes for Simultaneous Removal	7-25
8.	COMPARATIVE EVALUATION	8-1
	Differences Affecting Process Applications in the United States and Japan	8-1
	Wet Lime/Limestone Process	8-2
	Indirect Lime/Limestone Process	8-7
	Other Processes (Recovery Processes)	8-9
	By-Products and Wastewater	8-10
	Simultaneous Removal of SO _x and NO _x	8-13

LIST OF FIGURES

<u>Figure</u>		<u>Page</u>
1-1	Yearly Average SO ₂ Concentration in 15 Major Cities and Industrial Districts	1-2
1-2	SO ₂ Concentration in Kanagwa Prefecturs (Yearly Average in 1973 and 1977, PPM)	1-16
1-3	Increase of the Designated Patients	1-17
2-1	Flowsheet of Eureka Process	2-8
2-2	Material Balance for Residual Oil Decomposition Using Flexcoking Process	2-12
2-3	Flowsheet of Cherry Process	2-12
3-1	Schematic Flowsheet of Wet Lime/Limestone Gypsum Process	3-9
3-2	Production Capacity of Desulfurization	3-22
3-3	Price of By-Products	3-22
3-4	Demand for and Supply of Gypsum in Japan	3-23
3-5	FGD Capacity and Amount of Wastewater ⁶	3-28
3-6	Concentration of O ₂ and CL ⁻ in Solution and Stress Corrosion	3-28
4-1	One-Absorber System of MHI Process	4-7
4-2	Operation Data for Karita Plant, Kyushu Electric	4-11
4-3	Water Balance (Karita Plant, MHI Process)	4-12
4-4	Flowsheet of Mitsui-Chimico Process	4-14

LIST OF FIGURES (continued)

<u>Figure</u>		<u>Page</u>
4-5	Purge Water and Chloride Concentration	4-18
4-6	Flowsheet of Babcock-Hitachi Process	4-19
4-7	Relationship of pH of Slurry in Reactor to Solid Composition After Oxidation	4-24
4-8	Flowsheet of Kureha-Kawasaki Process	4-25
4-9	Flowsheet of Chiyoda Process	4-30
5-1	Performance of FGD Plant for No. 4 Sintering Machine	5-5
5-2	Flowsheet of Moretana Process (Kashima Plant, Sumitomo Metal)	5-8
5-3	Operation Data of No. 1 Train, Kashima Plant	5-11
5-4	Flowsheet of Cal Process	5-14
5-5	Flowsheet of SSD Process	5-18
6-1	Flowsheet of Kawasaki Magnesium-Gypsum Process	6-3
6-2	Dimensions of Jet Scrubber (MM) (120,000 Nm ³ /hr)	6-7
6-3	Flowsheet of MKK Jet-Scrubber Process (Naoshima Plant)	6-8
6-4	Flowsheet of Dowa Aluminum Sulfate-Limestone Process	6-9
6-5	Solubility Curves of SO ₂ in BAS Solution	6-11
6-6	Solubility Curves of SO ₂ at Various Temperatures of the Solution	6-11
6-7	Soluble Range of Aluminum Compound	6-11
6-8	Relationship of Al Loss to Concentration of Solution	6-11

LIST OF FIGURES (continued)

<u>Figure</u>		<u>Page</u>
6-9	Operation Data of Tamano Plant, Naikai	6-14
6-10	Flowsheet of Kurabo Ammonium Sulfate-Lime Process	6-16
6-11	Flowsheet of Kureha Sodium Acetate-Lime/Limestone Process	6-19
6-12	Flowsheet of Chemico-Mitsui Magnesium Process	6-21
7-1	Models of Combination of FGD and Denitrification	7-2
7-2	Simplified Flowsheet of Moretana Simultaneous Removal Process	7-8
7-3	MHI Simultaneous Removal Process	7-9
7-4	Schematic Flowsheet of IHI Simultaneous Removal Process	7-11
7-5	Flowsheet of Kureha Simultaneous Removal Process	7-17
7-6	Flowsheet of Mitsui Shipbuilding Process	7-17
7-7	Flowsheet of CEC Process	7-19
7-8	Flowsheet of Asahi Chemical Reduction Process	7-21
7-9	Flowsheet of Kawasaki Magnesium Process	7-24
7-10	Schematic Diagram of SO ₂ and NO _x Removal By Activated Carbon at Different Temperatures and Space Velocities	7-26
7-11	Flowsheet of Unitika Process	7-28

LIST OF TABLES

<u>Table</u>		<u>Page</u>
1-1	Supplies of Primary Energies in Japan	1-2
1-2	Power Generation Capacity	1-3
1-3	Power Generated	1-3
1-4	Domestic Demand For Heavy Oil	1-4
1-5	Consumption of Fuels By Power Companies	1-4
1-6	Cost of Fuels For Power Plants	1-5
1-7	Planned Imports of Liquefied Natural Gas	1-7
1-8	Long Term Energy Supply Plan	1-8
1-9	Ambient Air Quality Standards	1-9
1-10	Values of K Applicable To Locations In Japan	1-10
1-11	Relation of K Value To Maximum Ground-level Concentration of SO ₂	1-10
1-12	Allowable Sulfur Content of Oil - Existing and New Plants	1-14
1-13	NO _x Emission Standards	1-19
2-1	Hydrodesulfurization Systems Built by 1971	2-2
2-2	Hydrodesulfurization Systems Completed Between 1972 and 1976	2-3
2-3	Amount of Sulfur Recovered by HDS and FGD	2-5
2-4	Approximate Costs For HDS and FGD at Various Desulfurization Efficiencies	2-5
2-5	Typical Product Patterns	2-9

LIST OF TABLES (continued)

<u>Table</u>		<u>Page</u>
2-6	Economic Balance of Residual Oil Cracking Process	2-11
3-1	Numbers and Capacities of FGD Systems Expected to be Operational by End of 1977	3-4
3-2	Wet Lime/Limestone Process Units by MHI Process	3-5
3-3	Wet Lime/Limestone Process Units Using Scrubbers Developed in the United States	3-6
3-4	Wet Lime/Limestone Process Units Using Other Processes	3-7
3-5	Example of Operation Parameters of FGD Plants By-Producing Gypsum and Calcium Sulfite	3-10
3-6	Indirect Lime/Limestone Process Installations	3-13
3-7	FGD Installations By-Producing H_2SO_4 , S and $(NH_4)_2SO_4$	3-18
3-8	Wastewater From FGD Systems	3-27
3-9	Plant Cost In Battery Limits (\$1 = ¥300)	3-31
3-10	Examples of FGD Cost With Wet Lime-Gypsum Process	3-33
4-1	Capacities of Steam Power Generation and FGD of Power Companies	4-2
4-2	FGD Systems of Power Companies	4-4
4-3	Requirements at the Karita Plant	4-8
4-4	Main Equipment at Takasago Plant	4-16
4-5	Composition of Limestone	4-21

LIST OF TABLES (continued)

<u>Table</u>		<u>Page</u>
5-1	SO ₂ Removal Installation For Waste Gas From Iron-Ore Sintering Machines	5-2
5-2	Equipment Dimensions, FGD System At Kashima Plant	5-10
7-1	Processes For Simultaneous Removal of SO ₂ and NO _x	7-3

LIST OF PHOTOS

<u>Photos</u>		<u>Page</u>
2-1	Sodegaura Plant, Eureka	2-8
3-1	Handling of By-Product Gypsum (Chiba Plant, Showa Denko)	3-23
3-2	Calcium Sulfite Sludge Disposal (Omuta Plant, Mitsui Aluminum)	3-25
4-1	Karita Plant, Kyushu Electric (188 MW)	4-9
4-2	Owase Plant, Chubu Electric (2 Units Each 375 MW)	4-9
4-3	Takasago Plant, EPDC (250 MW) (Scrubber and Reactors)	4-15
4-4	Takasago Plant, EPDC (250 MW) (Gypsum Centrifuge and Storage)	4-15
4-5	Tamashima Plant, Chugoku Electric (500 MW)	4-20
4-6	Tamashima Plant, Chugoku Electric (500 MW)	4-20
4-7	Sakaide Plant, Shikoku Electric (450 MW) (Two Scrubbers in Parallel)	4-26
4-8	Sakaide Plant, Shikoku Electric (450 MW) (Oxidizers and Stripper)	4-26
4-9	Fukui Plant, Hokuriku Electric (350 MW)	4-32
4-10	Fukui Plant, Hokuriku Electric (350 MW)	4-32
5-1	Wakayama Plant, Sumitomo Metal	5-12

LIST OF PHOTOS (continued)

<u>Photos</u>		<u>Page</u>
6-1	Chiba Plant, Idemitsu Kosan (170 MW Equivalent, Scrubber and Kiln)	6-22
6-2	Chiba Plant, Idemitsu Kosan	6-22

1. FUEL USAGE AND REGULATIONS ON SO_x AND NO_x
SUPPLY AND USAGE OF ENERGY

The energy supply in Japan, after continuously growing at a yearly rate of more than 10 percent, has leveled off since 1974 because of economic depression caused by the international crisis in marketing of oil (Table 1-1). Supplies of imported oil, which accounts for more than 70 percent of Japan's total energy, decreased in 1974. Production of electric power in 1974 was slightly less than that in 1973, although the generating capacity increased (Tables 1-2 and 1-3).

Consumption of heavy oil, the major fuel in Japan, also decreased in 1974. Use of high-sulfur heavy oil (grade C) decreased markedly, whereas use of low-sulfur heavy oil (grade A) increased slightly (Table 1-4). Consumption of heavy oil by power companies also dropped, but use of sulfur-free fuels such as naphtha and LNG (liquefied natural gas) increased (Table 1-5). The average sulfur content of oils consumed by power companies decreased from 1.5 percent in 1970 to 0.54 percent in 1974. The price difference between low-sulfur and high-sulfur oils was about \$30/kl in 1974 and \$25/kl in 1976 (Table 1-6).

Table 1-1. SUPPLIES OF PRIMARY ENERGIES IN JAPAN¹

	1970	1971	1972	1973	1974
Hydroelectric power, 10^6 MWhr	80.0	86.8	87.9	71.6	84.8
Atomic power, 10^6 MWhr	4.5	8.0	9.5	9.7	19.7
Coal, 10^6 t					
Domestic	40.8	33.8	28.1	21.7	21.4
Imported	50.9	46.5	50.5	58.0	64.6
Oil, 10^6 kl					
Domestic	0.9	0.9	0.8	0.8	0.8
Imported	234.1	248.7	275.7	318.6	305.8
Natural gas, 10^9 m ³	2.8	2.9	2.9	2.9	2.8
LNG, 10^6 t	0.9	1.0	1.0	2.3	2.8
Other, 10^{13} kcal	1.0	1.4	1.2	0.6	1.0
Total, 10^{13} kcal	310.4	320.6	344.3	382.5	383.5

Table 1-2. POWER GENERATION CAPACITY

(1,000 MW)¹

	1970	1971	1972	1973	1974
Hydro	20.0	20.1	20.7	22.6	23.5
Thermal	47.0	54.9	62.7	70.6	76.9
Atomic	1.3	1.3	1.8	2.3	3.9
Total	68.3	76.3	85.2	95.5	104.2

Table 1-3. POWER GENERATED¹(10⁶ MWhr)

	1970	1971	1972	1973	1974
Hydroelectric	80.1	86.8	87.9	71.6	84.8
Thermal	274.8	290.8	331.1	388.8	354.6
Atomic	4.6	8.0	9.5	9.7	19.7
Total	359.5	385.6	428.5	470.1	459.0

Table 1-4. DOMESTIC DEMAND FOR HEAVY OIL¹(10⁶ kl)

	1970	1971	1972	1973	1974
Grade A (S=0.1-1.0)	11.1	13.3	16.2	19.3	19.6
Grade B (S=0.3-2.5)	12.7	12.7	12.9	12.8	11.9
Grade C (S=0.5-4.0)	90.0	92.4	95.6	105.0	86.6
Total	113.8	118.4	124.7	137.1	118.1

Table 1-5. CONSUMPTION OF FUELS BY POWER COMPANIES¹

	1970	1971	1972	1973	1974
Coal, 10 ⁶ t	18.8	13.9	10.7	8.3	7.3
Heavy oil, 10 ⁶ kl	34.5	35.3	38.1	42.8	34.7
Crude oil, 10 ⁶ kl	7.2	11.0	17.8	23.6	23.0
Naphtha, 10 ⁶ kl	0	0	0.2	2.2	3.5
LNG, 10 ⁶ t	0.7	0.7	0.6	1.4	2.5
Natural gas, 10 ⁶ Nm ³	0.07	0.13	0.17	0.22	0.24
Blast furnace and coke oven gases, 10 ⁶ Nm ³	15.6	18.1	22.9	32.4	32.4
Average sulfur content, of oil, %	1.50	1.31	1.03	1.75	0.54

Table 1-6. COST OF FUELS FOR POWER PLANTS

Fuel	s, %	1974		1976	
		\$/kl (t)	¢/M Btu	\$/kl (l)	¢/M Btu
Naphtha	0.02			110-115	245-255
Crude oil	0.3	86-90	210-220	96-100	234-244
Heavy oil	0.3	88-92	220-230	97-100	242-250
Heavy oil	1.6	72-75	180-188	80-84	200-210
Heavy oil	3.0	59-61	148-153	72-75	180-188
Coal	1-2.5	(19-29)	108-133		

Domestic coal has been used for fuel. Although coal is cheaper than oil, as shown in Table 1-6, consumption will not increase because of the limited capacity of domestic coal mines. So far, all of the coal imported to Japan has been used for coke production for the steel industry. Importing of coal for fuel is expected to start in a year or two and to increase. In recent years, great efforts have been made to increase the imports of LNG because it causes no emissions of SO_x and lower emissions of NO_x than other fuels. Many contracts are under way with several countries (Table 1-7). There is considerable uncertainty with respect to the amount and the costs of LNG to be imported.

It is expected that the Japanese economy will recover by the end of 1976 and will gain strength slowly, with energy consumption increasing at a rate of 5 to 6 percent yearly (Table 1-8).

EMISSION AND REGULATION OF SO_x

Japan today depends upon imported crude oil for more than 70 percent of its total energy supply. In 1974 and 1975, Japan imported nearly 300 million kiloliters of crude oil, which contained nearly 3 million tons of sulfur. Most of the sulfur in crude oil goes into heavy oil. More than 40 plants for hydrodesulfurization of heavy oil have been completed, and in 1975 about 30 percent of the oil was treated to produce 750,000 tons of elemental sulfur. Nearly

Table 1-7. PLANNED IMPORTS OF LIQUEFIED NATURAL GAS¹

(LNG, 1,000 t)

Source	Buyer	1973	1975	1977	1979	1981	1983	1985
Alaska	Tokyo Electric	720	720	720	720	720	720	720
	Tokyo Gas	240	240	240	240	240	240	240
Brunei	Tokyo Electric	580	2,920	3,450	3,450	960	960	960
	Tokyo Gas	370	710	1,060	1,060	1,060	1,060	1,060
	Osaka Gas	350	500	630	630	630	630	630
Abu Dhabi	Tokyo Electric			2,800	2,800	2,800	2,800	2,800
Indonesia	Kansai Electric			100	1,490	2,400	2,400	2,400
	Chubu Electric			500	1,350	1,700	1,700	1,700
	Kyushu Electric			300	1,340	1,500	1,500	1,500
	Osaka Gas			380	1,140	1,300	1,300	1,300
	Nippon Steel			380	590	600	600	600
Sarawak	Tokyo Electric					4,000	4,000	4,000
	Mitsubishi Shoji					2,000	2,000	2,000
Iran						2,500	2,500	2,500
Other							2,000	7,000
Total		2,260	5,090	10,560	14,850	24,900	26,900	31,900

Table 1-8. LONG TERM ENERGY SUPPLY PLAN¹

(MITI, August 1975)

	1973		1980		1985	
	Amount	%	Amount	%	Amount	%
Domestic energy						
Hydroelectric, 10 ⁶ MWhr	71.6	4.6	85.5	4.2 ^a	100.3 ^a	3.7
Geothermal, 10 ⁶ MWhr	0.1	0.0	2.1	0.1	14.7	0.5
Oil (gas), 10 ⁶ kl	3.7	0.9	6.4	1.2	14.0	1.8
Coal, 10 ⁶ t	21.7	3.8	20.0	2.5	20.0	1.9
Atomic power, 10 ⁶ MWhr	9.7	0.6	95.4	4.4	278.3	9.6
Imported energies						
LNG, 10 ⁶ t	2.4	0.8	20.6	5.2	42.0	7.9
Coal (for coke), 10 ⁶ t	58.0	11.7	87.3	12.7	87.8	9.6
Coal (for fuel), 10 ⁶ t	0	0	4.7	0.7	14.6	1.6
Oil, 10 ⁶ kl	318.0	77.4	393.0	68.9	485.0	63.3
Total		100.0		100.0		100.0

^a Including power to be generated by pumped storage power plants.

2 million tons of sulfur in heavy oil and crude oil was burned, constituting about 75 percent of the total emission of SO₂. About one-fourth of the heavy oil was burned in utility boilers of power companies and the rest by other industries. The ambient standard for SO_x concentration was changed from 0.05 ppm (yearly average) to 0.04 ppm (daily average) in May 1973. By the new standard, the hourly average should not exceed 0.1 ppm and the daily average should not exceed 0.04 ppm. The standard is much more stringent than those in the U.S.A. and West Germany (Table 1-9). The values of K applicable to specific existing and new plant locations in Japan are provided in Table 1-10. The relation of K-values to maximum allowable ground-level SO₂ concentrations is presented in Table 1-11.

Table 1-9. AMBIENT AIR QUALITY STANDARDS

(Daily or yearly average, converted to ppm)

	SO _x		NO ₂	
	Daily	Yearly	Daily	Yearly
Japan	0.04	0.016	0.02	0.008
United States		0.03	0.13	0.05
West Germany		0.05		0.05

The emission standard is given by the following equation:

$$Q = K \times 10^{-3} \text{ He}_2$$

Q: Amount of sulfur oxides, Nm³/hr
(1 Nm³/hr = 0.59 scfm).

K: The value shown in Table 1-10.

He: Effective height of stack, meters
(1 meter = 3.3 ft).

Table 1-10. VALUES OF K APPLICABLE TO LOCATIONS IN JAPAN

(For existing plants)

K = 3.0	K = 3.5	K = 4.67	K = 8.76	K = 14.6
Tokyo	Chiba	Sapporo	Hachinohe	Hakodate
Yokohama	Fuji	Kashima	Sendai	Miyako
Kawasaki	Handa	Shimizu	Niigata	Mobara
Yokkaichi	Himeji	Tokuyama	Okayama	Sasebo
Osaka	Mizushima	Omuta	Futuoka	Kagoshima

(For new plants)

K = 1.17	K = 1.75	K = 2.34
Tokyo, Yokohama	Chiba, Fuji	Kashima, Omuta
Kawasaki, Nogoya	Kitakyushu	Ube, Oita
Yokkaichi, Osaka	Handa, Himeji	Shimizu, Kyoto

Table 1-11. RELATION OF K VALUE TO MAXIMUM
GROUND-LEVEL CONCENTRATION OF SO₂

(ppm)

K	1.17	1.75	2.34	3.50	4.67	8.76	14.6
SO ₂	0.002	0.003	0.004	0.006	0.008	0.05	0.025

For a new plant with a capacity of 500 MW in the Tokyo and Osaka areas, the sulfur content of oil must be below 0.3 percent, even with a 200-meter stack. Through application of the regulations and such efforts as importing low-sulfur fuels, hydrodesulfurization of heavy oil, and flue gas desulfurization, ambient SO₂ concentrations in Japan have decreased, as shown in Figure 1-1.

TOTAL MASS REGULATION OF SO_x

The emission standard has not succeeded in keeping the ambient concentration below 0.04 ppm daily average (0.016 ppm yearly average) in large cities and heavy industrial areas. With the aim of attaining the ambient standard by March 1978, the central government promulgated a new regulation in November 1974 to restrict the total mass of SO₂ emissions from the following eleven polluted areas: (1) Tokyo (2) Chiba (3) Yokohama, Kawasaki (4) Fuji (5) Nagoya (6) Handa (7) Yokkaichi (8) Osaka, Sakai (9) Kobe, Amagasaki (10) Kurashiki, Mizushima (11) Kitakyushu. The new regulation applies to plants using more than 0.1 to 1.0 kiloliter of oil per hour (0.4 to 4.0 MW equivalent; a certain number between 0.1 and 1.0 is to be assigned to each prefecture by the Governor). The amount of allowable SO_x is calculated from one of the following formulas, to be selected by each prefecture:

$$Q = a \times W^b \quad (1)$$

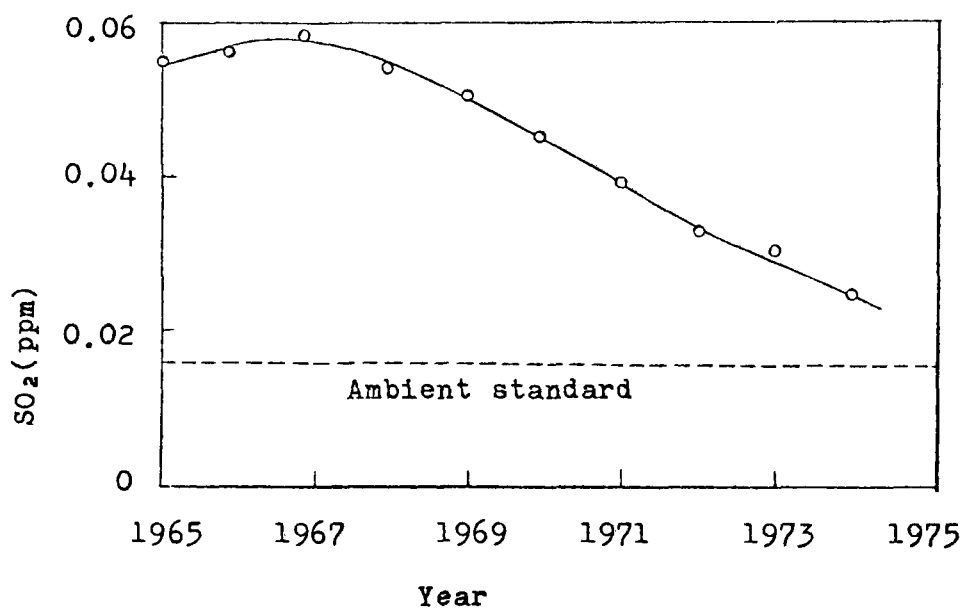


Figure 1-1. Yearly average SO_2 concentration in
15 major cities and industrial districts

- Q: Amount of allowable SO_x
- a: A constant to ensure SO_x abatement
- W: Amount of fuel used by each plant
- b: A constant between 1.00 and 0.80 to be selected by the prefectural governor.

$$Q = Q_o \times C_m / C_{m_o} \quad (2)$$

- Q: Amount of allowable SO_x
- C_m: Maximum ground level concentration to ensure SO_x abatement
- C_{m_o}: Maximum ground level concentration due to each plant
- Q_o: Amount of SO_x being emitted

Two prefectures, Mie (with Yokkaichi city) and Kanagawa (with Yokohama and Kawasaki cities), put the regulation in force recently. The regulations of Kanagawa are shown below.

$$Q = 1.5 W^{0.865} + 0.5 W_i^{0.865} \dots \dots \dots (I)$$

$$Q = 2.5 W^{0.865} + 0.8 W_i^{0.865} \dots \dots \dots (II)$$

Q: Amount of allowable SO_x, Nm³/hr

W: Fuel consumption by existing plants, kl/hr

W_i: Fuel consumption by new plants, kl/hr

The equations are applied to plants that consume more than 1 kl oil per hour. Equation (I) is for most polluted districts in Yokohama and Kawasaki cities, and (II) is for other districts of the cities and for all parts of Yokosuka city. The allowable sulfur content of fuel oil, as calculated from the equations for different fuel consumptions, is shown in Table 1-12.

Table 1-12. ALLOWABLE SULFUR CONTENT OF OIL -
EXISTING AND NEW PLANTS

(Percent)				
W	1	10	100	1,000
(MW equivalent)	(4.5)	(45)	(450)	(4,500)
Sulfur (I)	0.238	1.174	0.124	0.094
Sulfur (II)	0.397	0.290	0.207	0.155
W_i	1	10	100	1,000
(MW equivalent)	(4.5)	(45)	(450)	(4,500)
Sulfur (I)	0.079	0.058	0.041	0.031
Sulfur (II)	0.127	0.093	0.066	0.050

An existing 450-MW plant in district (I) is allowed to use oil with less than 0.124 percent sulfur. If the plant has two 225 MW units and one of them uses LNG with no sulfur, the other unit may use oil with 0.248 percent sulfur. New plants in district (I) need to use oil with less than 0.079 percent sulfur. Those plants will have to use naphtha, kerosene, or gas.

For plants consuming less than 1 kl/hr oil, sulfur content must be below 0.3 percent in district (I) and below 0.5 percent in district (II). The sulfur contents of fuel oils for diesel engine cars and ships are to be restricted to 0.2 and 0.5 percent, respectively. In districts other than (I) and (II), stationary sources are regulated by the national K-value control.

By those regulations, the total emission of SO_x in districts (I) and (II), which was $5,348 \text{ Nm}^3/\text{hr}$ in 1973, will decrease to $2,078 \text{ Nm}^3/\text{hr}$ by the end of 1977; all districts in the Kanagawa prefecture will have an SO_x concentration below the national environmental standard, 0.016 ppm yearly average (Figure 1-2).

POLLUTION-RELATED HEALTH DAMAGE COMPENSATION LAW

One of the driving forces for progress in SO_2 abatement has been the "Pollution-Related Health Damage Compensation Law" which has been in effect since 1972. By the law, certain regions with prevalent pollution are designated as polluted areas, and certain inhabitants who are diagnosed by nominated doctors to have pollution-related sickness such as chronic bronchitis are designated as pollution-related patients. Firms which emit more than $5,000 \text{ Nm}^3/\text{hr}$ of flue gas in those regions must pay a tax according to the amount of SO_2 ; and tax is used to provide medical care for the patients.

The number of designated patients increased remarkably, as shown in Figure 1-3. Most of their illnesses are considered due to air pollution, mainly by SO_2 . The total tax paid by the firms increased from \$11 million in 1974 to \$52 million in 1975; most of the tax was paid for SO_2 emissions. The tax rate, which changes each year, was $26¢/\text{Nm}^3 \text{ SO}_2$ in 1975. For example, a firm with a 100-MW boiler using 0.6

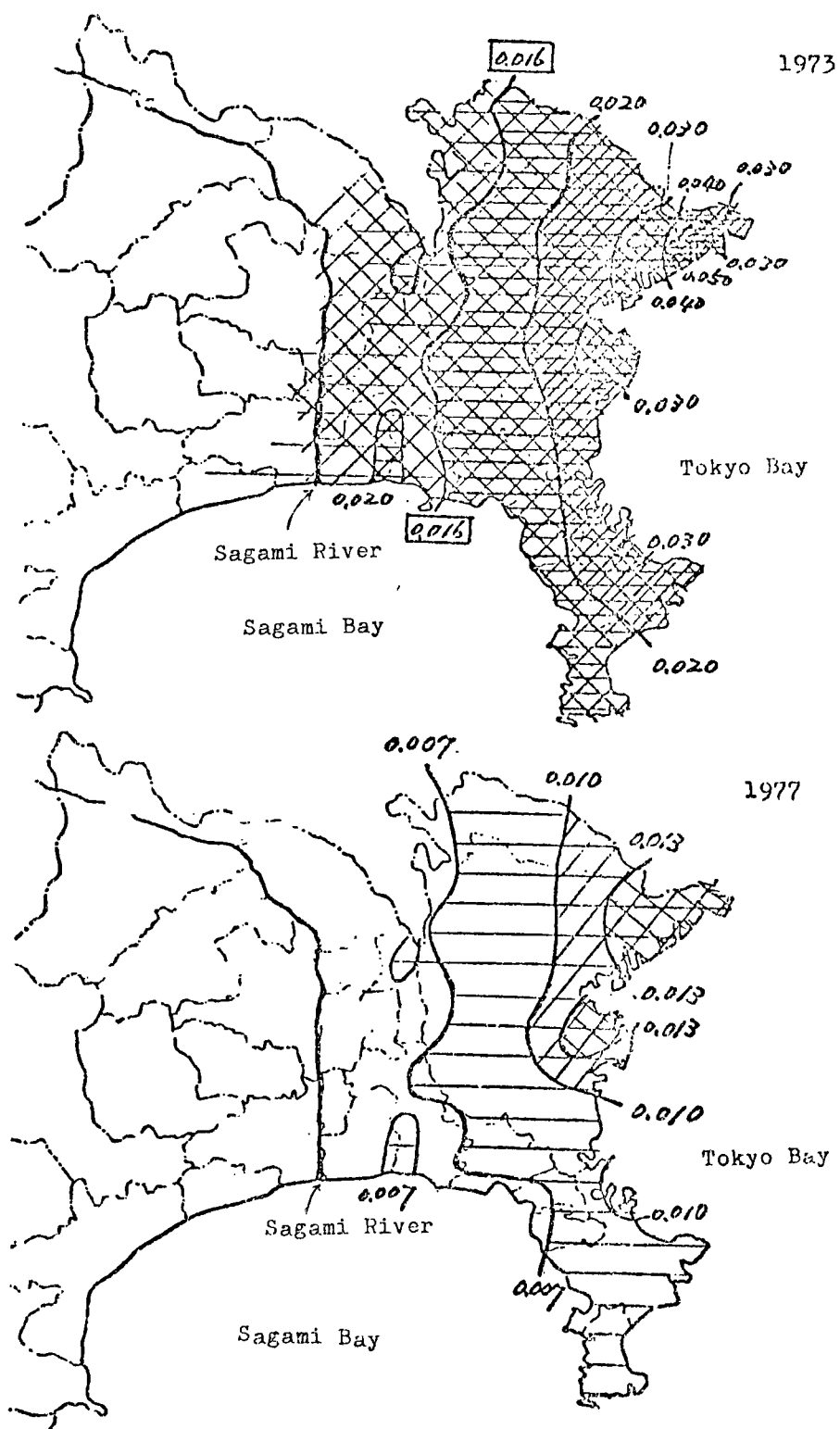


Figure 1-2. SO_2 concentration in Kanagawa prefecture
(yearly average in 1973 and 1977, ppm).

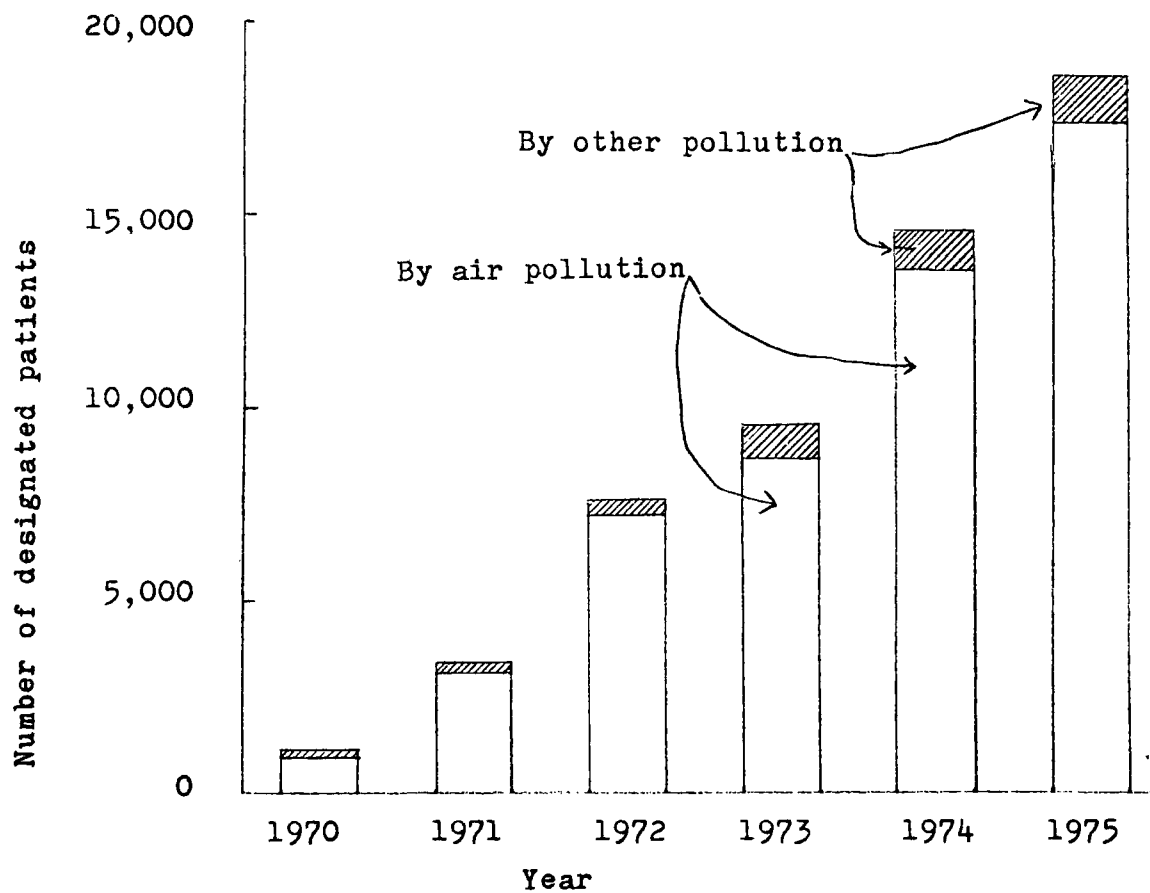


Figure 1-3. Increase of the designated patients.

percent sulfur oil paid about \$160,000 in 1975. The tax is equivalent to about 1 percent of the cost of the fuel oil.

Several organizations now claim that the system may not be working properly because the number of designated patients has increased markedly in spite of the remarkable decrease in ambient SO_2 concentrations. It is likely that the numbers of designated regions and patients will decrease in the future.

EMISSION AND REGULATION OF NO_x

Total emissions of NO_x in Japan are estimated at about 2 million tons yearly. More than 90 percent of the NO_x is caused by the burning of fuels, such as heavy oil and gasoline. About 40 percent of the total NO_x is derived from automotive exhausts, 20 percent from electric power generation, 30 percent from industry, and the rest from household heating, etc. In large cities such as Tokyo and Osaka, 60 to 70 percent of the NO_x is traced to automobiles.

The ambient air quality standard for NO_2 was set forth in 1973 at 0.02 ppm daily average, the most stringent standard in the world (Table 1-9). The present yearly average NO_2 concentration ranges from 0.02 to 0.03 ppm, and the daily average often reaches 0.04 to 0.07 ppm in many cities.

The NO_x emission standard for stationary sources was first set up in 1973 and revised in 1975. Table 1-13 shows

the standard for boilers larger than 100,000 Nm³/hr. Similar figures have been assigned to smaller boilers between 10,000 and 100,000 Nm³/hr since 1975. This standard also is more stringent than those in the U.S.A. and other countries.

Table 1-13. NO_x EMISSION STANDARDS
(ppm)

Fuel	For new boilers		For existing boilers	
	1973	1975	1973	1975
Gas	130	100	170	130
Oil	180	150	230	230
Coal	480	480	750	750

Even though combustion modifications and improvement of burners have been undertaken in efforts to meet the standard, the ambient air quality standard has not been attained, even with the stringent emission standard. More stringent regulations to restrict the total NO_x emissions from stationary sources are to be applied in several prefectures. The new regulations will require construction of many flue gas denitrification plants, which remove more than 80 percent of the NO_x.

2. HYDRODESULFURIZATION AND DECOMPOSITION OF OIL AND GASIFICATION DESULFURIZATION

STATUS OF HYDRODESULFURIZATION (HDS)

Eighteen heavy oil HDS systems have been constructed since 1967, as shown in Tables 2-1 and 2-2. HDS is accomplished by two methods. One is vacuum gas-oil HDS, by which the vacuum gas-oil obtained by vacuum distillation of heavy oil is desulfurized to about 0.2 percent sulfur. Although this treatment is relatively easy, it cannot desulfurize the residual oil from the distillation, which amounts to about 40 percent of the heavy oil and is rich in sulfur and metallic impurities. The second method is topped-crude HDS, by which heavy oil is treated directly. It is difficult to reduce sulfur content below 1 percent by this process. Since 1 percent sulfur oil has become unsatisfactory for use in many places, several oil companies, including Idemitsu Kosan, Seibu Oil, Asia Oil, and Maruzen Oil, have constructed new topped-crude HDS process plants to reduce sulfur to 0.3 percent or below by use of several reactors in series. Hydrogen consumption to decrease sulfur from 1.0 to 0.3 percent is about equal to that required to reduce it from 2.5 to 1.0 percent. About 700,000 tons of elemental sulfur

Table 2-1. HYDRODESULFURIZATION SYSTEMS BUILT BY 1971

Refiner	Plant site	Process	Completed	Capacity, per day	
				Oil, BPSD	Sulfur, tons
Idemitsu Kosan	Chiba	UOP ^a	1967	40,000	265
Fuji Oil	Sodegaura	CRC	1968	23,000	100
Toa Nenryo	Wakayama	ER & E	1968	25,000	180
Daikyo Oil	Umaokoshi	Gulf	1969	17,500	110
Nippon Oil	Negishi	CRC	1969	40,000	190
Showa Oil	Kawasaki	Shell	1969	16,000	66
Kyushu Oil	Oita	Shell	1969	14,000	55
Mitsubishi Oil	Mizushima	UOP	1969	30,000	100
Maruzen Oil	Chiba	Union	1969	35,000	165
Seibu Oil	Yamaguchi	Shell	1969	4,000	28
Nippon Mining	Mizushima	Gulf ^a	1970	27,760	165
Koa Oil	Marifu	CRC	1970	8,000	39
General Oil	Sakai	ER & E	1970	31,000	73
Kashima Oil	Kashima	UOP ^a	1970	45,000	265
Daikyo Oil	Umaokoshi	Gulf	1970	17,500	77
Kansai Oil	Sakai	ER & E	1971	20,000	88
Koa Oil	Osaka	CRC	1971	12,000	55
Toa Nenryo	Kawasaki	ER & E	1971	51,000	220
Total				465,260	2,241

^a Topped-crude HDS processes; all others are for vacuum gas-oil HDS.

Table 2-2. HYDRODESULFURIZATION SYSTEMS COMPLETED
BETWEEN 1972 AND 1976

Refiner	Plant site	Process	Year of completion	Capacity oil, BPSD
Nippon Oil	Negishi	CRC	1972	28,000
Idemitsu Kosan	Himeji	Gulf ^a	1972	40,000
Kyokuto Petroleum	Chiba	UOP	1972	60,000
Toa Nenryo	Kawasaki	ER & E	1972	9,000
Asia Kyoseki	Sakaide	CRC	1972	15,000
Kyushu Oil	Oita	UOP	1972	25,000
Showa Yokkaichi Oil	Yokkaichi	Shell	1972	35,000
Seibu Oil	Yamaguchi	Shell	1972	1,000
Nippon Oil	Muroran	CRC	1973	40,000
Toa Oil	Nagoya	CRC	1973	30,000
Nippon Mining	Mizushima	Gulf ^a	1974	3,240
Toa Oil	Nagoya	ER & E	1974	37,000
Kansai Oil	Sakai	ER & E	1974	2,000
Showa Yokkaichi Oil	Yokkaichi	Shell	1974	5,000
Mitsubishi Oil	Mizushima	UOP ^a	1974	45,000
Nippon Mining	Mizushima	UOP	1975	60,000
Idemitsu Kosan	Chiba	UOP	1975	34,000
Idemitsu Kosan	Tokuyama	UOP	1975	45,000
Idemitsu Kosan	Aichi	Gulf ^b	1975	50,000
Seibu Oil	Yamaguchi	Shell ^b	1975	45,000
Asia Oil	Yokohama	UOP ^b	1975	30,000
Asia Oil	Sakaide	Gulf ^b	1976	28,000
Toa Oil	Kawasaki	ER & E	1976	46,000
Fuji Oil	Sodegaura	CRC	1976	35,000
Maruzen Oil	Chiba	ER & E ^b	1976	60,000
Total				839,240

^a Topped-crude HDS.

^b New topped-crude HDS.

All others are for vacuum gas-oil HDS.

was recovered in 1973 by HDS; sulfur recovery has not increased much since then (Table 2-3), in spite of the completion of many new HDS plants. The slowdown in sulfur recovery was caused by economic depression; in 1975 operation of the HDS plants averaged only about 4,700 hours (54 percent of possible total of 8,760 hours).

About 25 HDS systems with a total capacity of nearly 1 million BPSD were planned for construction between 1976 and 1978. Most of the plans, however, were abandoned or postponed, partly because of the economic depression and partly because a great increase in the demand for heavy oil had become unlikely.

A rough estimate of HDS costs for heavy oil containing 3 percent sulfur at various desulfurization efficiencies is shown in Table 2-4, with FGD cost given for comparison. HDS is more expensive than FGD by the wet lime/limestone process but it is advantageous in that it produces elemental sulfur, which is a desirable by-product. The sulfur-by-producing FGD processes developed in Japan and other countries are more expensive, except for oil refineries that have Claus furnaces and hydrogen sulfide for sulfur production.

RESIDUAL OIL DECOMPOSITION AND COAL GASIFICATION

Outline

One of the major problems associated with HDS Technology is that its application is limited to heavy oil, containing

Table 2-3. AMOUNT OF SULFUR RECOVERED BY

HDS AND FGD

(1,000 tons)

	1973	1974	1975	1976
HDS	700	720	750	(800)
FGD ^a	100	200	350	(550)

^a Rough estimates including all sulfur compounds as converted into sulfur.

Table 2-4. APPROXIMATE COSTS FOR HDS AND FGD AT

VARIOUS DESULFURIZATION EFFICIENCIES

(\$/kl, at 6,000 hours operation per year)

Sulfur removal, %	70	80	90	97
Hydrodesulfurization	16	19	23	27
Flue gas desulfurization ^a	16	17	18	19

^a Wet lime/limestone process by-producing gypsum.

only small amounts of impurities because metallic impurities in the oil poison the HDS catalysts. It is estimated that only about 15 percent of the total heavy oil in Japan is suitable for the new topped-crude HDS process to reduce sulfur to 0.3 percent or below.

As demand for low-sulfur fuels increased, gasification desulfurization seemed promising in 1973, and many companies planned to construct gasification plants. Because of inflation following the oil crisis, however, most of the plans were given up. Ube Industries constructed a pilot plant to gasify 50 tons per day of heavy oil.² The Coal Research Center constructed a pilot plant to gasify 5 tons per day of coal. Activity at those works has been limited, however, by the economic situation.

Thermal decomposition of the vacuum residue of heavy oil (asphalt) with little gasification seems more promising. Two commercial plants have been constructed for this purpose. A plant with a capacity of treating 18,300 BPSD of asphalt was completed recently at Sodegaura Refinery, Fuji Oil, using the Eureka (Kureha) process. A plant capable of treating 50,000 BPSD of heavy oil is near completion at Kawasaki Refinery, Toa Oil, using the Flexicoking process.

Osaka Gas has developed a new process of asphalt decomposition and is operating a pilot plant.

Eureka Process (Kureha Process)⁴

Kureha Chemical Industry has developed a new process to decompose the residue from vacuum distillation of heavy oil (asphalt) using steam to produce a cracked oil (about 65%), gas (about 5 wt %) and pitch (about 30%). A commercial plant owned by Eureka Industry Co. (established by Kureha Chemical jointly with Fuji Oil, Arabia Oil, and Sumitomo Metal) with a capacity of treating about 1 million tons of the residual oil yearly was completed in February 1976 at Sodegaura, Chiba Prefecture, and has been operating smoothly since March 1976 (Photo 2-1).

A flowsheet of the process is shown in Figure 2-1. The residual oil containing 4 to 5 percent sulfur is treated with steam heated to above 700°C in reactors at 500°C for several hours. The steam carries heat and promotes distillation.

Typical product patterns are shown in Table 2-5. The cracked naphtha and gas oil contain few heavy metal impurities and are easily treated by hydrodesulfurization. The gas from the reactor contains about 15 percent H_2S , which is removed by a conventional process using an amine. The purified gas (about 16,000 kcal/m³) is used for fuel. The pitch, which contains 4 to 8 percent sulfur, is used by Sumitomo Metal as a binder for poor-coking coal in coke

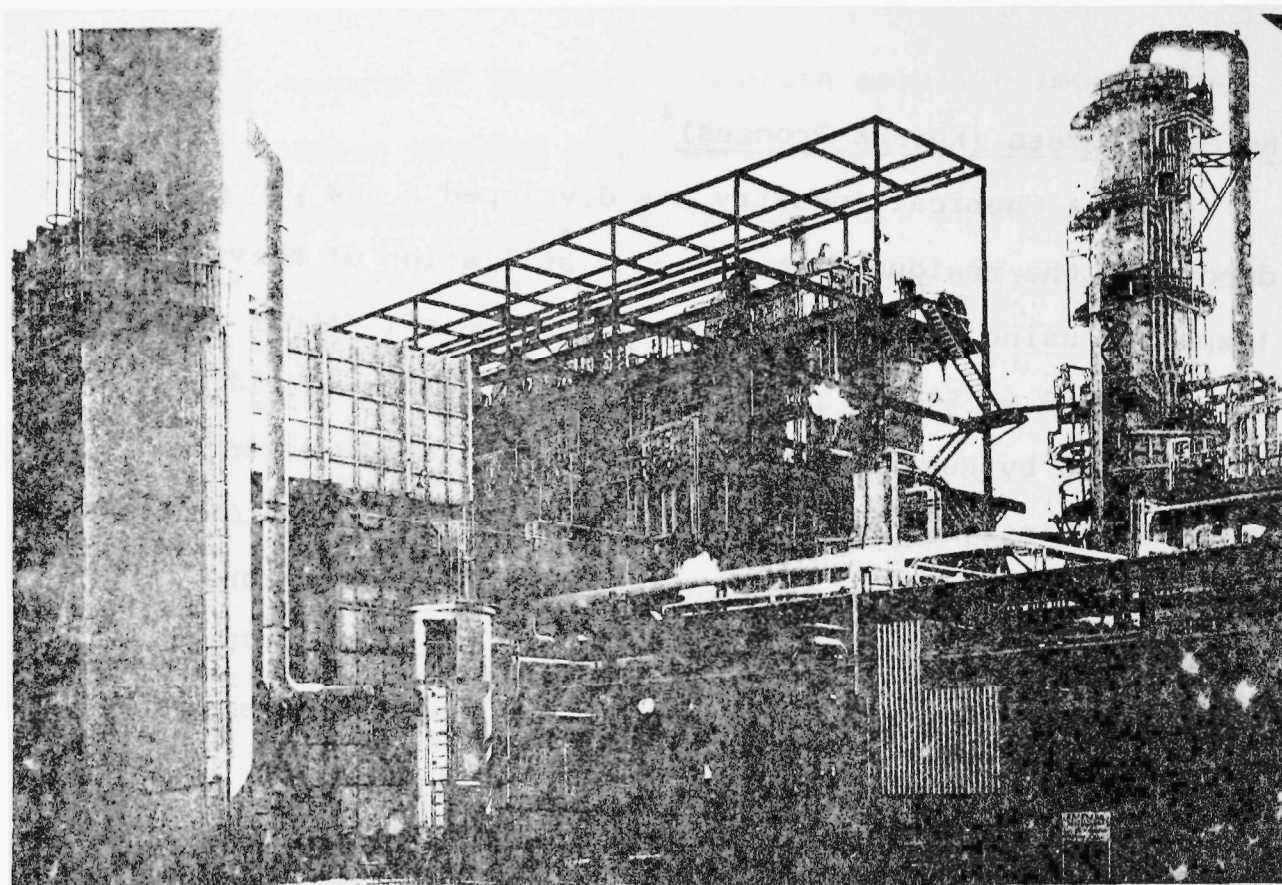


Photo 2-1. Sodegaura plant, Eureka.

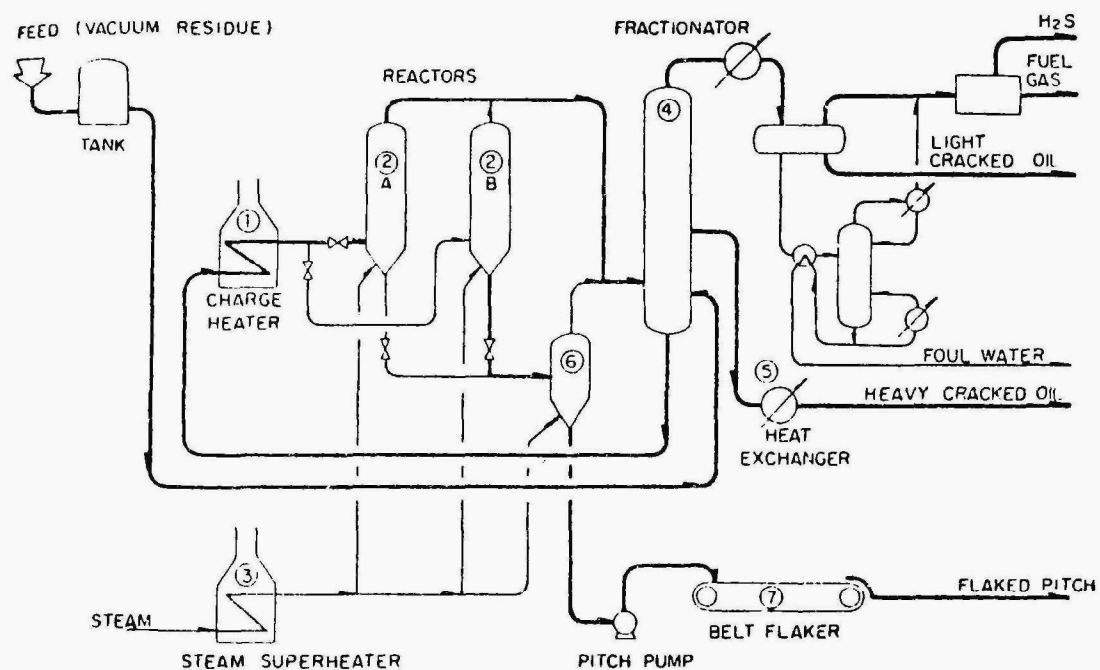


Figure 2-1. Flowsheet of Eureka process.

Table 2-5. TYPICAL PRODUCT PATTERNS

(Values in percent by weight)

Products	Khafji vacuum residue			Iranian Heavy vacuum residue		
	Yield	Sulfur content	Sulfur distribution	Yield	Sulfur content	Sulfur distribution
Gas	5.2	14.5	14.2	5.4	10.7	16.8
Cracked naphtha	8.3	1.9	3.0	11.3	1.7	5.6
Cracked gas oil	54.7	3.6	37.2	52.9	2.5	38.5
Pitch	31.8	7.6	45.6	30.4	4.4	39.1

NOTE: 1. Sulfur contents of Khafji vacuum residue and Iranian Heavy vacuum residue are 5.30 and 3.43 (% w), respectively.

2. Main components of gas are C₁-C₅ paraffins and olefins.

production. Kureha recently developed a process to produce a spherical activated carbon with the pitch.

The economic balance of the residual oil cracking process is shown in Table 2-6.

Flexicoking

A commercial plant with a capacity of treating 50,000 bbl/day heavy oil by vacuum distillation and 21,000 bbl/day residual oil from the distillator (asphalt) by Flexicoking is under construction and scheduled to be completed by summer 1976 at Kawasaki refinery, Toa Oil. A rough material balance is shown in Figure 2-2. Vacuum gas oil from the distillator and gas oil from the Flexicoker will be treated by HDS (Gofiner). High-calorie gas will be sold to an adjacent steel producer, low-calorie gas will be consumed by Toa Oil, and the coke will be sold.

Cherry Process

Osaka Gas Co., jointly with Mitsubishi Heavy Industries, has developed a new asphalt decomposition process to solve a common problem--coking on the reactor walls--by adding a small amount of pulverized coal to the asphalt. A pilot plant with a capacity of treating 26 t/day of asphalt was completed recently. A flowsheet is shown in Figure 2-3. Asphalt with a small amount of coal powder is heated to 400°C in a furnace, cracked and polymerized in a reactor,

Table 2-6. ECONOMIC BALANCE OF RESIDUAL OIL CRACKING PROCESS

(1 million tons of vacuum residue per year)

	Unit price		Cost	
	(yen/kg)	(¢/lb)	(million yen/year)	(million \$/year)
Raw material				
Vacuum residue, 1 million ton/yr	13.2	2	13,200	44
Processing cost				
Utilities			930	3.1
Fixed cost (30% of investment)			3,087	10.3
			17,217	57.4
Products cost				
Cracked oils 654 million ton/yr	19.8	3	12,930	43.1
Pitch 300 million ton/yr	14.3	2.17	4,287	14.3
			17,217	57.4

NOTE: 1. Investment including off-site is 10,290 million yen in Japan, 1975.

2. The price of coal tar pitch is considered between 3¢/lb and 5¢/lb in Japan, 1975.

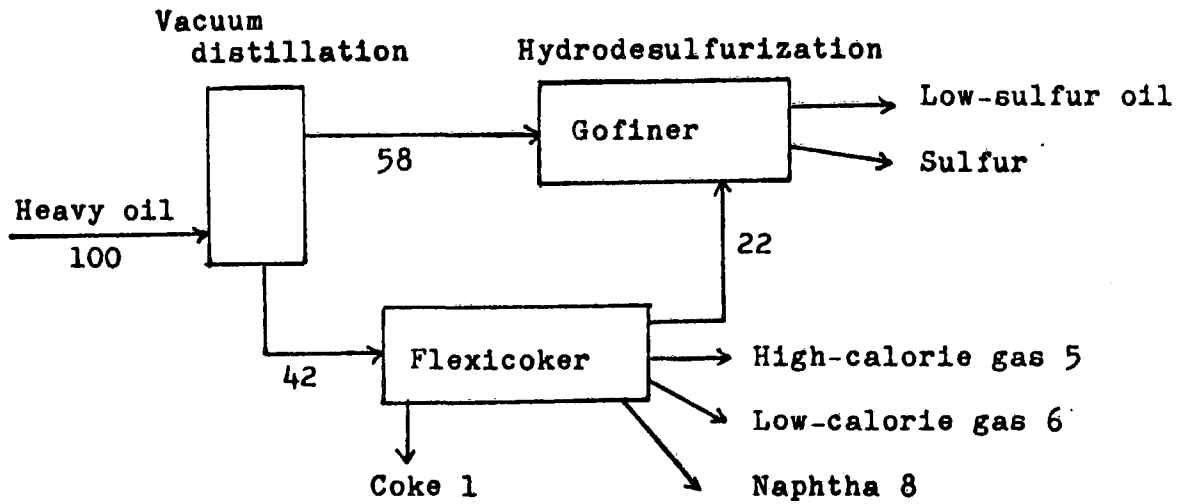


Figure 2-2. Material balance for residual oil decomposition using Flexcoking process.

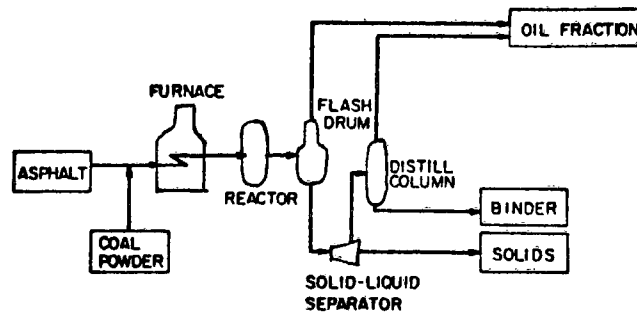


Figure 2-3. Flowsheet of Cherry Process.

then led into a flash drum, where lighter fractions are separated. The bottom product of the flash drum is centrifuged to separate liquids and solids. The liquid is sent to a distillation column to produce an oil and a pitch, which is used as a binder. The yield is 30 percent naphtha, 5 percent kerosene, 11 percent heavy fuel oil, 34 percent pitch binder, and 20 percent solids. The solid product is used as a coal substitute. A portion of the solids can be recycled to the system. No coking on the reactor walls has been observed.

Coal Gasification

The Coal Research Center has continued tests on coal gasification with a 5-t/day pilot plant at Yubari, Hokkaido. Coal is gasified in a fluidized bed under 10 atmospheres pressure at 800 to 900°C with air and steam to produce a low-calorie gas (1,200 to 1,500 cal/m³). A unit to desulfurize the gas by the Benfield process (a U.S. process) is under construction and is to start operation by the end of 1976. A plan to construct a 40-t/day test plant has been postponed.

Electric Power Development Co., which 2 years ago planned to construct a pilot plant for coal gasification, has postponed the plan and has joined the Coal Research Center in the development project.

3. GENERAL ASPECTS OF FLUE GAS DESULFURIZATION (FGD)

TRENDS

Japan has made remarkable progress in flue gas desulfurization since 1972. About 100 systems, including several large ones with a capacity of 250 to 500 MW, went into operation in 1975. The total FGD capacity, which was about 40 million Nm^3/hr (13,000 MW equivalent) at the end of 1974, will exceed 80 million Nm^3/hr by the end of 1976. The rapid growth is due to the economic advantage of FGD over the use of low-sulfur fuels and to the reliability of process and operation. The growth rate may decline after 1977, however, for the following reasons:

- (1) Ambient SO_2 concentrations in large cities and industrial districts have decreased to a range of 0.02 to 0.03 ppm, almost meeting the ambient standard.
- (2) The recent economic depression has prevented industry from building new plants.
- (3) Overproduction of FGD by-products and a decrease in the price difference between high-sulfur and low-sulfur oils (Table 1-6) have started an industry trend toward the use of low-sulfur fuels, because Japan has limited landspace available for discarding of useless by-products.
- (4) The stringent NO_x regulation is forcing industry toward flue gas desulfurization. Several processes for simultaneous removal of NO_x and SO_2 have been developed, and industry is waiting for the new technology.

MAJOR PROCESSES AND SYSTEMS

Table 3-1 lists major constructors of FGD units and numbers and capacities of the units to be operational by the end of 1977. The total number of the units will exceed 500 and the total capacity will reach 85,000,000 Nm³/hr, which is equivalent to 28,000 MW. About 80 percent of the units are completed (as of April 1976). About half of the total capacity is for utility boilers (mostly oil-fired) and the rest for industrial boilers, iron-ore sintering machines, nonferrous metals plants, sulfuric acid plants, and similar operations.

About half of the total capacity uses the wet lime/limestone process to by-produce gypsum; 16 percent use the indirect lime/limestone process (double alkali type); 13 percent use regenerable processes to by-produce sulfuric acid, ammonium sulfate and elemental sulfur; and 24 percent use sodium scrubbing to by-produce sodium sulfite or sulfate. The average system capacities are 427,000 Nm³/hr for wet lime/limestone, 291,000 Nm³/hr for indirect lime/limestone, 378,000 Nm³/hr for the regenerable processes, 59,600 Nm³/hr for the sodium scrubbing processes. About 80 percent of the sodium scrubbing units by-produce sodium sulfite for paper mills, and the rest oxidize the sulfite by air bubbling to by-produce sulfate, which is either used in the glass industry or purged in wastewater.

A recent survey by the Heavy Industry Newspaper Co. has revealed that in addition to the 335 sodium scrubbing units listed in Table 3-1, there are nearly 500 small commercial sodium scrubbing systems with an average capacity of about 20,000 Nm³/hr. The sodium scrubbing process is the least costly, and its operation is relatively easy.

WET LIME/LIMESTONE PROCESS

Outline

Wet lime/limestone process systems with a capacity larger than 60,000 Nm³/hr (20 MW) are listed in Tables 3-2, 3-3, and 3-4. The first commercial system using a wet lime-gypsum process was constructed in 1964 by Mitsubishi Heavy Industries (MHI), licensed by Japan Engineering Consulting Co. to treat 62,500 Nm³/hr of tail gas from a sulfuric acid plant, containing 2,200 ppm SO₂. Several problems, including scaling and corrosion, were encountered at the beginning of the operation. The problems have been gradually solved, and many FGD systems using wet lime/limestone processes have been constructed by MHI and other companies.

The MHI (or Mitsubishi-JECCO) process has been used most widely for oil-fired boilers, iron-ore sintering plants, etc., while the Chemico-Mitsui, Mitsui-Chemico, and Chemico-IHI processes have been applied to coal-fired boilers. Six other processes also have been used, mainly

Table 3-1. NUMBERS AND CAPACITIES^a OF FGD SYSTEMS EXPECTED TO BE
OPERATIONAL BY END OF 1977

Plant constructor	Wet lime limestone		Indirect lime limestone		Regenerable		Sodium scrubbing		Total	
Mitsubishi Heavy Industries (MHI)	33	(18,270)					3	(292)	36	(18,562)
Ishikawajima H.I. (IHI)	17	(4,445)					79	(4,351)	96	(8,796)
Hitachi - Babcock	13	(6,940)			2	(590)	15	(603)	30	(8,133)
Mitsubishi Kakoki (MKK)-Wellman	2	(256)			13	(6,478)	41	(913)	56	(7,643)
Kawasaki Heavy Industries	4	(756)	6	(5,450)			7	(256)	17	(6,380)
Tsukishima Kikai (TSK)	1	(80)	4	(398)	1	(88)	40	(4,042)	45	(4,608)
Chiyoda Chemical Eng. and Const.			14	(4,459)					14	(4,459)
Oji Koei							57	(4,280)	57	(4,280)
Fuji Kasui Engineering	7	(3,954)					6	(270)	13	(4,224)
Kurabo Engineering			5	(413)	1	(18)	106	(3,751)	112	(4,182)
Mitsui Miike-Chemico	4	(2,744)			1	(500)			5	(3,244)
Ebara Manufacturing			11	(1,914)			10	(1,167)	21	(3,081)
Nippon Kokan (NKK)	3	(245)	1	(150)	2	(1,990)	6	(62)	12	(2,447)
Kureha Chemical							8	(1,431)	8	(1,431)
Showa Denko							5	(1,372)	5	(1,372)
Gadeliuss							8	(1,291)	8	(1,291)
Sumitomo (SCEC)-Wellman					6	(1,288)			6	(1,288)
Mitsui Metal Engineering	4	(1,006)			2	(130)			6	(1,136)
Kobe Steel	5	(1,125)							5	(1,125)
Japan Gasoline	1	(330)			1	(125)			2	(455)
Dowa Engineering			4	(423)					4	(423)
Niigata Iron Works			1	(185)					1	(185)
Mitsui Shipbuilding							1	(160)	1	(160)
Sumitomo Heavy Industries					1	(150)			1	(150)
Total	94	(40,181)	46	(13,392)	30	(11,357)	335	(19,961)	505	(84,891)

^a Number of units followed by total capacity in parentheses; capacities are in thousands Nm³/Hr.

Table 3-2. WET LIME/LIMESTONE PROCESS UNITS BY MHI PROCESS

(larger than 60,000 Nm³/hr)

User	Plant site	Capacity, 1,000 Nm ³ /hr ^a	Source of gas	SO ₂ , ppm		Absorbent	Year of completion
				Inlet	Outlet		
Nippon Kokan	Koyasu	62.5	H ₂ SO ₄ plant	2,200	200	Ca(OH) ₂	1964
Kansai Electric	Amagasaki	100	Utility boiler ^b	700	70	Ca(OH) ₂	1972
Onahama Refining	Onahama	92	Copper smelter	20,000	100	Ca(OH) ₂	1972
Kawasaki Steel	Chiba	120	Sintering plant	600	60	Ca(OH) ₂	1973
Kansai Electric	Kainan	400	Utility boiler	550	60	Ca(OH) ₂	1974
Tokyo Electric	Yokosuka	400	Utility boiler	250	40	CaCO ₃	1974
Tohoku Electric	Hachinohe	380	Utility boiler	850	85	Ca(OH) ₂	1974
Kyushu Electric	Karita	550	Utility boiler	800	75	Ca(OH) ₂	1974
Kawasaki Steel	Mizushima	750	Sintering plant	830	40	Ca(OH) ₂	1974
Kansai Electric	Amagasaki	375	Utility boiler	500	50	Ca(OH) ₂	1975
Niigata Power	Niigata	530	Utility boiler	700	70	Ca(OH) ₂	1975
Kawasaki Steel	Mizushima	900	Sintering plant	500	40	Ca(OH) ₂	1975
Kawasaki Steel	Chiba	320	Sintering plant	800	60	Ca(OH) ₂	1975
Teijin	Ehime	270	Industrial boiler ^b	1,700	60	Ca(OH) ₂	1975
Mizushima Power	Mizushima	611	Utility boiler	1,050	40	Ca(OH) ₂	1975
Tohoku Electric	Niigata	420	Utility boiler	550	55	CaCO ₃	1976
Chubu Electric	Owase	1,200	Utility boiler	1,500	35	Ca(OH) ₂	1976 (March)
Chubu Electric	Owase	1,200	Utility boiler	1,500	35	Ca(OH) ₂	1976 (June)
Kawasaki Steel	Mizushima	750	Sintering plant	550	40	Ca(OH) ₂	1976
Toyobo	Iwakuni	200	Industrial boiler	1,400	50	Ca(OH) ₂	1976
Kashima Power	Kashima	431	Utility boiler	1,000	100	CaCO ₃	1976
Kyushu Electric	Karatsu	730	Utility boiler	550	70	CaCO ₃	1976
Kyushu Electric	Karatsu	570	Utility boiler	550	70	CaCO ₃	1976
Kyushu Electric	Ainoura	730	Utility boiler	880	110	CaCO ₃	1976
Kyushu Electric	Ainoura	730	Utility boiler	880	110	CaCO ₃	1976
Sakata Power	Sakata	1,100	Utility boiler	950	50	CaCO ₃	1976
Sakata Power	Sakata	1,100	Utility boiler	950	50	CaCO ₃	1976
Chugoku Electric	Shimonoseki	1,200	Utility boiler	1,600	50	CaCO ₃	1976
Confidential		475	Utility boiler	500	65	Ca(OH) ₂	1976
Confidential		1,200	Utility boiler	550	65	Ca(OH) ₂	1976
Confidential		530	Utility boiler			CaCO ₃	1976

^a 1,000 Nm³/hr = 590 scfm = 320 kW.^b All boilers are oil-fired.

Table 3-3. WET LIME/LIMESTONE PROCESS UNITS USING SCRUBBERS DEVELOPED

IN THE UNITED STATES

(larger than 60,000 Nm³/hr)

User	Plant site	Capacity, 1,000 Nm ³ /hr ^a	Source of gas	SO ₂ , ppm		Absorbent	Year of completion
				Inlet	Outlet		
Babcock-Hitachi Process							
Chugoku Electric	Mizushima	310	Utility boiler	1,500	60	CaCO ₃	1974
Asahi Chemical	Mizushima	481	Industrial boiler			CaCO ₃	1975
Kansai Electric	Osaka	500	Utility boiler			CaCO ₃	1975
Chugoku Electric	Tamashima	1,460	Utility boiler			CaCO ₃	1975
Kansai Electric	Osaka	500	Utility boiler			CaCO ₃	1975
Chugoku Electric	Tamashima	1,000	Utility boiler			CaCO ₃	1976
Showa Power	Ichihara	249	Utility boiler			CaCO ₃	1976
Showa Power	Ichihara	480	Utility boiler			CaCO ₃	1976
Maruzumi Paper	Kawanoe	342	Industrial boiler			CaCO ₃	1976
Confidential		500	Industrial boiler			CaCO ₃	1976
Electric Power Dev.	Takehara	852	Utility boiler ^b			CaCO ₃	1977
Ishikawajima Harima (IHI) - TCA Process							
Chichibu Cement	Kumagaya	104	Diesel engine	700	50	CaO	1972
Onahama Smelter	Onahama	120	Converter			CaO	1972
Furukawa Mining	Ashio	60	H ₂ SO ₄ plant			CaO	1972
Chichibu Cement	Kumagaya	106	Diesel engine			CaO	1973
Hibi Smelter	Hibi	300	Smelter			CaO	1974
Tokuyama Soda	Tokuyama	550 x 2	Industrial boiler				1975
Sumitomo Power	Niihama	450	Utility boiler			CaCO ₃	1975
Mitsui Alumina	Wakamatsu	300	Boiler, kiln			CaCO ₃	1975
Chemico - Mitsui and Mitsui - Chemical Processes							
Mitsui Aluminum	Omuta	512	Industrial boiler ^b	2,000	200	Ca(OH) ₂ ^c	1972
Mitsui Aluminum	Omuta	552	Industrial boiler	1,500	150	CaCO ₃	1975
Electric Power Dev.	Takasago	840	Utility boiler ^b	1,500	150	CaCO ₃	1975
Electric Power Dev.	Takasago	840	Utility boiler ^b	1,500	150	CaCO ₃	1976
Ishikawajima Harima (IHI) - Chemico Process							
Electric Power Dev.	Isogo	900 x 2	Utility boiler ^b	500	70	CaCO ₃	1976

^a 1,000 Nm³/hr = 590 scfm = 320 kW.^b Coal-fired boilers. Other boilers are oil-fired.^c Carbide sludge to by-produce throwaway calcium sulfite. Other plants by-produce gypsum.

Table 3-4. WET LIME/LIMESTONE PROCESS UNITS USING OTHER PROCESSES

(larger than 60,000 Nm³/hr)

User	Plant site	Capacity 1,000 Nm ³ /hr ^a	Source of gas	SO ₂ , ppm		Absorbent	Year of completion
				Inlet	Outlet		
Fujikasui-Sumitomo process (Moretana scrubber)							
Ide Paper	Fuji	60	Industrial boiler	1,340	20	CaCO ₃	1974
Sanyo Kokusaku Pulp	Onomichi	140	Recovery boiler	4,000	40	CaCO ₃	1975
Sumitomo Metal	Wakayama	370	Sintering machine	650	20	CaCO ₃	1975
Sumitomo Metal	Kokura	92	Heating furnace	820	40	CaCO ₃	1975
Sumitomo Metal	Kashima	880	Sintering machine	650	30	CaCO ₃	1975
Sumitomo Kainan Kokan	Wakayama	182	Heating furnace	680	35	CaCO ₃	1975
Sumitomo Metal	Kashima	1,000 x 2	Sintering furnace	650	30	CaCO ₃	1976
Sumitomo Metal	Kokura	750	Sintering furnace	650	20	CaCO ₃	1976
Nippon Kokan process (Spray tower absorber)							
Nippon Sheet Glass	Yokkaichi	120	Glass furnace	1,000	100	Ca(OH) ₂	1974
Nippon Sheet Glass	Maizuru	107	Glass furnace	1,550	200	CaCO ₃	1976
Nippon Kokan	Fukuyama		Incinerator	28,000	200	CaO	1976
Kobe Steel process (Cal process)							
Kobe Steel	Amagasaki	175 x 2	Sintering machine	500	30	Ca(OH) ₂	1976
Kobe Steel	Kobe	350	Sintering machine	500	30	Ca(OH) ₂	1976
Nakayama Steel	Osaka	375	Sintering machine	500	30	Ca(OH) ₂	1976
Chubu - MKK process (CM process)							
Ishihara Chemical	Yokkaichi	250	Industrial boiler	1,600	150	CaCO ₃	1974
Mitsubishi Gas Chem.	Yokkaichi	60	Industrial boiler	1,300	100	Ca(OH) ₂	1974
Kawasaki Heavy Industry process							
Jufo Paper	Akita	84	Recovery boiler			CaCO ₃	1973
Jufo Paper	Akita	90	Recovery boiler			CaCO ₃	1975
Unitika	Okazaki	200	Industrial boiler			CaO(MgO)	1975
Nippon Exlan	Saidaiji	300	Industrial boiler			CaCO ₃ (MgO)	1975
Nippon Steel process (SSD process)							
Nippon Steel	Tobata	200	Sintering machine			Slag	1974
Nippon Steel	Wakamatsu	1,000	Sintering machine			Slag	1976

^a 1,000 Nm³/hr = 590 scfm = 320 kW.

for flue gas from oil-firing boilers, and three others for sintering plants.

Many of the plants constructed before 1974 use lime at a stoichiometry of 0.95 to 1.0 to remove 93 to 98 percent of the SO_2 . Most of the plants constructed later use limestone ground to pass about 325-mesh screen at a stoichiometry of 1.0 to 1.05 to remove 90 to 96 percent of the SO_2 . Virtually all of the plants by-produce salable gypsum.

A schematic flowsheet common to most processes, except the Chemico-Mitsui and Mitsui-Chemico processes which use no cooler (prescrubber), is shown in Figure 3-1. Flue gas passes through an electrostatic precipitator, a cooler, a scrubber, a mist eliminator, and a reheater, before being emitted from a stack. Types of scrubbers and examples of operation parameters are listed in Table 3-5. Calcium sulfite formed by the reaction of SO_2 with lime or limestone slurry is oxidized by air bubbles into gypsum, which is then centrifuged.

The Omuta plant of Mitsui Aluminum is the only plant that produces a throwaway calcium sulfite sludge on a large scale. The plant uses a two-stage Chemico venturi scrubber and a carbide sludge as the absorbent. The plant has been operated in an unsaturated mode; gypsum is not formed, although about 10 percent of the calcium sulfite is oxidized in the scrubber. Operation has been trouble-free since

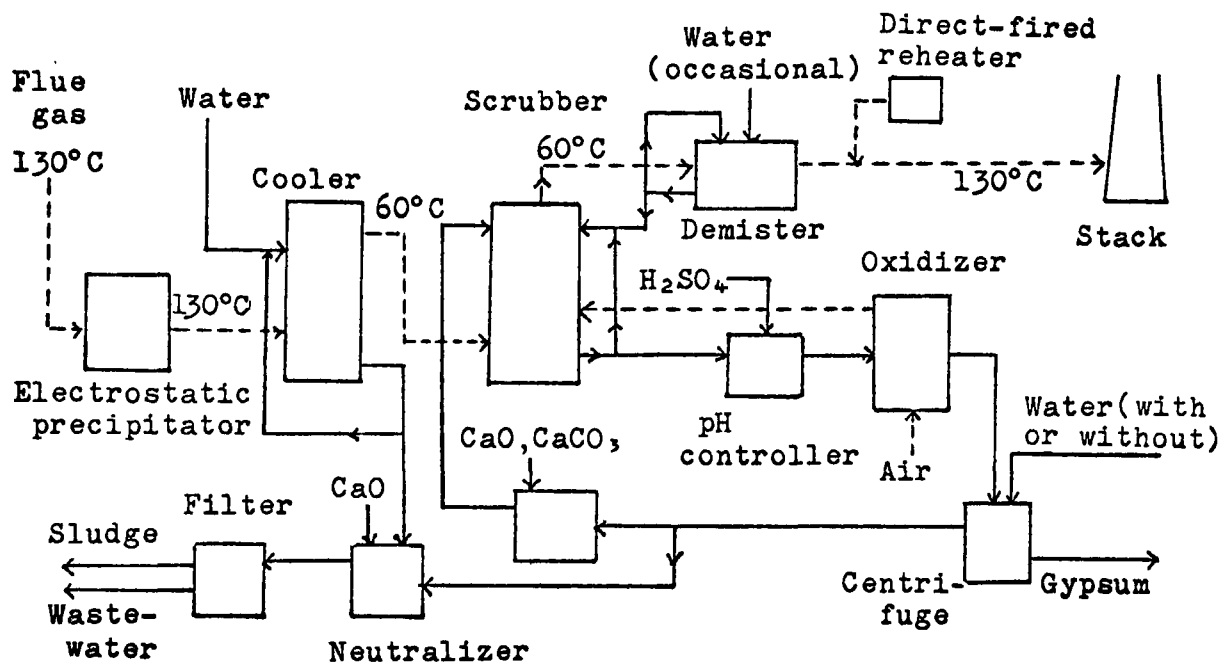


Figure 3-1. Schematic flowsheet of wet lime/limestone gypsum process.

Table 3-5. EXAMPLE OF OPERATION PARAMETERS OF FGD PLANTS
BY-PRODUCING GYPSUM AND CALCIUM SULFITE

Process developer	Absorbent, precipitant (stoichiometry)	Capacity, 1,000 Nm ³ /hr	Type of absorber	Slurry or solution,		L/G, liter/ Nm ³	Space velocity, m/sec	Pressure drop,* mm H ₂ O	SO ₂ , ppm		Moisture, % of gypsum
				pH	conc. %				In	Out	
Wet lime-limestone process											
MHI (Mitsubishi-JECCO)	CaO 1	1,200 ^d	GP ^a	6.6	10	10	3.5	200	1,600	40	8-10
MHI (Mitsubishi-JECCO)	CaCO ₃ 1	750	GP ^a	6	10	10	3.5	150	1,000	80	8-10
Chemico-Mitsui	Ca(OH) ₂ 1-1.05	385	venturi	7	3-5	10-15		400	2,900	150	CaSO ₃
Mitsui-Chemico	CaCO ₃ 1.0-1.05	840	venturi	6	5	10-15		200	1,500	130	10-15
Babcock-Hitachi	CaCO ₃ 1.05-1.15	1,460 ^c	PP ^b	6.1	20	10	3.2	850	1,500	60	8-10
Fuji Kasui-Sumitomo	CaCO ₃ 1.05-1.15	800	PP ^b	6	5-6	5	4.5	200	500	20	
Chubu-MKK	CaCO ₃	250	screen	6	10	10	4	120	1,500	150	10-12
Ishikawajima-TCA	CaO	100	TCA		2	7	3		700	50	10-15
Kobe Steel	CaO ^f 1.05	350	spray	6-8	30	3	3	190	300	20	10
Indirect lime-limestone process											
Kureha-Kawasaki	Na ₂ SO ₃ , CaCO ₃	1,260 ^d	GP ^a	6.2	20	10	2.5	150	1,070	5	6-8
Showa Denko	Na ₂ SO ₃ , CaCO ₃	500 ^c	cone	6.8	25	2		250	1,400	40	8-10
Nippon Kokan	(NH ₄) ₂ SO ₃ , CaO	150	screen	6	30	2	3	250	700	30	8-10
Chiyoda	dil. H ₂ SO ₄ , CaCO ₃	1,050	Tellerette	1	2-4	55-60	1		1,600	60	7-9
Kurabo	(NH ₄) ₂ SO ₄ , CaO	100	Tellerette	4	10	6-10	2	100	1,500	80	8-10
Dowa	Al ₂ (SO ₄) ₃ , CaCO ₃	150	Tellerette	4	10	3 ^e	1.5	100	600	20	10-12
Kureha	CH ₃ COONa, CaCO ₃	5	PP ^b	5.5	20	7-8 ^g	2-2.5	280	1,400	1-3	6-7

^a Grid packed.

^b Perforated plate.

^c Four scrubbers in parallel.

^d Two scrubbers in parallel.

^e For tail gas at 25°C. L/G 6-10 for flue gas.

^f In CaCl₂ solution.

^g Including limestone scrubbing.

* Including cooler, absorber, and mist eliminator.

start-up. For the recent FGD installations, Mitsui Aluminum selected a modified Mitsui-Chemico process to by-produce gypsum using limestone--because carbide sludge is getting short and the calcium sulfite sludge pond is becoming full.

Scale Prevention

In many of the plants by-producing gypsum, more than 20 percent oxidation of calcium sulfite occurs, resulting in formation of gypsum in the scrubber. Gypsum crystals are usually recycled to the scrubber as seed crystals to eliminate the formation of gypsum on the surface of structural materials, which tends to cause scaling. Maintaining a smooth surface on the materials and a continuous flow of slurry in all parts of the scrubbers and pipings also aids in scale prevention.

Mist eliminators are most susceptible to scaling. Washing the eliminator with fresh water dissolves the scale but increases the volume of wastewater. A recent trend is to wash eliminators with circulating liquor and only occasionally with fresh water when some scale forms.

Scaling of the mist eliminator apparently is much more serious in the United States than in Japan. In the U.S.A. a large excess of limestone is often used to remove 80 to 90 percent of the SO_2 . A mist containing a considerable amount of unreacted limestone adheres to the eliminator; passage of the gas containing 200 to 400 ppm of SO_2 and 2 to 4 percent

O₂ results in formation of gypsum on the mist eliminator. In Japan the excess of limestone is not so great, and the SO₂ concentration of the gas passing through the eliminator usually ranges from 20 to 150 ppm. Good reaction of limestone in the scrubber is attained by fine grinding and use of a higher L/G ratio.

INDIRECT LIME/LIMESTONE PROCESS

Outline

Processes developed to ensure scale-free stable operation, include double-alkali processes that use alkaline solutions for absorption and lime or limestone for precipitation, and also similar processes using acidic solutions for absorption. All of these processes are classified under the category of "indirect lime/limestone process." Operating parameters are shown in Table 3-5; major installations are listed in Table 3-6.

The scrubbing liquors are as follows: sodium sulfite for Kureha-Kawasaki, Showa Denko, and Tsukishima; ammonium sulfite for Nippon Kokan; dilute sulfuric acid containing ferric sulfate for Chiyoda; aluminum sulfate for Dowa; acidic ammonium sulfate for Kurabo; and sodium acetate for Kureha.

The pH values of the liquors are 6 to 7 for ammonium and sodium sulfites, 5 for sodium acetate, 3 to 4 for

Table 3-6. INDIRECT LIME/LIMESTONE PROCESS INSTALLATIONS

Process developer	Absorbent, precipitant	User	Plant site	Capacity, 1,000 Nm ³ /hr	Source of gas	Inlet SO ₂ , ppm	Year of completion
Chiyo-da	H ₂ SO ₄ , CaCO ₃	Nippon Mining	Mizushima	34	Claus furnace	9,300	1972
		Fuji Kōsan	Kainan	160	Industrial boiler	770	1972
		Mitsubishi Rayon	Otake	90	Industrial boiler	1,600	1972
		Daicel	Aboshi	99	Industrial boiler	1,400	1973
		Tochoku Oil	Sendai	14	Claus furnace	7,100	1973
		Mitsubishi Chem.	Yokkaichi	420	Industrial boiler	1,200	1974
		Aragasaki Coke	Kakogawa	36	Incinerator	11,300	1974
		Hokuriku Electric		750	Utility boiler	600	1974
		Mitsubishi Pet.	Yokkaichi	150	Industrial boiler	1,300	1974
		Mitsubishi Pet.	Yokkaichi	750	Industrial boiler	1,200	1974
		Gulf Power	Florida	85	Utility boiler	1,700	1975
		Denki Kagaku	Chiba	122	Industrial boiler	1,400	1975
		Hokuriku Electric	Fukui	1,050	Utility boiler	1,500	1975
		Toyama Power	Toyama	750	Utility boiler	1,500	1975
Showa Denko	Na ₂ SO ₃ , CaCO ₃	Showa Denko	Chiba	500	Industrial boiler	1,500	1973
		Kanagafuchi Chem.	Takasago	300	Industrial boiler	1,500	1974
		Showa Pet. Chem.	Kawasaki	200	Industrial boiler	1,400	1974
Showa Denko-Ebara	Na ₂ SO ₃ , CaCO ₃	Nippon Mining	Saganoseki	120	H ₂ SO ₄ plant		1973
		Yokohama Rubber	Hiratsuka	105	Industrial boiler		1974
		Nisshin Oil	Isogo	100	Industrial boiler		1974
		Poly Plastics	Fuji	212	Industrial boiler		1974
		Ajinomoto	Yokkaichi	82	Industrial boiler		1974
		Kyowa Pet. Chem.	Yokkaichi	150	Industrial boiler		1974
		Japan Food	Yokkaichi	100	Industrial boiler		1975
		Yokohama Rubber	Mie	100	Industrial boiler		1975
		Asia Oil	Yokohama	243	Industrial boiler		1975
Kureha-Kawasaki	Na ₂ SO ₃ , CaCO ₃	Tohoku Electric	Shinsendai	420	Utility boiler	420	1974
		Shikoku Electric	Sakaide	1,260	Utility boiler	1,500	1975
		Shikoku Electric	Anan	1,260	Utility boiler	1,500	1975
		Kyushu Electric	Buzen	730	Utility boiler	1,500	1977
		Tohoku Electric	Akita	1,050	Utility boiler	1,500	1977
Nippon Kokan	(NH ₄) ₂ SO ₃ , CaCO ₃	Nippon Kokan	Keihin	150	Sintering plant	400	1972
Tsukishima	Na ₂ SO ₃ , CaO	Kinura Utility	Nagoya	185	Industrial boiler		1974
		Daishowa Paper	Fuji	264	Industrial boiler		1975
Kurabo Eng.	(NH ₄) ₂ SO ₄ , CaO	Kuraray	Tamashima	100	Industrial boiler	1,500	1974
		Daicel	Aboshi	153	Industrial boiler	1,300	1975
		Bridgestone Tire	Tosu	60	Industrial boiler		1975
		Bridgestone Tire	Tochigi	80	Industrial boiler		1975
		Gujō Paper	Ishinomaki	200	Industrial boiler	1,200	1976
Dowa Mining	Al ₂ (SO ₄) ₃ , CaCO ₃	Tanaka Mining	Mobara	3,500	Kiln	7,500	1972
		Dowa Mining	Okayama	300	H ₂ SO ₄ plant	650	1974
		Naikai Engyo	Okayama	70	Utility boiler	1,500	1976
		Yahagi Iron	Nagoya	50	Sintering plant		1976
Kureha Chemical	CH ₃ COONa, CaCO ₃	Kureha Chem.	Nishiki	5	Utility boiler	1,500	1975

ammonium and aluminum sulfates, and 1 for sulfuric acid. The L/G ratios are 1/2 (7 to 14 gal/1,000 scf) for the solutions of pH 6 to 7, 3/10 for the solutions of pH 3 to 5, and 30/50 for the acid at pH 1. As acidity of the solution increases, the SO_2 absorption capacity and scaling are reduced and the reaction with limestone becomes easier. Limestone can be reacted with a sodium bisulfite solution, as in the Showa Denko and Kureha-Kawasaki processes, but the reaction occurs slowly and requires large reaction vessels. Lime is used for the Tsukishima, NKK, and Kurabo processes.

For the Chiyoda, Dowa, Kureha, and Kurabo processes, the liquors that absorb SO_2 are contacted with air to oxidize SO_3^{--} into SO_4^{--} . Limestone or lime is then added to precipitate gypsum. For other processes, limestone or lime is added first to precipitate calcium sulfite, which is then oxidized into gypsum. In the double alkali type process, gypsum usually grows in larger crystals than in the wet lime-limestone process. Moisture content of the by-product gypsum after centrifugalization ranges from 6 to 12 percent as compared with 8 to 15 percent for the wet lime-limestone process.

The liquor from the gypsum centrifuge is returned mainly to the scrubber system. Softening of the liquor, which is usually needed to prevent scaling, is not required when pH of the acidic solution is below 5.

At most plants, a small portion of the liquor is purged to maintain the concentrations of chloride, magnesium, and other impurities under a certain level.

Sodium scrubbing provides high SO₂ recovery but involves sodium sulfate formation because of the oxidation of sodium sulfite. The sulfate must be decomposed because it does not absorb SO₂.

Compared with sodium scrubbing, ammonia scrubbing is less expensive because ammonia is cheaper than sodium hydroxide and ammonium sulfate is readily decomposed by lime. Plume formation is the major problem in ammonia scrubbing.

Costs and Trends

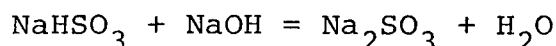
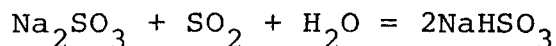
Generally speaking, scale-free, stable operation is more easily attained by indirect lime/limestone processes than by the wet lime/limestone process, but the capital and operating costs are 5 to 10 percent higher (Table 3-9).

As successful operation of wet lime/limestone process units continues, the indirect processes are losing their advantage. Sodium-based indirect processes may be used, however, when more than 98 percent SO₂ removal is required, even though the processes are fairly expensive. Other indirect processes may be used at plants where the operation is not readily controlled to prevent scaling with a wet lime/limestone process.

OTHER PROCESSES (RECOVERY PROCESSES)

Sodium Sulfite By-production

There are 335 sodium scrubbing systems with an average capacity of treating 60,000 Nm³/hr of flue gas (Table 3-1) and about 500 smaller units with an average capacity of 20,000 Nm³/hr, by-producing mainly sodium sulfite, with some sulfate. As sodium hydroxide absorbs not only SO₂ but also CO₂, sodium sulfite is generally used to absorb SO₂ only. The product sodium bisulfite is neutralized with sodium hydroxide to produce the sulfite solution, half of which is returned to the scrubber and the rest to a concentration step. The by-product sulfite, in the form of either a concentrated solution or crystal, is sold to paper mills.



The process is simple and the system costs are low; demand for the sulfite, however, is limited. In several units the sulfite in solution is air-oxidized into sulfate, which is used for glass production, etc. From some smaller units, the sodium sulfite solution is purged. The number of sodium sulfite scrubbing systems will not increase significantly because of the oversupply of the by-products and the increase in cost of the sodium hydroxide.

Wellman-Lord Process

Many Wellman-Lord process units have been constructed by Mitsubishi Kakoki Kaisha (MKK) and Sumitomo Chemical Engineering Co. (SCEC) (Table 3-7). The processes and the units were described in an earlier report.² Unit operation has been smooth. The major problem with the process has been treatment of wastewater to decompose reducing compounds. In addition to sodium thiosulfate $\text{Na}_2\text{S}_2\text{O}_3$, dithionate $\text{Na}_2\text{S}_4\text{O}_6$ is formed during heating of the sodium bisulfite solution, and the dithionate is not easily decomposed. MKK has succeeded in decomposing it by ozone oxidation at a pH below 1.5. The chemical oxygen demand (COD) is reduced enough to meet the regulation, but the treatment adds cost to the process. Most of the Wellman-Lord process units by-produce sulfuric acid; three units of oil companies by-produce elemental sulfur by feeding the recovered SO_2 into a Claus furnace.

Magnesium and Zinc Scrubbing

There are two magnesium scrubbing units in operation (Table 3-7). Onahama plant, Onahama Smelting Co., sends the recovered SO_2 to a sulfuric acid plant, as reported earlier.^{2,3} Chiba plant, Idenitsu Kosan, using the Chemico-Mitsui process, started operation recently, as described in Section 6.

Table 3-7. FGD INSTALLATIONS BY-PRODUCING H_2SO_4 , S AND $(\text{NH}_4)_2\text{SO}_4$

Process developer	Absorbent	User	Plant site	1,000 Nm ³ /hr ^a	Source of gas	Inlet SO ₂ , ppm	Year of completion	By-product
Wellman-MKK	Na_2SO_3	Japan S.R.	Chiba	200	Industrial boiler	2,000	1971	H_2SO_4
		Chubu Electric	Nishinagoya	620	Utility boiler	1,600	1973	H_2SO_4
		Kashima Oil	Kashima	30	Claus furnace	11,000	1973	S
		Japan S.R.	Yokkaichi	450	Industrial boiler	1,000	1973	H_2SO_4
		Toyo Rayon	Nagoya	330	Industrial boiler	1,500	1974	H_2SO_4
		J.N. Railways	Kawasaki	700	Industrial boiler	1,500	1975	H_2SO_4
		Mitsubishi Chem.	Mizushima	628	Industrial boiler	1,500	1975	H_2SO_4
		Kuraray	Okayama	410	Industrial boiler	1,500	1975	H_2SO_4
		Shindaikyowa Oil	Yokkaichi	400	Industrial boiler	1,500	1975	H_2SO_4
		Mitsubishi Chem.	Mizushima	628	Industrial boiler	1,500	1976	H_2SO_4
		Mitsubishi Chem.	Kurosaki	530	Industrial boiler	1,500	1976	H_2SO_4
		Tohoku Electric	Niigata	380	Utility boiler	1,000	1977	H_2SO_4
		Toa Nenryo	Kawasaki	67	Claus furnace	6,500	1971	S
		Sumitomo Chem.	Sodegaura	360	Industrial boiler	1,550	1973	H_2SO_4
Wellman-SCEC	Na_2SO_3	Toa Nenryo	Wakayama	17	Claus furnace	19,000	1974	S
		Fuji Film	Fujinomiya	150	Industrial boiler	1,300	1974	SO_2
		Sumitomo Chem.	Niihama	155	Industrial boiler	1,600	1975	H_2SO_4
		Sumitomo Chem.	Sodegaura	540	Industrial boiler	1,600	1975	H_2SO_4
		Onahama Smelting	Onahama	84	Copper smelter	20,000	1972	H_2SO_4
		Idemitsu Kosan	Chiba	500	Claus and boiler		1975	S
Chimico-Mitsui	MgO	Mitsui Mining	Hibi	80	H_2SO_4 plant		1971	H_2SO_4
Mitsui Mining	MgO	Mitsui Mining	Kamioka	50	H_2SO_4 plant		1975	H_2SO_4
Shell	ZnO	Showa Y.S.	Yokkaichi	120	Industrial boiler	1,500	1973	S
Sumitomo H.I.	CuO	Kansai Electric	Sakai	160	Utility boiler		1971	H_2SO_4
Hitachi Ltd.	Carbon	Tokyo Electric	Kashima	420	Utility boiler		1972	H_2SO_4
		Unitika	Oji	171	Industrial boiler	1,500	1975	H_2SO_4
		Nippon Kokan	Fukuyama	760	Sintering plant	400	1976	$(\text{NH}_4)_2\text{SO}_4$
Nippon Kokan	$(\text{NH}_4)_2\text{SO}_3$	Nippon Kokan	Ogishima	1,140	Sintering plant	400	1977	$(\text{NH}_4)_2\text{SO}_4$
Kurabo Engineering	$(\text{NH}_4)_2\text{SO}_4$	Taki Chemical	Befu	15	Industrial boiler	1,500	1976	$(\text{NH}_4)_2\text{SO}_4$
MHI-IFP	$(\text{NH}_4)_2\text{SO}_3$	Maruzen Oil	Shimozu	42	Claus furnace		1974	S
TEC-IFP	$(\text{NH}_4)_2\text{SO}_3$	Fuji Oil	Chiba	6	Claus furnace		1974	S

^a 1,000 Nm³/hr = 590 scfm = 320 MW.

Operation of a magnesium scrubbing system (80,000 Nm³/hr) at Hibi Works, Mitsui Mining² was discontinued. A lime scrubbing system of larger capacity (300,000 Nm³/hr) was constructed to replace it and is now in operation.

Kamioka plant, Mitsui Mining, started operation in 1975 to recover SO₂ from tail gas (50,000 Nm³/hr) in a sulfuric acid plant by zinc oxide scrubbing. Zinc sulfite formed by the reaction is calcined to regenerate zinc oxide and SO₂; the latter is returned to the sulfuric acid plant. The process is similar to the magnesium scrubbing process. Although the calcination temperature is much lower with zinc (about 350°C) than with magnesia (about 950°C), zinc is expensive and entails some environmental hazards.

Activated Carbon Process

Tokyo Electric Power operates an activated carbon process plant at Kashima (420,000 Nm³/hr).² The SO₂ absorbed on carbon is washed with water to produce a weak sulfuric acid, which is treated with limestone to produce good-quality gypsum. The system has been operated without trouble for over 2 years. Virtually no carbon has been lost during the period.

Unitika Co. recently constructed a carbon process system to by-produce a stronger sulfuric acid (Section 6).

Operation continues at the dry carbon process plant of Kansai Electric at Sakai, using a moving bed designed by

Sumitomo Heavy Industry (formerly Sumitomo Shipbuilding).² After SO₂ absorption, the carbon is heated in a reducing gas to recover a concentrated SO₂ gas, which is used for sulfuric acid production. Consumption of the carbon is fairly high and there is no plan to construct a new system with the process. Sumitomo has been testing simultaneous removal of SO₂ and NO_x in the process using ammonia (Section 7).

Ammonia Scrubbing

Nippon Kokan has developed a process to by-produce ammonium sulfate from SO₂ in flue gas and ammonia in coke oven gas² and is constructing two commercial plants (Table 3-7). To reduce formation of plume from the scrubber, which is a common problem with ammonia scrubbing processes, Nippon Kokan will use an afterburner to raise the temperature of the treated gas at its Fukuyama plant and a wet electrostatic precipitator at its Ogishima plant.

A small commercial plant to by-produce ammonium sulfate was completed recently by Kurabo Engineering for a fertilizer producer (Table 3-7, and Section 6).

Two relatively small units, with ammonia scrubbing, thermal decomposition of ammonium sulfate, and an IFP reactor to by-produce elemental sulfur, were constructed by Mitsubishi Heavy Industries and Toyo Engineering. Both have had problems, mainly in the decomposition step.

Shell Process

The SYS system using the Shell copper oxide process has continued operation (Table 3-7). Tests have been performed for simultaneous removal of NO_x by ammonia injection (Section 7).

BY-PRODUCTS OF FGD

Desulfurization efforts in Japan are oriented toward processes that yield salable by-products (Figures 3-2 and 3-3). The reasons are that domestic supplies of sulfur and its compounds are limited in Japan, as is land for disposal of useless by-products. About 60 percent of the SO_2 is converted into salable gypsum, 20 percent into sodium sulfite, 15 percent into sulfuric acid, and the rest into waste calcium sulfite and sodium and ammonium sulfates.

The process that first became popular is sodium scrubbing to by-produce sodium sulfite for paper mills. Although this is the easiest FGD process, by-production of sodium sulfite will not increase greatly because the supply has filled the demand and because sodium hydroxide has become expensive.

The by-product gypsum can be grown into fairly large crystals useful for wallboard production and as a retarder of cement setting (Figure 3-4, Photo 3-1). Although the demand for gypsum decreased last year because of the economic depression, gypsum will continue to be the major by-product of FGD.

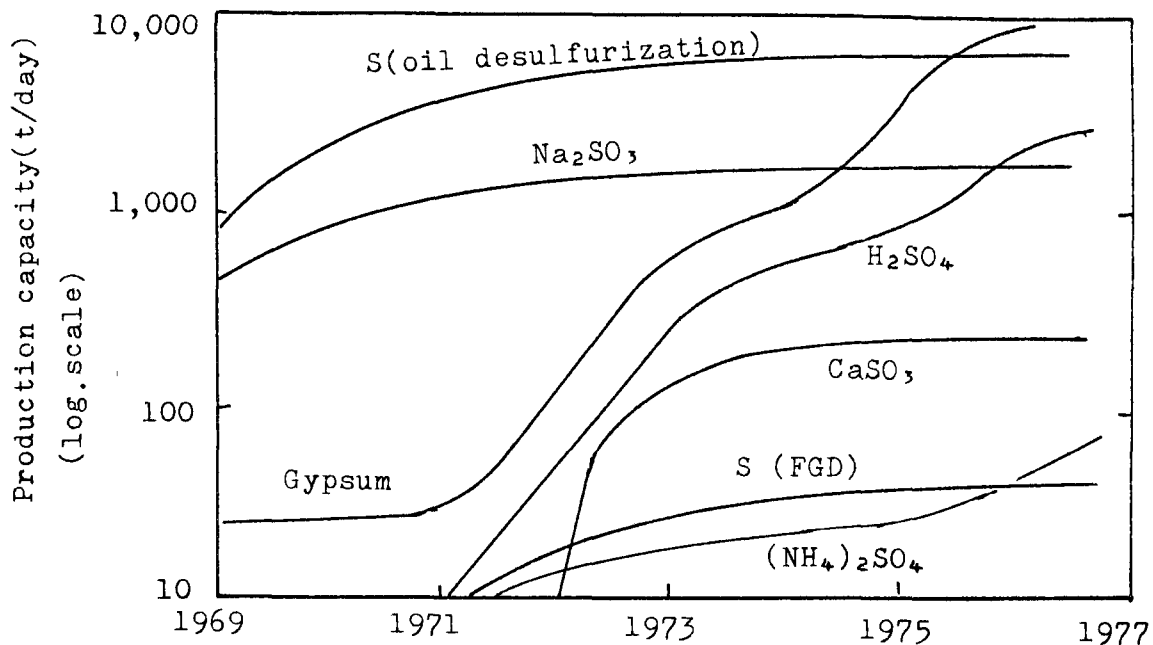


Figure 3-2. Production capacity of desulfurization.

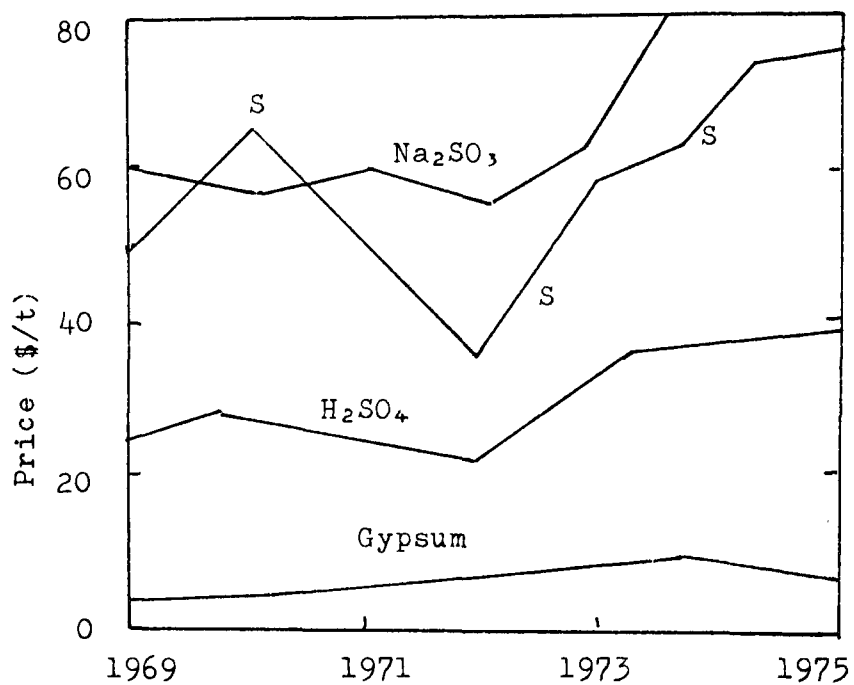
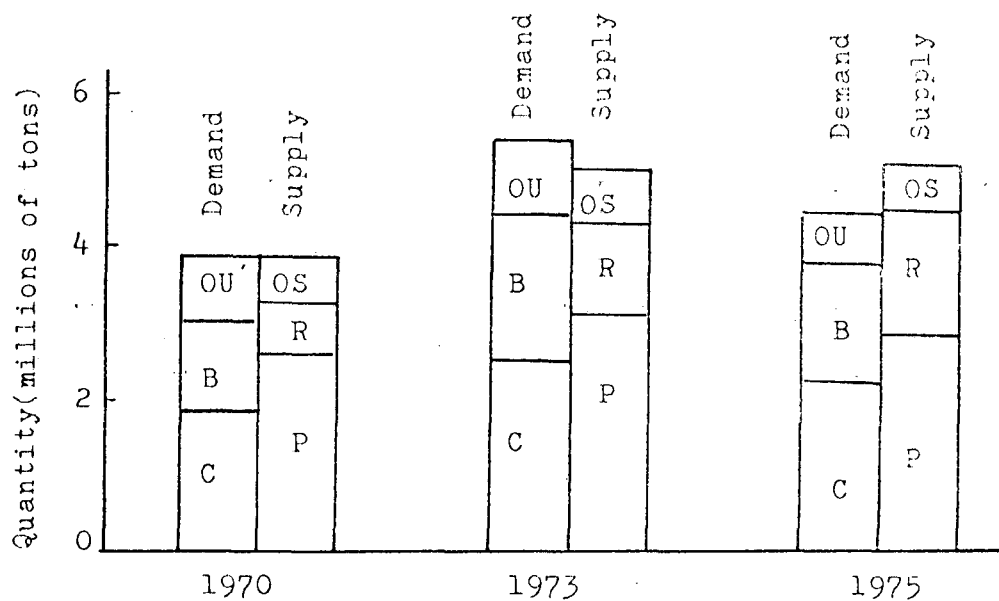


Figure 3-3. Price of by-products.



Demand: C:Cement B:Board OU:Other uses

Supply: P:Phosphogypsum R:Recovered OS:Other sources

Figure 3-4. Demand for and supply of gypsum in Japan.



Photo 3-1. Handling of by-product gypsum

(Chiba plant, Showa Denko).

Sulfuric acid has been produced by the Wellman-Lord and magnesium scrubbing processes. Elemental sulfur has been produced in relatively small units in oil refineries by the Wellman-Lord, Shell, and magnesium scrubbing processes using Claus furnaces.

Since FGD has developed rapidly and there is already a tendency toward oversupply of the by-products, it is desired to develop new applications. Efforts have been concentrated on establishing new uses of gypsum, mainly for new construction materials.

Throwaway calcium sulfite sludge has been produced at the Omuta plant, Mitsui Aluminum (385,000 Nm³/hr, Photo 3-2), where the sulfite slurry is discharged into a large pond and the supernatant is recycled to the scrubber. In two other smaller plants, the slurry is filtered and discarded.

Even for discarding into a pond, gypsum might be a better choice than calcium sulfite because it grows into much larger crystals and readily settles into a much smaller volume. The Omura plant, Mitsui Aluminum, that has been in operation since 1972 is going to change the process to by-produce gypsum because the pond is nearly full. The Wakamatsu plant, Nippon Steel, and Isogo plant, EPDC, are going to produce throwaway gypsum.

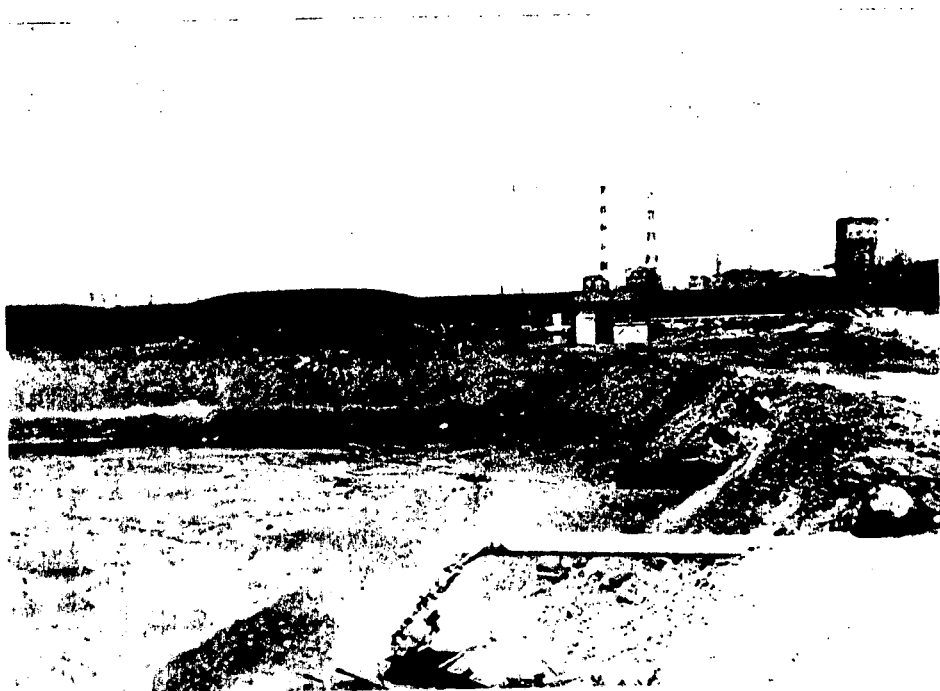


Photo 3-2. Calcium sulfite sludge disposal
(Omuta plant, Mitsui Aluminum).

WASTEWATER AND GAS REHEATING

Wastewater

Most Japanese FGD processes purge wastewater, as shown in Table 3-8 and Figure 3-5, mainly to prevent the accumulation of impurities, especially chloride, in the circulating liquor. Chloride, which is derived from fuel and process water, promotes corrosion, particularly when the liquor contains more than 1 ppm of oxygen, as shown in Figure 3-6.

Plants using the Kureha-Kawasaki process (Section 4) do not normally purge any water because the scrubber liquor is very low in oxygen. Chloride concentrations in the liquor at those plants has reached 4,000 ppm; the amount of chloride leaving the system with gypsum which contains 6 to 8 percent moisture has become equal to that going into the system. Another process free from wastewater is the Kobe Steel process (Section 5), which uses a calcium chloride solution dissolving lime as the absorbent and uses highly corrosion-resistant materials for construction of system components.

The dry processes are not free from wastewater, except for the Sumitomo Heavy Industry process, which uses carbon absorption and thermal regeneration. The Tokyo Electric - Hitachi and the Shell processes give relatively large amounts of wastewater because they use wet treatment in the regeneration steps.

Table 3-8. WASTEWATER FROM FGD SYSTEMS

Process	User	Plant site	MW	Inlet SO ₂ , ppm	Wastewater t/hr (A)	Gypsum, t/hr		Water ratio (A+C) (A+B+C)	Wastewater kg/MW hr
						Solid (B)	Moisture (C)		
Mitsubishi (MHI)	Kansai Electric	Kainan	150	270	1.5	0.9	0.1	0.64	10
	Kyushu Electric	Karita	188	600	3.7	2.2	0.2	0.64	20
	Chubu Electric	Owase ^a	750	1,480	14.0	29.0	2.9	0.37	19
Mitsui-Chemico	EPDC	Takasago ^b	250	1,500	5.0	10.0	1.1	0.38	20
Babcock-Hitachi	Chugoku Electric	Mizushima	105	400	1.5	0.9	0.1	0.64	14
	Chugoku Electric	Tamashima	500	1,500	5.0	19.5	1.9	0.26	10
Chubu-MKK	Ishihara Chemical	Yokkaichi	85	1,300	3.5	2.2	0.2	0.62	41
Showa Denko	Showa Denko	Chiba	150	1,400	3.5	5.2	0.5	0.45	23
Chiyoda	Hokuriku Electric	Toyama	250	610	15.0	7.5	0.7	0.73	60
	Hokuriku Electric	Fukui	350	1,540	24.0	14.0	1.4	0.64	68
Wellman-MKK	Chubu Electric	Nishinagoya	220	1,800	3.0				14

^a Designed value; the plant has just started operation.

^b Coal-fired boiler. All others are for oil-fired boilers.

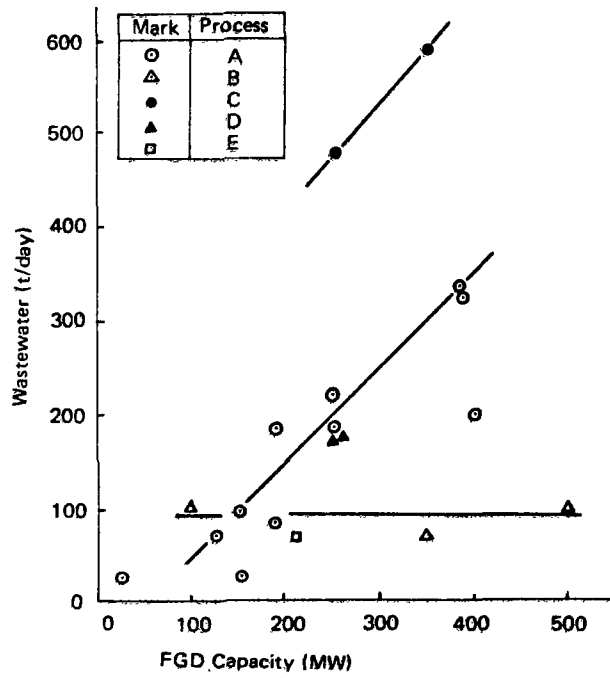


Figure 3-5. FGD capacity and amount of wastewater.⁶

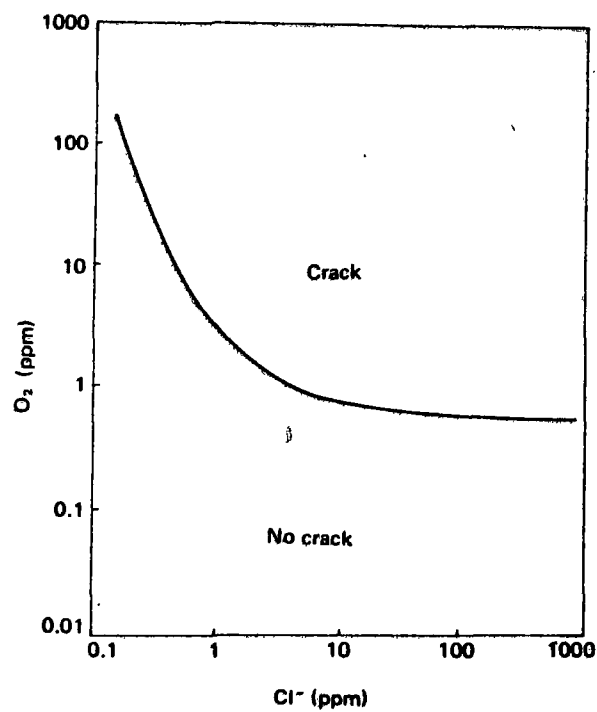


Figure 3-6. Concentration of O_2 and Cl^- in solution and stress corrosion.

Many states in the United States prohibit the discharge of wastewater but allow the discarding of calcium sulfite sludge after filtration; the sludge usually contains 50 to 60 percent water (water ratio 0.5 to 0.6). Although most Japanese processes purge some wastewater, the amount is about equal to or even less than that purged by sludge disposal. The Chiyoda process normally purges a larger amount of water to prevent corrosion because Chiyoda uses sulfuric acid saturated with oxygen as the absorbent. The volume of wastewater can be reduced by using a material with greater corrosion resistance.

Wastewater is treated to meet regulations. The treatment in most processes is simple, consisting principally of neutralization and filtration. Some processes, such as the Wellman-Lord process, require extensive treatment including ozone oxidation to decompose reducing compounds formed in the process. After being treated, the wastewater normally has a pH of 6 to 8.5, contains 5 to 20 mg/liter of SS (suspended solids), contains less than 10 mg/liter of COD (chemical oxygen demand), and does not adversely affect the environment.

Reheating

The temperature of boiler flue gas that has passed through an air preheater and electrostatic precipitator normally ranges from 130 to 150°C. Scrubbing in the wet

processes usually reduces the gas temperature to 50 to 60°C. In most systems the gas is heated to 110 to 150°C by afterburning of low-sulfur oil. Although afterburning is the easiest way to reheat stack gas, the oil requirement reaches 3 to 5 percent of that used for the boiler. Afterburning not only is somewhat costly but adds SO₂ and dust to the cleaned gas.

A few companies have made tests on reheating the gas using a gas-steam heat exchanger. Corrosion of the heat exchanger tubes is the major problem, particularly when the scrubber liquor is rich in chloride. Commercial use of such heat exchangers will start soon.

ECONOMIC ASPECTS OF FGD SYSTEMS

Among the various FGD processes, the sodium scrubbing process that by-produces sodium sulfite is the simplest and also the least expensive. The second cheapest, the throw-away wet-lime process, requires a large disposal pond. A system based on the wet lime-gypsum process costs about 25 percent more than one using the throwaway process, but does not require a pond.

Examples of plant cost in battery limits are shown in Table 3-9. The cost rose sharply until the middle of 1975 because of inflation and the active demand for FGD units, but has since lowered considerably. Generally speaking, a

Table 3-9. PLANT COST IN BATTERY LIMITS (\$1 = ¥300)

(The cost nearly tripled in late 1973 and has decreased considerably since late 1975)

Process	Absorbent	By-product	Capacity, MW	Plant cost		Year completed
				M \$	\$/kW	
Wellman-MKK	Na_2SO_3	H_2SO_4	70	2.6	37	1971
Sumitomo S.B.	Carbon	H_2SO_4	55	2.8	51	1971
Chemico-Mitsui	$\text{Ca}(\text{OH})_2$	Sludge	128	3.3	26	1972
Hitachi-Tokyo E.P.	Carbon	Gypsum	150	5.6	39	1972
Wellman-MKK	Na_2SO_3	H_2SO_4	220	7.0	32	1973
Shell	CuO	SO_2	40	3.3	83	1973
Chubu-MKK	CaCO_3	Gypsum	89	2.6	29	1973
Chemico-Mitsui	MgO	SO_2	180	13	72	1974
Mitsui-Chemico	CaCO_3	Gypsum	250	16	64	1974
Mitsubishi (MHI)	CaO	Gypsum	188	11.5	61	1974
Kureha-Kawasaki	Na_2SO_3	Gypsum	450	32	71	1975
Chiyoda	H_2SO_4	Gypsum	350	26	74	1975
Babcock-Hitachi	CaCO_3	Gypsum	500	35	70	1975
Wellman-MKK	Na_2SO_3	H_2SO_4	160	20	125	1975

wet lime-gypsum process unit (200-300 MW) now costs \$45 to \$60/kW in battery limits; system based on the indirect lime/limestone process costs 5 to 30 percent more, and one using sulfuric acid by-production processes costs 30 to 70 percent more than the wet lime-gypsum process.

Examples of FGD costs for removal of 90 to 95 percent of the SO_2 by the wet lime-gypsum process are shown in Table 3-10. The cost is about \$14 to \$17/k1 oil or 3.0 to 3.6 mil/kWh for different sizes of unit, based on 7 years depreciation and 7,000 hours yearly operation. As the fixed cost is greater than the running cost, the FGD cost is largely influenced by years of depreciation and by operating hours.

Capital cost of the wet limestone-gypsum process may be slightly higher than those of the lime-gypsum process, but operating costs are slightly lower. Indirect lime/limestone processes usually cost 5 to 20 percent more than does the lime-gypsum process. Sulfuric acid by-producing processes cost 20 to 30 percent more.

The current per-kiloliter price of heavy oil is about \$75 for high-sulfur oil ($S=3\%$) and \$100 for low-sulfur oil ($S=0.3\%$). Therefore, FGD is more economical than the use of low-sulfur oil (Table 1-6).

Table 3-10. EXAMPLES OF FGD COST WITH WET

LIME-GYPSUM PROCESS

(7 years depreciation, 7,000 hours full-load operation per year. Oil consumption: 150,000 kl/100 MW/year, S: 2.8%, 90% removal. Reheating to 110°C).

	100 MW	500 MW
Investment cost, \$1,000	6,600	27,000
Fixed cost, \$1,000/year		
Depreciation	940	3,860
Interest, Insurance	470	1,930
Total	1,410	5,790
Running cost, \$1,000/year		
Lime (at \$30/t)	230	1,140
Oil for reheating (at \$100/kl)	450	2,230
Power (at 3¢/kWh)	400	1,840
Labor	50	50
Maintenance	100	300
Other requirements	20	60
Gypsum (at \$5/t)	-120	-610
Total	1,130	5,010
Total annual cost, \$1,000	2,540	10,800
Desulfurization cost, \$/kl	16.9	14.4
Desulfurization cost, mills/kWhr	3.6	3.0

Flue gas desulfurization (FGD) is less expensive than hydrodesulfurization (HDS) at higher sulfur removal ratios (Table 2-4), but HDS is advantageous in that it produces elemental sulfur, which is a desirable by-product. In Japan and elsewhere, other processes yielding sulfur as a by-product seem much more expensive than HDS, except for oil refineries that have Claus furnaces for sulfur production.

4. MAJOR NEW FGD SYSTEMS FOR UTILITY BOILERS

STATUS OF FGD BY POWER COMPANIES

Table 4-1 lists power companies and their capacities for steam power generation and FGD. The nine major companies (Nos. 1 to 9 in the Table) have produced about 70 percent of the total steam power using mainly oil, with some LNG and a little coal. Electric Power Development Co. (EPDC, No. 10 in the Table), which was established by the nine major companies and the Central Government, has been the major consumer of domestic coal for power generation. Other power suppliers have relatively small capacities, burning mainly oil. Total capacity of the power generation plants, including those under construction and planned for construction, is 86,457 MW. The capacity of FGD systems in operation is 6,040 MW and that of systems under construction and being designed is 6,755 MW.

Among the major power companies, Tokyo Electric, Kansai Electric, and Chubu Electric, which supply power to the largest cities and industrial complexes in Japan, have relatively small capacities of FGD, with a B/A ratio (see Table 4-1) of only 1 to 7 percent. Those companies prefer

Table 4-1. CAPACITIES OF STEAM POWER GENERATION AND FGD OF POWER COMPANIES

No.	Power company	Power generation, MW			FGD, MW			B/A ^b (%)
		Existing	Under construction ^a	Total (A)	Existing	Under construction ^a	Total (B)	
1	Hokkaido	1,270	1,225	2,495	0	525	525	21.0
2	Tohoku	3,925	1,200	5,125	275	625	900	17.6
3	Tokyo	19,167	4,400	23,567	283	0	0	1.2
4	Chubu	9,935	3,800	13,733	970	0	970	7.1
5	Hokuriku	1,412	1,000	2,412	600	500	1,100	45.6
6	Kansai	10,672	1,200	11,872	360	469	838	7.1
7	Chugoku	3,777	1,800	5,777	950	1,100	2,050	36.8
8	Shikoku	2,687	450	3,137	900	0	900	12.5
9	Kyushu	4,500	2,700	7,200	188	1,438	1,626	22.6
10	EPDC	1,430	0	1,430	780	500	1,280	89.5
11	Niigata	350	350	700	175	175	350	50.0
12	Showa	550	0	550	150	250	400	72.7
13	Toyama	750	0	750	250	0	250	33.3
14	Mizushima	462	0	462	156	0	156	33.8
15	Sumitomo	368	250	618	0	218	218	35.3
16	Sekata	0	700	700	0	700	700	100.0
17	Fukui	0	250	250	0	250	250	100.0
18	Others	5,512	375	5,887	0	0	0	0.0
Total		66,775	19,700	86,475	6,040	6,755	12,795	14.8

^a Including those decided to be constructed.^b Capacity being scrubbed over total capacity.

use of low-sulfur fuels such as naphtha and LNG in polluted areas, because they believe that the regulation on SO_2 emission for those areas may become too stringent to be achieved by FGD. According to the recent regulation restricting total mass emissions of SO_2 , large power plants in the designated regions are required to keep SO_2 in flue gas below about 50 ppm, as described in Section 1. Although it is not difficult to reduce SO_2 from 1,500 ppm to 50 ppm by FGD, sulfur-free fuel is needed to reheat the treated gas. For plants to be constructed in regions to which much more stringent restriction is applied, FGD may entail some difficulty. On the other hand, Hokuriku Electric and Chugoku Electric, which have power plants remote from big cities, have larger B/A ratios (refer to Table 4-1).

FGD installations of power companies are listed in Table 4-2. Before 1973 power companies were not yet confident about the usefulness of FGD and therefore constructed test units to treat one-third to one-fourth of the gas from a boiler burning low-sulfur fuel; other industries that had difficulty in obtaining low-sulfur fuel constructed many FGD systems and demonstrated their reliability. The first commercial-scale FGD system for a utility boiler burning high-sulfur fuel was the Nishinagoya plant, Chubu Electric Power (200 MW, $620,000 \text{ Nm}^3/\text{hr}$) based on the Wellman-Lord

Table 4-2. FGD SYSTEMS OF POWER COMPANIES
(Oil-fired boilers unless otherwise indicated)

Power company	Power station	Boiler		FGD MW	Process developer	Absorbent, precipitant	By-product	Year of completion
		No.	MW					
Tohoku	Sinsendai	2	600	150	Kureha-Kawasaki	Na ₂ SO ₃ , CaCO ₃	Gypsum	1974
	Hachinohe	4	250	125	Mitsubishi H.I.	CaO	Gypsum	1974
	Niigata	4	250	125	Wellman-MKK	Na ₂ SO ₃	H ₂ SO ₄	1976
	Niigata H.	1	600	150	Mitsubishi H.I.	CaCO ₃	Gypsum	1976
	Akita	3	350	350	Kureha-Kawasaki	Na ₂ SO ₃ , CaCO ₃	Gypsum	1977
Tokyo	Kashima	3	600	350	Hitachi-Tokyo	Carbon, CaCO ₃	Gypsum	1972
	Yokosuka	1	265	133	Mitsubishi H.I.	CaCO ₃	Gypsum	1974
Chubu	Nishinagoya	1	220	220	Wellman-MKK	Na ₂ SO ₃	H ₂ SO ₄	1973
	Owase	1	375	375	Mitsubishi H.I.	CaO	Gypsum	1976
	Owase	2	375	375	Mitsubishi H.I.	CaO	Gypsum	1976
Hokuriku	Toyama	1	500	250	Chiyoda	H ₂ SO ₄ , CaCO ₃	Gypsum	1974
	Fukui	1	350	350	Chiyoda	H ₂ SO ₄ , CaCO ₃	Gypsum	1975
	Nanao	1	500	500	Not decided		Gypsum	1978
Kansai	Sakai	8	250	63	Sumitomo H.I.	Carbon	H ₂ SO ₄	1972
	Amagasaki	2	156	35	Mitsubishi H.I.	CaO	Gypsum	1973
	Amagasaki			121	Mitsubishi H.I.	CaO	Gypsum	1975
	Amagasaki	1	156	156	Mitsubishi H.I.	CaO	Gypsum	1976
	Osaka	3	156	156	Babcock-Hitachi	CaCO ₃	Gypsum	1975
	Osaka	2	156	156	Babcock-Hitachi	CaCO ₃	Gypsum	1975
	Osaka	4	156	156	Babcock-Hitachi	CaCO ₃	Gypsum	1976
	Kainan	4	600	150	Mitsubishi H.I.	CaO	Gypsum	1974
Chugoku	Mizushima	2	156	100	Babcock-Hitachi	CaCO ₃	Gypsum	1974
	Tamashima	3	500	500	Babcock-Hitachi	CaCO ₃	Gypsum	1975
	Tamashima	2	350	350	Babcock-Hitachi	CaCO ₃	Gypsum	1976
	Shimonoseki	2	400	400	Mitsubishi H.I.	CaCO ₃	Gypsum	1976
Shikoku	Anan	3	450	450	Kureha-Kawasaki	Na ₂ SO ₃ , CaCO ₃	Gypsum	1975
	Sakaide	3	450	450	Kureha-Kawasaki	Na ₂ SO ₃ , CaCO ₃	Gypsum	1975

Table 4-2 (continued). FGD SYSTEMS OF POWER COMPANIES

(Oil-fired boilers unless otherwise indicated)

Power company	Power station	Boiler No.	MW	FGD MW	Process developer	Absorbent, precipitant	By-product	Year of completion
Kyushu	Karita	2	375	188	Mitsubishi H.I.	CaO	Gypsum	1974
	Karatsu	2	375	188	Mitsubishi	CaCO ₃	Gypsum	1976
	Karatsu	2	500	250	Mitsubishi	CaCO ₃	Gypsum	1976
	Ainoura	1	375	250	Mitsubishi	CaCO ₃	Gypsum	1976
	Ainoura	2	500	250	Mitsubishi	CaCO ₃	Gypsum	1976
	Buzen	1	500	250	Kureha-Kawasaki	Na ₂ SO ₃ , CaCO ₃	Gypsum	1977
	Buzen	2	500	250	Kureha-Kawasaki	Na ₂ SO ₃ , CaCO ₃	Gypsum	1978
EPDC	Takasago	1	250 ^a	250	Mitsui-Chemico	CaCO ₃	Gypsum	1975
	Takasago	2	250 ^a	250	Mitsui-Chemico	CaCO ₃	Gypsum	1976
	Isogo	1	265 ^a	265	Chemico-IHI	CaCO ₃	Gypsum	1976
	Isogo	2	265 ^a	265	Chemico-IHI	CaCO ₃	Gypsum	1976
	Takehara	1	250 ^a	250	Babcock-Hitachi	CaCO ₃	Gypsum	1977
Niigata	Niigata	1	350	175	MHI	CaCO ₃	Gypsum	1975
Showa	Ichihara	1	150	150	Showa Denko	Na ₂ SO ₃ , CaCO ₃	Gypsum	1973
	Ichihara	5	250	250	Babcock-Hitachi	CaCO ₃	Gypsum	1976
Toyama	Toyama	1	250	250	Chiyoda	H ₂ SO ₄ , CaCO ₃	Gypsum	1975
Mizushima	Mizushima	5	156	156	Mitsubishi H.I.	CaO	Gypsum	1975
Sumitomo	Niihama	3	156	156	IHI	CaCO ₃	Gypsum	1975
Sakata	Sakata	1	350	350	Mitsubishi H.I.	CaCO ₃	Gypsum	1976
		2	350	350	Mitsubishi H.I.	CaCO ₃	Gypsum	1977
Fukui	Fukui	1	250	250	Not decided	CaCO ₃	Gypsum	1977

^a Coal-fired boilers.

process. The system has been operated smoothly at more than 97 percent operability. As regulations on wastewater have become increasingly stringent, however, wastewater treatment has posed a serious problem in the Wellman-Lord process.

Since 1973, power companies have constructed many full-scale FGD systems for utility boilers (250-500 MW) burning high-sulfur fuel (2.5 to 3 percent sulfur) using processes that produce gypsum. Processes and performance of several new FGD systems are described below.

In addition to the systems described in this section two units at Isogo Station, Electric Power Development Co., started operation very recently. These systems have a unit capacity of treating 900,000 Nm³/hr of flue gas from a coal-fired boiler using the Chemico-IHI process; no additional details are available.

PLANTS USING THE MHI LIME-GYPSUM PROCESS (MITSUBISHI-JECCO PROCESS)²

Karita Plant, Kyushu Electric⁸

The Karita plant, with a capacity of treating 550,000 Nm³/hr of flue gas from an oil-fired boiler (188 MW equivalent), went into operation in November 1974. The plant is based on the single-absorber system (Figure 4-1, Photo 4-1) and uses lime. Flue gas is first cooled to 55 to 60°C in a cooler and led into the plastic-grid-packed absorber. SO₂ concentration at the absorber inlet ranges from 400 to 600

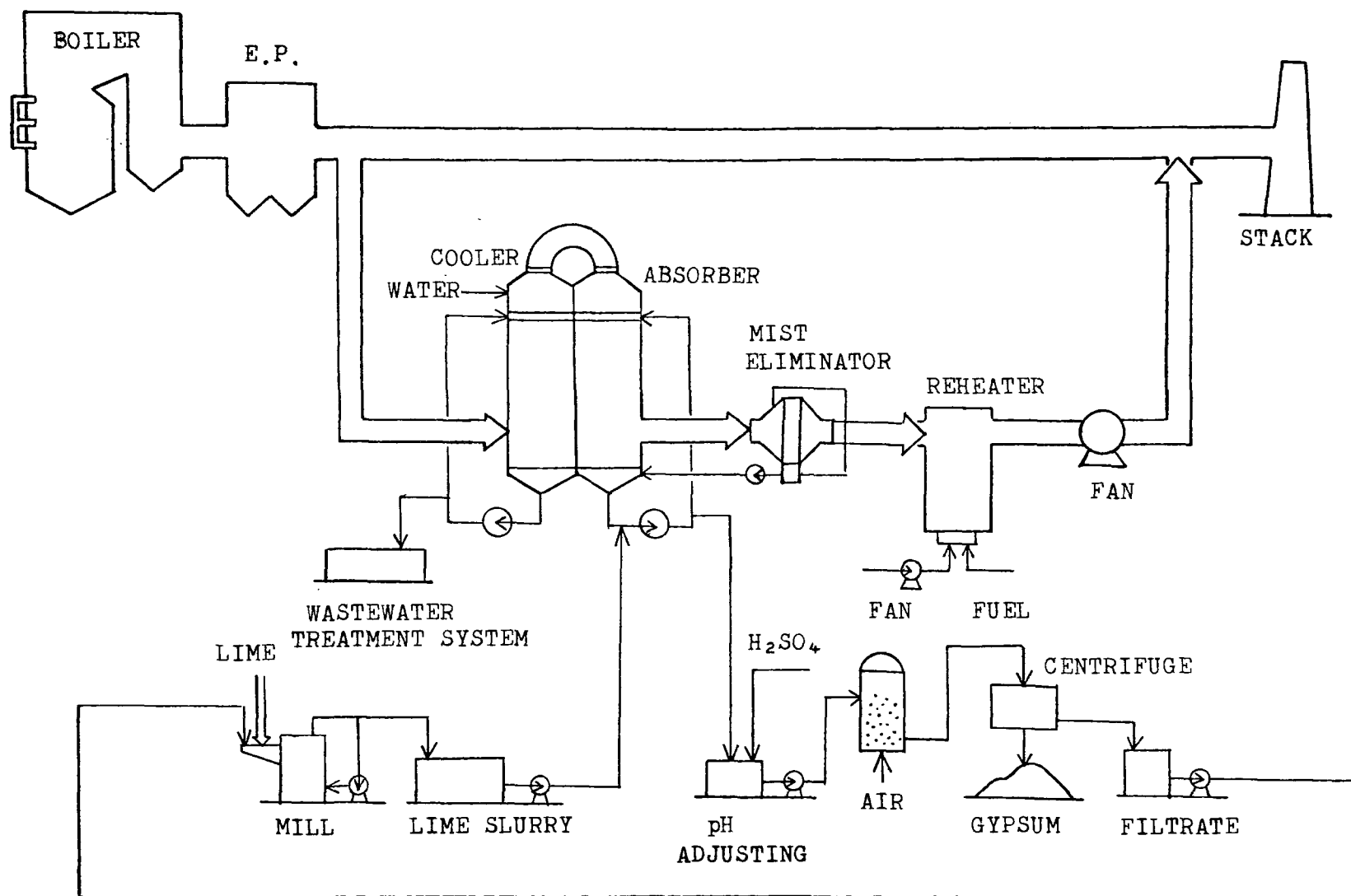


Figure 4-1. One-absorber system of MHI process.

ppm and at the outlet from 10 to 30 ppm. A lime slurry, at pH 6.4, 10 percent concentration is fed to the scrubber at an L/G ratio of 10. About 1.05 stoichiometry of lime is used. Space velocity (superficial gas velocity) of gas in the scrubber is 3.5 m/sec. Pressure drop through the cooler, absorber, and mist eliminator is 120 mm H₂O. Utility requirements at full load (550,000 Nm³/hr) are shown in Table 4-3.

Table 4-3. REQUIREMENTS AT THE KARITA PLANT

	Sulfur in fuel, %	
	1.2	0.8
Power, kW	3,200	3,200
Steam, t/hr	1.5	1.5
Industrial water, t/hr	3.0	3.0
Oil for reheating, t/hr	2.0	2.0
Lime, t/hr	1.1	0.8
Sulfuric acid, t/hr	0.2	0.13
Wastewater, t/hr	3.5	3.5
By-product gypsum, t/hr	2.9	2.0

As a little excess of lime is used to ensure high SO₂ removal efficiency by a single absorber, a considerable amount of sulfuric acid is needed to lower the pH of the calcium sulfite slurry to promote the oxidation of gypsum.

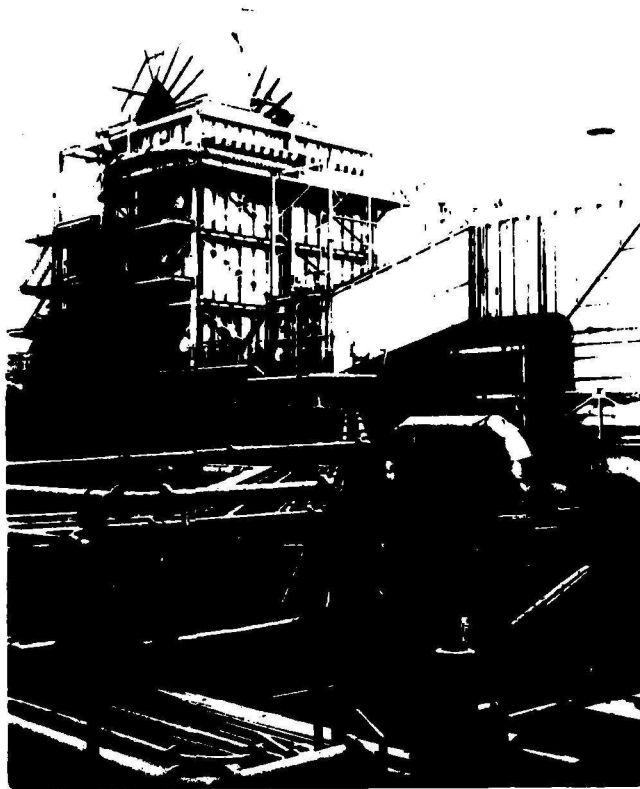


Photo 4-1. Karita plant, Kyusha Electric (188MW).



Photo 4-2. Owase Plant, Chubu Electric
(2 units each 375 MW).

The load fluctuates between 550,000 and 300,000 Nm³/hr every day. The flow rate of the slurry is kept constant, while the amount of lime is adjusted with the load.

The plant has been operated at 100 percent availability since start-up, except for a scheduled shutdown of the boiler from April 1 to May 14, 1975 (Figure 4-2). On February 24, 1975, a flow-rate-adjusting bulb was stopped up but was repaired without interrupting scrubber operation. In the inspection of April 1975, considerable scaling was found on the mist eliminators. The eliminators had been washed with circulating liquor; since May 1975, in an effort to reduce scaling, they have been washed alternately with the liquor and fresh water. Water balance is shown in Figure 4-3.

Other Plants

Two new units (175 and 250 MW) using the MHI process with limestone have started operation recently at the Karatsu station, Kyushu Electric, and two new units (375 MW each) using lime have started at the Owase station, Chubu Electric (Photo 4-2). Operating parameters are shown in Table 3-5.

The units at Owase have a two-tower system.² SO₂ concentration is reduced from 1,600 ppm to 30 ppm using slightly less than the stoichiometric amount of lime (Table

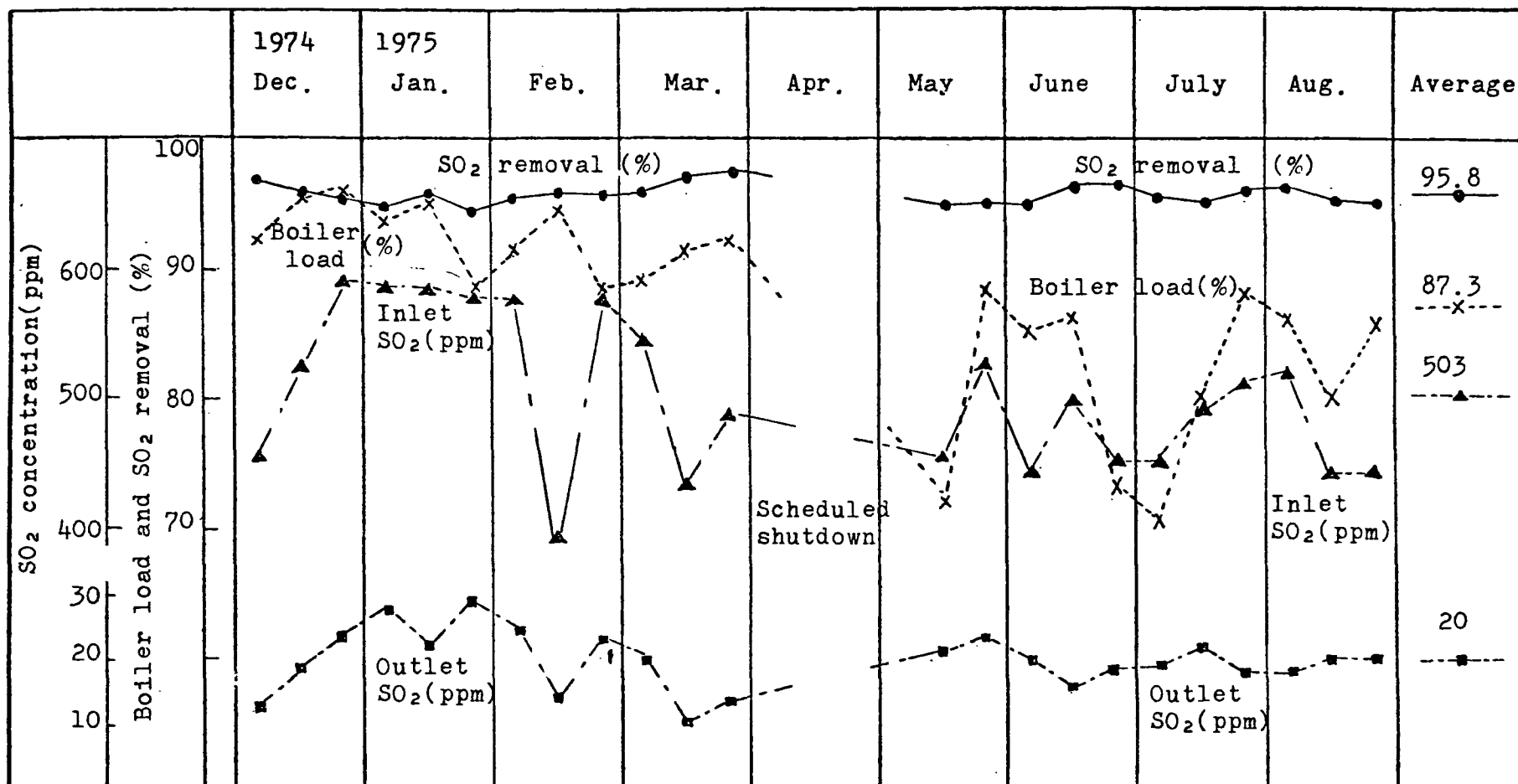


Figure 4-2. Operation data fo Karita plant, Kyushu Electric.

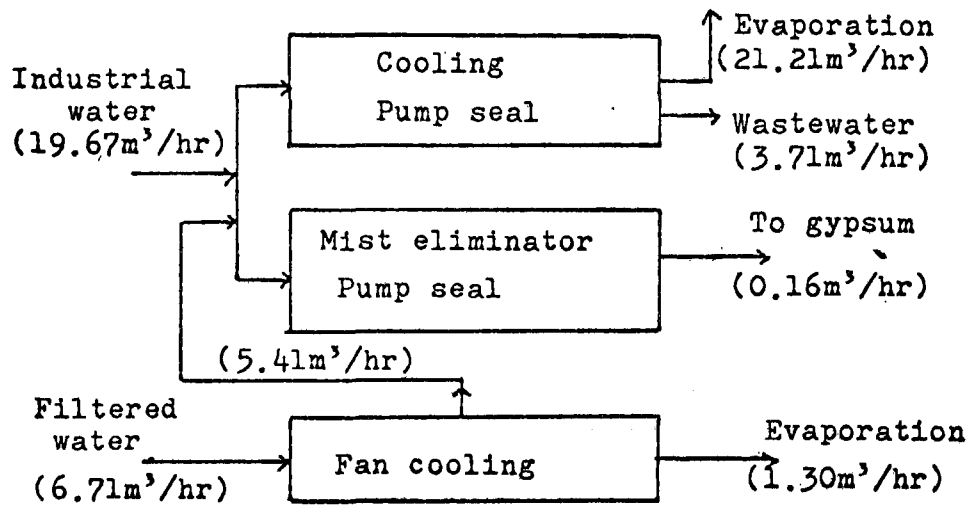


Figure 4-3. Water balance (Karita plant, MHI process).

3-5) at a pressure drop of 200 mm H₂O. No sulfuric acid is required because about 99 percent of the lime reacts and the pH of the calcium sulfite solution is lowered considerably by the use of two absorbers and also by the high concentration of SO₂ at the inlet.

MITSUI-CHEMICO LIMESTONE-GYPSUM PROCESS AT THE TAKASAGO PLANT, ELECTRIC POWER DEVELOPMENT CO.²

Process and Plant Design

This plant is based on the Mitsui-Chemico process developed by Mitsui Miike Machinery Co. It is the first Mitsui-Chemico process unit to be used commercially for a coal-fired utility boiler, and has a capacity of treating 840,000 Nm³/hr flue gas from a 250 MW boiler. The plant consists of two single-stage Chemico scrubbers placed in series, a pH controller to reduce the pH of the calcium sulfite slurry by introducing the flue gas, and two reactors for oxidation of the sulfite to gypsum by air (Figure 4-4, Photos 4-3 and 4-4). A catalyst is used to promote both SO₂ removal and oxidation. The flue gas containing 1,500 ppm SO₂ and 80 mg/Nm³ dust is passed through an electrostatic precipitator and the FGD unit to remove 90 percent of the SO₂ and 75 percent of the dust. Limestone, 95 percent under 325 mesh, is the absorbent. The operating parameters are shown in Table 3-5. Specifications of the principal equipment are shown in Table 4-4.

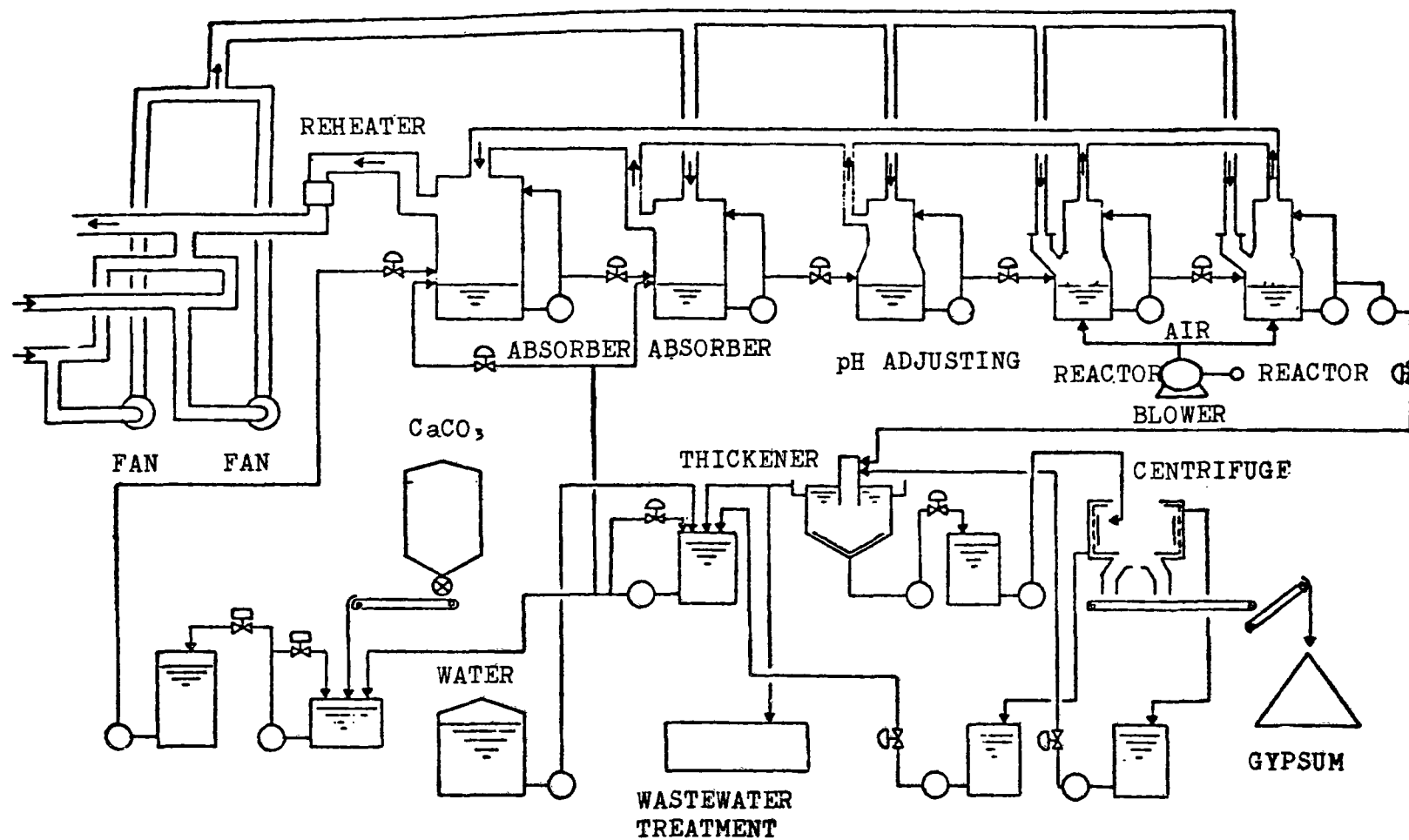


Figure 4-4. Flowsheet of Mitsui-Chemico process.

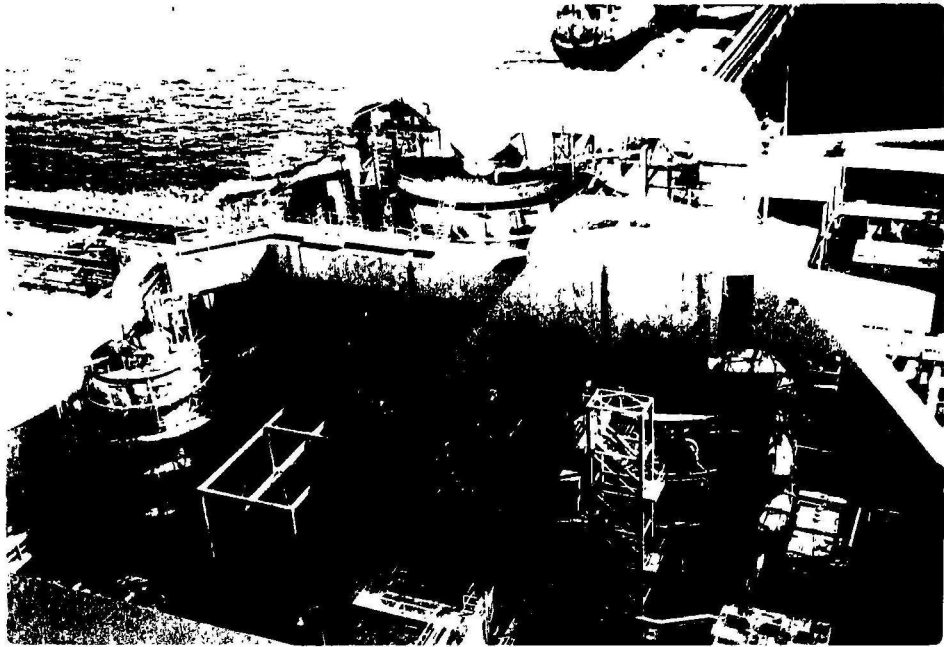


Photo 4-3. Takasago plant, EPDC (250 MW)
(scrubber and reactors).

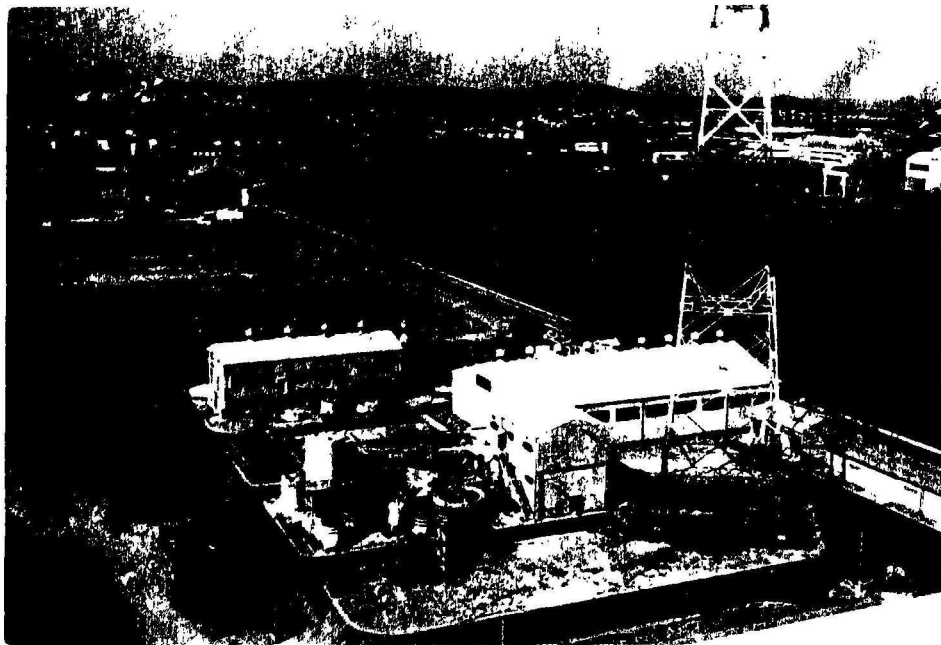


Photo 4-4. Takasago plant, EPDC (250 MW)
(Gypsum centrifuge and storage).

Table 4-4. MAIN EQUIPMENT AT TAKASAGO PLANT

Equipment	Type and dimensions
Limestone tanks	1,000 t x 2
Absorber (1st)	Venturi 11.5 m ϕ x 20 m
Absorber (2nd)	Venturi (double throat) 14 m ϕ x 23 m
Forced draft fans	(12,300 m ³ /min, 1,900 kW) x 2
Circulation pumps	
For 1st absorber	135 kW x 4
For 2nd absorber	200 kW x 5
pH adjusting tower	Venturi 6 m ϕ x 16 m
Reactors (oxidizers)	6 m ϕ x 18 m, 2 units
Blowers for oxidizers	(110 Nm ³ /hr, 1,000 mm H ₂ O, 250 kW) x 3
Centrifuges	(solid 1.5 t/hr) x 7
Reheating furnaces	(oil 625 liter/hr) x 4

Performance

The plant started up in January 1975 and has since operated well except for several brief shutdowns to clean the mist eliminators. The eliminators are washed with the circulating liquor and fresh water. System availability reaches 98 percent. The load fluctuates between 170 and 250 MW daily. The slurry flow rate in the scrubbers is kept constant.

The daily requirements for operation are as follows:

CaCO_3 - 118 tons

Power - 136,000 kWh

Oil - 41 liters (reheating)

Steam - 13 tons (heating of oil)

Industrial water - 1,200 tons

Catalyst - \$350

The coal contains about 500 ppm chloride. To keep the chloride concentration of the scrubber liquor below 7,000 ppm, about 5 ton/hr of wastewater is purged. The relationship between the amount of purge water and the chloride concentration of coal and equilibrated scrubber liquor is shown in Figure 4-5.

To prevent promotion of corrosion by chloride, a plastic or rubber lining and a high-grade stainless steel are used.

Another plant with the same capacity is near completion at Takasago. The new plant has one reactor instead of two, as at the plant just described.

BABCOCK-HITACHI PROCESS AT THE TAMASHIMA PLANT, CHUGOKU ELECTRIC⁸

Process Description²

Hitachi Ltd. constructed a plant (500 MW full scale) using Babcock-Wilcox scrubbers and an oxidizing system to by-produce gypsum (Figure 4-6, Photos 4-5 and 4-6). Limestone, more than 95 percent under 325 mesh as shown in Table

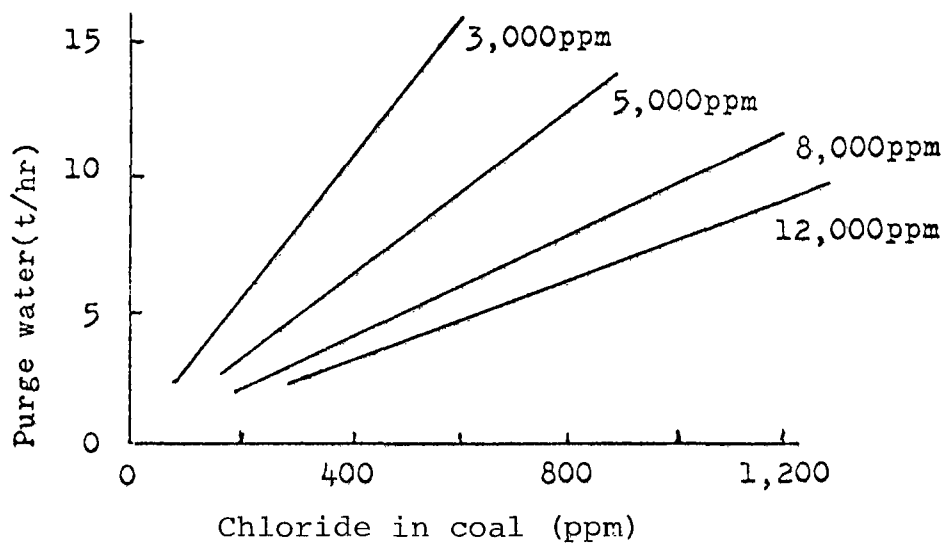


Figure 4-5. Purge water and chloride concentration.

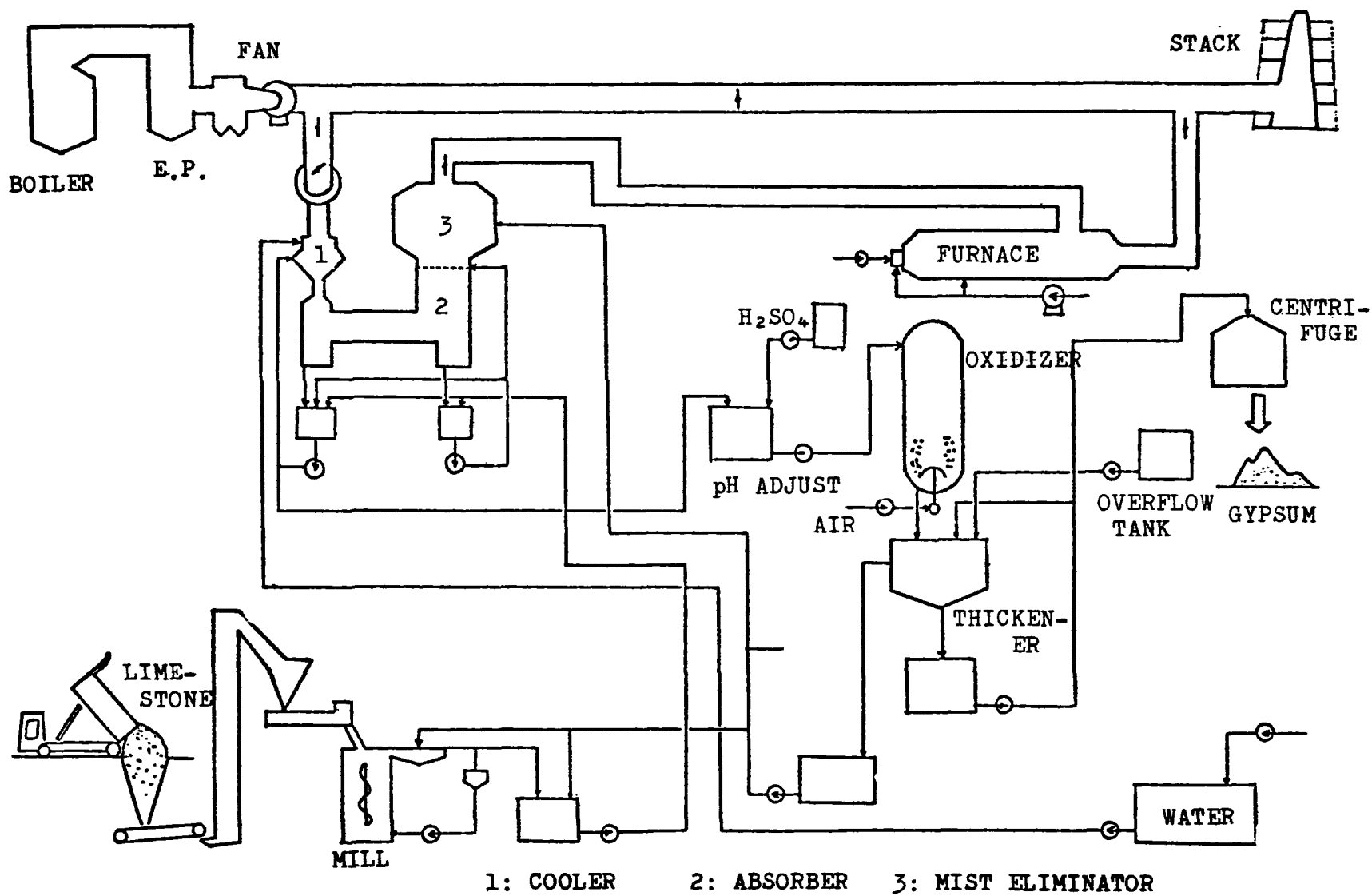


Figure 4-6. Flowsheet of Babcock-Hitachi process.

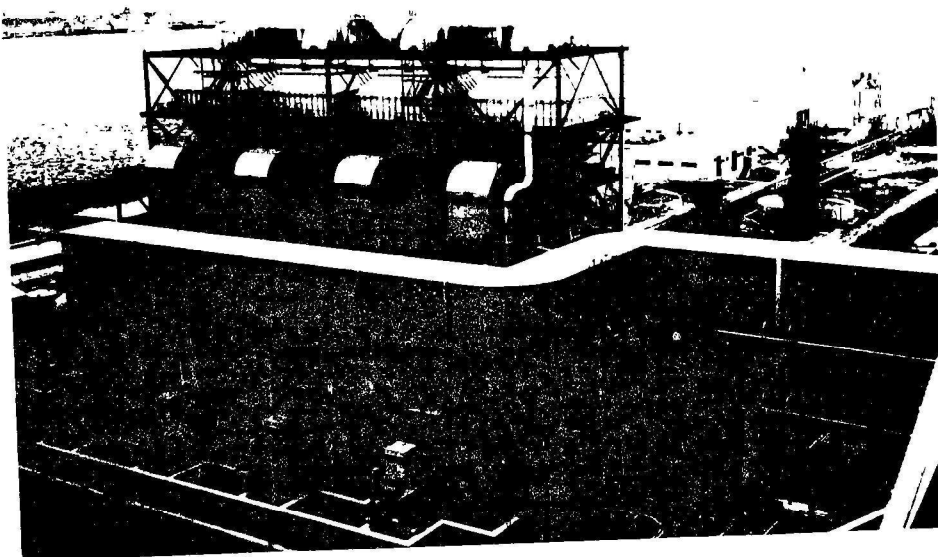


Photo 4-5. Tamashima Plant, Chugoku Electric (500 MW)

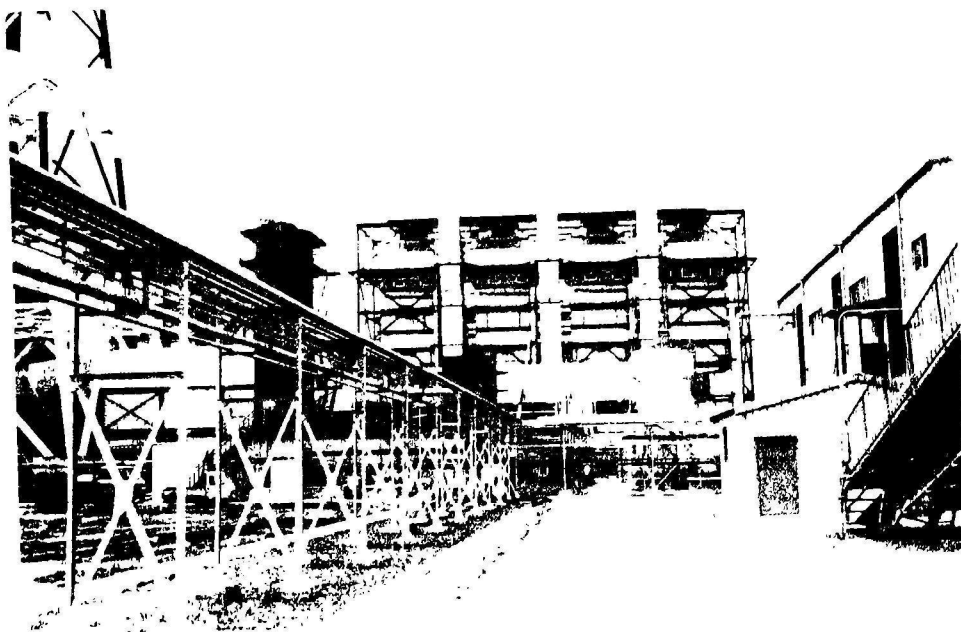


Photo 4-6. Tamashima Plant, Chugoku Electric (500 MW)

4-5, is used as the absorbent at a stoichiometry of 1.05 to 1.10. Four scrubbers were installed, of which three are in use and one is for standby. Flue gas from an oil-fired boiler, with an SO₂ concentration of 1,500 ppm, is first cooled to 55°C in venturi scrubbers and then led into absorbers with six stages of perforated plates, which remove about 95% of the SO₂.

The L/G ratio is 2.1 (liters/Nm³) in the venturi and 10 in the absorber. The pH of the slurry is 5.5 in the venturi and 6 to 6.2 in the absorber. The pressure drop is fairly heavy---230 mm H₂O in the venturi and 600 mm in the absorber. A calculation indicates that 99 percent SO₂ removal efficiency may be achieved at a total pressure drop of 1,300 ppm in the venturi and the absorber. Mist eliminators are of a new type designed by Hitachi for easy washing.

Table 4-5. COMPOSITION OF LIMESTONE

(Percent)

	CaO	SiO ₂	Al ₂ O ₃	Fe ₂ O ₃	MgO
Specification	>55	<0.3	<0.1	<0.1	<0.6
Actual	55	0.3	0.1	0.1	0.5

The calcium sulfite sludge discharged from the bottom of the venturi at pH 5.5 is sent to a reactor, where sulfuric acid is added to decompose the unreacted limestone and to lower the pH to 4.7. The slurry is sent to three oxidizers

in parallel and is oxidized by air bubbles generated by rotary atomizers. More than 97 percent of the calcium sulfite is oxidized to gypsum. The gypsum contains 7 to 8 percent moisture after being centrifuged and is used in the manufacture of wallboard and cement.

Performance

The plant came on-stream in July 1975 and has been used for base-load operation. Power consumption reaches 3.6 percent of the power generated. Small amounts of scale tend to form in some parts of the scrubber, which occasionally becomes dislodged and plugs the spray nozzles. The strainers of the circulation pumps have been improved, and operability (percentage of FGD operation hours to boiler operation hours) has reached 97 percent.

About 93 t/hr of industrial water is used, of which 13 tons are used for washing the mist eliminator. At the beginning of the operation, about 5 t/hr wastewater was purged to keep the chloride concentration of the circulating liquor below 1,000 ppm. Use of wastewater has been substantially reduced recently.

Composition of Slurry

Solids in the slurry in the absorber circulating tank consist of calcium sulfite, 50 to 60 percent; gypsum, 40 to 50 percent; and limestone 5 percent. The relation between

the pH of the slurry in the reactor (pH adjusting tank) and the composition of solids in the slurry after the oxidation is shown in Figure 4-7. When a slurry at pH 5.5 was oxidized without adding sulfuric acid, the oxidation ratio was about 80 percent. The product, consisting of about 80 percent gypsum and 20 percent calcium sulfite, has proved useful as a retarder of cement setting.

KUREHA-KAWASAKI SODIUM-LIMESTONE PROCESS AT THE SAKAIDE PLANT, SHIKOKU ELECTRIC

Process

The Sakaide plant went in operation in August 1975 with a capacity of treating 1,260,000 Nm³/hr of flue gas containing 1,050 ppm SO₂ from a 450 MW oil-fired boiler (Photos 4-7 and 4-8). The process is similar to that for Shinsendai plant, Tohoku Electric (Figure 4-8).³ Flue gas from the 450 MW oil-fired boiler, passed through an electrostatic precipitator by a forced-draft fan, is fed into a venturi-type precooler and then into a grid-packed-type scrubber, where the gas is washed with a sodium sulfite solution (about 20%) at pH 7.0 and an L/G ratio of nearly 10 (about 70 gal/1,000 scf). The gas is then passed through a mist eliminator, reheated by an afterburner, and ducted into a stack.

The liquor discharged from the scrubber at pH 6.5 is passed through a series of five reactors, where powdered limestone, ground in a vertical tower mill to pass 325 mesh, is reacted to precipitate calcium sulfite and regenerate sodium sulfite.

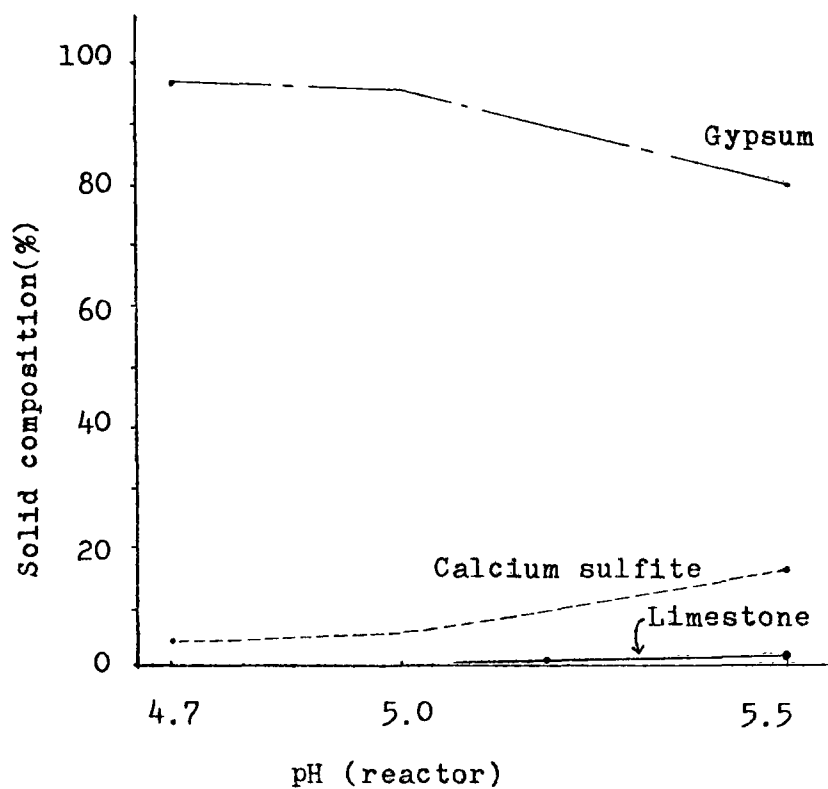


Figure 4-7 Relationship of pH of slurry in reactor to solid composition after oxidation

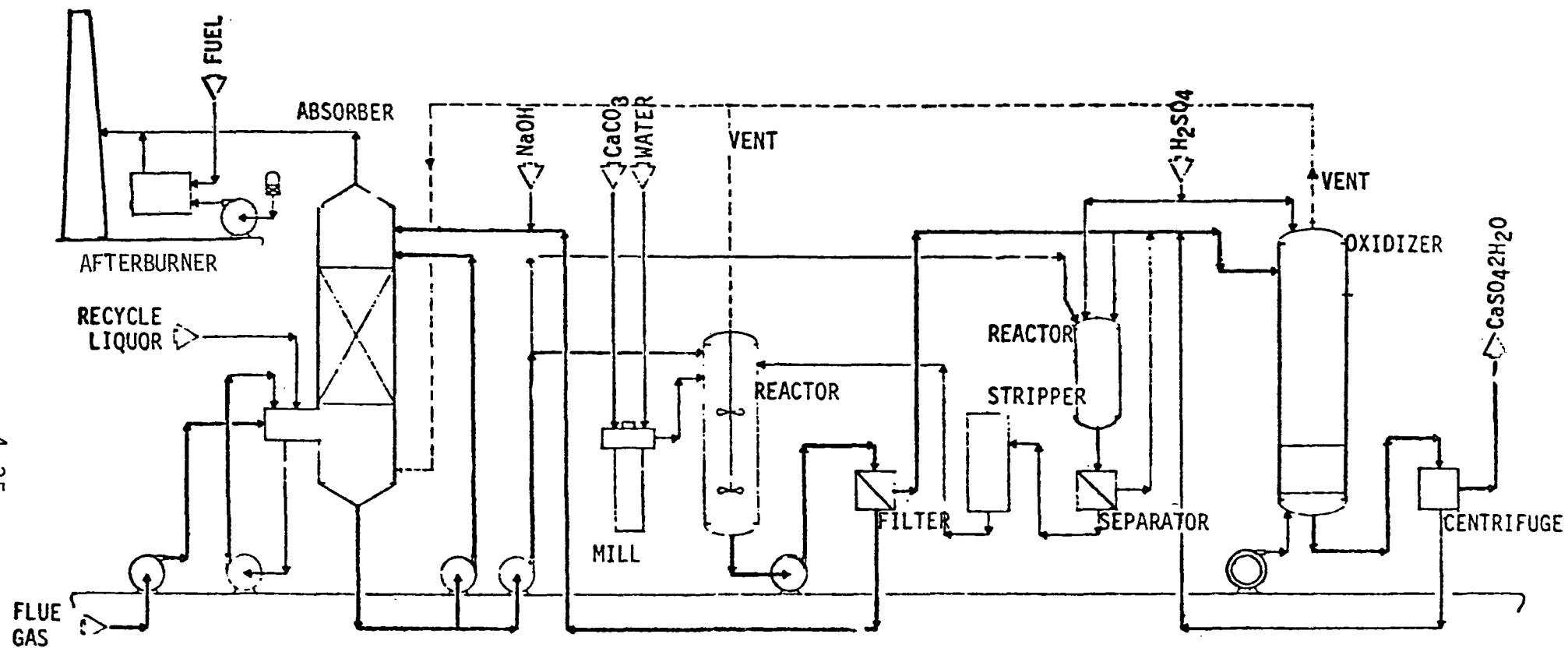


Figure 4-8. Flowsheet of Kureha-Kawasaki process.

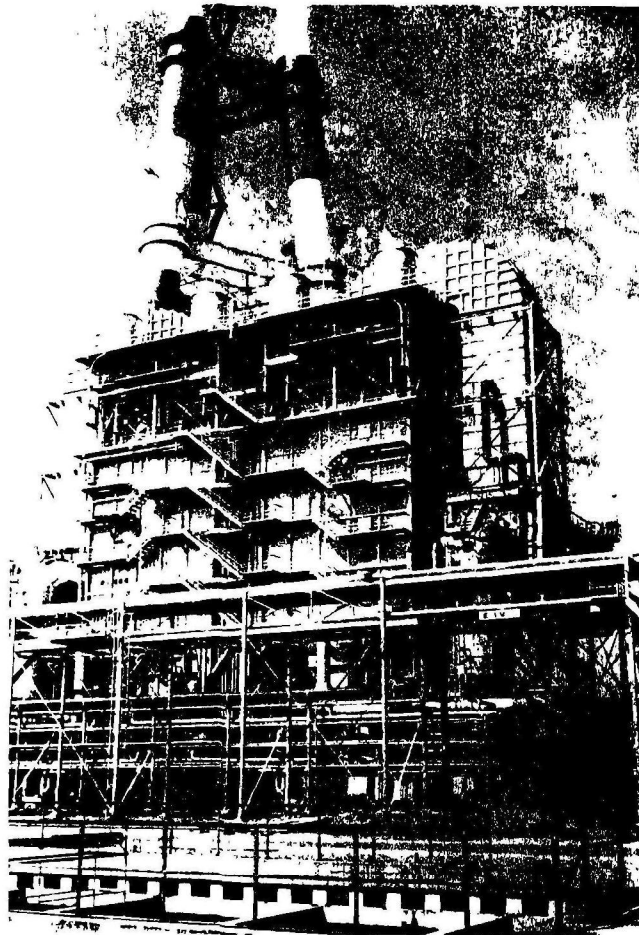


Photo 4-7. Sakaide plant, Shikoku Electric (450 MW)
(two scrubbers in parallel)

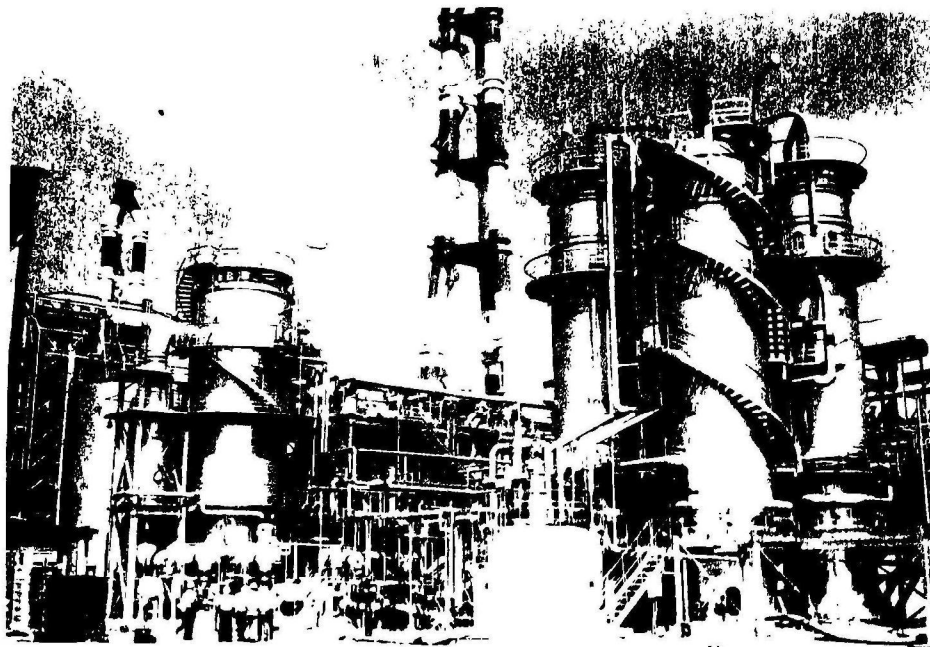
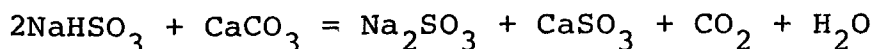
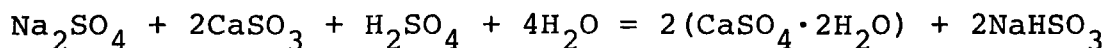


Photo 4-8. Sakaide plant, Shikoku Electric (450 MW)
(Oxidizers and stripper)



The pH of the slurry at the outlet is 7.3. The calcium sulfite (50% slurry) is separated on a vacuum filter and washed to remove sodium sulfite. The filter cake (about 60% water) is repulped to 10 percent slurry, treated with sulfuric acid to reduce pH, and oxidized by air bubbles in an oxidizer (at 2 atmospheres of pressure) developed by Kureha and Kawasaki. The gypsum slurry is centrifuged to less than 8 percent moisture; the separated liquid is recycled to repulp the calcium sulfite.

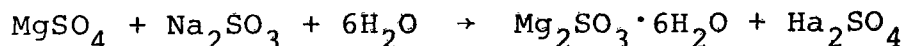
A portion of sodium sulfite is oxidized to sulfate in the scrubber by oxygen in the flue gas. A side stream of liquor from the scrubber is treated to decompose the sulfate by reaction with sulfuric acid and calcium sulfite.



The slurry from the desulfation unit is filtered, and the gypsum is sent to the oxidizer as seed to obtain well-grown gypsum crystals; the separated sodium bisulfite solution is passed through a steam-heated stripper to generate SO_2 , which is recycled to the desulfation step to reduce sulfuric acid consumption and then sent to the reactor.

A main difference from the Shinsendai plant is that the Sakaide plant incorporates a unit to remove magnesium from the circulating liquor. Magnesium derived from limestone tends to accumulate in the circulating liquor and

delay the reaction of limestone with sodium bisulfite. A portion of the liquor from the gypsum centrifuge is reacted with sodium sulfite solution from the stripper to precipitate magnesium sulfite, which is filtered off. The filtrate is mixed with the liquor discharged from the scrubber.



Another difference in the two plant operations is that here the gypsum is washed with water to reduce the sodium content as needed for use in wallboard production.

Performance⁸

Operation parameters are shown in Table 3-5. More than 99 percent of the SO_2 is removed by sodium sulfite scrubbing. Operation has been smooth since start-up. The load fluctuates daily between 1,260,000 and 534,000 Nm^3/hr . The sodium sulfate concentration of the circulating liquor has reached 11 to 12 percent, exceeding the design value of 8 percent, but a high SO_2 recovery ratio has been attained. This unit is characterized by the absence of water purging from the system. At full-load operation, 63.5 t/hr water is charged, of which 29.9 tons is used for gypsum wash. The same amount, 63.5 tons, leaves the system, of which 62.0 tons is evaporated and the rest is contained in the by-product gypsum. Chloride concentration of the circulating liquor has reached 3,500 ppm but has caused no corrosion problem because the

liquor contains little oxygen, which tends to cause stress corrosion in the presence of excess chloride. In the Kureha-Kawasaki process, oxidation is carried out with the calcium sulfite separated from the mother liquor, and therefore the oxygen content of the liquor is kept low. The chloride input from fuel and water and the output with gypsum containing 7 to 9 percent moisture appear to have equalized at the 3,500 ppm concentration level.

CHIYODA PROCESS AT THE FUKUI PLANT, HOKURIKU ELECTRIC

Process Description

A flowsheet of the Chiyoda process is shown in Figure 4-9. Flue gas is first treated by a prescrubber to eliminate dust and to cool the gas to 50°C. The cooled gas is led into a packed tower absorber containing 2-inch Tellerette. Dilute sulfuric acid (2 to 5% H_2SO_4), which contains ferric ion as a catalyst and is nearly saturated with oxygen, is fed to the packed tower. About 90 percent of the SO_2 is absorbed and partly oxidized into sulfuric acid.

The product acid is led to the oxidizing tower, into which air is bubbled from the bottom to complete the oxidation. Most of the acid at 50 to 60°C, saturated with oxygen, is returned to the absorber. Part of the acid is treated with powdered limestone (74% under 200 mesh) to produce gypsum. A special type of crystallizer has been

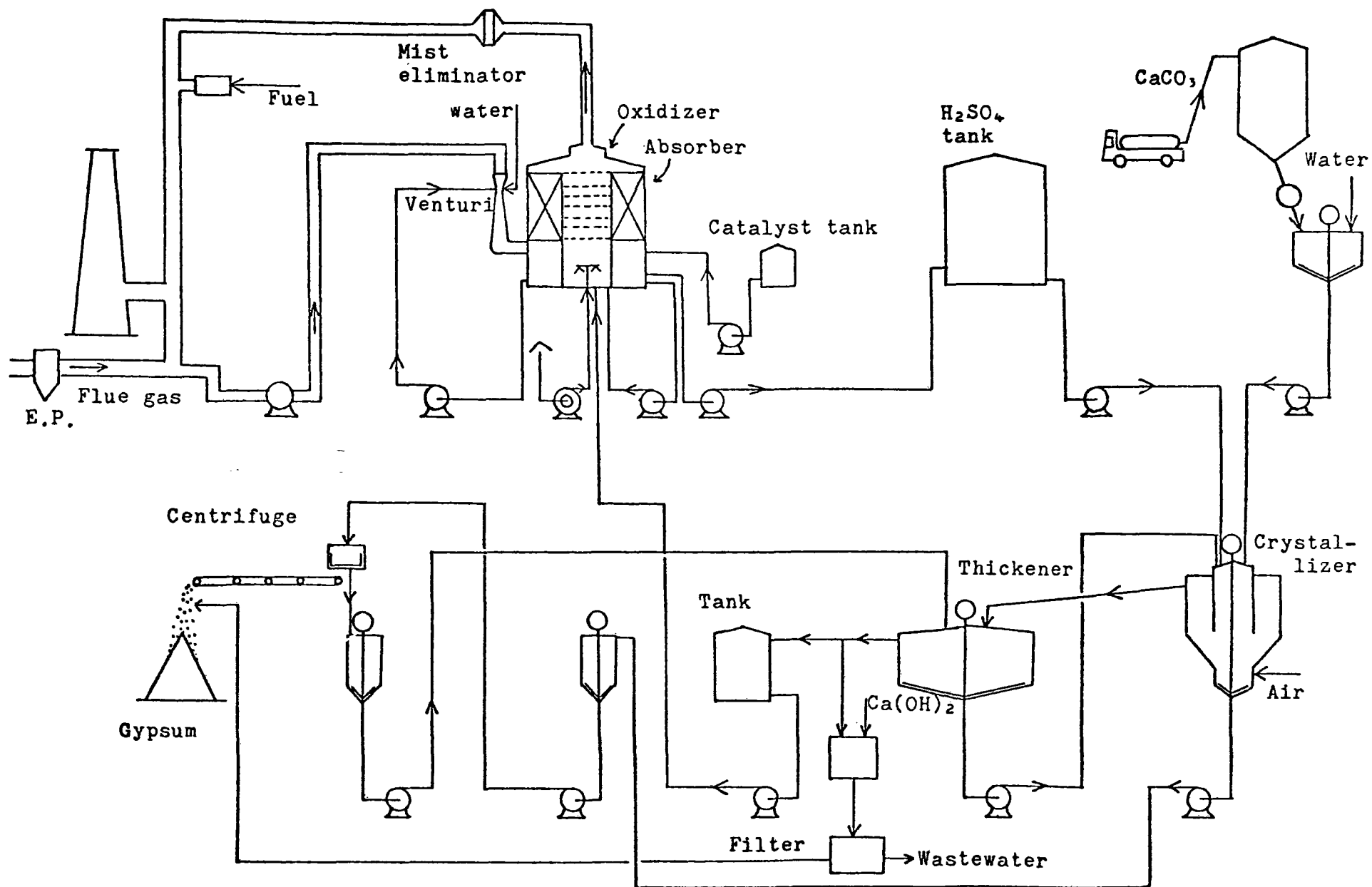


Figure 4-9. Flowsheet of Chiyoda process.

developed to obtain good gypsum with crystals ranging from 100 to 500 microns. The gypsum is centrifuged from the mother liquor and washed with water. The product gypsum is of good quality and salable.

The mother liquor and wash water are sent to the scrubber. A small amount of wastewater is discharged in order to prevent corrosion caused by the accumulation of chloride in the circulating liquor.

Performance

The Fukui plant (Photos 4-9 and 4-10) has operated smoothly since its start-up in the summer of 1975, treating 1,050,000 Nm³/hr of flue gas from a 350 MW oil-fired boiler. The plant has a double-cylinder-type absorber-oxidizer. Operating parameters are listed in Table 3-5. The requirements are 150 t/day of limestone, 3.8 m³/hr of oil for reheating, 11.8 MW of electric power, and 2,000 t/day of industrial water. About 580 t/day of wastewater is purged after being treated to keep the pH at 7.9, suspended solids below 5 mg/liter, and COD below 10 mg/liter. Plant operation requires only two operators per shift.

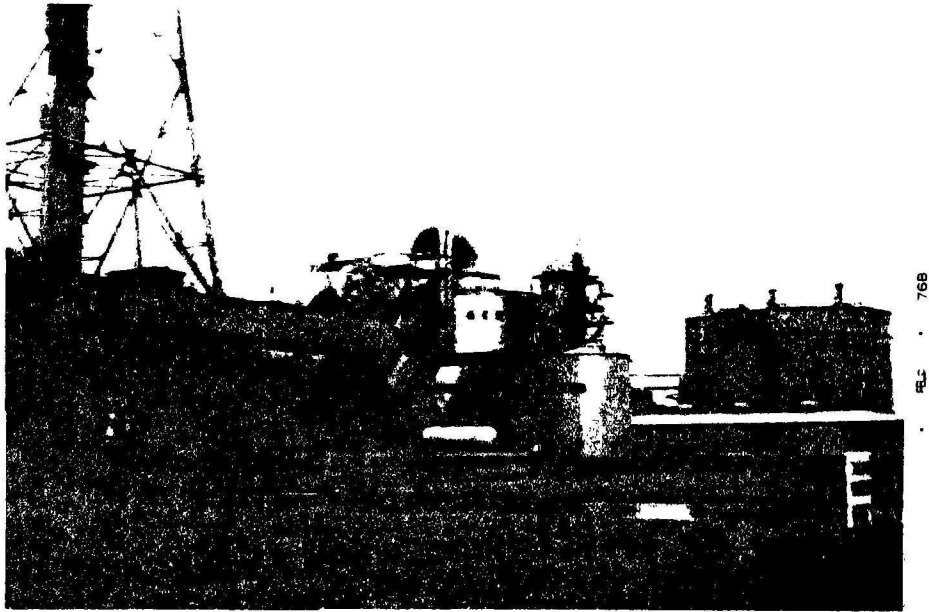


Photo 4-9. Fukui plant, Hokuriku Electric (350 MW)

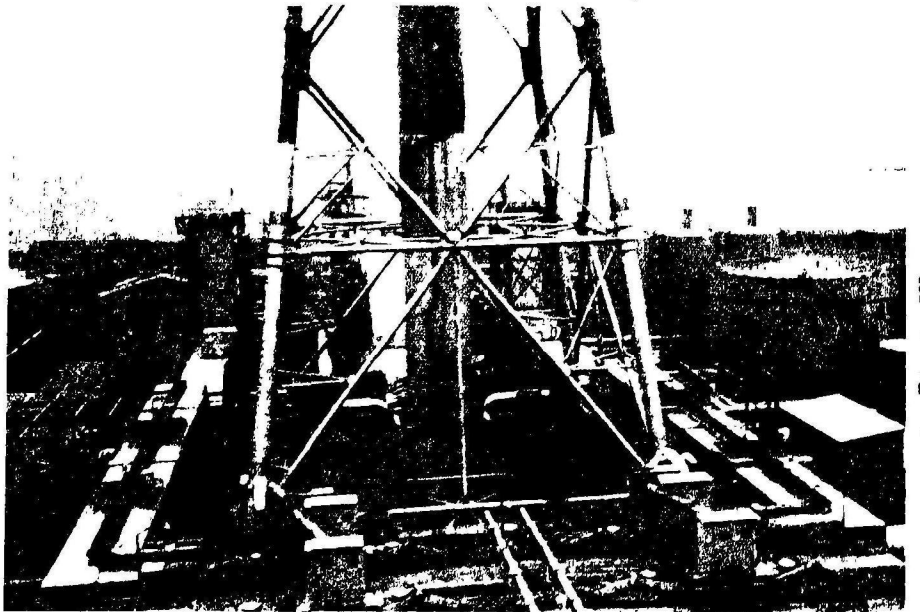


Photo 4-10. Fukui plant, Hokuriku Electric (350 MW)

5. FLUE GAS DESULFURIZATION IN THE STEEL INDUSTRY

INTRODUCTION

Many desulfurization units have been installed since 1971 to treat flue gas from iron-ore sintering plants, which constitute the major source of SO_2 emissions in the steel industry (Table 5-1). As the absorbent, a lime slurry is used by Kawasaki Steel (MHI process), a limestone slurry by Sumitomo Metal (Sumitomo-Fujikasui Moretana process), slurry of pulverized converter slag by Nippon Steel (SSD process), and a calcium chloride solution dissolving lime by Kobe Steel (Cal process). All of these systems by-produce gypsum. Nippon Kokan uses ammonia scrubbing to by-produce ammonium sulfate or gypsum by reacting lime with the sulfate.²

By 1977, 22 FGD systems will be in operation, with a total capacity of treating $13,800,000 \text{ Nm}^3/\text{hr}$ ($8,120,000 \text{ scfm}$) of flue gas, which is about half the total gas from all sintering plants in Japan.

Flue gas from sintering plants is characterized by a high O_2 concentration (12 to 16%), a relatively low SO_2 concentration (200 to 1,000 ppm), and a dust rich in ferric oxide. Oxidation of sulfite into sulfate occurs in the scrubbers much more readily than with flue gas from a boiler,

Table 5-1. SO₂ REMOVAL INSTALLATION FOR WASTE GAS FROM IRON-ORE SINTERING MACHINES

Steelmaker	Plant site	Gas treated 1,000 Nm ³ /hr	Process	Absorbent	Year of completion	Gypsum, t/year
Kawasaki Steel	Chiba	120	MHI	CaO	1973	3,600
	Chiba	320	MHI	CaO	1975	13,200
	Chiba	650	MHI	CaO	1976	26,500
	Mizushima	750	MHI	CaO	1974	27,600
	Mizushima	900	MHI	CaO	1975	32,400
	Mizushima	750	MHI	CaO	1977	27,600
Sumitomo Metal	Kashima	880	Moretana	CaCO ₃	1975	32,400
	Kashima	1,000	Moretana	CaCO ₃	1976	40,500
	Kashima	1,000	Moretana	CaCO ₃	1977	40,500
	Wakayama	370	Moretana	CaCO ₃	1975	14,400
	Kokura	720	Moretana	CaCO ₃	1976	26,500
Kobe Steel	Amagasaki	175 x 2	Cal	Ca(OH) ₂	1976	12,600
	Kobe	375	Cal	Ca(OH) ₂	1976	12,600
	Kakogawa	1,000 x 2	Cal	Ca(OH) ₂	1977	72,000
Nakayama Steel	Osaka	375	Cal	Ca(OH) ₂	1976	13,500
Nippon Steel	Tobata	200	SSD	Slag	1974	7,200
	Wakamatsu	1,000	SSD	Slag	1976	32,400
Nippon Kokan	Keihin	150	NKK	NH ₃ , CaO	1971	7,200
	Fukuyama	760	NKK	NH ₃ ^a	1976	12,000 ^b
	Ogishima	1,230	NKK	NH ₃ ^a	1977	20,000 ^b

^a Ammonia in coke oven gas.^b Ammonium sulfate.

because the oxidation is promoted by the high O_2/SO_2 ratio and also by the catalytic action of the ferric oxide.

This section will describe mainly the lime and limestone processes, discussing the dimensions and performance of the FGD systems.

MHI PROCESS AT THE MIZUSHIMA PLANT, KAWASAKI STEEL

FGD System For No. 4 Sintering Machine

Kawasaki Steel installed FGD systems first at its Chiba Works, using the lime-gypsum process developed by Mitsubishi Heavy Industries. Satisfied with operation of these systems, Kawasaki Steel introduced larger FGD systems at its Mizushima Works, where they operate four iron-ore sintering machines with a unit capacity of 8,000 to 15,000 t/day. The No. 4 machine gives 350,000 to 750,000 Nm^3/hr of waste gas at about 150°C containing 500 to 1,000 ppm SO_2 and about 13.5 percent O_2 .

The flowsheet of the FGD system is similar to that shown in Figure 4-1. The gas is first cooled to 57°C in a cooler with water sprays and led into a plastic-grid-packed absorber, where it is treated with a lime slurry at pH 6.4 to 7.5 at an L/G ratio of 7 liters/ Nm^3 (about 50 gallons/1,000 scf) to remove more than 90 percent of the SO_2 . The treated gas passes through a mist eliminator, is heated to about 140°C by afterburning, and sent to a stack. A calcium

sulfite slurry discharged from the absorber is acidified to pH 4 by addition of sulfuric acid, then is led into an oxidizer and oxidized into gypsum by air bubbles generated by a rotary atomizer. The gypsum slurry is sent to a thickener and then is centrifuged to less than 10 percent moisture. The by-product gypsum is sold as a retarder of cement setting. The liquor from the centrifuge is returned to the thickener; the thickener overflow is returned to the absorber after lime is added.

A portion of the circulating liquor of the cooler is neutralized with lime to recover low-grade gypsum. The liquor from the centrifuge is sent to a wastewater treatment system and reused.

As the calcium sulfite is oxidized to a considerable extent in the absorber because of high concentration of oxygen in the gas, the gypsum is recycled to the absorber as crystal seed in order to prevent scaling. An automatic system has been installed to shut down and restart the FGD unit with the sintering machine.

Performance

The FGD system for the No. 4 machine went into operation in November 1974. Performance characteristics are shown in Figure 5-1. The gas volume fluctuated from 350,000 to 850,000 Nm³/hr and inlet SO₂ concentrations ranged from

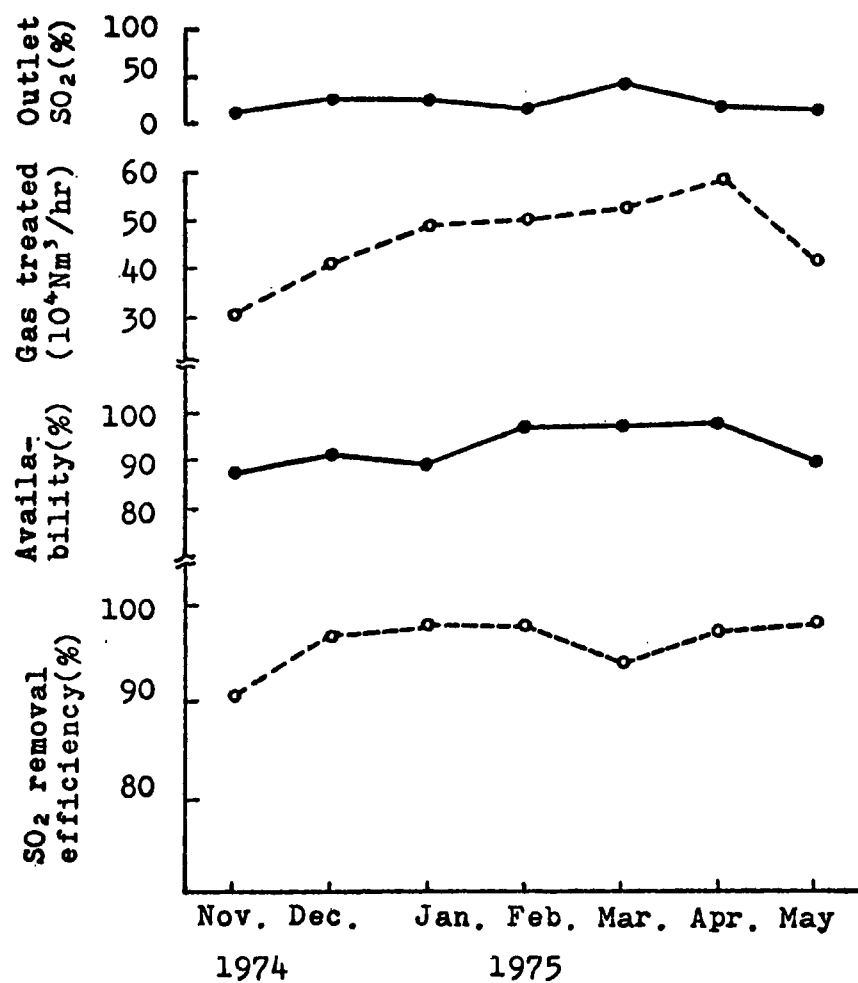


Figure 5-1. Performance of FGD plant for
No. 4 sintering machine.

400 to 1,100 ppm. The SO_2 removal efficiency was 91 to 98 percent and the SO_2 concentration at scrubber outlet ranged from 20 to 50 ppm. Availability (FGD operation hours as percent of total hours) was about 90 percent for the first 3 months because of several minor troubles, such as corrosion of the impeller of the cooler circulation pump, plugging of the lime-slurry pump, and breakage of a fire-brick in the furnace. Following the repairs, nearly 100 percent availability was obtained in the next 3 months. The low availability in May 1975 (about 90%) was due to a shutdown of the sintering machine.

On the average, the gas velocity in the scrubber is about 2.5 m/sec and the total pressure drop in the cooler, absorber, and mist eliminator is about 120 mm H_2O . Lime with less than 1 percent MgO has been used. The by-product gypsum contains about 7 percent moisture after being centrifuged and has an average crystal size of 40 microns. At the beginning of the operation, use of an excessive amount of lime to ensure high SO_2 removal efficiency (over 97%) resulted in consumption of a considerable amount of sulfuric acid. In later operation, slightly less than 1 mole lime to 1 mole inlet SO_2 has been used to obtain about 95 percent removal, and thus the consumption of sulfuric acid has been reduced.

FGD SYSTEMS OF SUMITOMO METAL (MORETANA PROCESS)

Moretana Process

Sumitomo Metal is operating two FGD systems and constructing three more (Table 5-1), all using the Moretana process developed by Sumitomo jointly with Fujikasui Engineering Co. The process is characterized by use of the Moretana scrubber fitted with four perforated plates made of stainless steel. The holes range from 6 to 12 mm diameter and the plate thickness from 6 to 20 mm. Both dimensions are varied depending on the specific scrubbing conditions. Free space in the cross section ranges from 25 to 50 percent. The bottom tray serves mainly as a gas distributor, and the upper three serve as absorbers. The gas and liquid flows are adjusted to maintain a liquor head of 10 to 15 mm on each plate. Gas velocity is higher than in usual scrubbers. The design gives extreme turbulence, producing foam layers 400 to 500 mm thick, and thus ensures a high ratio of SO₂ and dust removal. The mist eliminator is a set of vertical chevron sections mounted in a horizontal duct after the scrubber.

A flowsheet of the process is shown in Figure 5-2. Gas from a sintering machine is first treated with water in a Moretana scrubber for cooling and removal of more than 90 percent of the dust. Removal of ferric oxide dust is useful

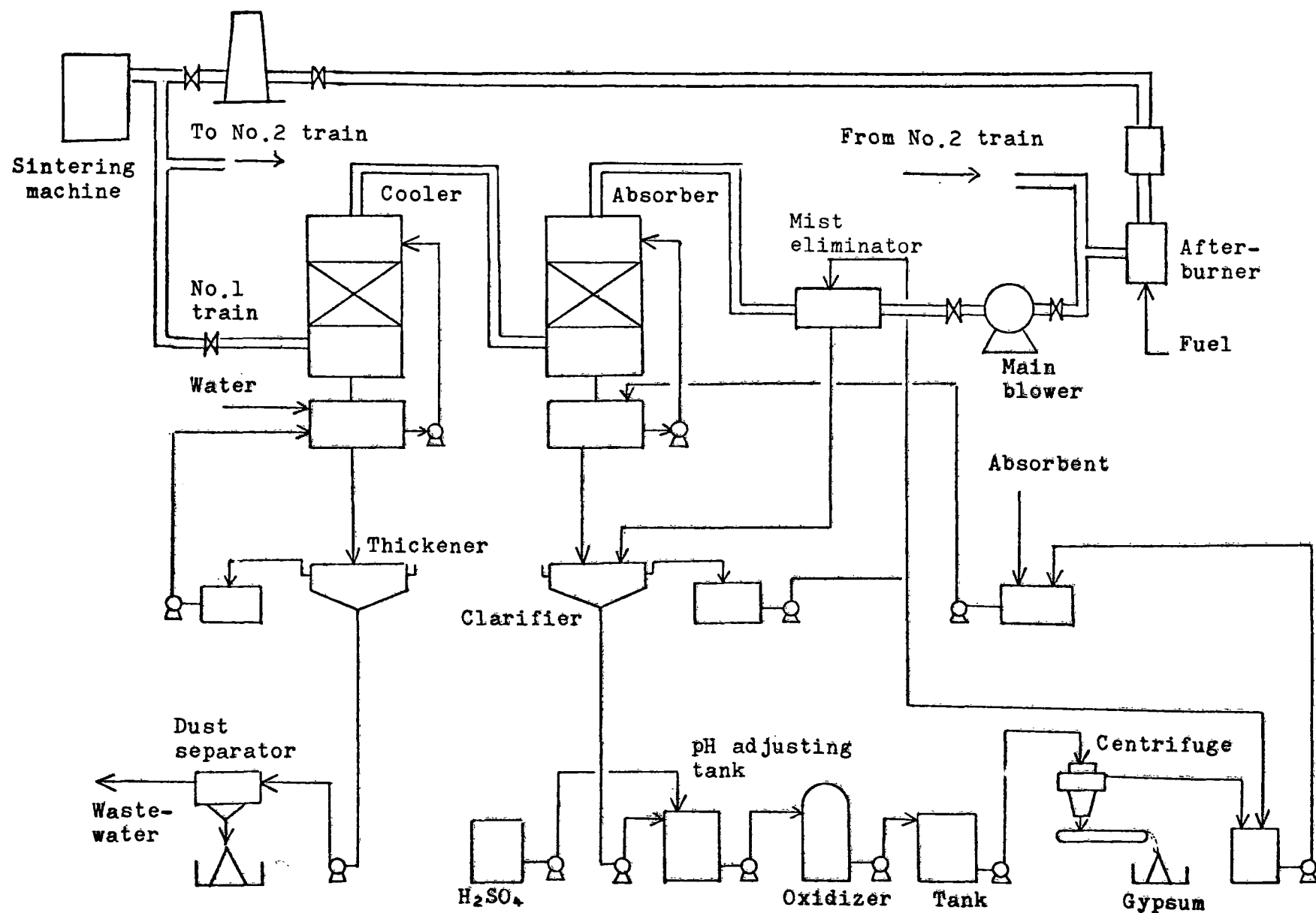


Figure 5-2. Flowsheet of Moretana process (Kashima plant, Sumitomo Metal).

in that it reduces oxidation in the absorber to allow scale-free operation. The gas is then treated with a limestone slurry (or a mixed slurry of lime and limestone), 10 to 20 percent in excess of stoichiometric amount, in a second Moretana scrubber to remove more than 95 percent of the SO_2 . The limestone contains less than 1 percent MgO and is ground to pass 325 mesh. The calcium sulfite slurry discharged from the scrubber is sent to a clarifier and then to a pH adjusting tank, where the pH is adjusted to about 4.0 by adding a small amount of H_2SO_4 . The slurry is then sent to an oxidizer developed by Fujikasui to convert calcium sulfite to gypsum. The gypsum slurry is centrifuged, and the filtrate is returned to the absorber.

The discharge from the cooler is sent to a thickener. The overflow is returned to the cooler; the underflow is filtered. The filter cake is returned to the sintering machine, and the filtrate is sent to a wastewater treatment system.

Kashima Plant, Sumitomo Metal

An FGD system at the Kashima plant, with capacity of treating $880,000 \text{ Nm}^3/\text{hr}$ of gas, was started up in September 1975 and has been in stable operation. All the gas from a sintering machine is treated, flow rates ranging from $350,000$ to $880,000 \text{ Nm}^3/\text{hr}$. The gas contains 200 to 450 ppm SO_2 , 14 to 15 percent O_2 , 6 to 8 percent CO_2 , 1 to 1.5

percent CO, 4 to 10 percent H₂O, and 0.15 to 0.23 g/Nm³ of dust at about 150°C. The scrubbing units consist of two trains, each with a capacity of treating 440,000 Nm³/hr of gas. The Moretana scrubber works normally with a gas velocity between 3 and 5 m/sec. When the gas flow rate is low, only one train is used. Equipment dimensions are shown in Table 5-2. Operational data on No. 1 train only are shown on Figure 5-3.

Table 5-2. EQUIPMENT DIMENSIONS, FGD SYSTEM

AT KASHIMA PLANT

Facility	Number	Size (Specification)
Cooler	2	6.5 m (dia.) 24.5 m (height)
Absorber	2	6.5 m (dia.) 20.5 m (height)
Mist eliminator	2	6 x 6 m, 2.4 m (length)
Oxidizer	2	2.8 m (dia.) 5.4 m (height)
Centrifuge	4	500 kg/hr each

Wakayama Plant, Sumitomo Metal

An FGD plant at Wakayama (Photo 5-1), with capacity of treating 375,000 Nm³/hr of waste gas from a sintering machine, started operation in May 1975 and has since operated well except for a defect in the plastic lining of a cooler, which was found early in the operation and was repaired. Operability of the plant has been about 98 percent. A scheduled shutdown of the sintering plant normally occurs every 2

5-11

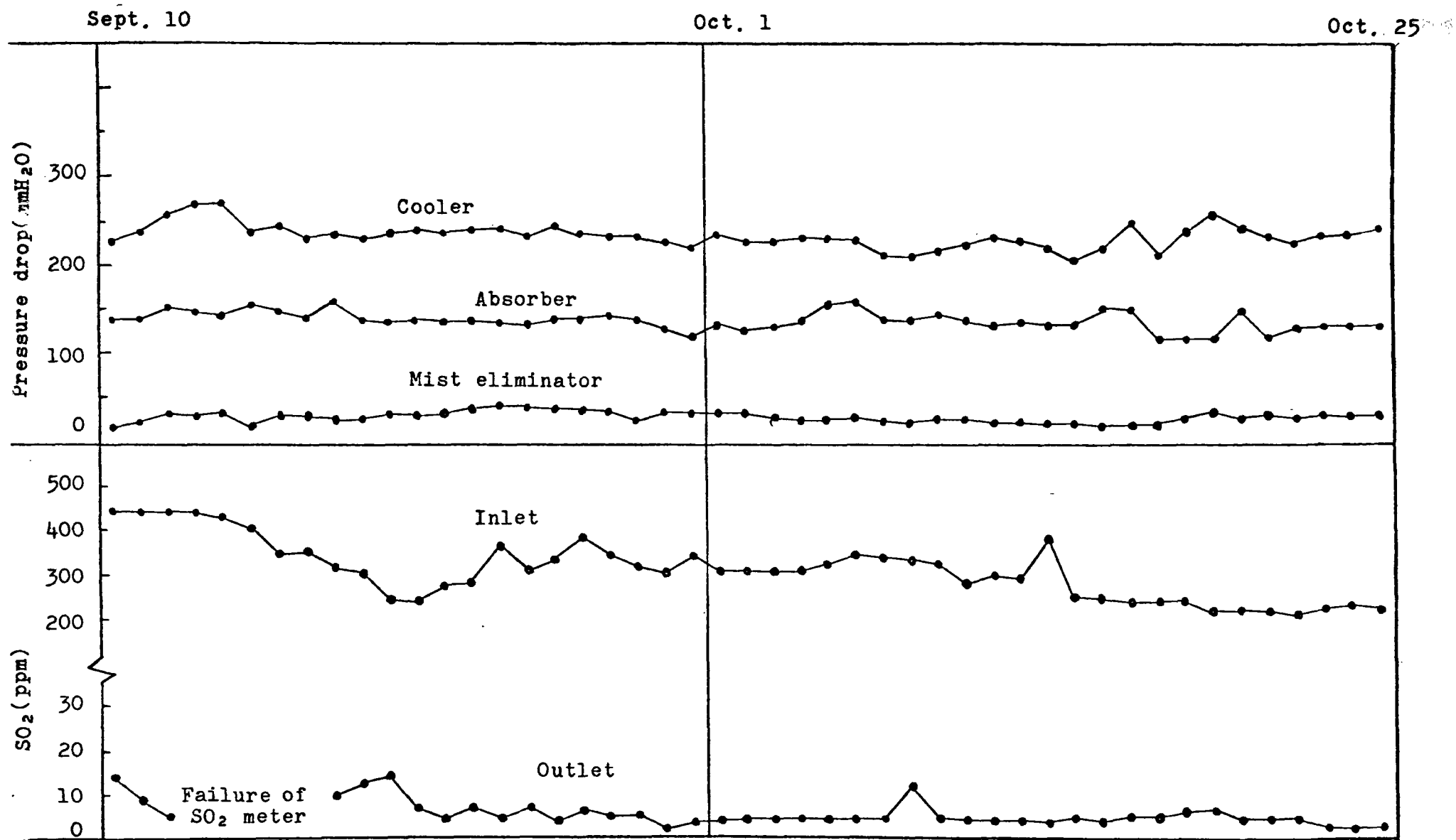


Figure 5-3. Operation data of No. 1 train, Kashima plant.

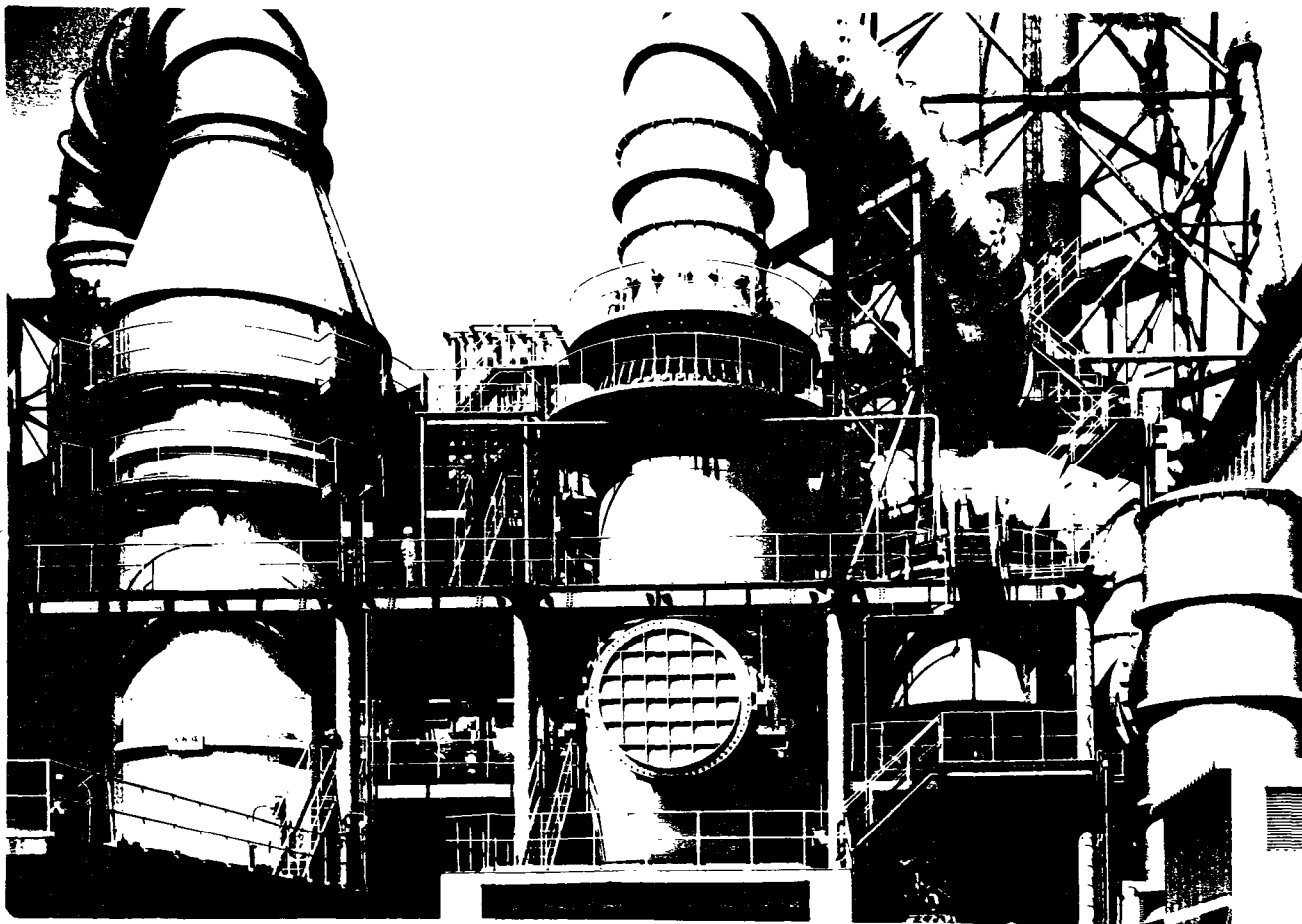


Photo 5-1. Wakayama plant, Sumitomo Metal (Absorber left, cooler right)

months. The mist eliminator is washed every 30 minutes with circulating liquor or fresh water. The pressure drop in the mist eliminator, which is 30 mm H₂O at start-up, gradually increases while the unit is washed with the circulating liquor. When the pressure drop reaches 50 mm, fresh water is used in place of the liquor until the pressure drop returns to 30 mm. The ratio of liquor to fresh water usage is about 80 to 20.

KOBE STEEL CALCIUM CHLORIDE PROCESS

Process Description

Kobe Steel has developed a new process using a 30 percent calcium chloride solution dissolving lime as the absorbent. A pilot plant (50,000 Nm³/hr) has been operated, and two commercial systems (Table 5-1) have just come on-stream to treat waste gas from iron ore sintering plants.

Calcium chloride solution dissolves 6 to 7 times as much lime as does water. High SO₂ recovery is attained at a low L/G ratio of 3 liters/Nm³. The flowsheet is shown in Figure 5-4.

Waste gas is first cooled in a cooler to which a calcium chloride solution (about 5%, from a gypsum centrifuge) is fed to cool the gas to 70°C and to remove most of the dust. The solution is concentrated here to about 30 percent and is sent to a scrubber system after dust removal

Figure 5-4. Flowsheet of Cal process.

by filtration. The gas then enters an absorber, in which a calcium chloride solution (about 30%, at pH 7 dissolving lime) is sprayed to remove more than 90 percent of the SO_2 . The gas is then passed through a mist eliminator to a stack. The liquor discharged from the absorber at pH 5.5 containing calcium sulfite is sent through a thickener to a centrifuge to separate most of the solution, which is sent to a tank where calcium hydroxide is dissolved to raise the pH to 7. The calcium sulfite sludge from the centrifuge is repulped with water and some sulfuric acid to produce a slurry at pH 4. The slurry is oxidized by air bubbles into gypsum, which is then centrifuged. The liquor from the centrifuge, containing about 5 percent calcium chloride, is returned to the cooler. The system gives no wastewater.

Since vapor pressure of the liquor is low, the temperature of the gas after the scrubbing reaches 70°C . Thus less energy is required for reheating than in the usual wet processes with gas temperatures of 55 to 60°C at the scrubber exit. The mist eliminator is washed with the circulating liquor. The solubility of gypsum in the liquor is very low (nearly $1/100$ of that in water), and evaporation of the liquor does not cause scaling.

In continuous operation of the pilot plant for about 6 months, a soft deposit formed on the wall of the absorber

when the L/G ratio was less than 1; the deposit could be removed by use of an L/G ratio greater than 2. A highly corrosion-resistant material is required for the cooler; the lower part at the hot gas inlet is made of titanium.

Amagasaki Plant

The FGD system at the Amagasaki plant has two trains, each with a capacity of treating 175,000 Nm³/hr of flue gas at 120°C containing 240 to 400 ppm SO₂, 0.05 to 0.2 g/Nm³ dust and 14 to 16 percent O₂. The plant began test operation in February 1976. The following problems were encountered during a 2-month test run:

Unusual vibration of a centrifuge.

Wearing of a control valve.

Scaling of pH meter electrode.

Breakage of rubber lining in a reducer.

Those problems have been solved, and the system went into commercial operation in April 1976. SO₂ removal efficiency ranges from 91 to 94 percent. Dust removal efficiency is about 50 percent. Gas velocity in the absorber is 3 m/sec. Total pressure drop in the cooler, absorber, and mist eliminator is 190 mm H₂O. The L/G ratios are 4.0 in the cooler and 3.0 in the absorber. More than 50 percent of the calcium sulfite is oxidized in the absorber. The by-product gypsum has an average crystal size of 40 microns and contains about 8 percent moisture and 0.1 percent chloride after being centrifuged.

NIPPON STEEL SLAG PROCESS (SSD PROCESS)

Nippon Steel has developed an FGD process that uses converter slag as the absorbent (Figure 5-5). The slag contains about 40 percent CaO, 16 percent SiO₂, 3 percent MgO, 3 percent Al₂O₃, and 35 percent FeO and Fe₂O₃; the slag is otherwise useless. Nippon Steel has operated a prototype system with a capacity of treating 200,000 Nm³/hr of waste gas from a sintering plant since 1974. A commercial unit (1,000,000 Nm³/hr) has just started operation.

The process is similar to other lime/limestone-gypsum processes except that it uses no oxidizer. The gas is cooled and led into two absorbers in series to remove 95 percent of the SO₂. The slag is fed to the second absorber to produce a calcium sulfite slurry; the slurry then goes to the first scrubber, where it is entirely oxidized into gypsum due to a low pH and large amounts of iron compounds, which act as a catalyst. The by-product gypsum contains about 40 percent impurities and is discarded. Scaling encountered in the prototype system must be reduced to ensure long-term continuous operation. The system may be useful for steel processes that normally yield large amounts of useless slag.

Figure 5-5. Flowsheet of SSD process.

6. NEW FGD PROCESSES

STATUS OF NEW DEVELOPMENTS

Since 1974 several FGD systems have been newly developed in order to improve the wet lime/limestone process (Kawasaki magnesium-gypsum process and MKK jet scrubber process), to ensure stable operation at low cost (Dowa aluminum sulfate process), to attain more than 99 percent SO₂ removal (Kureha sodium acetate process), or to obtain as by-products elemental sulfur (Chemico-Mitsui process in combination with Claus furnace), sulfuric acid (Hitachi-Unitika activated carbon process), or ammonium sulfate (Kurabo process). These processes are described in this section.

Mitsubishi Heavy Industries and Toyo Engineering have operated pilot plants for ammonia scrubbing followed by thermal decomposition of ammonium sulfate and production of elemental sulfur using the IFP reactor. These operations may be abandoned, however, because of problems encountered, mainly in the decomposition step. A few other companies have made small-scale tests on wet processes that yield elemental sulfur, but the results have not yet been disclosed.

KAWASAKI MAGNESIUM-GYPSUM PROCESS

Process Description

Kawasaki Heavy Industries completed a lime-gypsum process plant in 1973. Scaling occurred in the early operations but was reduced by addition of magnesium. Recently Kawasaki constructed two commercial plants using a magnesium-gypsum process, both of which went into operation in January 1976. A flowsheet of the process is shown in Figure 6-1. Flue gas is treated in a multi-venturi type scrubber with a slurry containing calcium and magnesium sulfites. More than 90 percent of the SO_2 is absorbed to form bisulfites. A portion of the sulfite-bisulfite slurry is sent to an oxidizer and oxidized by air bubbling to produce gypsum and magnesium sulfate.

The gypsum slurry is treated in a thickener and then centrifuged. The filtrate is returned to the thickener. Most of the thickener overflow, containing about 5 percent magnesium sulfate, is returned to the absorber. A portion of the overflow is sent to a reactor and reacted with lime to precipitate gypsum and magnesium hydroxide. The resulting slurry is sent to the absorber; gypsum in the slurry works as seed crystal.

The pH of the slurry in the scrubber ranges from 5 to 5.5, lower than that in usual lime/limestone scrubbing.

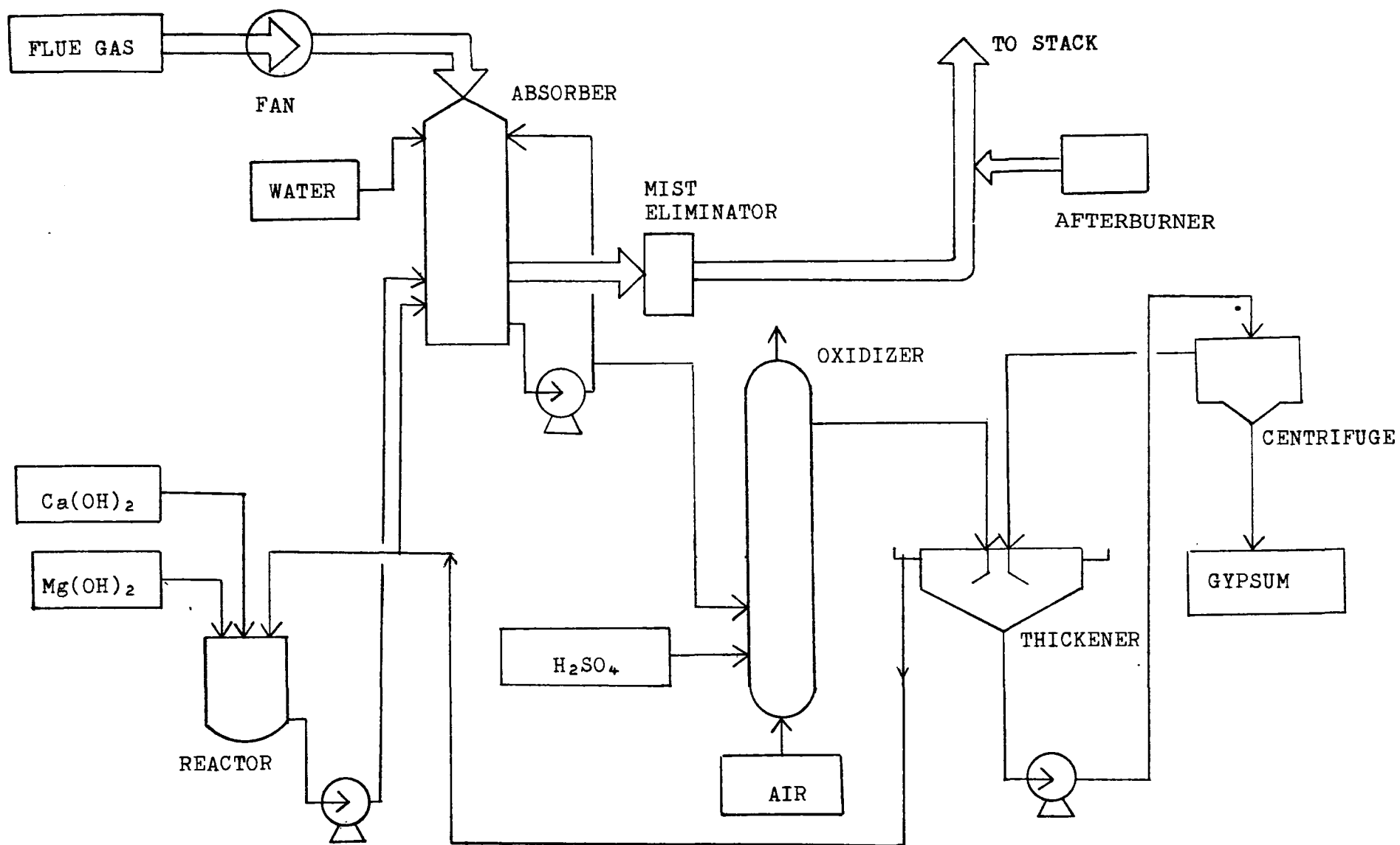


Figure 6-1. Flowsheet of Kawasaki magnesium-gypsum process.

Scale-free operation may be achieved more easily at the low pH, but the SO₂ removal ratio is fairly high because of the effect of magnesium.

Plant Operation

The Saidaiji plant, Nippon Exlan, with a capacity of treating 200,000 Nm³/hr of flue gas from an oil-fired boiler, started operation on January 17, 1976. Formation of soft scale in a pipe from the reactor forced a plant shutdown on January 28 for four days. An additional pipe was installed for use during cleaning of the original pipe. A 10-day shutdown, starting February 26, 1976, was carried out to clean build-up scale from a mist eliminator, which had been washed with a circulating liquor. Fresh water wash has been used since then to eliminate scaling. Because a large amount of water is evaporated in the scrubber, no wastewater has been discharged from the plant, even with the use of fresh water for washing. Nearly trouble-free operation has been achieved since April 1976. SO₂ concentration of the gas is about 1,400 ppm at the inlet and 100 ppm at the outlet. The by-product gypsum contains 9 to 12 percent moisture after being centrifuged.

The Okazaki plant, Unitika Co., with a capacity of treating 220,000 Nm³/hr of flue gas from an oil-fired boiler, went into operation on January 6, 1976. In this plant,

limestone is added to the scrubber and less lime is added to the reactor. The ratio of limestone to lime is 3.5 to 1. Problems similar to those at the Saidaiji plant were encountered at the beginning of the operation but were solved fairly easily. Operability of the plant (FGD system operating hours as a percent of boiler operating hours) from start-up until the beginning of May 1976 was 95 percent. SO_2 concentration is about 1,300 ppm at the inlet and 100 ppm at the outlet.

In both plants, small amounts of sulfuric acid have been added to the slurry prior to the oxidation to adjust the pH of the slurry to 5. Sulfuric acid may not be needed when the SO_2 concentration is over 2,000 ppm at the inlet and over 200 ppm at the outlet (as in the U.S.), because the slurry pH can be reduced to below 5 by the scrubbing. The multi-venturi scrubber has limited application. For a plant larger than $400,000 \text{ Nm}^3/\text{hr}$, Kawasaki may use the Bischoff scrubber developed in Germany.

MKK LIME-GYPSUM PROCESS USING JET SCRUBBER

Mitsubishi Kakoki Kaisha (MKK) has constructed two lime-gypsum process systems using a screen-type scrubber. Operation is not yet completely successful. MKK recently completed a new lime-gypsum process system at Naoshima Smeltery, Mitsubishi Metal, with a capacity of treating

120,000 Nm³/hr of tail gas from a sulfuric acid plant using a jet scrubber, which is a simple structure, as shown in Figure 6-2. The inside wall is kept washed with the absorbent slurry so that no scaling takes place. The tail gas, containing 1,000 to 3,500 ppm SO₂, is cooled in a cooler and led to the scrubber. A lime slurry is sprayed from the top by a specially designed nozzle at an L/G ratio of 12 to 15 liters/Nm³; the slurry removes 90 to 95 percent of the SO₂. Calcium sulfite formed by the reaction is oxidized to gypsum in the scrubber by oxygen in the gas; therefore, there is no need for an oxidizer. Gypsum is centrifuged. Most of the filtrate is returned to the scrubber after lime is added. A small portion of the filtrate is sent to a wastewater treatment system together with the discharge from the cooler. The pressure drop in the scrubber is about 80 mm H₂O. A flow-sheet of the MKK lime-gypsum jet-scrubber process installed at the Naoshima Smelter is provided in Figure 6-3.

DOWA ALUMINUM SULFATE PROCESS

Process Description⁹

Dowa Mining Co. has developed an indirect limestone process using an aluminum sulfate solution at about pH 4 as the absorbent. A flowsheet of the process used at the Okayama plant, Dowa Mining, is shown in Figure 6-4. The plant has two units to treat tail gas from two sulfuric acid plants. The capacity of each unit is 150,000 Nm³/hr.

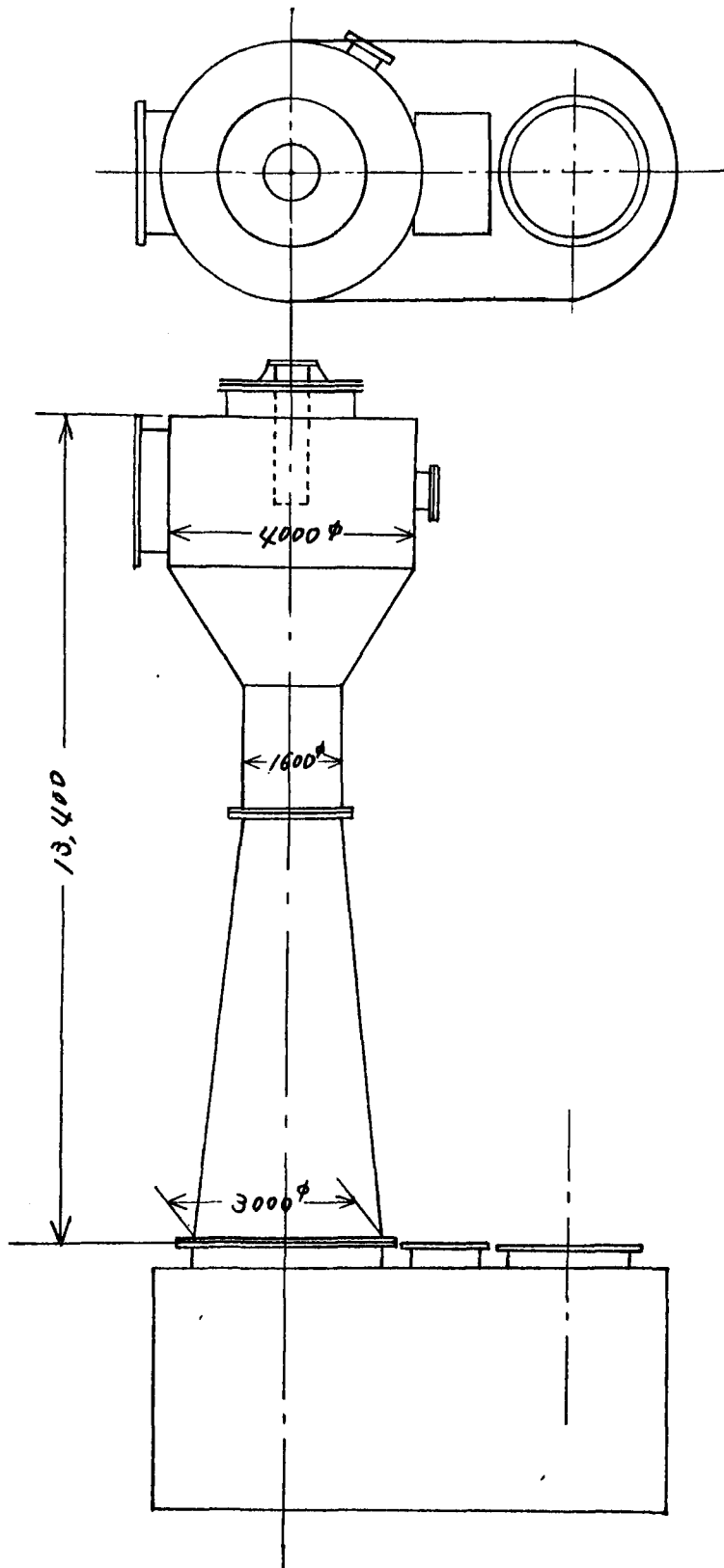


Figure 6-2 Dimensions of jet scrubber(mm)
(120,000 Nm³/hr)

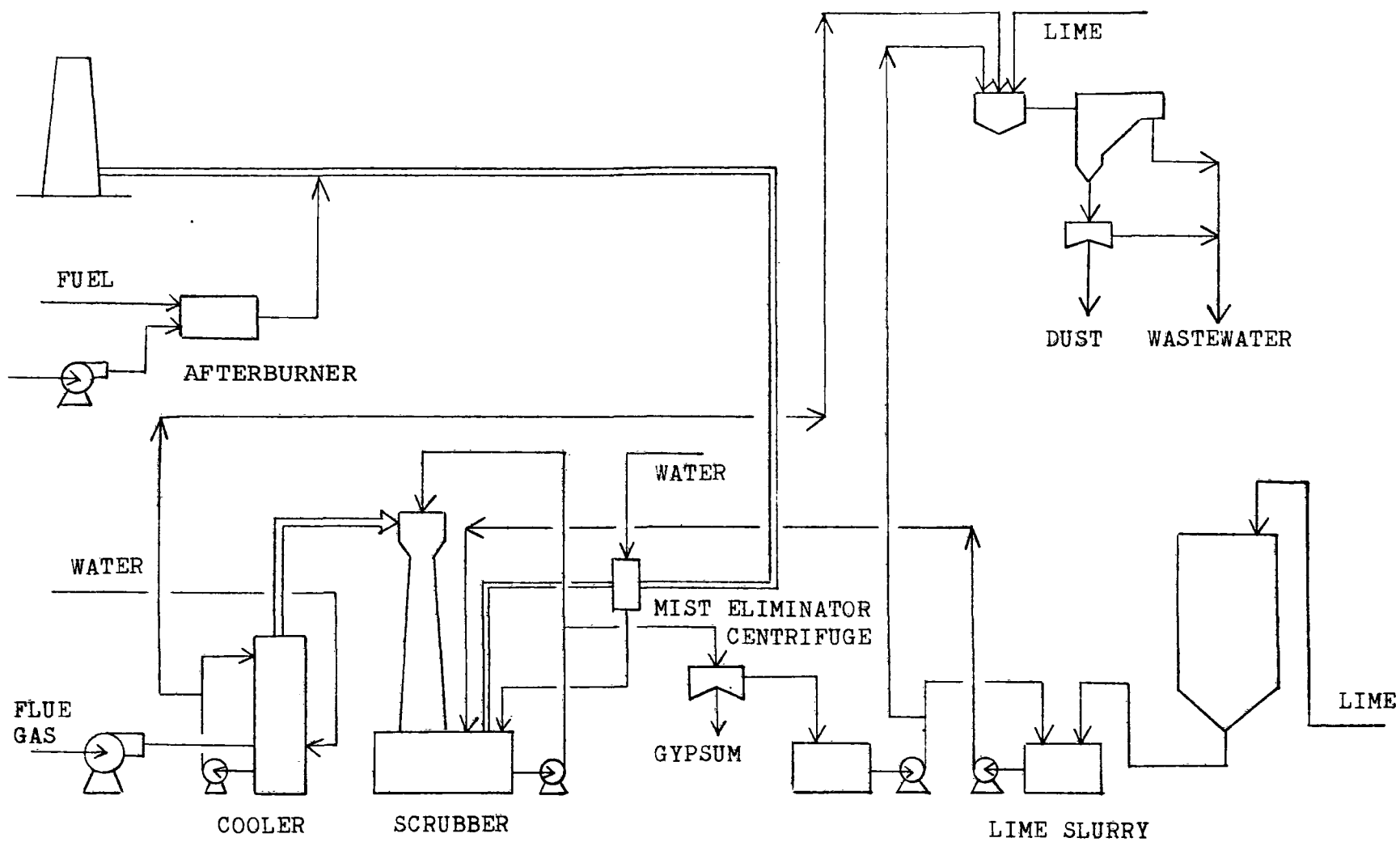


Figure 6-3 Flowsheet of MKK jet-scrubber process (Naoshima plant)

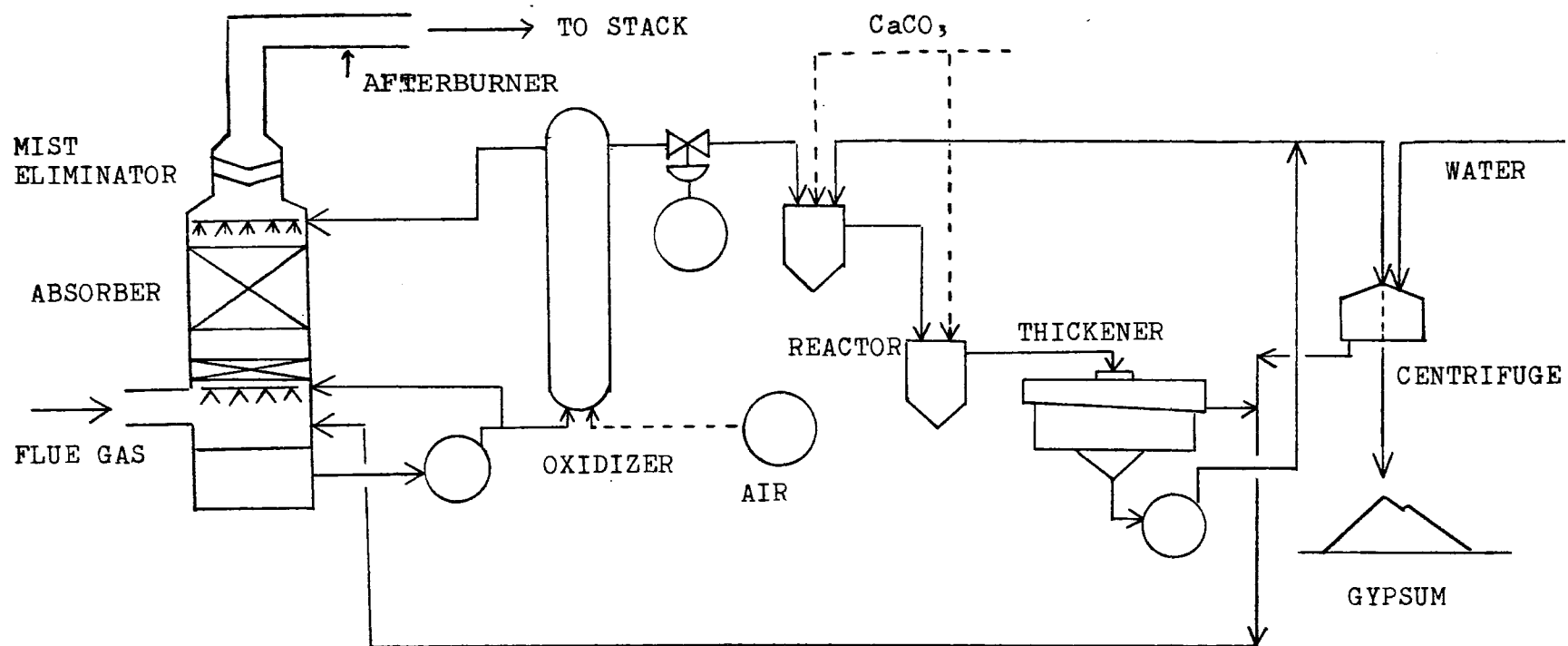
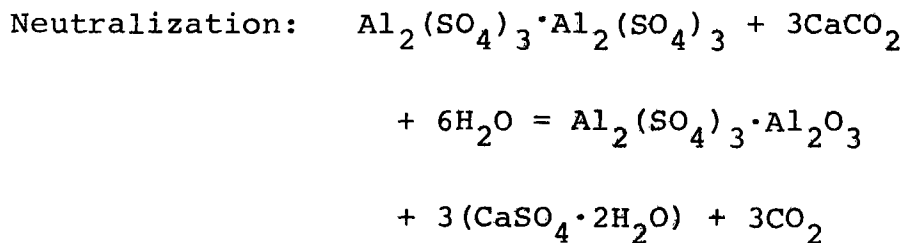
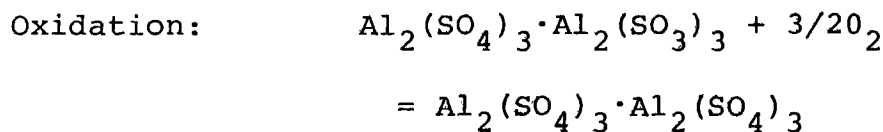
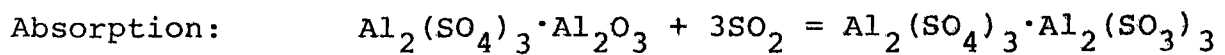


Figure 6-4 Flowsheet of Dowa aluminum sulfate-limestone process

Waste gas is led into a packed-tower absorber. SO_2 is absorbed in a solution of basic aluminum sulfate, $\text{Al}_2(\text{SO}_4)_3 \cdot \text{Al}_2\text{O}_3$, of pH 3 to 4 to form $\text{Al}_2(\text{SO}_4)_3 \cdot \text{Al}_2(\text{SO}_3)_3$. The liquor is oxidized with air into $\text{Al}_2(\text{SO}_4)_3$, which is then treated with powdered limestone to precipitate gypsum and to regenerate the basic aluminum sulfate solution. The gypsum is centrifuged, and the liquor and wash water is returned to the absorber.



The nature of the absorbing liquor is indicated in Figures 6-5 to 6-8. The basicity is the ratio of uncombined Al_2O_3 to total Al_2O_3 . An optimum concentration as well as basicity of the absorbing liquor is selected according to the SO_2 concentration of the gas and the removal efficiency required. Four plants are in operation (Table 3-6).

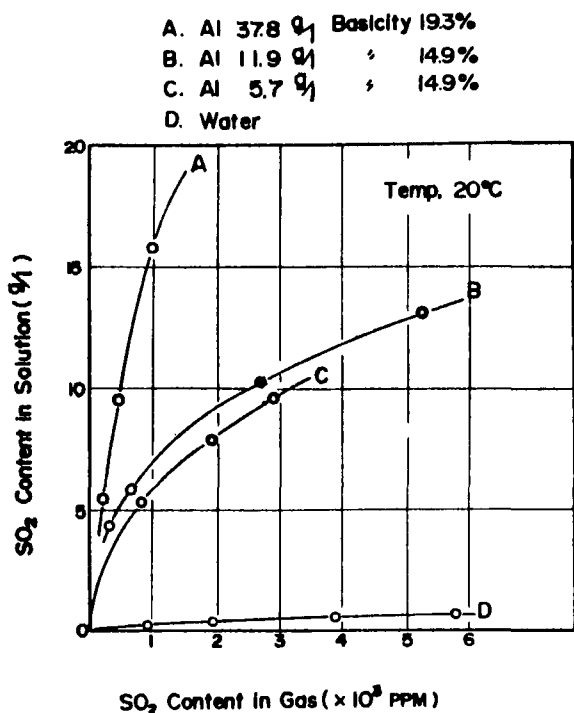


Figure 6-5 Solubility curves of SO₂ in BAS solutions

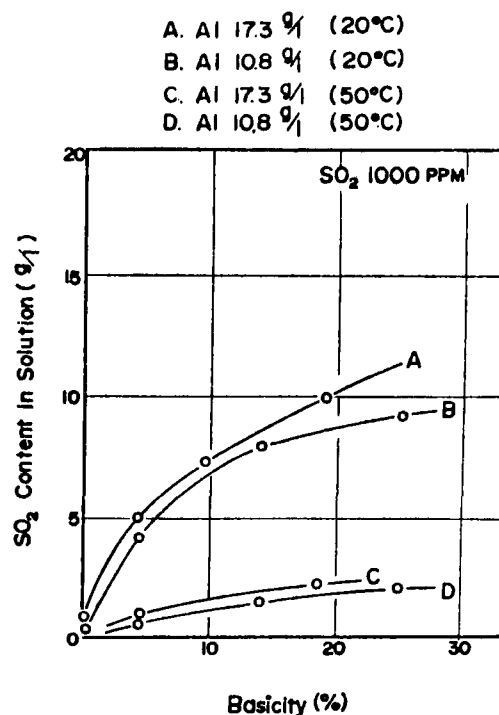


Figure 6-6 Solubility curves of SO₂ at various temperatures of the solution

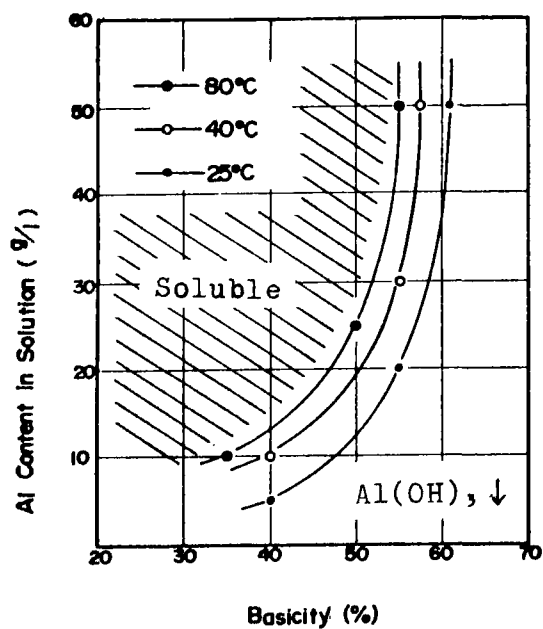


Figure 6-7 Soluble range of aluminum compound

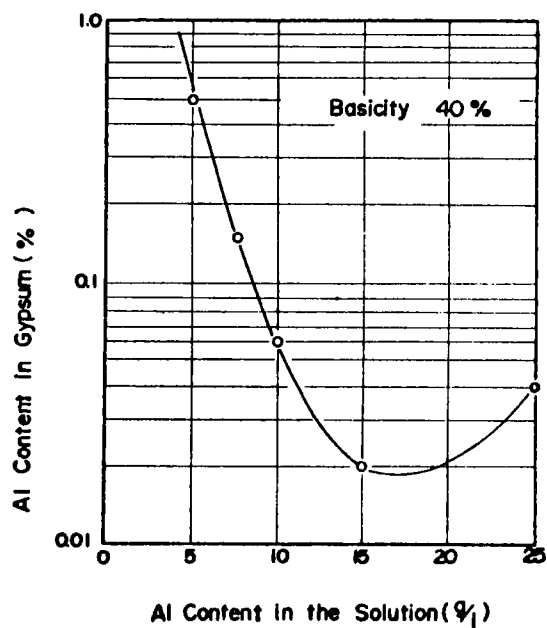


Figure 6-8 Relationship of Al loss to concentration of solution

Okayama Plant

The operation parameters of the Okayama plant are shown in Table 3-5. An L/G ratio of 2.5 to 3 is used to reduce SO₂ from 500 to 600 to 10 to 20 ppm with a packed height of 2 to 4 meters. The plant has been in continuous operation since its start-up in 1974, except for the period of shut-down of the acid plant.

No wastewater was purged for over a year and magnesium accumulated in the liquor. Although magnesium does not interfere with SO₂ removal, its concentration should be kept below a certain level to yield good-quality gypsum for wallboard production. A small additional unit has been installed to eliminate the magnesium. A small portion of the absorbing liquor is neutralized with an excessive amount of limestone to precipitate aluminum hydroxide and is then sent to a settler. Overflow from the settler, containing the magnesium, is discarded. The underflow, containing aluminum hydroxide and limestone, is sent to the reactor.

Requirements for the Okayama plant for treating 280,000 Nm³/hr of tail gas containing 650 ppm SO₂ are as follows:

CaCO ₃	0.81 t/hr
Al ₂ (SO ₄) ₃ solution (Al ₂ O ₃ 8%)	15 kg/hr
Water	7 t/hr
Electric power	1,000 kW
Operator	One per shift

A small amount of aluminum is contained in the gypsum but this does not affect the quality of wallboard or cement produced from the gypsum. Consumption of aluminum is about 0.5 kg (as Al) for every ton of gypsum.

Tamano Plant, Naikai Salt Production Co.

The Tamano plant, Naikai Salt Production Co., has a capacity of treating 80,000 Nm³/hr of flue gas from an oil fired boiler containing 1,500 ppm SO₂. This plant went into operation in March 1976 and has been in trouble-free continuous operation since then. The system has a gas cooler before the absorber. A portion of the liquor discharged from the absorber is sprayed in the cooler. It has been found that about 70 percent of the SO₂ is removed in the cooler. Operation data for the first month are shown in Figure 6-9.

At the beginning of operation, the SO₂ removal ratio ranged from 83 to 93 percent. A small amount of a soluble metallic catalyst was then added to improve the removal ratio. Since two weeks after start-up, when the gas volume increased to nearly full load and the SO₂ concentration to nearly 1,500 ppm, the removal ratio has been kept above 95 percent. The L/G ratio has been 10 for the scrubber. Pressure at the cooler inlet was about 100 mm H₂O at full load.

No wastewater has been purged. In case of wastewater purge, the catalyst can be recovered by neutralizing the liquor.

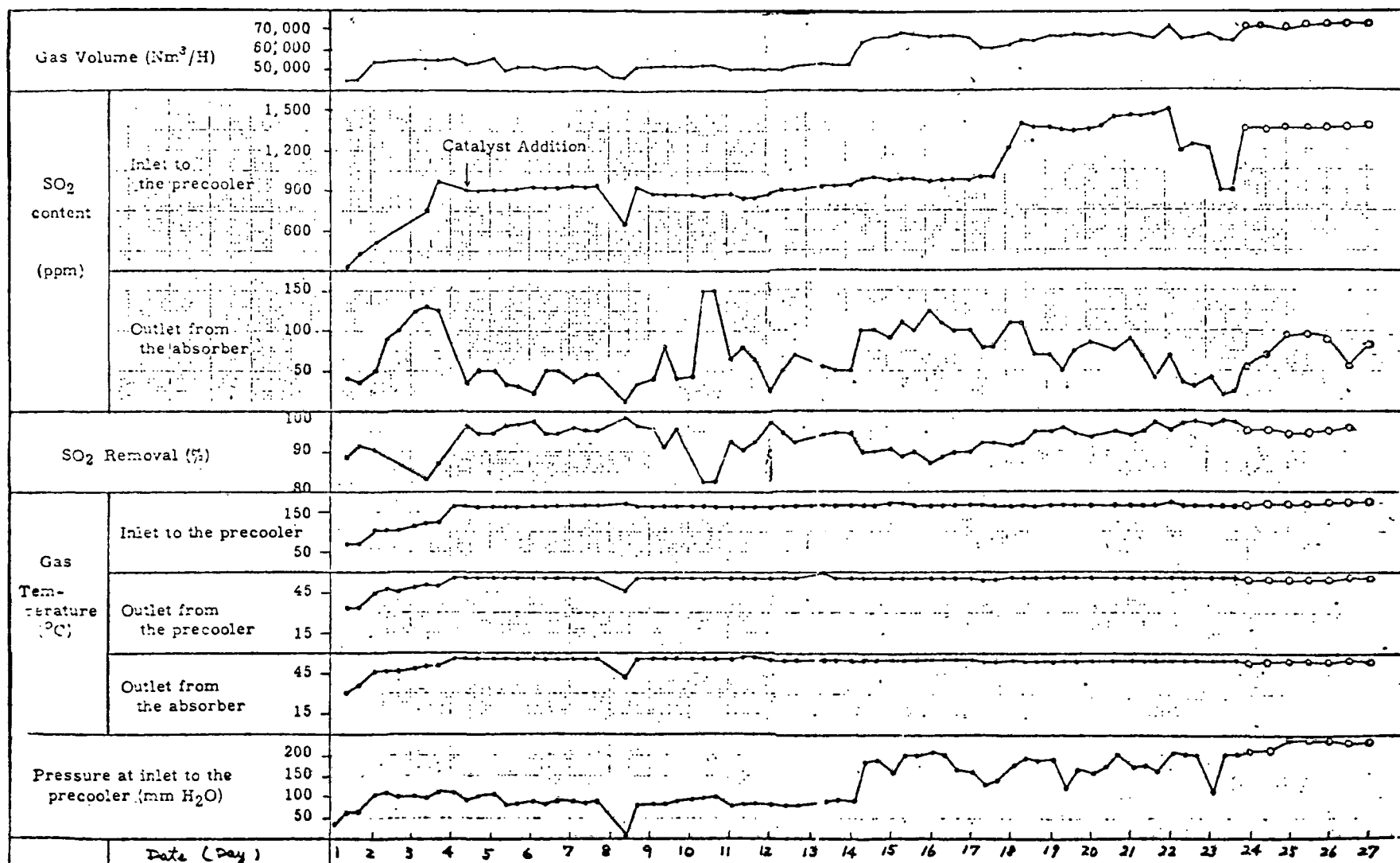
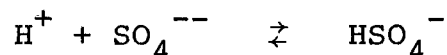


Figure 6-9. Operation data of Tamano plant, Naikai.

KURABO AMMONIUM SULFATE-LIME PROCESS²

In order to prevent plume formation, which is a common problem in ammonia scrubbing, Kurabo Industries has developed a process using as the absorbent a slightly acidic ammonium sulfate solution at pH 3 to 4. Plume formation can be eliminated by the use of the acidic absorbing liquor because the vapor pressure of NH_3 is less than 1 ppm equivalent with a solution at a pH lower than 4. The acidic ammonium sulfate solution has a greater capacity for SO_2 absorption than plain water or a saturated calcium sulfate solution because of the smaller pH drop due to the following equilibrium in ammonium sulfate solution:



A flowsheet of the process is shown in Figure 6-10. Flue gas is first led into a KBCA scrubber and then into a packed tower absorber. The main function of the KBCA scrubber is to cool the gas to 60°C and to concentrate the absorbing liquor (ammonium sulfate solution). More than 90 percent of the SO_2 is removed. The liquor from the packed tower absorber is sent to the KBCA unit, concentrated, and then led into an oxidizer. The pH of the liquor in the oxidizer is adjusted to 3 or 4 by adding dilute aqua ammonia; the sulfite in the liquor is oxidized to sulfate by small bubbles of air formed by introducing a jet stream of circulating liquor accompany air into a pool of the liquor.

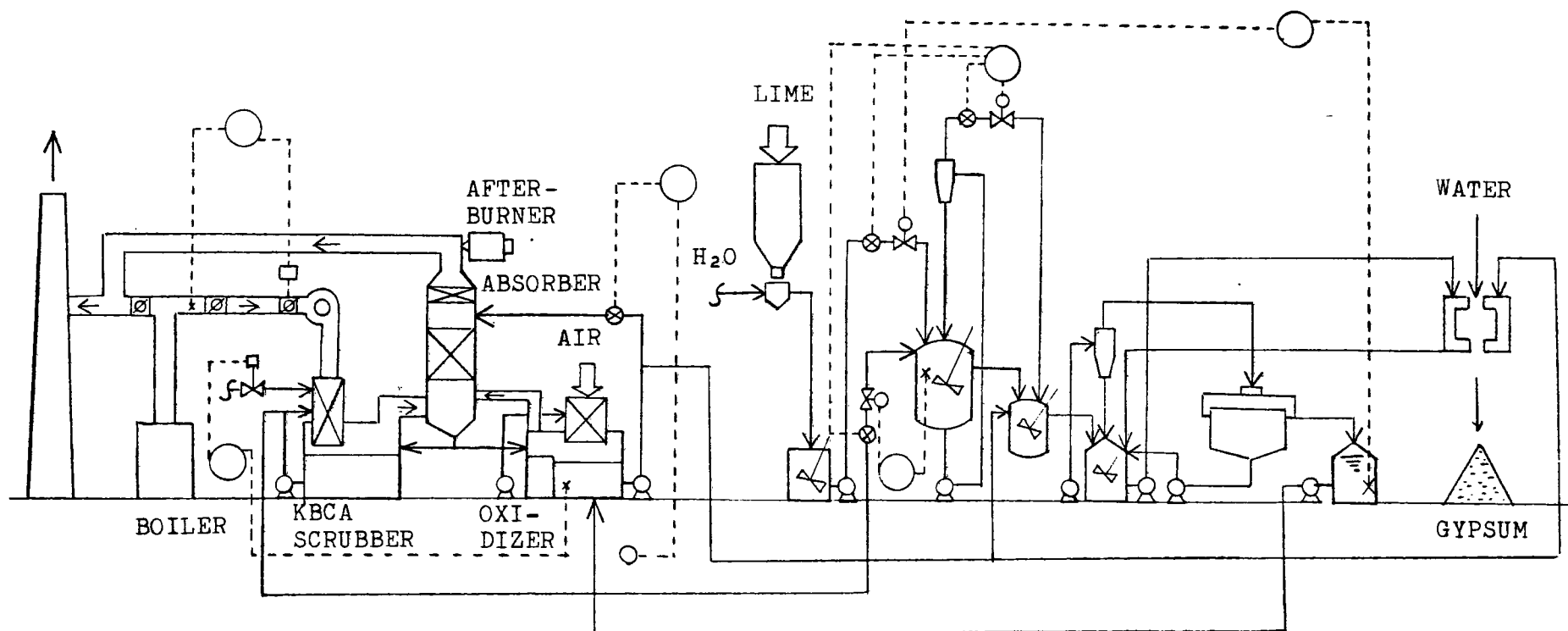


Figure 6-10 Flowsheet of Kurabo ammonium sulfate-lime process

About five times the stoichiometric amount of air is used.

Most of the liquor from the oxidizer is returned to the absorber, and a portion is sent to a set of three reactors, in which the liquor is treated with milk of lime to precipitate gypsum. The liquor from the centrifuge and the wash water are sent to an aqua ammonia tank, and the aqua ammonia is sent to the oxidizer.

Five commercial plants are in operation (Table 3-6). Operation parameters are shown in Table 3-5. The plume is almost invisible.

Ammonium sulfate solution (about 15% concentration) has been produced in the small commercial plant of Taki Chemical and used as the promoter of gypsum hardening for wallboard production. The plant has a capacity of treating 15,000 Nm³/hr of flue gas containing 1,000 ppm SO₂ from an oil-fired boiler. More than 98 percent SO₂ removal has been attained with a 4.5 m packed height and an L/G ratio of about 10. The plant has been in operation since early April 1976 with 100 percent operability.

In by-production of solid ammonium sulfate, a 1.0 to 1.5 mole/liter solution may be used for absorption and concentrated in the KBCA scrubber to about a 3 mole/liter solution (nearly saturation).

KUREHA SODIUM ACETATE PROCESS¹⁰

Kureha Chemical has developed an indirect lime/limestone process using a sodium acetate solution as the absorbent and has operated a pilot plant with a capacity of treating 5,000 Nm³/hr of flue gas from an oil-fired boiler. The reason for the use of the acetate is to eliminate the problem of sodium sulfate, an undesirable compound that forms in sodium scrubbing processes and must be treated. In this process the acetic acid reacts with calcium to form calcium acetate, which is soluble and readily reacts with sodium sulfate to precipitate gypsum and to regenerate sodium acetate. A flow-sheet of the process is shown in Figure 6-11.

SO₂ in flue gas is absorbed by a sodium acetate solution to form sodium bisulfite and acetic acid. Acetic acid is vaporized and caught, together with the remaining SO₂, by a limestone slurry at the upper part of the absorber to form a calcium acetate solution. Sodium sulfite is oxidized by air into sodium sulfate, which is then reacted with calcium acetate as mentioned above. Operation parameters are shown in Table 3-5.

Plant operation is easy, with more than 99 percent SO₂ recovery. Losses of acetic acid and sodium can be kept very small. On the other hand, the absorber and reactors are large, and the L/G ratio is fairly high.

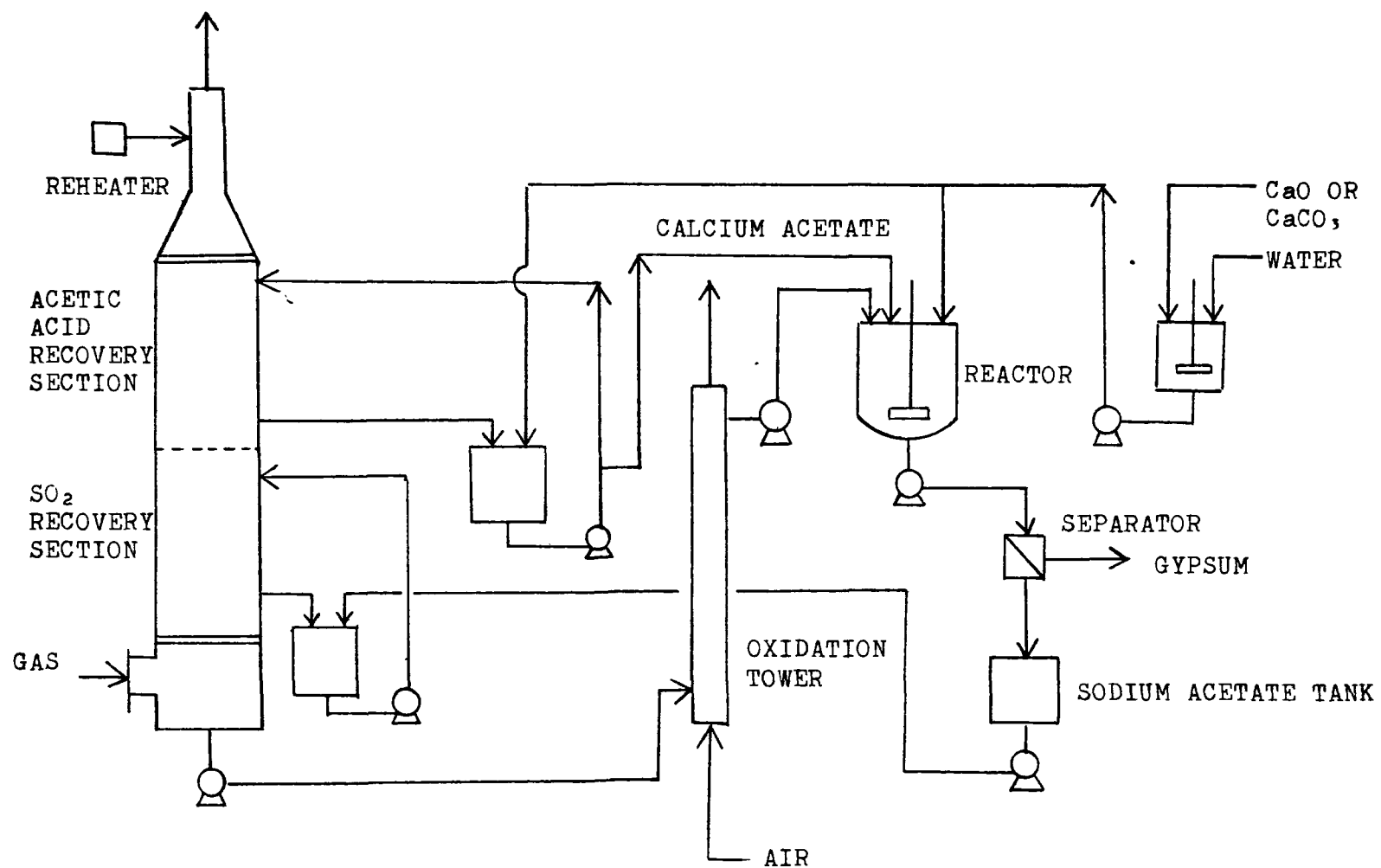


Figure 6-11. Flowsheet of Kureha sodium acetate-lime/limestone process

Kureha recently started to use lime in place of limestone in order to reduce the size requirements and the L/G ratio substantially. Process economy is improved by the use of lime. For plants which do not need very high SO₂ removal efficiencies, 70 to 80 percent of the gas may be treated by the process and then mixed with the untreated hot gas to eliminate reheating of the gas. The sodium acetate process is useful also for simultaneous removal of NO_x (Section 7).

mitsui-chemico magnesium process

Mitsui Miike Machinery Co. constructed a magnesium scrubbing system with a capacity of treating 500,000 Nm³/hr of waste gas from industrial boilers and Claus furnaces at Chiba refinery, Idemitsu Kosan. The unit went into operation in late 1974. The flowsheet is shown in Figure 6-12.

Two Chemico venturi scrubbers in the same shell are used (Photo 6-1) as at the Omuta plant, Mitsui Aluminum. The magnesium sulfite slurry discharged from the scrubber is pH-adjusted and filtered. The sulfite cake is dried in a rotary drier with countercurrent flow. The dried sulfite is calcined in an oil-fired rotary kiln (Photo 6-2). As 10 to 15 percent oxidation of the sulfite occurs in the scrubber and regeneration section, coke is added to the kiln to reduce the sulfate. The regenerated MgO goes to a slaker and the slaked product is wet-milled to increase activity.

Figure 6-12 Flowsheet of Chemico-Mitsui magnesium process



Photo 6-1. Chiba plant, Idemitsu Kosan
(170 MW equivalent, scrubber and kiln)

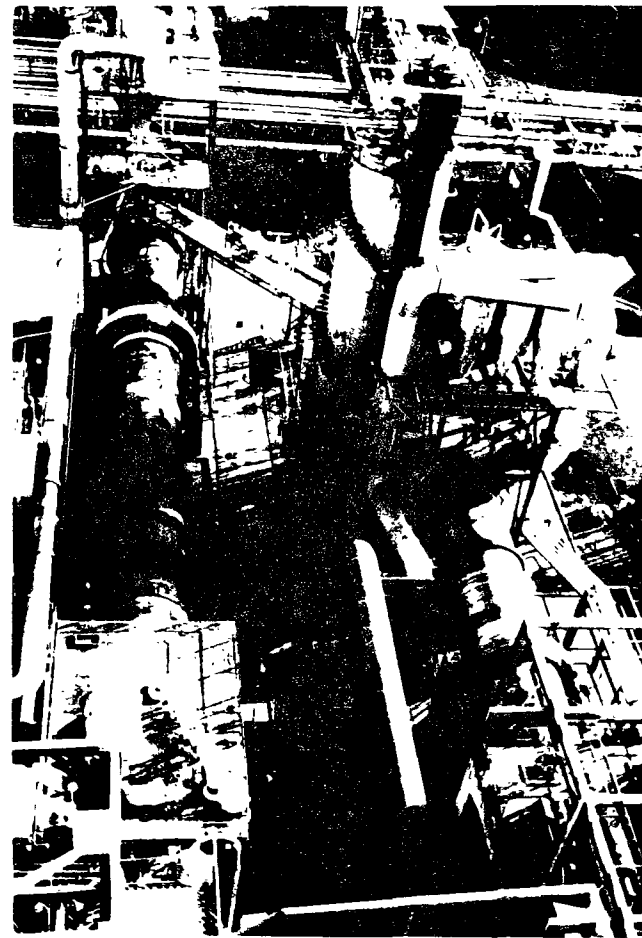


Photo 6-2. Chiba plant, Idemitsu Kosan
(View from the top of the scrubber,
drier at right, kiln at left)

Gas from the kiln goes through a cyclone, wet venturi, and wet precipitator. The cleaned gas containing 10 to 12 percent SO_2 is fed to the Claus furnace, where H_2S recovered in the refinery is reacted with SO_2 to produce elemental sulfur. SO_2 concentration at the scrubber inlet is about 2,500 ppm, and the removal efficiency is 95 to 97 percent.

The form of magnesium sulfite, trihydrate or hexahydrate, is an important key to plant operation. Usually hexahydrate is preferred because it grows in much larger crystals than does trihydrate. Mitsui Miike has found that mixing the two forms gives the best results. There has been no problem with drier operation. The major problem was dislodging of the firebricks in the rotary kiln, caused possibly by the cooling of the kiln by rain. To prevent corrosion, the chloride concentration in the liquor is maintained below 2,300 ppm. Usually no wastewater is purged because the inlet chloride concentration is low.

HITACHI-UNITIKA ACTIVATED CARBON PROCESS

Unitika Co. has installed at Uji an activated carbon process with a capacity of treating $170,000 \text{ Nm}^3/\text{hr}$ of flue gas from an industrial boiler burning oil with 2.5 percent sulfur. The adsorption-desorption unit was designed by Hitachi Ltd, which had constructed a larger unit at Kashima Station, Tokyo Electric, where the dilute sulfuric acid obtained by water wash of the carbon is reacted with limestone

to produce gypsum of good quality. The Kashima plant is fairly costly, but operation has been trouble-free for 3 years with virtually no loss of carbon.

At the Uji plant, Unitika, which went into operation in December 1975, dilute sulfuric acid (6 to 7 percent H_2SO_4) is sprayed into the incoming hot flue gas at 170°C. The gas is cooled to 80 to 90°C, and the acid is concentrated to about 60 percent. The acid is somewhat dirty and too dilute for commercial use. Unitika has used the acid in its own chemical plant and hopes to sell it locally for wastewater treatment. The acid concentration unit was designed by Unitika. There is no mist eliminator between the concentrator and the absorber. The acid mist is caught by the carbon in a fixed bed.

7. SIMULTANEOUS REMOVAL OF SO_2 AND NO_x

OUTLINE

In Japan, NO_x removal technology has developed rapidly since 1973, when a stringent ambient standard for NO_2 (0.02 ppm in daily average) was set forth. Among many processes, selective catalytic reduction of NO_x by ammonia at about 400°C has been considered a most feasible means to remove 85 to 95 percent of the NO_x in flue gas. Several commercial plants are treating flue gas from oil-fired industrial boilers (200,000-450,000 Nm^3/hr), and many plants are under construction using the reduction process.¹¹ This dry denitrification process, however, is not suited for use in conjunction with FGD (for which a wet process is economical), because it requires a large heat exchanger and a considerable amount of energy for heating (No. 3, Figure 7-1) or an expensive hot electrostatic precipitator (No. 4, Figure 7-1). Moreover, many existing boilers have not enough space for installation of both FGD and denitrification units. For these reasons, many dry and wet processes have been developed for simultaneous removal of SO_2 and NO_x (No. 1 and 2, Figure 7-1), and many plants are in operation, as shown in Table 7-1.

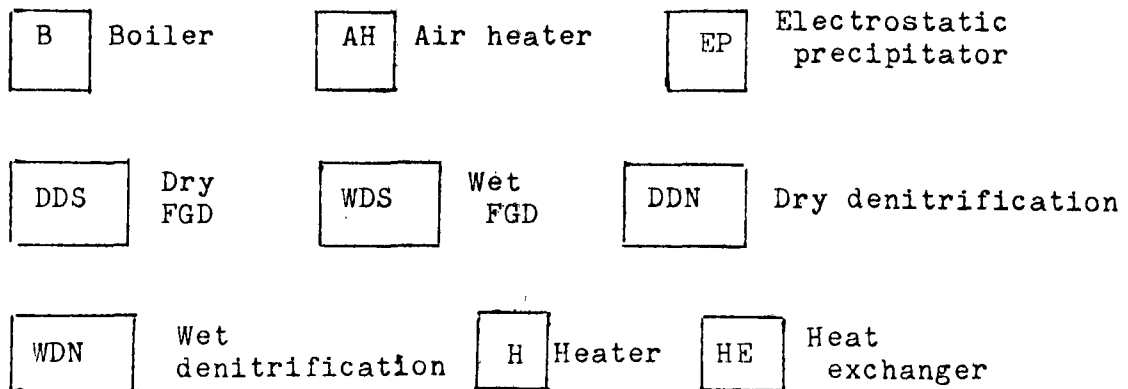
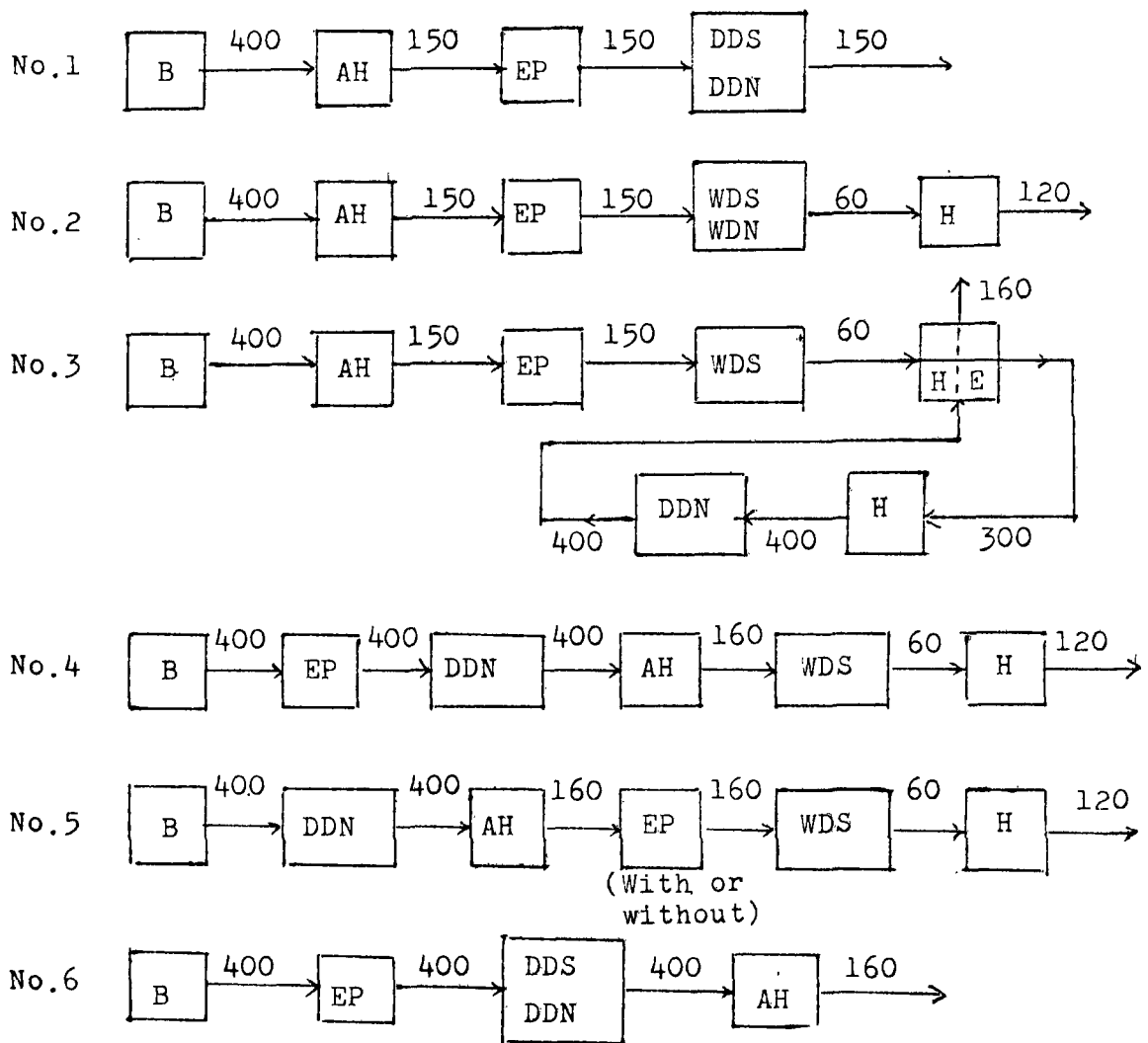


Figure 7-1 Models of combination of FGD and denitrification
(Figures show gas temperature)

Table 7-1. PROCESSES FOR SIMULTANEOUS REMOVAL OF SO₂ AND NO_x

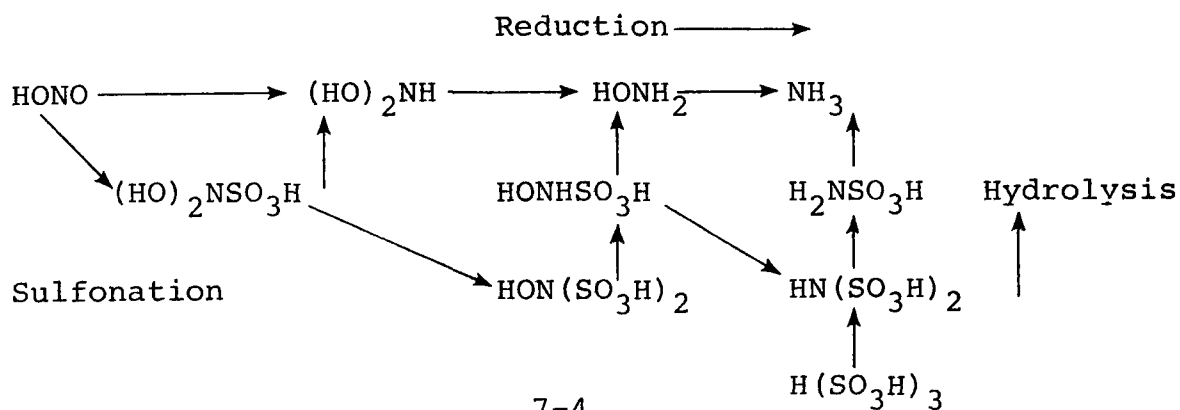
Process developer	Type of process	Plant owner	Plant site	Capacity, Nm ³ /hr	Source of gas	Completion	By-product
(Sumitomo Metal Fujikasui)	Oxidation reduction	Sumitomo Metal	Amagasaki	62,000	Boiler ^a	1973	(NaNO ₃ , NaCl, Na ₂ SO ₄)
(Sumitomo Metal Fujikasui)	Oxidation reduction	Toshin Steel	Fuji	100,000	Furnace	1974	(NaNO ₃ , NaCl, Na ₂ SO ₄)
(Sumitomo Metal Fujikasui)	Oxidation reduction	Sumitomo Metal	Osaka	39,000	Boiler ^a	1974	(NaNO ₃ , NaCl, Na ₂ SO ₄)
Osaka Soda	Oxidation reduction	Osaka Soda	Amagasaki	60,000	Boiler ^a	1976	(NaNO ₃ , NaCl, Na ₂ SO ₄)
Shirogane	Oxidation reduction	Mitsui Sugar	Kawasaki	48,000	Boiler ^a	1976	(NaNO ₃ , NaCl, Na ₂ SO ₄)
Chiyoda	Oxidation reduction	Chiyoda	Kawasaki	1,000	Boiler ^a	1973	Gypsum, Ca(NO ₃) ₂
Mitsubishi H.I.	Oxidation reduction	Mitsubishi H.I.	Hiroshima	2,000	Boiler ^a	1974	Gypsum, NH ₃
Ishikawajima H.I.	Oxidation reduction	Ishikawajima	Yokohama	5,000	Boiler ^a	1975	Gypsum, N ₂
Kureha Chemical	Reduction	Kureha Chem.	Nishiki	5,000	Boiler ^a	1975	Gypsum, N ₂
Chisso Eng.	Reduction	Chisso P.C.	Goi	300	Boiler ^a	1974	(NH ₄) ₂ SO ₄
Mitsui S.B.	Reduction	Mitsui S.B.	Chiba	150	Boiler ^a	1974	H ₂ SO ₄ , N ₂
Asahi Chemical	Reduction	Asahi Chem.	Mizushima	600	Boiler ^a	1974	Gypsum, N ₂
Kawasaki H.I.	Magnesium	EPDC	Takehara	5,000	Boiler ^b	1975	Gypsum, Mg(NO ₃) ₂
Unitika Co.	Carbon ^c	Union Glass	Hirakata	5,000	Furnace	1975	Conc. SO ₂ , N ₂
Sumitomo H.I.	Carbon ^c	Sumitomo Metal	Kokura	150	Furnace	1975	Conc. SO ₂ , N ₂
Shell	Copper ^c	Showa Y.S.	Yokkaichi	120,000	Boiler ^a	1975	Conc. SO ₂ , N ₂
Ebara-JAERI	Electron beam ^c	Ebara	Fujisawa	1,000	Boiler ^a	1974	Mist, dust

^a Oil-fired boiler.^b Coal-fired boiler.^c Dry process.

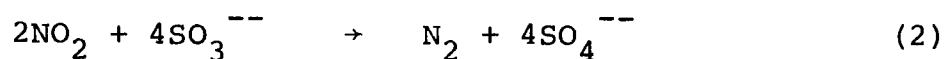
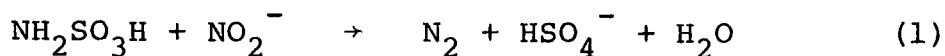
CHEMISTRY AND PROBLEMS OF WET PROCESSES

Most of the NO_x in combustion gas is in the form of NO, which has little reactivity and does not readily dissolve in solutions. NO can be absorbed in a solution containing an oxidizing agent such as potassium permanganate, hydrogen peroxide, or calcium hypochlorite; an absorption oxidation process may be too expensive for commercial use on a large scale, however, particularly when the gas contains much SO_2 , which consumes the oxidizing agent. Moreover, treatment of the by-product liquor containing a nitrate, nitrite, and sulfate will present a problem.

NO is slowly oxidized into NO_2 in air. Gaseous oxidizing agents, ozone and chlorine dioxide, oxidize NO very rapidly, within 1 second, but they hardly oxidize SO_2 into SO_3 . Since 1972 several oxidation reduction processes have been developed in Japan, by which NO is first oxidized to NO_2 by the oxidizing agent and is absorbed in a solution or slurry together with SO_2 . Various reactions occur in the solution or slurry as shown below, resulting in the reduction of NO_x by SO_2 (or sulfite) to N_2 or NH_3 .¹²



The reactions to form N_2 are more complex but may be described as in equation (1) or more simply as in equation (2).



In some of the processes a considerable portion of NO_x remains in the resulting liquor as a nitrite and nitrate. For a large-scale operation, it is desirable not to have those nitrogen compounds in solution because they present problems in wastewater treatment.

In addition to the oxidation reduction processes, several reduction processes have been developed using solutions containing ferrous ion and EDTA (ethylenediamine-tetraacetic acid, a chelating compound whose present cost in Japan is about \$2,700/t), which absorbs NO fairly well. The absorbed NO reacts with sulfite and is converted to N_2 or $(NH_4)_2SO_4$. Otherwise, the NO can be regenerated in a concentrated form.

The wet simultaneous processes have not yet been commercialized on a large scale; five relatively small commercial plants are in operation, as shown in Table 7-1. The major problem for the oxidation reduction processes has been the high cost of ozone (\$1.2 to \$1.4/kg). Although chloride dioxide is less expensive, it brings chloride into the system and complicates the process. Another problem has

been the formation of considerable amounts of nitrate and nitrite in the absorbing liquor. Improvements have been made to prevent the nitrogen compounds from remaining in the liquor as well as to produce ozone at lower cost. The major problems for the reduction process have been the requirements of expensive EDTA, a large absorber, and a high L/G ratio. Efforts have been made to minimize the consumption of EDTA.

On the positive side, the wet processes offer the following advantages: (1) They are not affected by particulates in the gas as are the dry processes using a catalyst. (2) Their NO_x removal efficiency exceeds 80 percent, whereas that of the ammonia injection process without catalyst may not reach 70 percent on a large scale. (3) The wet processes do not consume ammonia. Some of the processes can produce ammonia from NO_x . For future worldwide use, the wet processes may be practicable if the process economy can be considerably improved, because the dry processes consume large amounts of ammonia and may cause shortages of nitrogen fertilizers and foods. (4) A high SO_2/NO_x ratio is favorable to the reduction of NO_x to N_2 or NH_3 . The wet processes may be useful for treatment of flue gas burning of high-sulfur coal.

OXIDATION REDUCTION PROCESSES

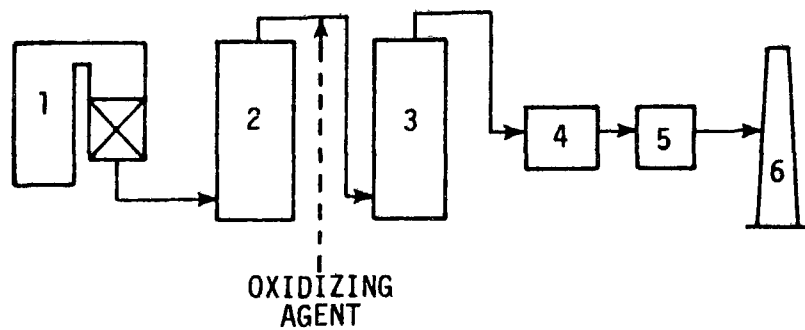
Fujikasui-Sumitomo Process (Moretana Process)

Fujikasui Engineering, jointly with Sumitomo Metal Industries, has developed a sodium scrubbing process for removal of SO_2 and NO_x (Figure 7-2) and has constructed three plants (Table 7-1). Gaseous ClO_2 is added to the gas just before the scrubber and oxidizes NO into NO_2 within 0.5 second. The gas is then introduced into a Moretana scrubber and is reacted with a sodium hydroxide solution. More than 98 percent of the SO_2 is absorbed to produce sodium sulfite. About 90 percent of the NO_x in the gas is removed. About half of the removed NO_x is converted into N_2 by the reaction with sodium sulfite and the rest into sodium nitrate and nitrite.

Capital cost is in the range of \$60 to \$90/kW. Operating cost including depreciation (7 years) is roughly \$32/k1 oil or 7 mil/kWh.

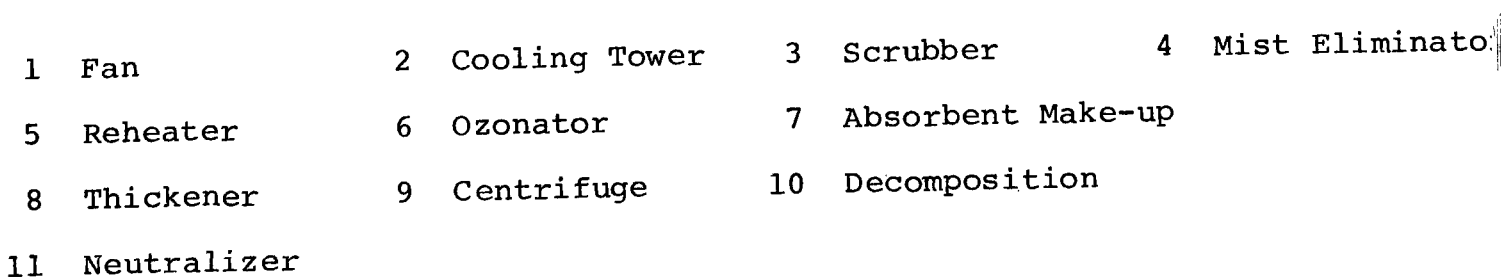
The liquor from the scrubber, which contains sodium sulfate, chloride, nitrate, and nitrite, is concentrated to separate most of the sodium sulfate in a crystal form. The remaining liquor is sent to a wastewater treatment system.

Fujikasui recently started tests on ozone oxidation followed by lime scrubbing to reduce NO_x to N_2 or NH_3 and to by-produce gypsum.



1: BOILER 2: COOLER
3: SCRUBBER 4: MIST ELIMINATOR
5: AFTERBURNER 6: STACK

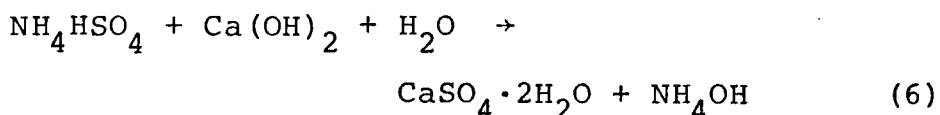
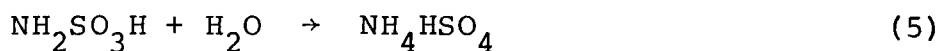
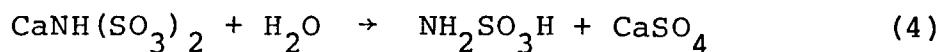
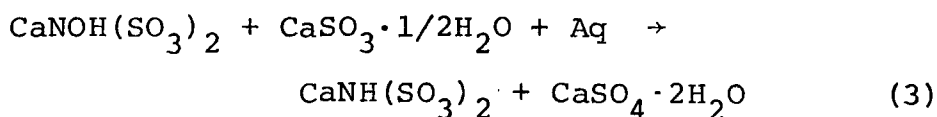
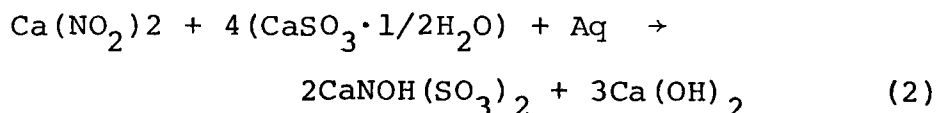
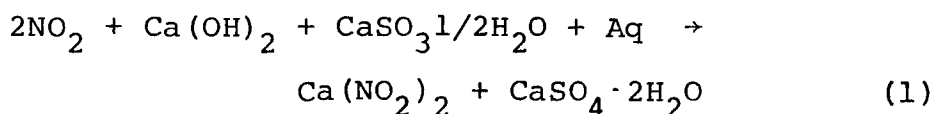
Figure 7-2. Simplified flowsheet of Moretana simultaneous removal process.



7-9

MHI Process

Mitsubishi Heavy Industries (MHI) has modified the wet lime/limestone FGD process for simultaneous removal of NO_x and has operated a pilot plant (Table 7-1, Figure 7-3). Ozone is introduced into the flue gas prior to scrubbing. A water-soluble inorganic catalyst is added to a lime/limestone slurry to promote the reaction of NO_2 . About 80 percent of the NO_x is removed with more than 90 percent of the SO_2 when more than 3 moles of SO_2 per mole of NO_x are present in the gas. The scrubber liquor contains essentially no nitrate or nitrite. A portion of the liquor is treated to decompose N-S compounds to NH_4HSO_4 , which is treated with lime to generate ammonia. The pilot plant has been operated smoothly. Cost of the simultaneous removal is estimated at about 40 percent higher than that of desulfurization only.



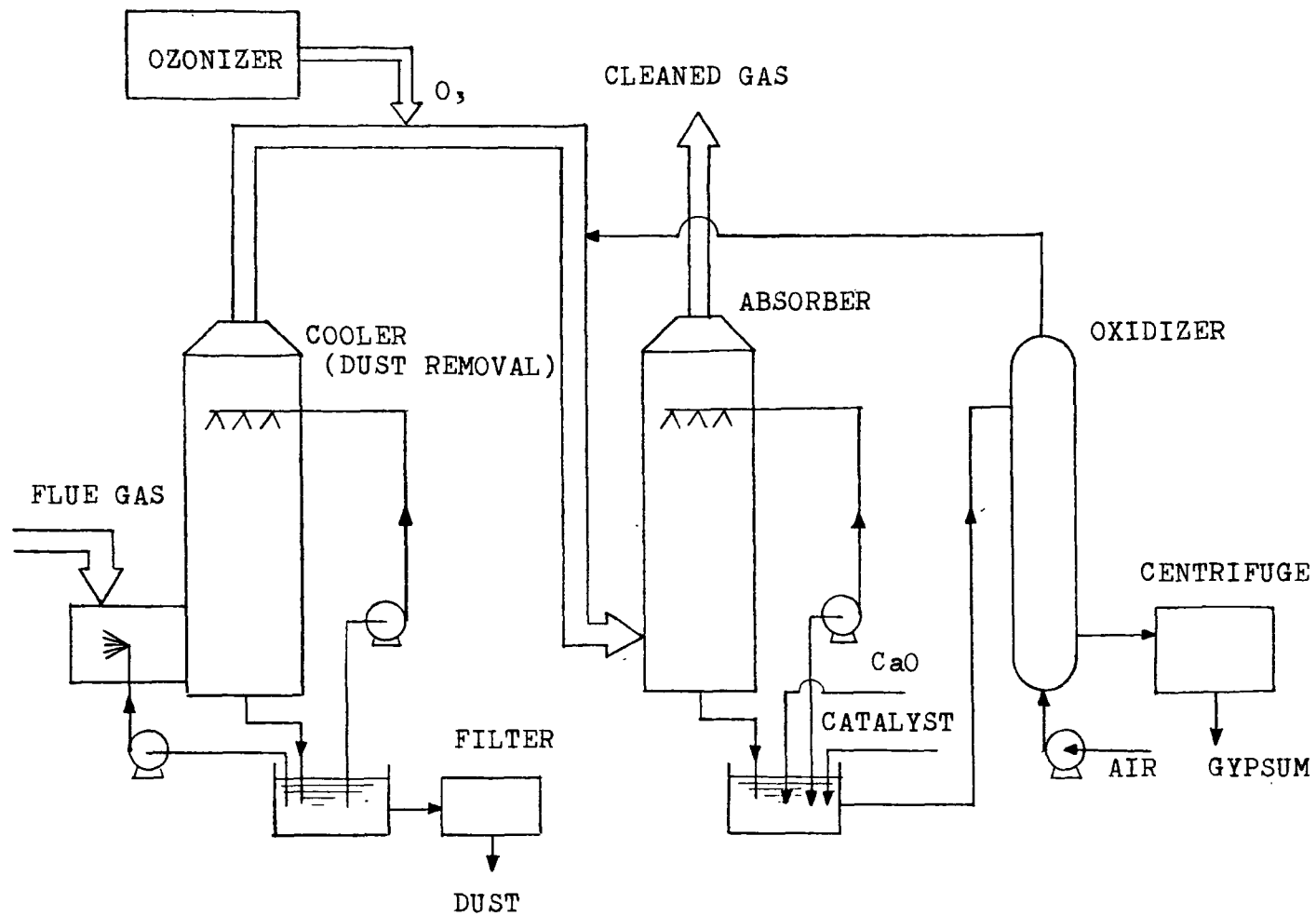
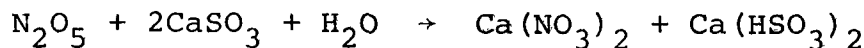
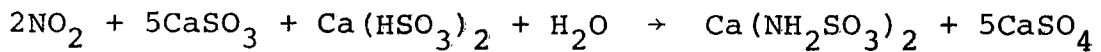
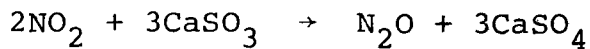
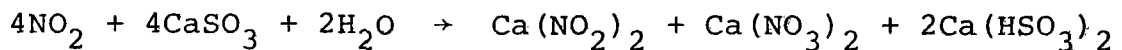
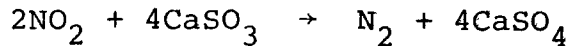


Figure 7-4 Schematic flowsheet of IHI simultaneous removal process

Ishikawajima-Harima Heavy Industries (IHI) has been testing an oxidation reduction process at a pilot plant with a capacity of treating 5,000 Nm³/hr of flue gas from an oil-fired boiler containing about 1,000 ppm SO₂ and 200 ppm NO_x (Figure 7-4).

The flue gas is cooled, injected with ozone to oxidize NO to NO₂, and treated in a scrubber with a lime/limestone slurry at pH 5 to 6, containing small amounts of CuCl₂ and NaCl as catalysts for NO_x absorption. A lower pH is favorable to NO_x absorption. More than 80 percent of the NO_x and 90 percent of the SO₂ are absorbed, resulting in various reactions in the liquor. The following reactions are assumed to take place:



About a half of the NO_x is reduced to N₂, and the rest stays in the liquor as nitrate and other compounds. Tests are in process to achieve further reduction of NO_x to N₂.

Chiyoda Thoroughbred 102 Process

Chiyoda has made a simple modification of the Thoroughbred 101 process to remove NO_x . Ozone is added to the gas prior to scrubbing. More than 60 percent of the NO_x is removed along with about 90 percent of the SO_2 . A portion of the removed NO_x is converted into nitric acid, which forms calcium nitrate, and the rest is converted into N_2 and N_2O . Wastewater treatment is required to remove the nitrate.

Other Oxidation-Reduction Processes

Osaka Soda, a chemical company, has developed a process similar to the Fujikasui-Sumitomo process and has constructed a prototype unit (Table 7-1). Tests on wastewater treatment are in progress.

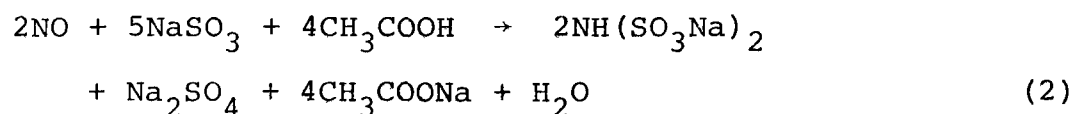
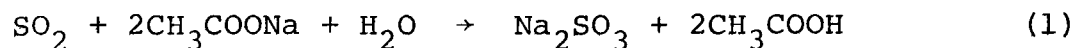
Shirogane Co., an engineering company, has built a unit (Table 7-1) using a process similar to the Fujikasui-Sumitomo process except that ozone is substituted for chloride dioxide. The wastewater containing sodium sulfate and nitrate is sent to a wastewater treatment system along with other wastewaters.

REDUCTION PROCESSES

Kureha Process

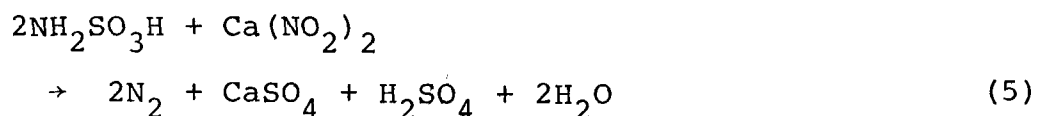
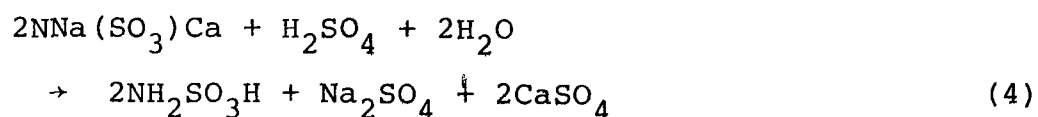
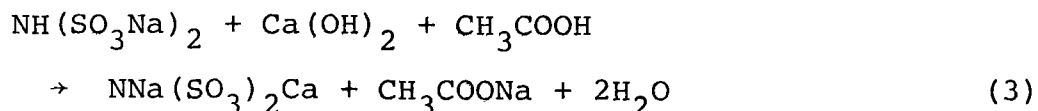
Kureha Chemical has developed a process to remove NO_x in combination with the sodium acetate FGD process (Section 6) (Figure 7-5).

SO₂ is absorbed by a sodium acetate solution to produce sodium sulfite and acetic acid. NO is absorbed by a sodium sulfite solution in the presence of acetic acid and a soluble metallic catalyst to produce sodium imidodisulfonate [reaction (2)].



The remaining sodium sulfite is air-oxidized into sulfate. The sulfate is treated with calcium acetate, as in the FGD process.

Sodium imidodisulfonate is reacted with slaked lime to precipitate and separate sodium calcium imidodisulfonate [reaction (3)], which is then hydralized in the presence of sulfuric acid into sulfamic acid [reaction (4)]. The sulfamic acid is treated with calcium nitrite to release nitrogen [reaction (5)].



Kureha has been operating a pilot plant with a capacity of treating 5,000 Nm³/hr of flue gas from an oil-fired boiler. The process seems fairly complicated, since it includes many reaction steps. Recently the sodium imidodisulfonate has been found useful as a builder of detergents to replace sodium tripolyphosphate, which has been causing eutrophication problems. Tests have been in progress on the effect of the disulfonate on the environment. Possible commercial use of the disulfonate will make the process useful.

Mitsui Shipbuilding Process¹¹

Mitsui Shipbuilding, one of the largest engineering and construction companies, has developed a simultaneous removal process which by-produces concentrated SO_2 which can be used in sulfuric acid production (Figure 7-6).

Flue gas is treated with a ferrous compound solution containing ethylenediaminetetraacetate, which absorbs both SO_2 and NO .

A portion of the ferrous ion is converted to ferric ion by the oxygen in the flue gas. The absorbed liquor is sent to a reduction step, where the ferric ion is reduced to ferrous ion by electrolysis.

The liquor is then sent to a stripper, where it releases concentrated SO_2 and NO by steam distillation. The NO is reduced to N_2 ; the SO_2 is used in sulfuric acid production. The liquor from the scrubber is returned to the absorber. In tests with a pilot plant ($150 \text{ Nm}^3/\text{hr}$) about 95% of the SO_2 and 85 percent of the NO_x were removed at an L/G ratio of 1 liter/ Nm^3 .

It is estimated that the cost is \$80 million for a 67 MW system, EDTA consumption per year is 300 to 400 tons (\$500,000 to 600,000), and simultaneous removal cost is \$34/kl oil, including fixed costs.

By use of H_2S in the reduction step, elemental sulfur may be produced. Tests with a larger plant are required for further evaluation.

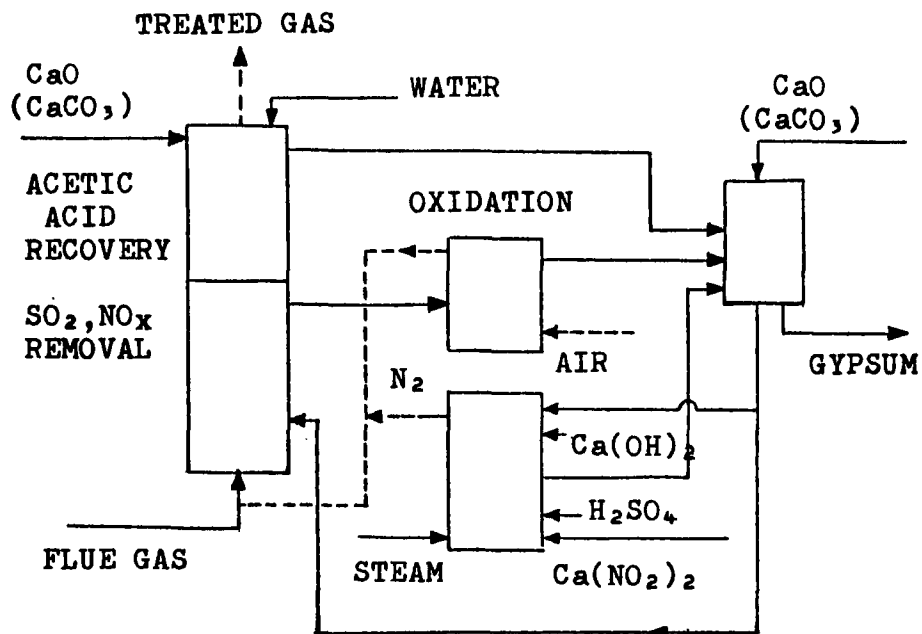


Figure 7-5 Flowsheet of Kureha simultaneous removal process

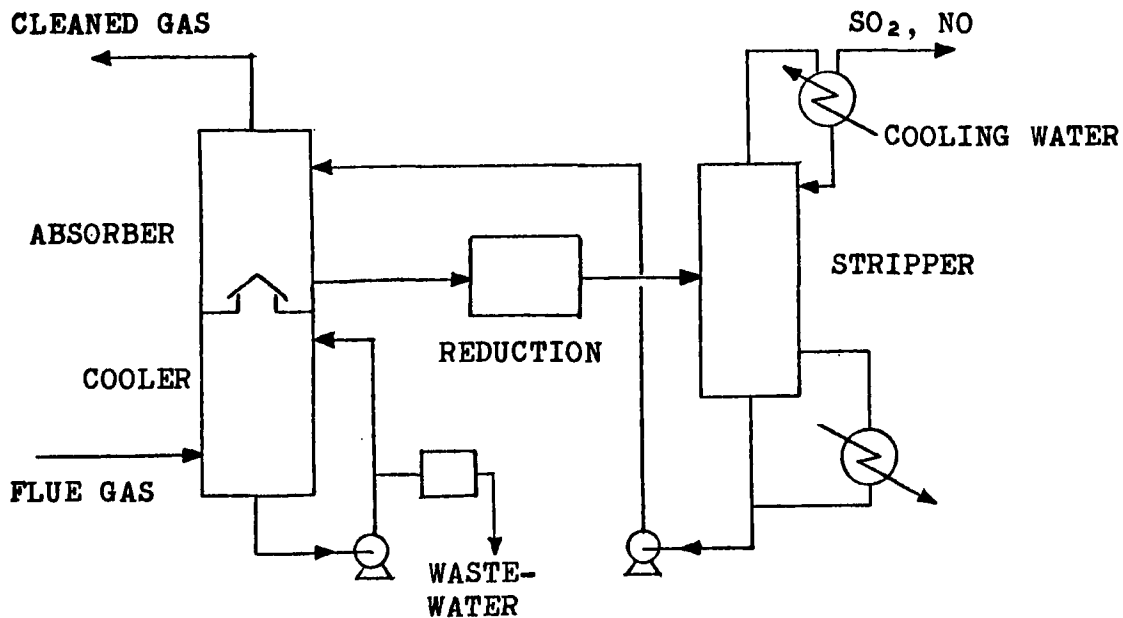


Figure 7-6. Flowsheet of Mitsui Shipbuilding process

Chisso Engineering Process (CEC Process)¹¹

Chisso Engineering, a subsidiary of Chisso Corporation, has developed a process for simultaneous removal of SO₂ and NO_x from flue gas by ammonia scrubbing using a catalyst (chelating compound) to by-produce ammonium sulfate. A pilot plant (300 Nm³/hr of flue gas from an oil-fired boiler) has been operated (Figure 7-7).

Flue gas containing SO₂ and NO_x is absorbed with an ammoniacal solution containing a soluble catalyst to reduce the absorbed NO_x into NH₃ by ammonium sulfite and bisulfite, which are formed from SO₂ and ammonia. Most of the catalyst is separated from the product solution containing ammonium sulfate and sulfite and intermediate compounds. The solution is oxidized by air and then heated to convert the intermediate compounds into ammonium sulfate. The product solution is concentrated in an evaporator to crystallize ammonium sulfate, which is separated by a centrifuge. The mother liquor, which contains a small amount of the catalyst, is returned to the catalyst separation step. The overall reaction may be expressed simply in the following way:



In order to recover 80 percent of 300 ppm NO_x it is desirable to have more than 1,200 ppm SO₂ in the flue gas.

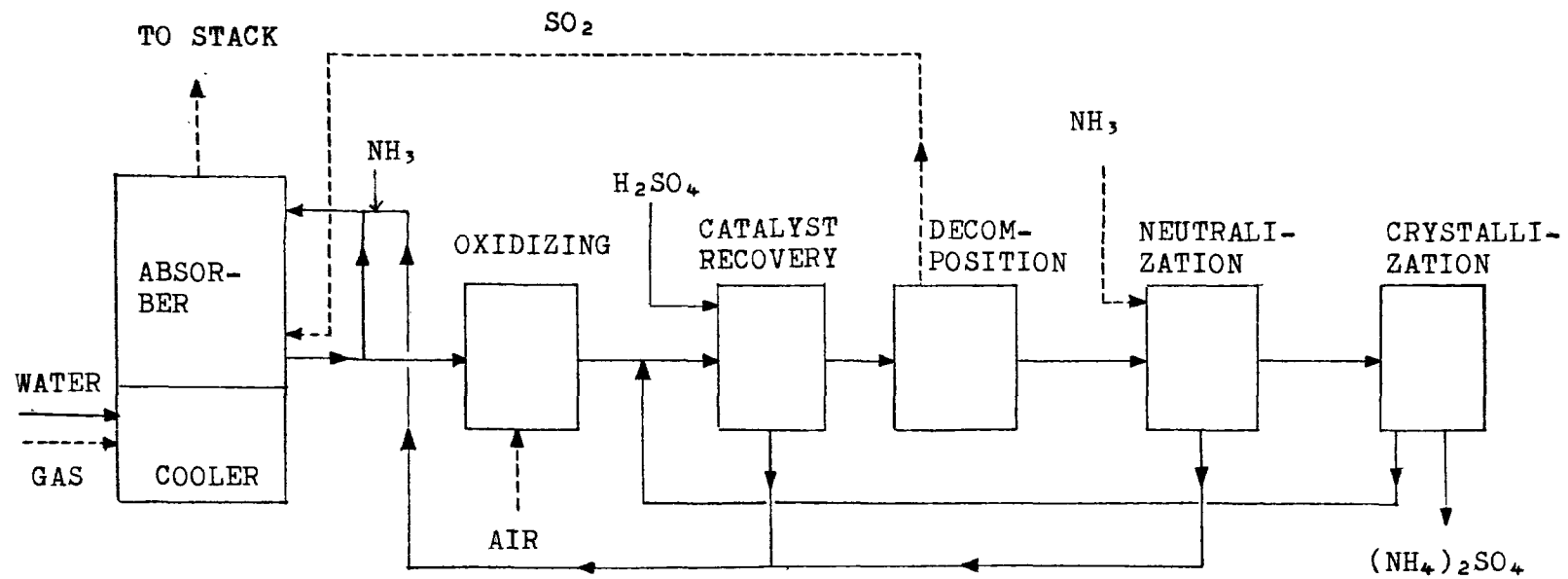


Figure 7-7 Flowsheet of CEC process

Reaction of NO_x with the sulfite liquor is not rapid, and the removal ratio has ranged from 70 to 80 percent. The use of a dilute liquor is favorable for the absorption but necessitates much evaporation for ammonium sulfate recovery. Therefore, NO_x removal of about 70 percent by use of a moderate concentration may be suitable. The catalyst is not affected by nickel and vanadium derived from flue gas. Chisso estimates the cost for simultaneous removal of SO_2 and NO_x to be about 40 percent more than for SO_2 removal only.

The process has the advantage of producing ammonium sulfate, utilizing both SO_2 and NO_x . Plume formation common to ammonia scrubbing processes might be a problem for this process.

Asahi Chemical Process

Asahi Chemical Co. has been testing a reduction process with a pilot plant ($600 \text{ Nm}^3/\text{hr}$). A flowsheet of the process is shown in Figure 7-8. A flue gas containing 1,250 ppm SO_2 , 3 to 4 percent O_2 , and 200 ppm NO_x is led to a sieve-tray absorber and treated with a sodium sulfite solution at pH 6.3 containing EDTA and ferrous ion. More than 80 percent of the NO_x is absorbed, forming an adduct with ferrous ion and EDTA, while more than 90 percent of the SO_2 is absorbed reacting with the sulfite. The NO adduct reacts with the sulfite to form sodium sulfate and nitrogen by the following reaction:

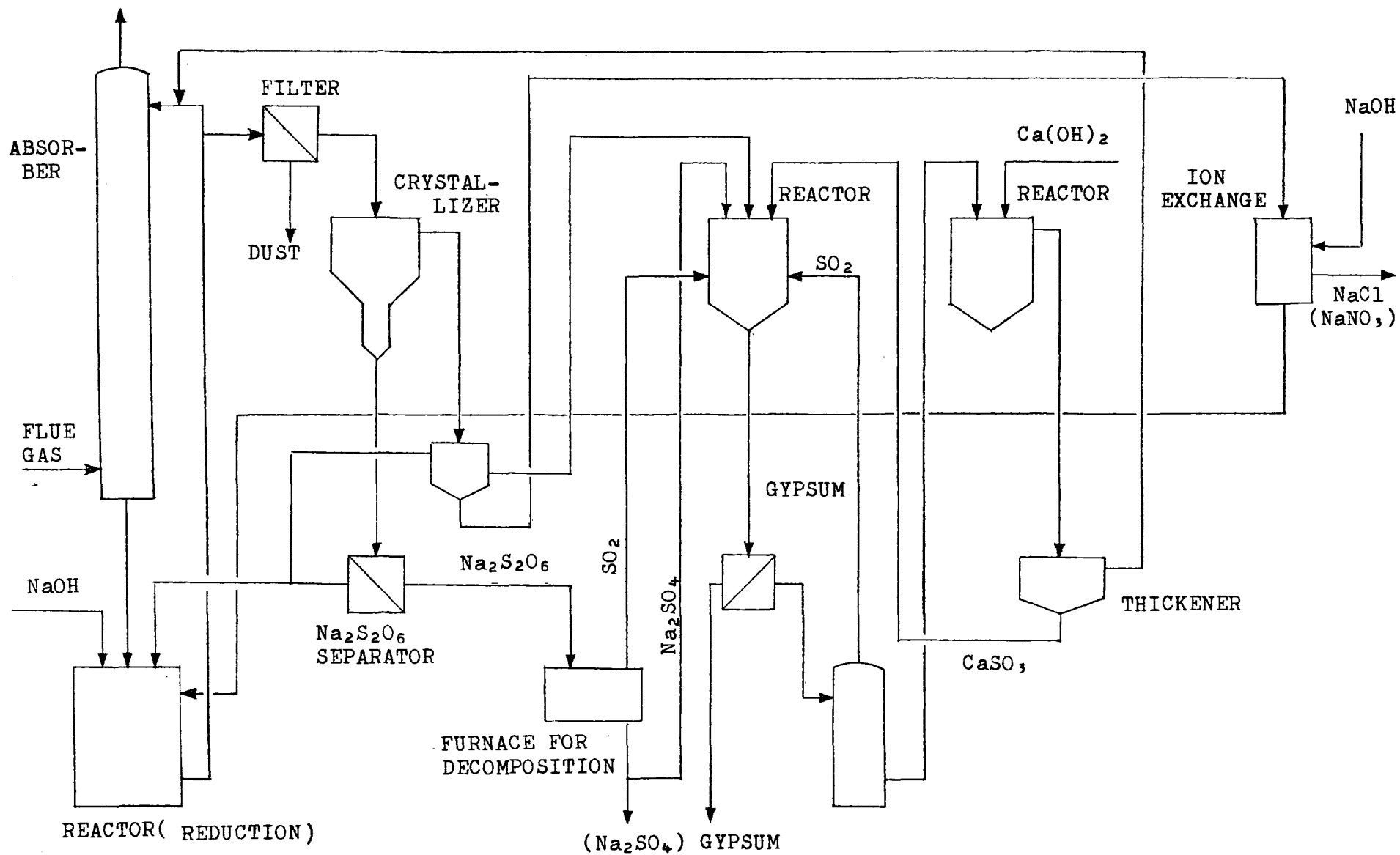
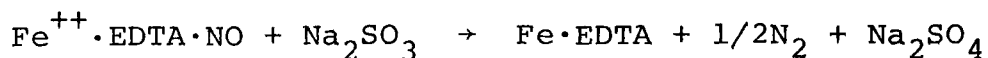
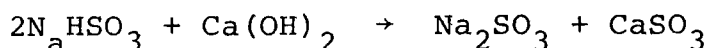
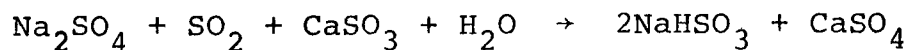
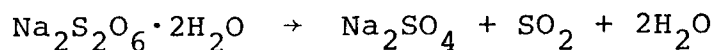


Figure 7- 8 Flowsheet of Asahi Chemical reduction process



Most of the resulting liquor is returned to the absorber. A portion is sent to a crystallizer, where sodium dithionate $\text{Na}_2\text{S}_2\text{O}_6 \cdot 2\text{H}_2\text{O}$ is crystallized. The dithionate is separated and heated at 300°C to be decomposed to Na_2SO_4 and SO_2 , both of which are sent to a reactor and reacted with calcium sulfite to precipitate gypsum.



Sodium bisulfite solution formed by the reaction is treated with lime to precipitate calcium sulfite and to regenerate sodium sulfite; the former is sent to the reactor and the latter is recycled to the absorbing system. Chloride, which is derived from the fuel and accumulates in the scrubbing liquor, can be eliminated by ion exchange, with which Asahi Chemical has had much experience.

Asahi Chemical estimates that the cost for a 500,000 Nm^3/hr unit (160 MW equivalent) is \$16 million and that requirements for treatment of 10,000 Nm^3 of gas containing 2,000 ppm SO_2 are as follows:

$\text{Ca}(\text{OH})_2$	6.7 kg
$\text{FeSO}_4 \cdot 7\text{H}_2\text{O}$	1.0 kg
EDTA	1.0 kg

NaOH	4.2 kg
Oil (gas reheating)	30 kg
Oil (thermal decomposition)	5 kg
Steam	60 kg
Cooling water	6 tons
Power	15 kWh

The system is not simple but represents a combination of feasible unit processes. Operation data of a larger pilot plant may be needed for a reliable evaluation of commercial feasibility.

Kawasaki Magnesium Process

Kawasaki Heavy Industries has been operating a pilot plant with magnesium scrubbing with a capacity of treating 5,000 Nm³/hr of flue gas from a coal-fired boiler (Table 7-1, Figure 7-9).

The gas, containing about 1,000 ppm SO₂ and 400 ppm NO_x, is mixed with NO₂ gas to adjust the NO₂/NO ratio to 1 and is treated with a magnesium hydroxide slurry to form magnesium sulfite and nitrite. The nitrite is separated and decomposed by adding sulfuric acid to produce NO, which is oxidized to NO₂ and returned to the absorber. The magnesium sulfite is oxidized into sulfate and reacted with calcium nitrate to precipitate gypsum, which is centrifuged. The separated magnesium nitrate solution is reacted with

Figure 7-9. Flowsheet of Kawasaki magnesium process.

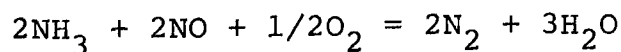
calcium nitrate to precipitate gypsum, which is centrifuged. The separated magnesium nitrate solution is reacted with calcium hydroxide to precipitate magnesium hydroxide, which is returned to the absorber. Part of the calcium nitrate liquor formed by the reaction is returned to the system for the reaction with magnesium sulfate and the rest is obtained as by-product.

The process has the advantage of removing both SO_2 and NO_x while by-producing gypsum and calcium nitrate. The process is not simple, however, and the demand for by-product calcium nitrate is limited.

DRY PROCESSES FOR SIMULTANEOUS REMOVAL

Reaction of Activated Carbon

Activated carbon has been used commercially as the absorbent of SO_2 . Although it also absorbs NO_x , the absorbing capacity is not sufficient to treat a large amount of gas. Takeda Chemical has produced an activated carbon containing metallic components or with a special structure to promote the reaction of NO_x with ammonia to form N_2 . Higher temperature is favorable to the reaction but decreases the SO_2 absorbing capacity (Figure 7-10). Optimum temperature for simultaneous removal by this process is about 250°C .



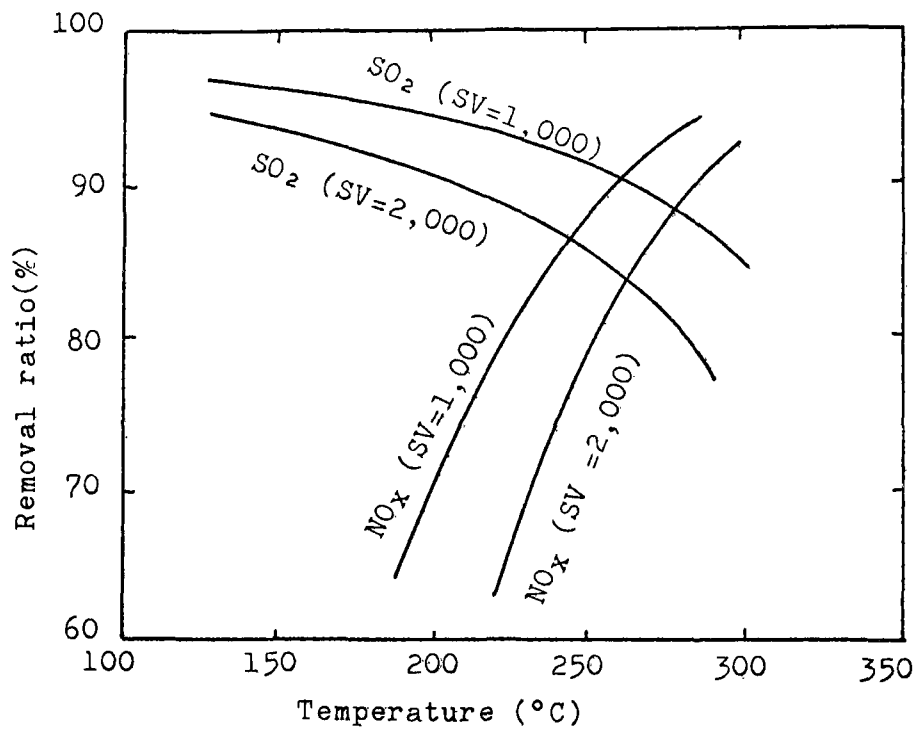


Figure 7-10. Schematic diagram of SO₂ and NO_x removal by activated carbon at different temperatures and space velocities.

Carbon for simultaneous removal of SO_2 and NO_x costs about 8,000/t, whereas carbon used commercially for FGD costs \$3,000/t. Simultaneous removal for flue gas from a 300 MW boiler will require about 1,000 t of the carbon, which may be too expensive. The price will be substantially lowered through mass production.

Unitika Activated Carbon Process

Unitika Co. recently started operating a pilot plant with a capacity of treating 4,500 Nm^3/hr of flue gas from a glass melting furnace; the gas contains about 400 ppm SO_2 and 500 ppm NO_x (Figure 7-11). The plant has a tower with four compartments, all of which have a fixed carbon bed. About 600 ppm NH_3 is added to the gas at about 230°C, and the mixed gas is led into three compartments. About 90 percent of the NO_x and SO_2 is removed. The carbon that has absorbed SO_2 is heated to 350°C in a reducing hot gas to release concentrated SO_2 for sulfuric acid production. Ammonium sulfate and sulfite, which tend to form on the carbon, are decomposed to SO_2 and N_2 in the regeneration step.

Design and operating parameters are as follows:

Tower height	17 m
Carbon bed thickness	1 m
Pressure drop	100 mm H_2O

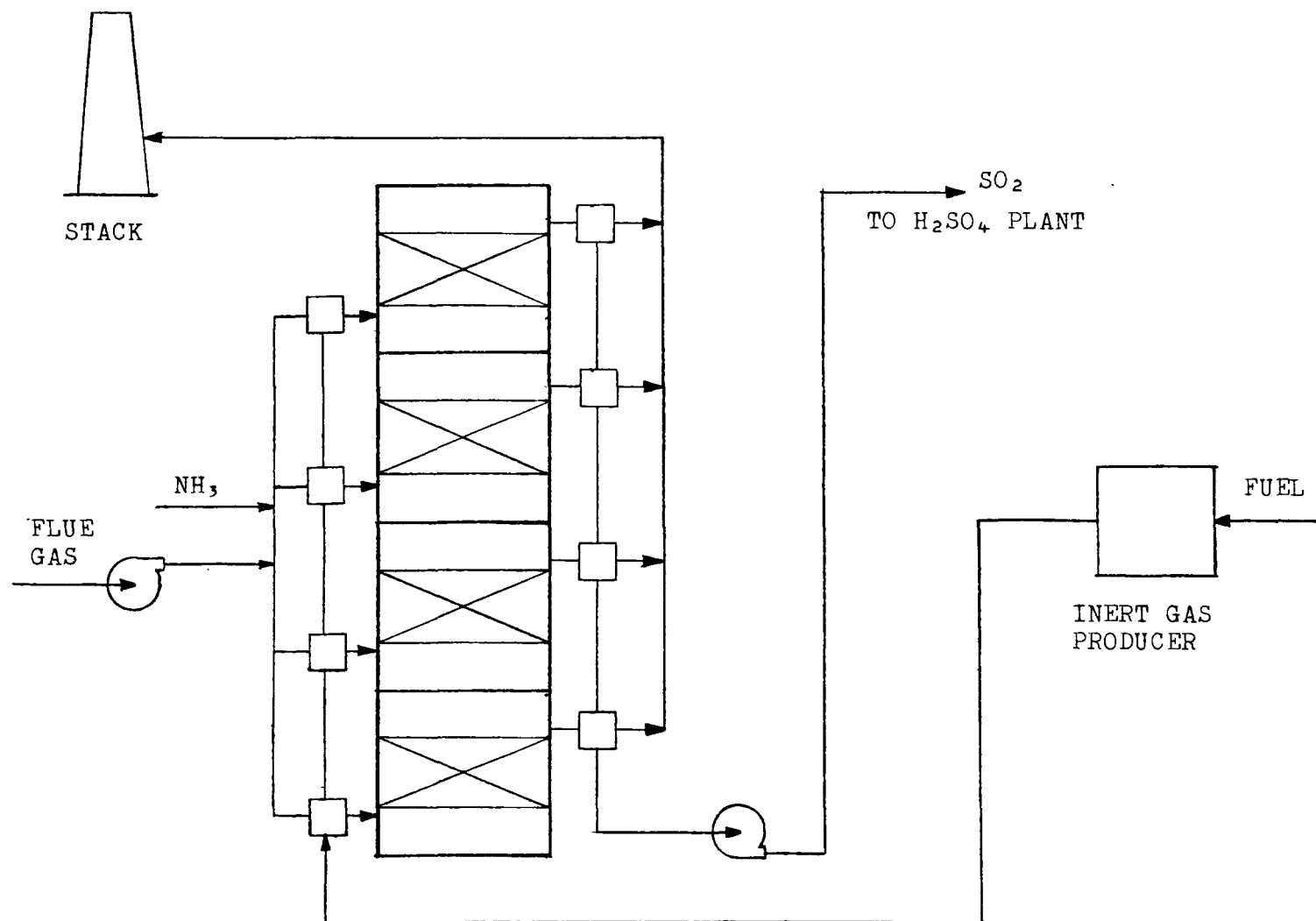


Figure 7-11 Flowsheet of Unitika process

Absorption time for one cycle 3 days

Regeneration time for one cycle 12 hours

SO₂ in gas from regeneration step 5-10%

A superficial gas velocity of about 700 is used. The carbon consumption is estimated to be less than 10 percent of a charge per year. In the 6-month operation, the loss has been only 1 to 2 percent. The gas from the regeneration step, containing 5 to 10 percent SO₂, may be used for sulfuric acid production.

Other Activated Carbon Processes

Sumitomo Heavy Industries has constructed a prototype FGD system (175,000 Nm³/hr) using moving beds of absorbent activated carbon, which is regenerated by heating a reducing gas.² With this unit they have studied simultaneous removal and are constructing a test unit with a capacity of treating 1,500 Nm³/hr flue gas using moving beds.

Hitachi Ltd. has found that activated carbon treated with ammonium bromide is effective even at 100°C for NO_x reduction by ammonia.¹⁴ The low-temperature activity may result in energy saving (Figure 7-1, No. 4), but deposition of ammonium sulfate and bisulfate on the carbon may present a problem.

Shell Copper Oxide Process

Copper oxide used as an absorbent of SO_2 in the Shell process works as catalyst in the reaction of NO_x with ammonia. The Yokkaichi plant, SYS, treating $120,000 \text{ Nm}^3/\text{hr}$ of flue gas from an oil-fired boiler by the Shell process, has introduced ammonia into a reactor at 400°C since 1975. Up to about 70 percent of the NO_x can be removed. Copper sulfate formed by SO_2 absorption is reacted with hydrogen to generate concentrated SO_2 , which is sent to a Claus furnace to produce sulfur.

Ebara Electron Beam Process¹⁵

Ebara Manufacturing Co. has developed a unique process for simultaneous removal by electron beam radiation. Flue gas is introduced into a reactor and exposed to the beam. About 80 percent of the SO_2 and 90 percent of the NO_x can be removed, forming a sulfuric acid mist and a powdery product containing S, N, O, and H, which are caught by an electrostatic precipitator. A pilot plant ($1,000 \text{ Nm}^3/\text{hr}$) has been operated and a larger plant ($3,000 \text{ Nm}^3/\text{hr}$) is to be constructed. Investment cost and power consumption seem high.

8. COMPARATIVE EVALUATION

DIFFERENCES AFFECTING PROCESS APPLICATIONS IN THE UNITED STATES AND JAPAN

Significant differences in circumstances in the United States and Japan can affect the selection of an FGD process and unit design. The major differences are as follows:

- (1) In the United States, gypsum and sulfur from natural sources are plentiful and inexpensive, whereas these are limited in Japan. By-products from desulfurization, including sodium sulfite and gypsum, can be sold in Japan, a fact conducive to the development of recovery processes.
- (2) In the United States, most plants have enough space for disposal of waste products, whereas in Japan space limitations necessitate maximum utilization of by-products.
- (3) In the United States, about 60 percent of the electric power is generated by burning coal, which gives much fly ash. In Japan, most power plants burn oil, which gives little dust--an advantage in recovering by-products with high purity.
- (4) In the United States, many power plants are located far from chemical plants. In Japan, power and chemical plants are usually close to each other; hence it is easy for chemical plants to utilize desulfurization by-products and for power plants to use chemicals.
- (5) SO₂ concentration of flue gas usually ranges from 400 to 1,500 ppm in Japan, whereas it reaches 2,000 to 3,000 ppm in the United States.

- (6) In Japan, many plants are close to cities. More than 90 percent removal of SO_2 or less than 100 ppm SO_2 in emitted gas is usually required. In the United States, about 85 percent desulfurization or 300 ppm SO_2 in the gas is usually acceptable.
- (7) Regulations of the purge of wastewater in many states in the United States are more stringent than in Japan.
- (8) Compared with flue gas from oil burning, that from coal is richer in NO_x and dust. It is not easy to reduce NO_x concentration in flue gas from coal combustion to less than 400 ppm; moreover, the dust contaminates the catalyst for NO_x removal. Simultaneous removal of SO_2 and NO_x by wet processes may be useful for treatment of flue gas from coal combustion.

WET LIME/LIMESTONE PROCESS

System Cost and Operation

The plant cost for the Japanese lime/limestone-gypsum process is considerably higher than that for the United States throwaway sludge process because of the requirement of a pH controller (or additional absorber), oxidizer, and centrifuge, which account for about 30 percent of the plant cost. Where there is enough space for sludge discarding, the throwaway process may be most economical.

Most of the lime/limestone-gypsum process systems have a gas cooler before the scrubber, and the scrubber is designed to ensure high SO_2 removal efficiency as well as high utilization of the absorbent, both above 95 percent in many cases. These factors make the system more costly

but free from serious scaling problems. The gas cooler humidifies the gas and prevents local drying in the scrubber, thus helping to prevent formation of scale. The high SO₂ removal and absorbent utilization in the scrubber may reduce scaling of the mist eliminator, as will be discussed below.

Japanese units are usually provided with spare pumps but seldom with a standby scrubber because long-term continuous operation can be carried out without severe scaling problems.

Saturated or Unsaturated

"Unsaturated mode operation" has been carried out successfully at the Paddy's Run plant, Louisville Power and Electric, and at the Omuta plant, Mitsui Aluminum. The operation prevents gypsum formation by keeping the oxidation of calcium sulfite below 20 percent and thus permits scale-free operation.

Japanese processes producing gypsum are usually operated in the "saturated mode", circulating a slurry containing gypsum as a crystal seed, and might be more widely applicable to various gas compositions. Unsaturated mode operation is not suitable for gases with a low SO₂/O₂ ratio such as that from a low-sulfur coal, because the low ratio encourages oxidation. Even with a gas from high-sulfur coal, which normally contains 2,000 to 3,000 ppm SO₂ and 4 to 5 percent

O₂, there is a possibility of scaling in the scrubber because of gypsum formation due to a temporary increase in the O₂ concentration and a temporary decrease in pH of the slurry as the operation load fluctuates.

Scaling on Mist Eliminator

Scaling tends to occur most readily on mist eliminators where, in addition to wetting and drying, lime or limestone from the mist reacts with SO₂. The gas passing through the eliminator usually contains 200 to 400 ppm SO₂ and 4 to 5 O₂ in the U.S. and 20 to 100 ppm SO₂ and 1 to 2 percent O₂ in Japan. In both cases gypsum would form by the reaction with lime or limestone because of the low SO₂/O₂ ratio. Washing the eliminator with fresh water can dissolve the gypsum but will increase the wastewater load. The scaling may be reduced by washing with a supernatant of gypsum slurry containing fine crystals of gypsum, which can serve as crystal seed, as has been done in Japan.

It has been argued that scale-free operation of wet lime/limestone process systems in Japan may be due largely to the low SO₂ concentration of inlet gas, which is below 600 ppm in many units. However, the inlet SO₂ concentration may not be very important because new full-scale systems for oil- or coal-fired utility boilers (250-500 MW) with SO₂ concentrations of 1,500 to 1,800 ppm in the gas have been in

scale-free operation even with frequent changes of operation load. For prevention of scale on the mist eliminator, the SO_2 concentration in the scrubber outlet gas and the utilization of lime or limestone in the scrubber may be important. The presence of much unreacted lime or limestone in the mist and much SO_2 in the gas flowing through the eliminator would increase formation of gypsum on the eliminator.

In Japan, the outlet SO_2 concentration is low because of the stringent regulation, and the utilization of lime or limestone usually exceeds 95 percent or 90 percent because of the necessity to produce high-quality gypsum. Such a situation may have reduced the scaling on the mist eliminator. The high utilization of lime or limestone is obtained by a good gas-liquid contact resulting from suitable scrubber design, by increasing the liquid/gas ratio, and by the fine grinding of limestone in a wet mill to pass 325 mesh. The high utilization of lime and limestone tends to reduce the pH of the slurry, increase the oxidation, and encourage the formation of gypsum.

Many of the Japanese companies once used a zigzag baffle-type mist eliminator placed horizontally at the top of the scrubber, such as is widely used in the United States, because it is relatively inexpensive; now they seldom use this type because performance is poor. Many now

place the mist eliminator vertically in a separate vessel in the exit duct outside the scrubber because this configuration provides better drainage and better over-all performance.

Effect on Fly Ash and Wastewater on Scaling

The effect of fly ash on scaling is not yet clear. The ash from coal consists of fine, spherical, glassy particles having little reactivity. The ash would increase erosion when present in large amounts in the slurry but not always increase scaling. It might help prevent scaling because of its nonreactive, erosive nature, as pointed out by R.H. Borgwardt, EPA. At present, six FGD systems for coal-fired utility boilers (160 to 300 MW) are in operation in Japan. One of them uses the unsaturated mode and the others use the saturated mode. Operation of all six plants is virtually trouble-free. They are equipped with electrostatic precipitators which remove most of the fly ash, and thus the effect of fly ash has not been clearly demonstrated.

It has been stated that Japanese FGD units purge wastewater to add large amounts of fresh water to the system and thus help to reduce scaling. This view may not be correct, because the sum of wastewater and moisture in the by-product gypsum is no larger than in the United States practice, where the sludge normally contains 60 percent or more water. As shown in Table 3-8, the water ratio or the equivalent water

ranges from 26 to 64 percent for the plants using wet lime/limestone processes.

INDIRECT LIME/LIMESTONE PROCESS

Sodium Scrubbing

In the United States the sodium scrubbing double alkali processes are simple, as they use lime and by-produce calcium sulfite sludge; in Japan the processes are fairly complex, as they use limestone and are equipped with sulfate decomposition and gypsum recovery units. Limestone is cheaper than lime but limestone processes require larger reactors.

In both processes a portion of the sodium sulfite is oxidized to form sulfate ion, which must be removed from the system. In the U.S. systems, sulfate ion is removed in two ways: (1) by entering calcium sulfite crystals replacing sulfite ion and (2) as sodium sulfate solution contained in the sludge. The oxidation of the sulfite must be maintained under a certain level to prevent accumulation of sodium sulfate in the liquor. The process suits high-sulfur coal but may not suit low-sulfur coal, because it needs a flue gas with a relatively high SO_2/O_2 ratio.

The Japanese processes, Kureha-Kawasaki and Showa Denko, incorporate a sulfate decomposition unit. This makes them costly, but they are applicable to any kind of gas.

Gypsum grows into much larger crystals than does calcium sulfite and is washed well. Sodium content in the by-product gypsum is less than 0.1 percent, whereas in the U.S. systems the content in the sulfite sludge usually exceeds 2 percent.

Other Indirect Processes

The Chiyoda and Dowa processes are much simpler than the above-mentioned Japanese processes. Compared with the U.S. systems, they are more complex because of the requirement for an oxidizer; moreover, they need higher L/G ratios because of the low pH of the absorbing liquor. They do, however, offer the following advantages: (1) use of limestone, (2) much smaller sizes of thickener and filter, (3) smaller loss of absorbent, (4) less possibility of scaling, and (5) applicability to any kind of gas, including that from low-sulfur coal. They may be most applicable at plants that do not require very high SO₂ removal efficiency.

The Kurabo ammonium sulfate and Kureha sodium acetate processes use lime, as do the U.S. processes. Although they are less simple than the U.S. processes, they provide some advantages. The Kurabo process gives virtually no plume, which has been a common problem for ammonia scrubbing processes. The Kureha process can recover more than 99 percent of the SO₂. Although such a high removal ratio is not

usually needed, in certain districts it may be a good idea to treat 70 to 80 percent of flue gas by this process and then mix the treated gas with untreated hot gas to eliminate reheating.

OTHER PROCESSES (RECOVERY PROCESSES)

Four types of FGD processes are in commercial use in Japan to by-produce concentrated SO_2 , which is used to produce sulfuric acid or is sent to a Claus furnace to produce elemental sulfur. These are the Wellman-Lord, magnesium (or zinc) scrubbing, activated carbon, and the Shell processes.

The Wellman-Lord process has been most widely used because of smooth operation of the systems and recovery of virtually all of the SO_2 , which permits substantial reduction of the size of the sulfuric acid plant relative to a conventional plant using 7 to 9 percent SO_2 gas obtained by burning pyrite or sulfur. The process, however, is becoming increasingly costly as regulations concerning wastewater become increasingly stringent.

The magnesium scrubbing process has the advantage of giving virtually no wastewater. But operation of a rotary kiln may not be willingly accepted by power companies. The recovered SO_2 , having a low concentration of 7 to 8 percent, can be used in producing sulfuric acid or in charging into a Claus furnace; it is not suitable for reduction to H_2S to produce elemental sulfur. It may be said that magnesium

scrubbing better suits chemical plants and oil refineries, and the Wellman-Lord process better suits power plants.

Carbon absorption with thermal regeneration gives a moderate concentration of the recovered SO_2 (about 20 percent) which can be used for production of both sulfuric acid and sulfur. Consumption of carbon increases when a moving bed is used and makes the process costly, particularly in Japan where good-quality and expensive (about \$2,700/t) carbon is used. Cheaper carbon is useful but has a lower absorbing capacity and tends to burn during the operation.

The carbon process using a fixed bed and water wash has proved to consume very little carbon. Although the by-produced sulfuric acid is weak, it can be concentrated to about 60 percent by the heat of flue gas, as has been done by Unitika. The process may be suitable for certain chemical plants that can use the acid.

The Shell process seems more costly than the other recovery processes because hydrogen is needed for regeneration. The process is advantageous in that it can effect simultaneous removal of NO_x up to about 70 percent.

BY-PRODUCT AND WASTEWATER

Gypsum and Calcium Sulfite

Oversupply of gypsum has presented a problem in Japan where land for disposal is scarce. Because per capita

consumption of gypsum in Japan is still about half of that in the United States, a considerable increase in future demand for construction material is expected. Some oversupply may continue, however. Gypsum is not suitable for land filling of ground to support large buildings because it has a little water solubility. A stabilized calcium sulfite sludge, which has been produced in the United States, may be better for land filling. Because fly ash is scarce in Japan, industrial slags may be used instead.

In contrast with current practice, gypsum may be produced in the future in certain districts in the United States. Gypsum by-produced from desulfurization of flue gas from coal has proved useful for cement and wallboard production, as recently practiced in Japan. It may be possible to substitute the gypsum for a portion of that now imported to the United States which amounts to 7 million tons yearly. Gypsum can be piled up to heights of more than 100 feet, as is done at phosphoric acid plants in Florida, thus allowing a great saving of land space as compared with the use of calcium sulfite sludge ponds.

Elemental Sulfur

Generally speaking, less effort has been made in Japan than in the United States to develop FGD processes that yield elemental sulfur as a by-product. Several oil refineries

have installed FGD units to recover concentrated SO_2 , which is sent to an existing Claus furnace to recover sulfur. Ammonia scrubbing with the IFP reactor to by-produce sulfur has been tested by Mitsubishi Heavy Industries and Toyo Engineering; this effort will be abandoned, however, because of operational problems including the sulfate decomposition step. Except for these, active research to develop new processes has not been made.

This may be due largely to the use of oil as the major fuel in Japan. Elemental sulfur can be recovered from oil at lower cost by hydrodesulfurization than by FGD. An economical process to by-produce sulfur is being sought, however, for the following two reasons: (1) all other by-products of FGD are in oversupply, and (2) heavy oil suitable for hydrodesulfurization to reduce sulfur below 0.5 percent is limited by the content of metallic impurities that poison the catalyst.

Chloride and Wastewater

When no wastewater purge is allowed, chloride in the flue gas accumulates in the scrubber liquor and tends to increase corrosion and decrease SO_2 removal efficiency. In the throwaway sludge process, the chloride concentration in the liquor may be kept within an acceptable level because a considerable amount of water is removed from the system

with the sludge, which normally contains more than 60 percent water. In a process by-producing gypsum, which normally contains less than 10 percent moisture, chloride concentration in the liquor will exceed 5 percent when the coal burned is relatively rich in chloride. Plant cost will be increased by the need for highly corrosion-resistant material, such as in the Kobe Steel calcium chloride process.

Chloride may be removed in a prescrubber by water wash of the gas; the liquor from the prescrubber may be neutralized with lime or limestone, and the resulting sludge may be discarded. In a sodium scrubbing process, a portion of the scrubber liquor may be concentrated to crystallize and separate sodium chloride. All such treatments add considerably to the cost of the process. The author believes that regulations should permit purging of a limited amount of wastewater after treating it thoroughly to remove any impurities that may be harmful to the environment.

SIMULTANEOUS REMOVAL OF SO_x AND NO_x

The present denitrification efforts in Japan have been motivated by the quite stringent regulation. NO_x concentrations in large cities in the United States, such as Los Angeles and Chicago, are much higher than those in Tokyo, Osaka, and other Japanese cities. Although a larger portion of NO_x in crowded cities is derived from automobiles,

NO_x from coal-fired boilers will become significant as consumption of coal increases, because the flue gas is relatively rich in NO_x (from 500 ppm to 1,000 ppm) and it is not easy to reduce the concentration below 400 ppm by combustion control.

Among the denitrification processes, the Exxon process injecting ammonia into flue gas at about 800°C without catalyst would require the least cost. The process has a narrow range of optimum reaction temperatures, and for large-scale applications about 50 percent NO_x removal is expected. Catalytic reduction processes using ammonia can attain over 90 percent removal but entail the problem of catalyst contamination by dust in the gas. The activated carbon process can remove about 90 percent of the SO₂ and NO_x simultaneously but requires a large amount of carbon and also involves a dust problem. The Shell process can remove about 70 percent of the NO_x together with about 90 percent of the SO₂ with a smaller dust problem due to the use of the parallel passage reactor. Selection of a process should be decided according to the NO_x removal ratio required.

The dry processes using ammonia entail some common problems: (1) Formation of ammonium bisulfate in the gas, which condenses anywhere at lower temperature, for example, in heat exchangers. (2) Possibility of secondary pollution

by emission of ammonia, which may occur to a considerable extent when the operation load fluctuates. (3) Widespread use of the ammonia processes on a large scale could cause a worldwide shortage of nitrogen fertilizer.

For these reasons, it is desired to develop not only ammonia-using processes but also wet processes for simultaneous control. The wet process by-producing ammonia or ammonium sulfate is desirable from the standpoint of fertilizer supply and will possibly be used extensively in the future if the process economy can be substantially improved. For plants where the transportation of ammonia or ammonium sulfate may be inconvenient, a combination of the Exxon process with a wet process by-producing ammonia might be useful. The by-produced ammonia may be returned to the boiler to reduce a portion of the NO_x to N_2 and thus promote simultaneous removal, for which a low NO_x/SO_2 ratio is favorable.

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The process descriptions in this report are based primarily on Ando's visits to the desulfurization plants, his discussions with the users and developers of each process, and data made available by them. In addition, the following publications were used and are cited as references.

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