

Estimating Total Sulfuric Acid Emissions from Stationary Power Plants

2018 Update

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Technical Update, March 2018

EPRI Project Manager

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ABSTRACT

The U. S. Environmental Protection Agency's (EPA's) Toxics Release Inventory (TRI) program requires electric utilities to estimate and report annual emissions of 682 chemicals and chemical categories. Sulfuric acid (H₂SO₄) is one of the chemicals included in the TRI reporting requirement, when it is present in an aerosol form. The EPA defines sulfuric acid aerosols as "includ[ing] mists, vapors, gas, fog, and other airborne forms of any particle size." The objective of this report is to provide a method for predicting the sulfuric acid manufactured during fuel combustion, removed by air pollution control equipment or flue gas treatment, and released to the environment in stack gases. The estimation method is an empirical model, whereby emissions are predicted from factors derived from sulfuric acid measurements at full-scale power plants. The power generation units considered in the report include coal, oil and natural gas-fired boilers, as well as both simple and combined cycle gas turbines. The procedures presented here can be used to estimate sulfuric acid manufacture from combustion, operation of nitrogen oxide reduction control equipment and flue gas conditioning, and the removal of SO₃ or H₂SO₄ from the flue gas stream by air heaters, particulate control devices such as electrostatic precipitators (ESPs) and fabric filters, sulfur dioxide control equipment, and nitrogen oxide control equipment.

Keywords

Air emissions Coal combustion Natural gas combustion Sulfuric acid Toxics Release Inventory (TRI)



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PRIMARY AUDIENCE: Power company environmental staff responsible for Toxics Release Inventory reporting and air permitting.

SECONDARY AUDIENCE: Air quality agencies and other organizations with an interest in power plant emissions.

KEY RESEARCH QUESTION

The U. S. Environmental Protection Agency's (EPA's) Toxics Release Inventory (TRI) program requires owners of some coal-, oil-, and natural gas-fired power plants to estimate and report annual emissions of 682 chemicals and chemical categories. Sulfuric acid (H_2SO_4) is one of the chemicals included in the TRI reporting requirement, when it is present in an aerosol form. The objective of this report is to provide a method for predicting the sulfuric acid manufactured during fuel combustion, removed by air pollution control equipment or flue gas treatment, and released to the environment in stack gases.

RESEARCH OVERVIEW

The sulfuric acid estimation method is an empirical model, whereby emissions are predicted from factors derived from sulfuric acid measurements at full-scale power plants. The power generation units considered in the report include coal, oil, and natural gas-fired boilers, as well as both simple and combined cycle gas turbines. The procedures presented here can be used to estimate sulfuric acid manufacture from combustion, operation of nitrogen oxide reduction control equipment and flue gas conditioning, and the removal of SO₃ or H_2SO_4 from the flue gas stream by air heaters, particulate control devices such as electrostatic precipitators (ESPs) and fabric filters, sulfur dioxide control equipment, and nitrogen oxide control equipment. The 2018 update to this report incorporates modifications to the methodology for estimating emissions from coal plants with hot-side electrostatic precipitators (ESPs), and plants that add magnesium oxide to inhibit conversion of sulfur dioxide to sulfur trioxide.

KEY FINDINGS

- The principal sources of sulfuric acid emissions from fossil fuel-fired power plants are 1) oxidation of sulfur during fuel combustion, 2) injection of sulfur trioxide for flue gas conditioning to improve the efficiency of ESPs, and 3) increased conversion of sulfur dioxide to sulfur trioxide when flue gas passes through selective catalytic reduction (SCR) NOx control equipment.
- Sulfuric acid is removed from flue gas when it reacts with ammonia to form solids that are not reportable to TRI, and are also removed when the flue gas passes through wet or dry scrubber equipment.
- Measurements of sulfuric acid before and after power plant process equipment were used to develop
 factors representing the impact of fuel type and pollution control devices on formation and removal of
 sulfuric acid.



WHY THIS MATTERS

Sulfuric acid is emitted from power plants that burn coal, and to a lesser extent, those that burn oil or natural gas for electricity production. For some U.S. power plants, the quantities emitted must be reported annually to the Environmental Protection Agency TRI program, as well as to state agencies. This report provides power plant owners with a methodology to estimate the impact of fuel changes or new equipment on sulfuric acid emissions.

HOW TO APPLY RESULTS

The equations in this report provide a complete methodology for estimating emissions from most common fossil fuel-fired power plant configurations. The equations may be calculated by entering them into a spreadsheet. The Electric Power Research Institute (EPRI) also incorporates the equations into the Toxics Release Inventory for Power Plants software, designed to assist power plant owners with annual TRI reporting.

LEARNING AND ENGAGEMENT OPPORTUNITIES

• Toxics Release Inventory for Power Plants User's Group

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PROGRAM: Program 59: Power Plant Multimedia Emissions Characterization

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ACRONYMS AND ABBREVIATIONS

APH	air preheater
ABS	ammonium bisulfate
AS	ammonium sulfate
Btu	British thermal units
CCS	controlled condensation system
CEMS	continuous emissions monitoring system
EDTA	ethylenediamine tetraacetic acid
EGU	electric generation unit
EPA	U.S. Environmental Protection Agency
EPCRA	Emergency Planning and Community Right-To-Know Act
EPRI	Electric Power Research Institute
ESP	electrostatic precipitator
FGC	flue gas conditioning
FGD	flue gas desulfurization
lb	pound
HV	heating value
MBtu	million Btu
Mcf	thousand cubic feet (of natural gas)
mmBtu	million Btu (of natural gas)
mol	mole
MW	molecular weight
PCD	pollution control device
ppm	parts per million
ppmv	parts per million by volume
PRB	Powder River Basin
scf	standard cubic feet
SBS	sodium bisulfate
SCR	selective catalytic reduction
SCS	Southern Company Services
SNCR	selective non-catalytic reduction

TRI	Toxics Release Inventory
TSAM	total sulfuric acid manufactured
TSAR	total sulfuric acid release
WAF	wall adjustment factor
yr	year

SI CONVERSION FACTORS

	English (US) units	X	Factor	=	SI units
Area:	1 ft ²	X	9.29×10^{-2}	=	m ²
Flow Rate:	1 gal/min	Х	6.31 × 10 ⁻⁵	=	m ³ /s
	1 gal/min	Х	6.31 × 10 ⁻²	=	L/s
Length:	1 ft	Х	0.3048	=	m
201801	1 in	X	2.54	=	cm
	1 yd	Х	0.9144	=	m
	•				
Mass:	1 lb	Х	454	=	g
	1 lb	Х	0.454	=	kg
	1 gr	Х	0.0648	=	g
	1 ton	Х	0.907		tonne
Volume:	1 ft^3	Х	28.3	=	L
	1 ft^3	Х	0.0283	=	m ³
	1 gal	Х	3.785	=	L
	1 gal	Х	$3.785 imes 10^{-3}$	=	m ³
Temperature:	°F-32	Х	0.556	=	°C
Energy:	Btu	Х	1055.1	=	joule

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1 INTRODUCTION

Background

Estimation of sulfuric acid emissions from power plants is a topic of increasing importance to the U.S. electric utility industry. Most significantly, Section 313 of the Emergency Planning and Community Right-to-Know Act (EPCRA), also known as the Toxics Release Inventory (TRI), requires facilities that "manufacture," "process," or "otherwise use" a listed chemical above certain threshold amounts to report their annual releases of the chemical to EPA and state agencies. For sulfuric acid, the TRI reporting requirements are triggered if a facility "manufactures" or "processes" more than 25,000 pounds of this chemical or "otherwise uses" more than 10,000 pounds of it in a given calendar year.

In June 1995, the U. S. Environmental Protection Agency (EPA) modified the list of chemicals subject to the EPCRA § 313 reporting requirements so that only aerosol forms of sulfuric acid would be subject to the TRI reporting requirements (EPA, 1995a). EPA defines sulfuric acid aerosols as "includ[ing] mists, vapors, gas, fog, and other airborne forms of any particle size." Although initially it appeared that only liquid droplets of sulfuric acid needed to be reported, the present method estimates the sulfuric acid emissions regardless of the physical state of the molecules.

Beginning on July 1, 1999, certain coal- and oil-fired electric power plants have been required to report annual releases of TRI chemicals that they manufacture, process or otherwise use above threshold amounts. Under EPA's EPCRA § 313 regulations, coal- and oil-fired electric utilities are deemed to "manufacture" sulfuric acid. Thus, electric utilities have to submit TRI reports on sulfuric acid aerosol releases if they "manufacture" more than 25,000 pounds of the chemical in a given reporting year (EPA, 1997).

Emissions of sulfuric acid received considerable attention in with the broad application of selective catalytic reduction (SCR) NOx control technology. The use of SCR unavoidably contributes to production of sulfur trioxide (SO₃), the precursor of sulfuric acid. Several notable incidents have been witnessed where an increase in sulfuric acid emissions, as manifested by an increase in stack plume visibility, was attributed to the addition of SCR process equipment to an existing power plant. The deployment of approximately 115 GW of selective catalytic reduction (SCR) from 1997 through 2009 prompted power plant owners to conduct extensive measurements documenting the fate of SO₃ and sulfuric acid in power plants.

The methodology described in this report was developed by Southern Company Services (SCS). Between 2000 and 2005, SCS developed several versions of the estimation methodology and released draft reports informally (Hardman, 1998; Hardman, 1999; Monroe, 2001, Harrison, 2004, and Harrison, 2005). The SCS model was widely used in the electrical power generation industry for TRI reporting and other emissions estimation purposes. In 2007, EPRI received permission from Southern Company Services to modify the SCS estimation method to include additional data and improvements, and to publish this method under EPRI sponsorship. The 2008

version of the model added the ability to estimate emissions based on a specified sulfuric acid flue gas concentration at the stack, which can be used when alkali injection is employed to reduce sulfuric acid emissions. The 2010 version updated several of the "technology impact factors" (F2 factors, used to reflect the impact of pollution control devices on sulfuric acid releases) with new SO₃ measurements. The 2012 version of the model (EPRI, 2012) made minor changes to the calculation procedure to more accurately represent flow through from the boiler and control devices, changed the approach used to account for sulfuric acid from flue gas conditioning (FGC), and made several corrections and clarifications to technology factors, including a new technology factor for units with a wet electrostatic precipitator (ESP) and the addition of a new sulfuric acid source through the oxidation of sulfur dioxide across a carbon monoxide catalyst in combined cycle units.

The current version of the model makes minor changes to the application of the sulfuric acid removal factors (Technology Impact Factors, F2) for configurations that include a hot-side ESP or magnesium oxide addition to the fuel or furnace. This version also incorporates methodology for the calculation of releases when SCR or SCNR is only injected on a partial-year basis (previously published in 2014 as an erratum to the 2012 report).

Objective

The objective of this report is to present a methodology for estimating sulfuric acid emissions from power generating facilities, for use in Toxics Release Inventory reporting and other applications. The scope of units considered includes coal- and oil-fired steam generating units, as well as simple cycle and combined cycle natural gas- and oil-fired combustion turbines. To date, the most authoritative and documented data describe sulfuric acid emissions from coal-fired steam generators. The focus on this category of units is due to the increased availability of measurements describing the fate of sulfuric acid attributable to the retrofit of SCR process equipment on coal-fired steam generators.

Report Organization

Section 2 of this report addresses the uncertainty in measuring SO₃ concentration using the preferred controlled condensation system (CCS) method, and the accuracy and representativeness of the available data. Section 3 outlines the general methodology adopted in this model while Section 4 describes the details of the methodology, focusing on data that can be used to predict sulfuric acid manufacture by combustion. This section also summarizes data describing the removal of sulfuric acid across air heaters, ESPs, and wet flue gas desulfurization (FGD) process equipment. Section 5 outlines how to conduct the calculation for steam boilers firing a mixture of fuels. Section 6 addresses simple cycle and combined cycle combustion turbine applications. Example calculations for various power plant configurations are presented in Appendix A. A chronology of changes to the sulfuric acid model is presented in Appendix B.

2 MEASUREMENT UNCERTAINTY

A predictive method for sulfuric acid emissions requires knowledge of sulfuric acid production and the fate of emissions from specific boilers and combustion turbines, to an acceptable degree of measurement accuracy. In addition, a large database of measurements from many electric generating units (EGUs) is desired to obtain a robust predictive method. Both the size and the representativeness of the measurement database are critical to the accuracy of a predictive method.

The predictive correlations presented in this report are based on data obtained from field tests conducted predominantly within the last 10 years, with some estimates conducted in the 1990s. The quality of the data is thought to vary widely. It was not possible to obtain quality control information for all of the available measurements; thus, it was not possible to critically review all data used in this model.

The technique used to measure SO_3/H_2SO_4 is not known for all data sources, but most measurements were conducted using the controlled condensate system (CCS), a widely used technique that is generally considered the most accurate method for measuring SO_3/H_2SO_4 in stack gases from stationary combustion sources. EPA Method 8 is sometimes used for this purpose, but that method has a known positive bias from oxidation of SO_2 to SO_3 in the impinger solution.

CCS Measurement Uncertainty

A series of field and laboratory trials conducted by EPRI evaluated the measurement bias of the CCS technique by comparing results from both laboratory tests and field trials (EPRI, 2001). SO₃/H₂SO₄ measurements at similar gas compositions were compared in a 1) clean, ash-free laboratory environment; 2) simulated ash environment, and 3) actual field duty. The findings of the EPRI tests conducted under high-ash conditions are applicable to much of the field data used to derive removal factors in the current estimation model. The results showed that, when fly ash was present in the flue gas stream, the technique imposed a low bias (e.g., indicated lower SO₃/H₂SO₄ than measured in the ash-free laboratory environment), due to reactions of SO₃ or sulfuric acid with fly ash in the filter thimble holder upstream of the CCS condenser. Conversely, the EPRI tests indicated that use of CCS downstream of the particulate control device produces unbiased measurements, as long as appropriate procedures are followed.

The extent of bias observed in the EPRI study depended on the sampling location, which could be the air heater inlet, air heater outlet, or flue gas desulfurization outlet. Furthermore, the bias depended on the quantity of ash collected, the alkalinity of the ash, and the coal sulfur content (e.g., SO₂ content in the sample gas). Measurement bias in the simulated ash environment was observed for both high sulfur coal and low sulfur Powder River Basin (PRB) coal; however biasing in the PRB coal tests can generally be considered non-significant in light of already low SO₃ concentrations and the expected absorption of sulfuric acid with the alkaline ash in actual field tests. Data for high sulfur coal suggested a low bias of 20-25% could occur in measurement

locations upstream of a particulate control device with typical ash loadings, and no significant bias downstream of the particulate control device.

The precision of the CCS data used in this estimation model can be estimated from the variability among duplicate measurements believed to be conducted under the same process conditions. Data from recent field trials with experienced test crews suggest a precision of up to \pm 20% is realistic for most conditions. The repeatability of the CCS measurements depends on avoiding stratification within the gas stream (traversing is very difficult), maintaining correct temperatures in the sampling train, and having well trained sampling personnel. Therefore, the precision of the entire data set used in this model, which includes data from older studies of unknown quality, is expected to be closer to \pm 50%.

Representativeness

The accuracy of the correlations presented in this report depends on the representativeness of the constituent data points. For some power plant configurations, only a few measurements are available, and the correlations provide only rough approximations of the flue gas SO₃/H₂SO₄ releases for similar units. The category with the largest number of data points is dry-bottom boilers, firing low sulfur, eastern bituminous coal. In general, the number of SO₃ measurements for various power plant configurations are proportional to the occurrence of those configurations among the total population of U.S. EGUs. The number of measurements in each power plant category should be considered when applying this model; factors based on one measurement have much higher uncertainty than factors based on dozens of measurements.

As a consequence of the uncertainty in SO_3/H_2SO_4 measurements and the limited number of data points that comprise the relevant correlations, the predictive technique should be assumed to provide estimates within a \pm 50% relative accuracy.

3 BACKGROUND AND METHODOLOGY OVERVIEW

This section provides a brief overview of sulfuric acid production mechanisms in power plants and presents the basic equations for estimating sulfuric acid releases. A detailed review of this topic has been provided by Srivastava (2004) and more recently by Monroe (2006).

The production or manufacture of SO₃, and ultimately sulfuric acid, is determined initially by processes occurring directly within the flame zone of a boiler, the convective heat transfer sections, or environmental control components. These production or manufacturing processes can be gas phase or can be induced by catalysis on the surface of fly ash particles or heat removal surfaces.

Mechanisms that remove sulfuric acid from the flue gas also can occur both in the gas phase and on heat transfer surfaces, usually promoted by ash-derived deposits. The amount of sulfuric acid released from the stack is the aggregate result of production mechanisms, generally occurring at temperatures above 650°F, and removal mechanisms, including alkali-based sorbent injection, occurring at lower temperatures.

Description of General Methodology

The method estimates the manufacture and the subsequent release of sulfuric acid from the power generation process. The *manufacture* of sulfuric acid is defined as any process step that increases the flue gas content of sulfuric acid, regardless of its ultimate fate. To estimate the *release* of sulfuric acid, the manufacture estimate is corrected by a factor or series of factors to account for sulfuric acid removal within the system.

Three process steps ultimately lead to the manufacture of sulfuric acid in a fossil fuel-fired electricity generation unit: the combustion process, selective catalytic reduction (SCR) for NOx control, and flue gas conditioning. All remaining steps in the power generation process reduce sulfuric acid. Therefore, the total sulfuric acid released (TSAR) is the cumulative sum of the total sulfuric acid manufactured (TSAM), adjusted by sulfuric acid losses.

Sulfuric Acid Manufacture from Combustion

A detailed analytical study by Senior (2002) explored details of SO₂ oxidation to SO₃. Specifically, the author modeled the relevant reaction sets employing the temperature-time history of a typical utility boiler, considering gas phase events only and ignoring catalytic effects. The results showed that insignificant SO₃ formed in the early stage of the flame, but SO₃ production increased as the gases exited the flame zone and cooled. The author reported production of SO₃ to be essentially complete prior to flue gas entering the economizer section. The observation that most SO₃ forms during cooling from post-flame temperatures (2,900-3,100°F) to temperatures typical of the economizer entrance was also noted by Buckley (2002). Senior further quantified the relationship between coal sulfur content, excess air, and SO₃ production that flame four seconds of residence time, SO₃ production could range from between 0.3 - 0.6% of the flue gas SO₂ concentration. In the absence of SCR NOx control or FGC, the boiler or turbine combustion process comprises the only source of sulfuric acid. The sulfuric acid manufacture by the combustion process (EM_{Comb}) is expressed as follows:

$$EM_{Comb} = K \bullet F1 \bullet E2$$
 Eq. 3-1

Where K is a constant, F1 is a Fuel Impact Factor describing the SO₃ production associated with combustion for a specific fuel type, and E2 is the annual rate of SO₂ production. Further details of this relationship and the derivation of constants and other terms are presented in Section 4. There are two situations where the TSAR equations must be modified to reflect reduction or removal of sulfuric acid from combustion: 1) the use of a hot-side ESP and 2) magnesium oxide (Mag-Ox) addition to the boiler or fuel. All other sources of sulfuric acid (SCR or FGC) are manufactured downstream of these two technologies and require no modifications. The derivation and use of technology impact factors for hot-side ESP and Mag-Ox addition are discussed in Section 4.

Sulfuric Acid Manufacture from SCR/SNCR NOx Control

The SCR process increases the production of SO₃, and therefore, H_2SO_4 , as a percentage of SO₂. SO₃ production can range from as low as 0.3%, for the most recent catalyst technology, to as high as 3% of the SO₂ concentration.

The production of SO₃ from SO₂ is a well-known consequence of SCR, and the degree of SO₂ oxidation is a performance variable that should be part of catalyst procurement specifications. To a degree, SO₃ production can be mitigated by reducing the content of vanadium pentoxide and compensating with other catalyst oxides such as tungsten or molybdenum, which offer improved selectivity (e.g., lower SO₂ oxidation) but also lower activity for NOx removal. Alternatively, advanced catalyst formulations can be used that concentrate the vanadium-containing active ingredients on the catalyst surface and minimize content within the substrate, lowering SO₂ oxidation. To achieve an equivalent level of NOx and residual ammonia slip, a low SO₂ oxidation catalyst may require a larger volume of catalyst than one for which SO₂ conversion is not constrained. In general, the degree of SO₂ oxidation ranges from about 0.2% to 1.5% for most SCR process reactors designed for bituminous coals. If SCR is applied to extremely low sulfur, high alkalinity coals such as PRB, higher SO₂ oxidation is acceptable. Depending on how the measurement of SO₂ oxidation is performed, the oxidation can be 3% or higher.

In many instances, owners and catalyst suppliers prefer that measurements to quantify SO₂ oxidation for the purpose of commercial guarantees be conducted not with authentic flue gas but in a laboratory-scale test facility, without fly ash present. This approach has the advantage of improved accuracy and repeatability of the SO₃ concentration. The measurements are generally conducted using CCS. However, particularly for PRB coals, this method will over-predict SO₃ manufactured by the SCR, as the mitigating role of high fly ash alkalinity is not considered. Consequently, the role of fly ash alkalinity must be considered in predicting SO₃ manufactured by SCR.

The observation that calculated laboratory-scale SO₃ values at the exit of an SCR reactor exceed measurements in the field has been witnessed on numerous occasions. Specifically, Lindenhoff (2004) reported that the measured SO₂ oxidation rate for PRB-generated flue gas of 1.25% was below the 1.6% value predicted based on the boiler and SCR catalyst specifications. Further,

tests conducted by this catalyst manufacturer noted significant absorption of SO₃ generated within the SCR reactor prior to the air heater. Sufficient data was accumulated to derive both a mathematical model and to project a graphical relationship describing SO₃ removal within an SCR reactor as a function of both ash content and the number of alkaline sites available in the flue gas stream.

An alternative NOx control process, selective non-catalytic reduction (SNCR), does not use a catalyst and does not promote the oxidation of SO₂ to SO₃. Accordingly, there is no manufacture of sulfuric acid attributable to SNCR.

The following relationship describes the total H₂SO₄ manufactured from an SCR equipped utility boiler or steam generator:

$$EM_{SCR} = K \bullet S2 \bullet f_{sops} \bullet E2 \bullet F3_{SCR}$$
 Eq. 3-2

Where K is a constant, S2 is the catalyst SO₂ oxidation rate, f_{sops} is the fraction of the coal burn in which flue gas is directed through the SCR, E2 is the annual rate of SO₂ production, and F3_{SCR} is the Technology Impact Factor for SCR used to adjust for the presence of alkaline fly ash. Further details of this relationship and the derivation of constants and other terms are presented in Section 4.

Sulfuric Acid Manufacture from Flue Gas Conditioning

Flue gas conditioning (FGC) is a process that is typically used in power plants to assist in particulate control in an ESP or baghouse. The conditioning additives can be any of the following: SO₃, SO₃ plus NH₃, or NH₃ alone. In SO₃-based FGC, SO₃ is introduced into the flue gas either preceding or following the air heater. The SO₃ is typically produced on-site from sulfurous fuel that is burned to produce SO₂, which is then catalytically oxidized to SO₃ with a conversion rate typically exceeding 95%. When injected into the flue gas, the SO₃ immediately reacts with water vapor to create sulfuric acid, thus resulting in the manufacture of sulfuric acid. Estimating the manufacture source requires knowledge of the concentration of SO₃ injected and the associated oxygen content of the flue gas.

Equation 3-3 estimates the manufacture of sulfuric acid from FGC, where K_e is a constant, B is the coal burn, f_e is the FGC operating factor, and I_s the SO₃ injection rate. F3_{FGC} is the Technology Impact Factor for FGC that is used to adjust for the presence of alkaline fly ash in PRB coals. Further details and derivation of the constant are provided in Section 4.

$$\mathsf{EM}_{\mathsf{FGC}} = \mathsf{K}_{\mathsf{e}} \bullet \mathsf{B} \bullet \mathsf{f}_{\mathsf{e}} \bullet \mathsf{I}_{\mathsf{s}} \bullet \mathsf{F3}_{\mathsf{FGC}} \qquad \qquad \mathsf{Eq. 3-3}$$

Methodology to Estimate Release

Sources of sulfuric acid manufactured upstream of the air preheater are calculated separately (modified for hot-side ESP or Mag-Ox if necessary), then summed. The combined SO₃ is modified to reflect reactions of SO₃ with residual equivalent NH₃ slip from SCR/SNCR equipment and/or FGC ammonia injection to form ABS, and then adjusted by removal in applicable downstream equipment such as the air preheater, electrostatic precipitator or other particulate control device (PCD) and flue gas desulfurization (FGD) equipment. In the calculations to determine the amount of ABS formation, ammonia slip is subtracted from sulfuric acid. However, the 1:1 ratio of H₂SO₄ to NH₃ is on a molar basis. Therefore, all the following

calculations of ammonia slip are actually the amount of ammonia slip that is equivalent to sulfuric acid based on a 1:1 molar ratio, as determined through the use of the constant K. The adjustments are made using Technology Impact Factors (F2) which describe the fraction of sulfuric acid that penetrates each component.

The methodology employs the follow steps, combined into a single release equation, which is shown in Equation 3-4, and illustrated schematically in Figure 3-1:

- 1. Calculate the sources of sulfuric acid upstream of the APH from combustion, SCR and FGC
- 2. If applicable, apply the technology factor for sulfuric acid removal to the sulfuric acid manufactured from combustion only, for hot-side ESP and/or Mag-Ox addition
- 3. Sum the resulting sources of sulfuric acid upstream of the APH from combustion, SCR and FGC
- 4. Calculate residual equivalent ammonia slip from the SCR/SNCR and FGC and subtract this ammonia from the sum of the sources upstream of the APH to account for ABS formation which is not reportable
- 5. If the result of Steps 1) and 2) is a positive value, apply the technology factor for APH removal of SO₃ (F2_{APH})
- 6. Add downstream sources of sulfuric acid (from FGC, if injected downstream), and subtract downstream ammonia injection (from FGC, if injected downstream) to account for ABS, which is not reportable
- 7. Apply all remaining relevant downstream technology factors

In equation form, the methodology is expressed as shown in Equation 3-4.

 $TSAR = \{[((EM_{Comb} * F2_{Hotside}_{ESP} * F2_{Mag-ox}) + EM_{SCR/SNCR} + EM_{FGC_{beforeAPH}}) - (NH3_{SCR} + NH3_{FGC_{beforeAPH}})] \bullet F2_{APH} + (EM_{FGC_{afterAPH}} - NH3_{FGC_{afterAPH}})\} \bullet F2_{x}$ Eq. 3-4

where,

TSAR	=	Total sulfuric acid released
EM _{Comb}	=	Sulfuric acid manufactured through combustion
EMscr/sncr	=	Sulfuric acid manufactured from SCR or SNCR
EM _{FGC_beforeAP}	Ή	= Sulfuric acid manufactured from FGC injected upstream of the air preheater (APH)
EMFGC_afterAPH	=	Sulfuric acid manufactured from FGC injected downstream of the APH
NH ₃ Slip	=	Calculated equivalent ammonia slip that reacts with H ₂ SO ₄ to form ammonium bisulfate (ABS) from SCR/SNCR and/or FGC
F2 _{арн}	=	Technology Impact Factor for APH
$F2$ Hotside_ESP	=	Technology Impact Factor for hot-side ESP

	= 0.63 if hot-side ESP is present
	= 1.0 if hot-side ESP is <u>not</u> present (optional)
F2 _{Mag-Ox}	= Technology Impact Factor for magnesium oxide addition in fuel or furnace
	= 0.25 (or custom) if Mag-Ox is sprayed into furnace
	= 0.50 (or custom) if Mag-Ox is mixed with fuel
	= 1.0 if Mag-Ox is <u>not</u> applied (optional)
F2 _x	= Technology Impact Factors, all others that apply (downstream of APH)

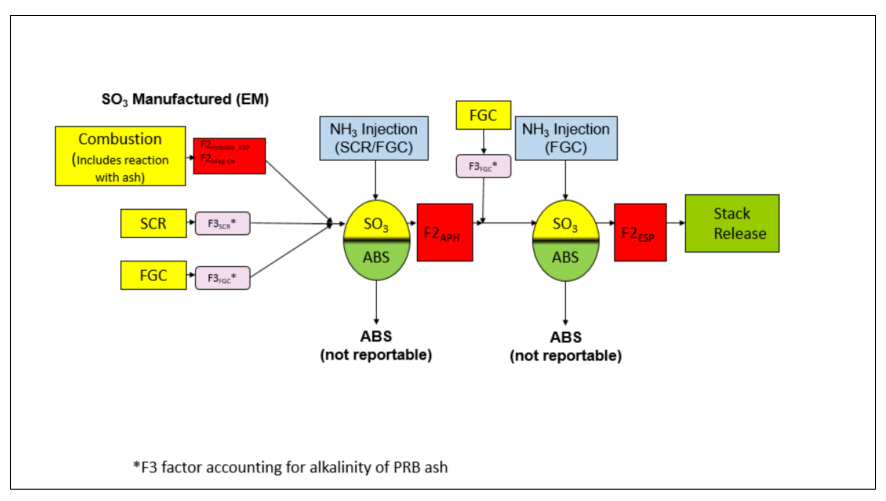


Figure 3-1 Schematic of Methodology Used to Estimate Sulfuric Acid Releases

Ammonia from SCR/SNCR NOx Control

Both SCR and SNCR can introduce residual ammonia (NH₃), also called ammonia slip, into the flue gas. This residual ammonia can react with SO₃ or sulfuric acid to form ammonium sulfate and/or ammonium bisulfate, thereby removing from the flue gas some of the SO₃ that is a precursor to sulfuric acid, and reducing the amount released.

For an SCR, ammonia slip is expected to usually range between 0 and 2 parts per million (ppm) for bituminous coal cases. Higher values of ammonia slip are possible if the unit has not been recently tuned, the boiler or SCR process operates under upset, or as the catalyst ages. Higher ammonia slip may be tolerable in PRB coal applications, since ammonia is not readily absorbed by the highly alkaline PRB ash and adverse operational impacts (air preheater pluggage and ash contamination by ammonia) are avoided. Reactions between the residual ammonia and SO₃ or H₂SO₄ occur in the air preheater and result in a solid product that may deposit or accumulate on the surface of the fly ash. Any SO₃ or H₂SO₄ that participates in these reactions is no longer present as sulfuric acid and is not required to be reported to the EPA TRI program as a release of sulfuric acid.

SNCR employs ammonia or urea injection in the upper furnace for NOx reduction. The higher temperatures characteristic of the upper furnace zone (1,800 to 2,400°F, as compared to 700°F for SCR) do not require reduction catalyst so there is no additional manufacture of SO₃ or sulfuric acid. Typically, SNCR systems operate with higher levels of residual ammonia (5 to 10 ppm), so any sulfuric acid that is present from combustion is reduced through reaction with the ammonia. Consequently, SNCR systems will always *reduce* the overall amount of sulfuric acid released, while SCR systems can *reduce* but will likely *increase* the amount of sulfuric acid released.

The reactions between SO₃ and/or H₂SO₄ and ammonia produce ammonium sulfate [(NH₄)₂SO₄] and/or ammonium bisulfate (NH₄HSO₄). While both chemicals are solids, the bisulfate pairs one molecule of ammonia with one of sulfuric acid, while the sulfate requires two ammonia molecules for each sulfuric acid. Which reaction product is present is usually determined by the stoichiometry, or the relative amount of each substance on a mole basis, that is present in the flue gas. When ammonia is present in an amount over twice the mole content of sulfuric acid, the reaction product will always be ammonium sulfate (AS). Conversely, when sulfuric acid is present in an amount on a mole basis greater than ammonia, the product will be ammonium bisulfate (ABS). Between these two extremes, a mixture of ammonium sulfate and bisulfate is produced.

A SCR-equipped unit firing bituminous coals with low-to-medium sulfur content will always produce an excess of sulfuric acid over ammonia slip. Accordingly, ammonium bisulfate is the primary byproduct anticipated. For subbituminous and lignite coals, any sulfuric acid produced is typically adsorbed by the ash and it is likely that residual ammonia will exceed sulfuric acid content on a mole basis. Under these conditions, ammonium sulfate is the likely product.

For SNCR-equipped units, ammonia slip levels of 5 ppm and possibly higher will favor ammonium sulfate, particularly for western U.S. coals where the alkaline fly ash will reduce the SO₃ content. For eastern bituminous coals, which have generally higher sulfur content and lower ash alkalinity, ammonium bisulfate will likely predominate.

For the purpose of predicting sulfuric acid emissions, these distinctions are not important. A single molecule of sulfuric acid will capture a single molecule of ammonia, producing ammonium bisulfate. If additional ammonia is available, the ammonium bisulfate can react with another ammonia molecule to form ammonium sulfate. It can be assumed that all sulfuric acid forms ammonium bisulfate before any further reaction to the ammonium sulfate form occurs. This assumption leads to the calculation strategy that ammonia captures all of the sulfuric acid it can as ammonium bisulfate, and since the bisulfate form is no longer reportable, the sulfuric acid disappears from the release calculation. Additional ammonia reacting with the ammonium bisulfate is of no consequence to the sulfuric acid calculation – although this reaction is important when estimating ammonia releases.

The total sulfuric acid released is reduced by the residual equivalent ammonia, or ammonia slip, from SCR or SNCR. This equivalent ammonia slip is calculated as follows, and is added to any ammonia injection from FGC before it is subtracted from the sum of the sulfuric acid manufactured.

$$NH3_{SCR} = K_s \bullet B \bullet f_{sreagent} \bullet S_{NH3}$$
 Eq. 3-5

Where K_S is a constant, B is the coal burn in TBtu/yr, $f_{sreagent}$ is the fraction of SCR operation with reagent injection, and S_{NH3} is the NH₃ slip for the SCR/SCNR. This calculation and factors are described further in Section 4.

Ammonia from Flue Gas Conditioning

Flue gas conditioning may involve injection of NH₃ alone or NH₃ with SO₃. Thus, FGC can result in sulfuric acid manufacture or may result in sulfuric acid removal, if NH₃ is injected alone or in quantities greater than the sulfuric acid. The following equation describes the calculation of equivalent ammonia that is used to reduce sulfuric acid. If ammonia is injected upstream of the APH, this value would be added to the residual ammonia from SCR/SNCR. If added downstream of the APH, the value would be subtracted from any downstream SO₃ injection introduced by FGC.

$$NH3_{FGC} = K_e \bullet B \bullet f_e \bullet I_{NH3}$$
 Eq. 3-6

Where K_e is a constant, B is the coal burn in TBtu/yr, f_e is the fraction of coal burn with FGC operation, and I_{NH3} is the NH₃ injected. This calculation and factors are described further in Section 4.

Estimating Release with Alkali Injection

Many utility operators inject alkali into the flue gas to control SO₃. A wide variety of alkali materials based on calcium, sodium, or in some cases magnesium compounds, have been used with varying degrees of effectiveness to remove SO₃. These alkali materials include sodium bisulfate, trona, and various types of lime (conventional, hydrated, and magnesium enhanced).

Sodium bisulfate (SBS) is the most widely used reagent for control of SO₃. SBS is introduced into the flue gas, usually at the air heater exit, but in some units at the air heater inlet, to maximize residence time and improve SO₃ removal. SBS can be highly effective in removing SO₃. Additional sodium can be added to the flue gas to compensate for any compromise in ESP

performance due to increased ash resistivity associated with SO₃ reduction. A detailed summary of experience with SBS usage has recently been published (Moser, 2007).

Trona is another sodium-containing reagent that is used to control SO₃. Trona can be injected at several locations within the boiler, but most applications introduce trona at the air heater exit, prior to the particulate collector (Ritzenthaler, 2006). As with SBS, additional sodium introduced into flue gas is advantageous to compensate for the reduction in ESP performance.

Several forms of calcium-based sorbents can be utilized. These are injected at various locations in the flue gas, including the economizer inlet, economizer outlet (e.g., air heater inlet), and the air heater outlet. A survey of recent experience with lime-derived sorbents addressed the advantages of this approach (Benson, 2006a). Hydrated lime has been shown to be an effective calcium-based reagent for SO₃ control (Gale, 2006), as has lime supplemented with magnesium hydroxide (Benson, 2006b). Calcium–based sorbents can be very effective in reducing SO₃, but may compromise ESP performance, as the reduced SO₃ content in flue gas increases the electrical resistivity of the ash on the collecting plate. For this reason, some operators have proposed using both calcium- and sodium-based sorbents, the latter to both augment SO₃ removal and promote ESP performance. In summary, any of several alkali materials can be used to control SO₃, with the least cost solution depending on the plant and access to low cost sorbent.

Plant units that deploy alkali injection will generally have conducted field tests to determine the SO_3 concentration in flue gas for a specified sorbent injection rate. In most cases, these tests will be conducted at the stack. The alkali injection system usually is operated to reduce SO_3 emissions to between 5 and 15 ppm, an optimal range to prevent formation of a visible plume. A procedure to convert measured sulfuric acid concentration in flue gas to a mass emission rate is summarized below and described in more detail in Section 4. An example application of this procedure is provided in Appendix A.

$$ER_{ALKINJ} = K_{alkali} \bullet B \bullet S_{SO3} \bullet F2_x \bullet F_{alkali}$$
 Eq. 3-7

Where K_{alkali} is a constant, B is the coal burn, S_{SO3} is the measured SO₃ content with injection, F_{alkali} is the fraction of time alkali injection is in operation, and F2 are any applicable downstream Technology Impact Factors.

Users of this approach should be cautioned that the current EPA-approved test method for SO₃ (EPA Method 8) has a significant positive bias, and that corrections for this bias may be needed to accurately assess flue gas concentrations. A more accurate method is the controlled condensate system (CCS). Although no general-purpose, EPA-approved CCS method is currently available, it is widely used by stack test contractors for sulfuric acid measurement and is more accurate than Method 8, if performed correctly.

Some units will not have measured stack emissions. This version of the model adds an alternate approach to estimate releases with alkali injection if stack SO₃ measurements are not available. Instead of using Equation 3-7 for ER_{ALKINJ}, the total sulfuric acid release is calculated by estimating total releases using Equation 3-4. An alkali injection factor, F3_{ALKINJ}, is then applied to the total release as shown in Equation 3-8. This F3_{ALKINJ} is either the expected fractional reduction in SO₃ (generally as guaranteed by the vendor) or a default of 0.2. The basis of the 0.2 value is test results indicating that 80% removal is easily achievable (EPRI, 2010b). This method is explained in further detail in Section 4.

To estimate $SO_3 \underline{manufacture}$ with alkali injection, plant owners should use the standard approaches given in Section 4 for the appropriate plant configuration. At this time, EPRI does not have sufficient information to estimate the impact on SO_3 formation of adding alkali at various points in the fuel combustion and stack gas treatment process. Assuming that there is no reduction of SO_3 formation is conservative for a <u>threshold</u> determination for TRI reporting purposes.

Estimating Sulfuric Acid with Partial-Year SNCR or SCR Operation

When ammonia or urea is employed in SCR or SNCR during only part of a year, the methodology outlined in this report may not provide an accurate estimate of total sulfuric acid released through the stack. The reason is that the model calculates the pounds of ammonia injected on an annual basis, and uses that quantity to subtract an equivalent quantity of sulfuric acid as non-reportable ammonia salt. However, if there is an excess of ammonia slip present during SCR/SNCR operation above the amount of SO₃ available to react, the model is in effect borrowing SO₃ from periods when the SCR/SNCR is not operating. As a result, the annual release of sulfuric acid will be underestimated for that unit. Underestimation of sulfuric acid releases is most likely to when the following conditions are met:

- SNCR (more likely) or SCR (less likely) operated partial-year
- Sulfuric acid emissions are calculated from fuel sulfur, not CEMS
- The operating factor for reagent addition in the SCR/SNCR system (fsreagent) is small
- The unit burns a fuel that inherently manufactures low levels of SO₃ from combustion, such as PRB coal
- The unit has high ammonia slip levels when ammonia is injected

To determine whether the model is underestimating releases for a unit with SCR or SNCR partial-year operation, the user should perform separate calculations for the SCR/SNCR "Ammonia On" condition and SCR/SNCR "Ammonia Off" condition, allocating the amount of fuel burned during the two conditions appropriately. If the sum of the sulfuric acid releases for these two conditions is greater than for the full-year calculation, separate calculations should be employed going forward.

4 ESTIMATING GUIDELINE: STEAM BOILERS

This section describes an estimation procedure for calculating the manufacture and release of sulfuric acid from coal-fired steam generators. The topics addressed are (a) formation within the furnace; (b) the role of SCR, (c) flue gas conditioning (FGC), and (d) alkali injection; and (e) removal by downstream equipment such as air heaters, ESPs or other particulate control devices, and FGD process equipment. Each of the subsequent sections in this chapter provides the information to conduct this stepwise calculation.

Sulfuric Acid Manufacture

Sulfuric Acid Manufactured by Combustion (EMcomb)

The premise of the methodology is that the amount of sulfuric acid manufactured by the boiler is a function of the amount of SO_2 produced – which may be determined either from coal usage (amount burned and sulfur content) or from continuous emission monitoring system (CEMS) output. Units equipped with FGD equipment or other methods of SO_2 control are required to estimate emissions either from coal data or from the output of a CEMS positioned ahead of the scrubber. As described below, it may be necessary to correct SO_2 CEMS data for non-ideal stack flow conditions.

The following relationship is used to estimate the sulfuric acid manufactured from combustion in utility sources:

$$EM_{Comb} = K \bullet F1 \bullet E2 Eq. 4-1$$

where,

EM_{Comb}	=	total H ₂ SO ₄ manufactured from combustion, lbs/yr
Κ	=	Molecular weight and units conversion constant 98.07 / 64.04 • 2000 = 3,063
		$98.07 = Molecular weight of H_2SO_4$
		$64.04 = Molecular weight of SO_2$
		2000 = Conversion from tons per year to pounds per year.
F1	=	Fuel Impact Factor
E2	=	Sulfur dioxide (SO ₂) emissions, either: (1) recorded by a continuous emissions monitor, tons/yr, or (2) calculated from coal burn data, tons/yr.

The Fuel Impact Factor (F1) is a numerical value or a mathematical relationship reflecting the conversion of SO_2 to SO_3 in the boiler for a specific coal type and boiler type. In the derivation of Equation 4-1, the following assumptions are made:

- SO₃ concentrations are proportional to SO₂ concentrations.
- The grade of coal being burned impacts the rate of conversion from SO₂ to SO₃.
- All SO₃ that forms is converted to H₂SO₄.
- The rate of SO₃ formation is independent of the boiler firing rate (unit load).

The estimates of sulfur dioxide emissions (E2) and F1 factors are further described in the following sections.

Sulfur dioxide (SO2) Emissions from Combustion (E2)

Estimating the sulfuric acid production from Equation 4-1 requires knowledge of the mass rate of SO_2 generated. This rate is designated as E2 in Equation 4-1. As noted previously, the value of E2 can be estimated from either (a) the EPA CEMS data, or (b) calculated from the coal burn data (EPA, 1995a).

If the CEMs data are used to directly quantify the annual SO2 production rate in tons/year of SO_2 , it may be necessary to correct for stack geometry, depending on the measurement method used. The user should check with their CEMs operator to obtain the bias correction factors, or the bias corrections may already be taken care of in the instrument software.

If a facility used EPA Methods 1 and 2 for CEM flow monitor setup and validation under the guidelines described in 40 CFR Part 60, Appendix A, flow bias and wall effects corrections should be made to the CEMs SO2 measurements to accurately estimate sulfuric acid releases. This correction can be made using one of the following two equations:

$$E2 = E\left[1 - \frac{(C1 \bullet R^{2} + C2 \bullet R + C3)}{100}\right]$$
Eq. 4-2

where,

- $E2 = Corrected SO_2$ mass rate, tons per year
- E = CEMS-derived SO₂ mass rate, tons per year
- C1 = 0.0264 (non-axial flow bias factor)
- C2 = 0.183 (non-axial flow bias factor)
- C3 = 1.5 (wall effects bias factor) default value for cylindrical stacks
- R = Stack/duct average resultant angle (or swirl angle) from site verification tests, degrees

The average resultant angle, R, is determined using Method 1. The wall effects bias factor, C3, is the percent difference between flow rates calculated using standard Method 1 sampling and flow rates calculated using measured near-wall velocity data. Many utilities do not have bias data

available, and as such, an average default value of 1.5, derived from actual data, can be used. Measured bias factors were within the range of 0.9 to 2.2% (Noble, 1998).

If a facility used EPA Method 2H (EPA, 1999) for round stacks, or CTM-041 (EPA, 2003) for rectangular stacks to determine a Wall Adjustment Factor (WAF), then this value can be used instead and the correction equation can be simplified to the following:

where,

 $E2 = Corrected SO_2$ mass rate, tons per year

E = CEMS-derived SO₂ mass rate, tons per year

WAF= Wall Adjustment Factor, determined from Method 2H

The WAF correction factor determined by Method 2H is applicable to stack flow rates determined by Methods 2, 2F and 2G. A default value of 0.995 can be used, although actual data should be used if available.

As an alternative to using CEMS data, the following relationship based on coal burn data can be used to estimate the rate of SO₂ emissions:

$$E2 = K1 \bullet K2 \bullet C1 \bullet S1 \qquad \qquad Eq. 4-4$$

where,

 $E2 = SO_2$ mass rate, tons/yr

- C1 = Dry coal burn, tons/yr. The dry coal can be calculated from wet coal through the following relationship: Dry coal (tons/yr) = wet coal (tons/yr) • (1-moisture(%)/100%)
- S1 = Coal sulfur weighted average, %, dry
- K1 = Molecular weight and units conversion constant= (64.04)/(100 • 32.06) = 0.02 64.04 = molecular weight of SO₂ 32.06 = molecular weight of S 100 = conversion of % S to a fraction

K2 = Sulfur conversion to SO₂, implicit from EPA AP-42 (EPA, 1995b)

= 0.95 for bituminous coals

= 0.875 for subbituminous coals

- = 0.55 to 0.85 for lignite, based on the Na content
- = 1.0 for oil

When any source uses FGD equipment or another technology to control SO₂ emissions, either the fuel basis must be used for the manufacturing and release calculations, or CEMS data can be used, but only when the CEMS precedes the FGD or SO₂ control equipment. <u>Data from a CEMS</u> located after a flue gas desulfurization system cannot be used, because the measured SO₂ has already been decreased by the control equipment, and therefore is not an accurate predictor of the SO₃ emissions rate.

<u>Fuel Impact Factor (F1).</u> Figure 4-1 depicts the fraction of H_2SO_4 produced as a function of flue gas SO₂ content for several coal sources and boiler types. The coal ranks and boiler types consist of (a) high sulfur (>2.5%) eastern bituminous coal, fired in a dry bottom boiler (wall-fired or cell-fired), (b) low sulfur eastern bituminous coal, fired in a dry bottom boiler, (c) PRB coal, fired in both a cyclone and dry bottom boilers. Data are also shown for one unit that fires 75% lignite with the balance PRB and one unit that fires 100% lignite.

Figure 4-1 shows that a wide range in SO₃ production is observed for all coals and boiler types. This range exceeds the theoretical predictions by Senior (2000), suggesting that the role of ash in either catalyzing SO₃ production or absorbing/neutralizing SO₃ is not fully accounted for. The only consistent results in Figure 4-1 are for SO₃ from PRB-fired units, regardless of boiler type, in that less than 1 ppm was observed.

The current model uses a linear relationship to estimate SO₃ production for all eastern bituminous, dry bottom boiler data, as shown in Figure 4-1. The equation expressing this relationship is shown in Equation 4-5. This relationship is significant at the 95% confidence level, although the correlation coefficient (R2) is low (0.13) due to scatter in the data. The current model retains F1 factors based on average fraction of SO₂ converted to SO₃ for all other coal types.

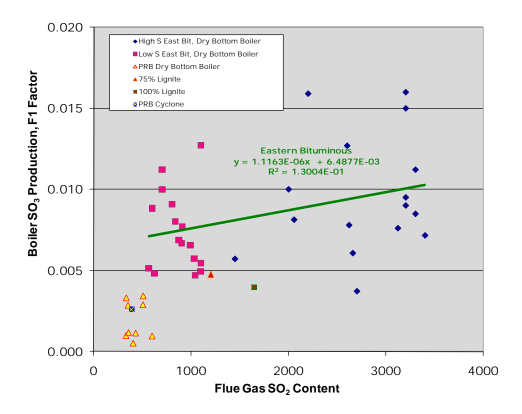


Figure 4-1 Relationship between Boiler SO₃ Production and Flue Gas SO₂ (corrected to 3% O₂)

$$F1_{ebit} = 1.12E-6 \bullet SO_2 + 0.0065$$
 Eq. 4-5

where,

F1_{ebit} = Fuel Impact Factor for all Eastern Bituminous coals burned in a drybottom boiler

 SO_2 = Boiler SO_2 concentration (ppm), derived from fuel sulfur content (%)

In order to use this approach, it is necessary to relate coal sulfur (%) to the SO₂ concentration in the boiler. Equation 4-6 is used to calculate that relationship:

$$SO_2 = S1 \bullet \frac{K_{F1}}{HV}$$
 Eq. 4-6

where,

 $SO_2 = Boiler SO_2$ concentration (ppmvd, 3% O_2 , dry) derived from fuel sulfur content (%)

S1 = Coal sulfur weighted average, %, dry

 K_{F1} = Conversion factor = 10,003,602

HV = Coal heating value, Btu/lb, dry

The conversion factor K_{F1} considers all relevant constants to yield the result in ppm of SO₂. The derivation of this constant is presented in Text Box A.

<u>Text Box A: Derivation of Conversion Factor, K_{Fl} .</u> The U.S. Code of Federal Regulations 40, Part 60, Table 19-1 "F Factors for Various Fuels" states that 1 million Btu of heat input for bituminous or subbituminous coal will produce 10,640 wet standard cubic feet of flue gas, defined at 0% oxygen and on a wet basis at 20°C and 760 mm Hg. Correcting this volume to 3% O₂ on a dry basis (typical of the SO₂ measurement data used in the Fuel Impact Factors correlation) yields a volume of 11,419 scf. The standard volume of one pound mole of any gas is 359 scf, defined at 0°C and 760 mm Hg. Converting this to the English units standard of 20C (68°F), one pound mole occupies 385.5 standard cubic feet. The value 0.95 is the AP-42 (EPA, 1995b) K2 factor for sulfur conversion to SO₂ for a bituminous coal.

Equation 4-4 above can then be expanded as follows:

S1%	0.95	385.5 scf S	1 lb mol S	1 MBtu	1 lb fuel	10 ⁶ Btu	10 ⁶ ppm
	100%	1 lb mol S	32.06 lbs S	11,419 scf	HV Btu	1 MBtu	

Grouping terms,

 $SO_2 = S1 \bullet 10,003,602 / HV$

Therefore, the value of K_{F1} is equal to 10,003,602 [ppmvd SO₂ (3% O₂, dry) • Btu/lb]/%S.

Table 4-1 summarizes the F1 factors. There are no changes to these factors from the previous version of the model (EPRI, 2012).

Table 4-1 Summary of Fuel Impact (F1) Factors for Steam Generating Units

Fuel	Equipment	F1	Comment
E. Bituminous (all)1	Dry Bottom Boiler	Slope: 1.1163E-6 Intercept: 0.0064877	F1 = slope SO2 + intercept 32 data points
Med-High S Eastern Bituminous (>2.5%)	Cyclone	0.016	One data point.
W. Bituminous	Dry Bottom Boiler	0.00111	One data point.
W. Bituminous	Cyclone	0.0022	One data point.
Subbituminous/PRB	All Boilers	0.0019	Average of 8 units
Lignite	Dry Bottom Boiler	0.0044	Two data points.
Lignite	Cyclone	0.00112	One data point.
Petroleum coke	Boiler	0.04	One data point.
Natural gas	Boiler	0.01	
#2 Fuel oil	Boiler	0.01	
#6 Fuel oil	Boiler	0.025	
Used oil	Boiler	0.0175	
Natural gas	СТ	See Table 6-1	
#2 Fuel oil	СТ	See Table 6-1	
Natural gas	CC	0.0555	
#2 Fuel oil	CC	0.0555	
Other Alternative Fuels	Any	0.04	
Other Alternative Fuels, co-fired w/coal, >75% heat throughput	NA		Use Coal F1, in absence of any applicable data.

 $^1\!For$ eastern bituminous coal, a linear relationship between SO_2 and SO_3 is used instead of an average F1 factor.

Sulfuric Acid Manufactured from SCR (EMscr)

This section describes a method to estimate impacts of SCR on sulfuric acid emissions. As discussed in Section 3, SCR produces SO₃ (and ultimately sulfuric acid), while SNCR removes or reduces it due to the reaction with residual ammonia. As such, only the SCR process will be addressed in this section. The removal equations will be discussed in subsequent sections, and will address both SCR and SCNR, as residual ammonia from either process contributes to sulfuric acid removal.

The following relationship estimates the total H₂SO₄ manufactured from an SCR equipped utility boiler or steam generator:

$$EM_{SCR} = K \bullet S2 \bullet f_{sops} \bullet E2 \bullet F3_{SCR}$$
 Eq. 4-7

where,

 $EM_{SCR} = Total H_2SO_4$ manufactured from SCR, lbs per year

- K = Molecular weight and units conversion constant = $98.07 / 64.04 \cdot 2000 = 3,063$
 - 98.07 =Molecular weight of H₂SO₄
 - $64.04 = Molecular weight of SO_2$
 - 2000 = Conversion from tons per year to pounds per year.
- S2 = SCR catalyst SO₂ oxidation rate (specified as a fraction, typically from 0.001-0.03)
- f_{sops} = Operating factor of SCR system, or the fraction of coal burn when the flue gas is directed through the SCR, whether NH₃ reagent is injected to derive NOx reduction or not. This value should reflect the hours the SCR reactor processed flue gas, which will be site-specific but can be approximated by generally 0.8 for year-round peaking operation, 0.98 for year-round baseloaded operation, or 0.43 for seasonal operation
- E2 = SO_2 produced, tons per year
- $F3_{SCR}$ = Technology Impact Factor, for SCR

The Technology Impact Factor for SCR ($F3_{SCR}$) is required to adjust SO₃ content for the presence of alkaline fly ash, as described in Section 3, unless direct measurements exist for the subject unit documenting SO₃ increase across the SCR reactor.

It is important to select the correct SO_2 oxidation rate, S2, for use in Equation 4-7. Typically, the SO_2 oxidation rate from SCR catalysts can range from as low as 0.3% of flue gas SO_2 content, up to 3% for low sulfur, highly alkaline coals (e.g., PRB). There are two means by which SO_2 oxidation is specified and measured for SCR process equipment, and each has significantly different implications for SO_3 produced. One method – based on actual field tests in commercial units – does not require adjustment and can be used as reported to predict sulfuric acid. The other method is based on laboratory tests of catalyst samples in the absence of ash and alkaline

materials, and must be adjusted to account for these latter factors. The consequences of each of these methods on estimates of sulfuric acid emissions are described as follows:

<u>Full-Scale Catalyst Performance Tests.</u> Ideally, SO₂ conversion data will be determined with commercial testing of full-scale equipment. These data – measured on commercial plants under actual operating conditions – are the most authentic in reflecting actual SO₃ content, as the impact of ash alkalinity is taken into account. This approach reflects commercial process operation, and the resulting fractional oxidation rate can be used as the S2 value in Equation 4-7 without adjustment by the F3_{SCR} Technology Impact Factor.

Laboratory Bench-Scale Tests. Some owners and process suppliers prefer to specify and conduct guarantee measurements for SO₂ oxidation on laboratory-scale equipment using synthetic flue gases without fly ash. There are good reasons to select this methodology; evidence suggests SO₃ measurements in the environment of the test laboratory are more accurate and reproducible than field tests. However, this method does not expose the catalyst being tested to ash, so there is no alkaline component to mitigate or absorb any SO₃ generated. Accordingly, SO₂ oxidation reported by this method exceeds that measured in commercial practice, particularly for PRB coals. SO₂ oxidation data from bench-scale tests can be used, but must be adjusted for the role of ash alkalinity. This adjustment is provided by the Technology Impact factor F3scR.

Table 4-2 shows the $F3_{SCR}$ factors. It should be noted that the 0.17 factor for PRB coals when using laboratory-scale SO₂ conversion (S2) data without fly ash is based on measurements at two PRB-fired units. SO₃ emitted from these units was lower than specified in the catalyst guarantee, which was based on laboratory test data. At present, there are no data to support F3_{SCR} factors for other coals, so a factor of 1 (full penetration) is recommended. Additionally, if full-scale catalyst performance SO₂ conversion data is used, the F3_{SCR} factor is 1 for all coal types.

Coal Type	F3 _{scr}			
Using Laboratory-Scale SO ₂ Conversion Data (S2), Without Fly Ash				
PRB	0.17 (n = 2)			
Other Coals	1 (no data available)			
Using Full-Scale Catalyst Performance SO ₂ Conversion Data (S2)				
All Coals	1			

Table 4-2 F3_{SCR} Technology Impact Factors for SCR

SNCR does not result in the manufacture of sulfuric acid; thus, EM_{SNCR} for SNCR is by definition zero.

Sulfuric Acid Manufactured from FGC (EM_{FGC})

In order to moderate ash resistivity, most SO₃–based FGC systems attempt to maintain a fixed concentration of SO₃ in the flue gas, typically between 5 and 7 ppm by volume (ppmv). Calculations to estimate the sulfuric acid manufactured during FGC use the setpoint of the FGC system as the key input. The FGC system is assumed to operate during most of the plant's duty, except for perhaps startup and low load operation. The sulfuric acid manufactured by SO₃-based FGC equipment (EM_{FGC}) is calculated as follows:

where,

- $EM_{FGC} = Total H_2SO_4$ manufactured from FGC, lbs per year, either upstream of APH ($EM_{FGC_beforeAPH}$) and/or downstream of APH ($EM_{FGC_afterAPH}$)
- K_e = Conversion factor = 3,799, see Text Box B.
- B = Coal burn in TBtu/yr
- fe = Operating factor of FGC system: the fraction of coal burn when the FGC system operates. This value is site-specific, must be determined for each unit, but generally will be about 0.8.
- $I_s = SO_3$ injection rate in ppmv at 6% O₂, wet; generally,
 - = 7 ppmv if before the APH
 - = 5 ppmv if after the APH
- $F3_{FGC}$ = Technology Impact Factor for FGC
 - = 0.17 for PRB coal
 - = 1 for all other coals

A Technology Impact Factor for FGC (F3_{FGC}) is used to adjust the flue gas SO₃ manufacture from FGC injection to account for the presence of alkaline fly ash. The principle is the same as for the F3_{SCR} factor that is used to adjust the flue gas SO₃ produced by SO₂ oxidation rate on the SCR catalyst.

The method first reduces the SO₃ that is injected into a PRB-fired boiler using the F3_{FGC} factor to account for sorption by the alkaline fly ash. Subsequently, SO₃ from FGC is treated the same as all other sources of sulfuric acid with respect to ABS formation or reduction through control devices. The F3_{FGC} factor (0.17) is assumed to be equal to the F3_{SCR} factor. A F3_{SCR} of 1.0 is used for eastern bituminous coals (zero reduction), as those coals produce an acidic ash that does not adsorb appreciable amounts of SO₃.

The conversion factor Ke, equal to 3,799, considers all relevant constants to yield the result in pounds per year of sulfuric acid. The derivation of this constant, for the case where residual SO₃ is reported in terms of 6% oxygen and "wet" flue gas at 8.1% H₂O, is presented in Text Box B.

<u>Text Box B: Derivation of Conversion Factor, K_{e} .</u> The U.S. Code of Federal Regulations 40, Part 60, Table 19-1 "F Factors for Various Fuels" lists that 1 million Btu of heat input for bituminous or subbituminous coal will produce 10,640 wet standard cubic feet of flue gas, defined at 0% oxygen and on a wet basis at 20°C and 760 mm Hg. Correcting this volume to 6% O₂ (typical at ESP conditions) yields a volume of 14,925 scf. The standard volume of one pound mole of any gas is 359 scf, defined at 0°C and 760 mm Hg. Converting this to the English units standard of 20°C (68°F), one pound mole occupies 385 standard cubic feet. Using these in the equation above,

 $(K_e \bullet B \bullet f_e \bullet I_S) = lbs H_2SO_4 per yr =$

B TBtu	<i>Is(S_{NH3})</i> scf NH ₃	1 lb mol SO ₃	1 lb mol H ₂ SO ₄	98 lbs H ₂ SO ₄	14925 scf fg	10 ⁶ MBtu
year	10 ⁶ scf flue gas	385 scf SO ₃	1 lb mol NH ₃	1 lb mol H ₂ SO ₄	1 MBtu	TBtu

Grouping terms,

 $= \mathbf{B} \bullet \mathbf{f}_{e} \bullet \mathbf{I}_{S} \bullet 3,799$

Therefore, the value of K_e is equal to 3,799 lbs $H_2SO_4/(TBtu \bullet ppmv SO_3 @ 6\% O_2$ and wet).

Equation 4-8 specifies the concentration of SO₃ in flue gas (Is) in terms of ppmv at 6% O₂, wet flue gas basis. If the concentration of SO₃ is reported at different conditions, the value of the constant K_e will change. The following formula is used to adjust the value of the constant K_e :

$$\begin{aligned} \mathsf{K}_{\mathrm{e}} \left(\mathsf{O}_{2},\,\mathsf{H}_{2}\mathsf{O}\right) &= 3,799 \,(6\%\,\mathsf{O}_{2},\,8.1\%\,\mathsf{H}_{2}\mathsf{O}) \bullet \left[(100\text{-}8.1)/(100\text{-}\mathrm{new}\,\mathsf{H}_{2}\mathsf{O})\right] \\ &\quad \bullet \left[(20.9-6.0)/(20.9-\mathrm{new}\,\mathsf{O}_{2})\right] & \qquad \mathsf{Eq} \end{aligned}$$

. 4-9

For example, if the SO_3 concentration is quantified at a value (in ppm) that is defined at 0% oxygen and a dry basis, the value of K_e that should be used is:

 $\mathsf{K_{e}}\ (\mathsf{O}_{2},\,\mathsf{H}_{2}\mathsf{O})=3,799\ (6\%\ \mathsf{O}_{2},\,8.1\%\ \mathsf{H}_{2}\mathsf{O})\bullet [(100\text{-}8.1)/(100\text{-}0)]\ \square\ [(20.9-6.0)/(20.9-0)]$

$K_e = 2,489$

The sulfuric acid manufacture from FGC should be noted as either upstream of the APH, $\rm EM_{FGC_beforeAPH}$, or downstream of the APH, $\rm EM_{FGC_afterAPH}$. These values must be kept separate for the release equations.

FGC equipment that employs NH₃ injection alone does not manufacture sulfuric acid and therefore this amount, EM_{FGC}, would be zero. However, the injected NH₃ will reduce the total

release of sulfuric acid by reacting with SO₃ or the resultant H₂SO₄, and therefore should be used in the release equation, as discussed in subsequent sections.

Total Manufacture from All Sources

The total manufacture of sulfuric acid is the sum of the individual manufacture estimates, prior to any adjustments or modifications due to fuel conditioning or particulate deices. Therefore, if a site burns coal and uses both SCR and FGC, the amount of sulfuric acid manufactured – by combustion, SCR, and/or FGC – is summed to determine the total amount.

Accordingly, total sulfuric acid manufacture (TSAM) is estimated for a generating unit equipped with SCR and flue gas conditioning by the following equation:

Total Sulfuric Acid Manufacture (TSAM) is described by the following equation:

TSAM = EM_{Comb} + EM_{SCR} + (EM_{FGC_beforeAPH} + EM_{FGC_afterAPH}) Eq. 4-10

Sulfuric Acid Release

Both SCR and/or ammonia-based FGC may have a negative effect on sulfuric acid release, as the ammonia slip can combine with some or all of the sulfuric acid generated from combustion to form a non-reportable ammonia salt (ABS or AS). Under these conditions, ammonia-based FGC can serve to only reduce the release of sulfuric acid. Therefore, sources of ammonia are calculated in terms that are equivalent on a 1:1 molar basis as sulfuric acid, summed in relation to the APH, and used to adjust the sulfuric acid manufactured. Once adjusted by the equivalent ammonia, the remaining sulfuric acid is then adjusted by the removals in applicable downstream equipment such as the APH, ESP or other particulate control device and FGD equipment by applying Technology Impact Factors (F2) which describe the fraction of sulfuric acid that penetrates each component.

Total Sulfuric Acid Release (TSAR) is described by the following equation:

$$TSAR = \{[((EM_{Comb} * F2_{Hotside}_ESP * F2_{Mag-ox}) + EM_{SCR} + EM_{FGC_beforeAPH}) - (NH_{3SCR} + NH_{3FGC_beforeAPH})] \bullet F2_{APH} + (EM_{FGC_afterAPH} - NH_{3FGC_afterAPH})\} \bullet F2_{x} \qquad Eq. 4-11$$

where,

TSAR	= Total Sulfuric Acid Release, lbs per year
EM _{Comb}	= Total sulfuric acid manufactured from combustion, lbs per year
F2Hotside_ESP	= Technology Impact Factor for hot-side ESP
	= 0.63 if hot-side ESP is present
	= 1.0 if hot-side ESP is <u>not</u> present (optional)
F2 _{Mag} -Ox	 Technology Impact Factor for magnesium oxide addition in fuel or furnace
	= 0.25 (or custom) if Mag-Ox is sprayed into furnace

		= 0.50 (or custom) if Mag-Ox is mixed with fuel
		$= 1.0 \text{ if Mag-Ox is } \underline{\text{not}} \text{ applied (optional)EM}_{SCR} = \text{Total}$ sulfuric acid manufactured from SCR, lbs per year
$EM_{FGC_beforeAPH}$	=	Total sulfuric acid manufactured from FGC upstream of the APH
$EM_{FGC_afterAPH}$	=	Total sulfuric acid manufactured from FGC downstream of the APH
NH3scr	=	Total equivalent ammonia slip produced from SCR/SNCR, lbs per year
NH3FGC_beforeAPH	=	Total equivalent ammonia produced from FGC upstream of the APH, lbs per year
NH3FGC_afterAPH =	=	Total equivalent ammonia produced from FGC downstream of the APH, lbs per year
F2арн =	=	Technology Impact Factor for APH, applied only if subtotal for releases upstream of the APH is non-negative
F2 _x =	=	Technology Impact Factors for processes downstream of the APH, all that apply

Examples in Appendix A detail the use of these calculations. Ammonia produced from SCR/SNCR and FGC are calculated using equations from the sections below.

The application of the hot-side ESP and Mag-Ox F2 factors to the sulfuric acid manufactured from combustion are a change from all previous versions of the report where these factors were applied to the total sulfuric acid release, which also included sulfuric acid from flue gas conditioning and SCR/SNCR. However, as a hot-side ESP removes sulfuric acid upstream of the APH, and therefore prior to any subsequent manufacture of sulfuric acid via SCR or FGC, the hot-side ESP F2 factor should only be applied to EM_{COMB}. Furthermore, application of magnesium oxide compounds into the furnace or fuel has been shown to only affect sulfuric acid produced from combustion, and does not affect any downstream formation due to SCR oxidation (Blythe, 2004). These two factors are discussed in more detail in subsequent sections, but it should be noted that as with other devices, these F2 factors only need to be included if the technology or device is present, and are optional (i.e. equal to 1.0) if the technology is not present.

The F2 factor for the APH (F2_{APH}) should only be applied if the subtotal of the sum of the sulfuric acid manufactured minus the ammonia slip upstream of the APH [(($EM_{Comb} * F2_{Hotside_ESP} * F2_{Mag-ox}$)+ $EM_{SCR} + EM_{FGC_beforeAPH}$) – (NH3_{SCR} + NH3_{FGC_beforeAPH})] is a positive value. If this subtotal is negative, this would indicate that excess ammonia slip is still present after all the sulfuric acid from combustion, SCR and FGC has been consumed to form ABS. In this instance, the resulting ammonia slip should not be reduced by the F2_{APH} factor, as all the ammonia is expected to penetrate the APH. This negative ammonia slip value can then be added to any downstream FGC injection, as summarized in Equation 4-12:

 $TSAR_{ExcessNH3} = \{[((EM_{Comb} * F2_{Hotside}_ESP * F2_{Mag-ox}) + EM_{SCR} + EM_{FGC_beforeAPH}) - (NH_{3SCR} + NH_{3FGC_beforeAPH})] + (EM_{FGC_afterAPH} - NH_{3FGC_afterAPH})\} \bullet F2_x \qquad Eq. 4-12$

Text Box C summarizes calculations for a unit equipped with a FGD process that employs partial flue gas bypass, to account for the fact that the entire flue gas flow is not subject to sulfuric acid removal by the FGD process. This bypass factor is pertinent only to units that employ flue gas bypass.

 $\begin{array}{ll} \underline{Text\ Box\ C:\ Flue\ Gas\ Desulfurization\ Bypass\ Calculation.}}_{Those units\ equipped\ with\ scrubbers\ where\ some\ of\ the\ flue\ gas\ bypasses\ the\ scrubber\ should\ take\ this\ into\ account\ in\ their\ total\ release\ calculations.\ No\ credit\ for\ sulfuric\ acid\ removal\ should\ be\ taken\ for\ the\ fraction\ of\ the\ flue\ gas\ that\ bypasses\ the\ scrubber.\ Therefore,\ this\ amount\ of\ the\ flue\ gas\ should\ be\ taken\ for\ the\ fraction\ of\ the\ flue\ gas\ that\ bypasses\ the\ scrubber.\ However,\ the\ flue\ gas\ volume\ should\ still\ be\ multiplied\ by\ the\ F2\ factor\ for\ the\ scrubber.\ However,\ the\ flue\ gas\ volume\ should\ still\ be\ multiplied\ by\ the\ F2\ factor\ for\ the\ scrubber.\ However,\ the\ flue\ gas\ volume\ should\ still\ be\ multiplied\ by\ the\ F2\ factor\ for\ the\ scrubber.\ However,\ the\ flue\ gas\ volume\ should\ still\ be\ multiplied\ by\ the\ F2\ factor\ for\ the\ scrubber.\ However,\ the\ flue\ gas\ volume\ should\ still\ be\ multiplied\ by\ the\ F2\ factor\ for\ the\ scrubber.\ However,\ the\ flue\ gas\ volume\ should\ be\ used\ where\ appropriate.\ TSAR_{bypass} = \{[(EM_{Comb} + EM_{SCR/SNCR} + EM_{FGC_beforeAPH}) - (NH3_{SCR} + NH3_{FGC_beforeAPH})] \bullet F2_{APH} + (EM_{FGC_afterAPH} - NH3_{FGC_afterAPH}) \} \bullet F2_x$ $TSAR = [SB_f + (1 - SB_f) \bullet F2_s] \bullet TSAR_{bypass}\ where,\ SB_f = fraction\ of\ scrubber\ bypass,\ as\ a\ decimal factor\ f$

 $F2_x$ = All other applicable F2 factors except for scrubber.

Ammonia Slip from SCR and SNCR (NH3scr)

A key assumption in the calculation methodology is that residual ammonia from either a SCR or SNCR captures all available sulfuric acid in the (ammonium) bisulfate form. The bisulfate form is not reportable under the Toxics Release Inventory rules and thus is not part of the manufacture calculation. If any additional ammonia reacts with bisulfate to form ammonium sulfate, that reaction is of no consequence to the sulfuric acid calculation – although it needs to be taken into account when estimating ammonia releases.

The total sulfuric acid released from combustion, SCR, and FGC is determined by subtracting from the sulfuric acid manufactured the amount removed by the residual ammonia, or equivalent ammonia slip, as calculated on a 1:1 molar basis with sulfuric acid. For SCR/SNCR, the ammonia slip is calculated using the following:

$$NH3_{SCR} = K_s \bullet B \bullet f_{sreagent} \bullet S_{NH3}$$
 Eq. 4-13

where,

NH3scr	=	Total equivalent ammonia slip produced from SCR/SNCR, lbs per year
Ks	=	Conversion factor = 3799, which is equivalent to K_e (See Text Box B)
В	=	Coal burn in TBtu/yr
fsreagent	=	fraction of SCR operation with reagent injection, when residual NH ₃ is produced that will remove SO ₃ . The value of $f_{sreagent}$ will be similar to, but slightly less than, the value of f_{sops} , defined for Equation 4-6.
$\mathbf{S}_{\mathrm{NH3}}$	=	NH ₃ slip from SCR/SNCR, ppmv at 6% O ₂ , wet:

- SCR averages 0.75 ppmv over catalyst guarantee period
- SNCR averages 5 ppmv
- Note: actual NH₃ slip data should be used if available

The conversion factor K_s , equal to 3,799, is equivalent to the factor K_e used for SO₃ injected from FGC, and considers all relevant constants to yield the result in pounds per year of equivalent ammonia. The derivation of this constant, in the form of K_e , was presented previously in Text Box B.

Equation 4-13 specifies the concentration of NH₃ in flue gas (S_{NH3}) in terms of ppmv at 6% O₂, wet flue gas basis. If the concentration of NH₃ is reported at different conditions, the value of the constant K_s will change. As with K_e, Equation 4-9, described previously and repeated again below for reference, is used to adjust the value of the constant K_s:

$$\begin{split} \mathsf{K}_{s} \left(\mathsf{O}_{2},\,\mathsf{H}_{2}\mathsf{O}\right) &= 3799 \,(6\% \,\,\mathsf{O}_{2},\,8.1\% \,\,\mathsf{H}_{2}\mathsf{O}) \bullet \left[(100\text{-}8.1)/(100\text{-}\mathsf{new} \,\,\mathsf{H}_{2}\mathsf{O})\right] \\ &\bullet \left[(20.9-6.0)/(20.9-\mathsf{new} \,\,\mathsf{O}_{2})\right] \end{split}$$

The operating factor of the SCR describes the portion of the coal burn that reflects the period of SCR operation, based on whether the unit operates seasonally (0.43), annually in a base-load duty (0.98), or annually in peaking duty (0.80).

The coal burn rate in TBtu/yr can be obtained from coal use records, such as those reported to EIA in Form 767, or can be calculated from the annual coal burn (tons/yr) and the heating value of the coal, as described in the equation below:

$$B = C1 \bullet HV \bullet K_B \qquad \qquad Eq. 4-14$$

Eq. 4-9

where,

- B = Coal burn in TBtu/yr
- C1 = Dry coal burn, tons/yr. The dry coal can be calculated from wet coal through the following relationship: Dry coal (tons/yr) = wet coal (tons/yr) • (1-moisture(%)/100%)
- HV = Coal heating value, Btu/lb, dry
- K_B = Conversion factor = 2E-9. The factor is calculated from the conversion of tons per year to pounds per year and Btu to TBtu (2000/10¹² = 2E-9)

Ammonia Injection from FGC (NH3_{FGC})

Similar to reactions with ammonia slip from SCR or SNCR, any ammonia injected from FGC will react with sulfuric acid manufactured to form ABS. The following equation is used to calculate equivalent ammonia from FGC. Care must be taken to note the location of the ammonia injection, upstream or downstream of the APH, such that the ammonia can be utilized correctly in the release equation.

$$NH3_{FGC} = K_e \bullet B \bullet f_e \bullet I_{NH3}$$
 Eq. 4-15

where,

NH3fgc =	Total equivalent ammonia produced from FGC, lbs per year, either upstream
	of APH, NH3FGD_beforeAPH, or downstream of APH, NH3FGC_afterAPH

- K_e = Conversion factor = 3,799, as described in Text Box B
- B = Coal burn in TBtu/yr
- fe = Operating factor of FGC system, or the fraction of fuel burn when the FGC system operates. This value will be the same as fe described previously and in many cases can be approximated by 0.8.
- I_{NH3} = NH₃ injection for dual flue gas conditioning, ppmv at 6% O₂, wet; generally 3 ppmv NH₃ if operating, or zero (0) if no ammonia is used.

Total Release Calculation for Sources Employing Alkali Injection

As described in Section 3, sources that employ alkali injection to mitigate stack opacity or a visible plume will typically have access to field measurements describing SO₃ concentration in flue gas, most likely at the stack. The concentration of SO₃ measured at the stack can be converted to a mass emission rate using Equation 4-16.

$$ER_{ALKINJ} = K_{alkali} \bullet B \bullet S_{SO3} \bullet F2_x \bullet F_{alkali}$$
 Eq. 4-16

where,

ERALKIN	J	= Total H ₂ SO ₄ released from unit equipped with alkali injection, lbs per year
Kalkali	=	Constant, equal to 3,799, which is equivalent to K _e , as described in Text Box B.
В	=	Coal burn in TBtu/yr
Sso3	=	SO ₃ content, as measured in the stack or particulate collector exit, ppmv at 6% O ₂ , wet
Falkali	=	Operating factor for the alkali injection system, fraction of coal burn when unit is operated with alkali addition to remove SO ₃
F2x	=	Technology Impact Factors, to be applied only if SO_3 measurements are conducted at a location preceding the stack. The value of $F2_x$ is 1 if measurements are conducted at the stack, and there are no control components downstream of the measurement location. If the control device is an SO_2 scrubber, $F2x$ is denoted as $F2s$, but is handled the same way.

If the SO₃ content (S_{SO3}) is known at conditions other than 6% O₂ and wet, Equation 4-8 can be used to calculate a new value K_{alkali} at the new conditions.

The Technology Impact Factors selected should represent the control equipment between the point of SO₃ measurement and the stack. For example, if a unit is equipped with only an ESP,

and the SO₃ measurement is conducted at the ESP inlet, then the F2 factor for an ESP (Table 4-5, below) should be selected.

If the utility employing alkali injection does not have access to measured data, the total sulfuric acid releases may be calculated as described by Equation 4-11 (i.e., by estimating and summing the manufacture and applying the technology Impact Factors), and an alkali injection factor, F3_{ALKINJ}, is then applied to the total release. This F3_{ALKINJ} is either the expected fractional reduction in SO₃ (generally as guaranteed by the vendor), a default value of 0.2, as available data indicates 80% removal is easily achievable (EPRI, 2010b).

where,

ER _{ALKINJ} =	Total H ₂ SO ₄ released from unit equipped with alkali injection, lbs per year
TSAR _{Comb+SCR+FGC}	= Total H ₂ SO ₄ released from combustion, SCR and FGC, as calculated using Equation 4-10.
Falkali =	Operating factor for the alkali injection system, fraction of coal burn when unit is operated with alkali addition to remove SO_3
F3alkinj =	Technology Impact Factor for alkali injection; the expected fractional reduction in SO ₃ (generally as guaranteed by the vender). If no vendor information is available, use a default value of 0.2.

Partial Year Injection

If alkali injection is not utilized for the entire operating year (F_{alkali} does not equal 1), then Equation 4-16 (or Equation 4-17) must be added to the fraction of the total sulfuric acid release (TSAR) calculated from combustion, SCR/SNCR and/or FGC for the remainder of the operating time (Equation 4-11). Furthermore, in the calculation of TSAR for partial-year alkali injection, if SO₃ measurements also exist for operation without alkali injection, these data should be used with Equation 4-16 instead of the standard estimation equations, as shown below:

$$ER_{NOINJ} = K_{alkali} \bullet B \bullet S_{SO3} \bullet F2_x \bullet (1-F_{alkali}) Eq. 4-18$$

where,

 $ER_{NOINJ} = Total H_2SO_4$ released from unit without alkali injection, lbs per year

- K_{alkali} = Constant, equal to 3,799, which is equivalent to K_e, as described in Text Box B.
- B = Coal burn in TBtu/yr
- S₅₀₃ = SO₃ content without alkali injection, as measured in the stack or particulate collector exit, ppmv at 6% O₂, wet

- F_{alkali} = Operating factor for the alkali injection system, fraction of coal burn when unit is operated with alkali addition to remove SO₃
- $F2_x$ = Technology Impact Factors, to be applied only if SO₃ measurements are conducted at a location preceding the stack. The value of $F2_x$ is 1 if measurements are conducted at the stack, and there are no control components downstream of the measurement location. If the control device is an SO₂ scrubber, F2x is denoted as F2s, but is handled the same way.

Accordingly, total sulfuric acid release (TSAR) for a unit employing partial-year alkali injection is estimated by one of the following equations:

TSAR = ER _{ALKINJ} + (TSAR _{Comb+SCR+FGC}) ● (1-F _{alkali})	Eq. 4-19
TSAR = ERALKINJ + ERNOINJ	Eq. 4-20

Technology Impact Factors (F2)

The sulfuric acid estimating methodology employs empirically-derived Technology Impact Factors (F2). These F2 factors describe sulfate removal observed over the air heater, the ESP or other particulate control device, and FGD process equipment. Background information on these processes and the derivation of these F2 factors are addressed in this section.

Air Heater

The removal of SO₃ and/or H₂SO₄ within the air heater is due to the condensation of sulfuric acid and its removal as discrete individual particles (along with the fly ash) on the surface of this heat exchanger. The conventional Ljungstrom-type air heater has been documented to provide a removal sink for sulfuric acid (Saranuc, 1999). In fact, the largest supplier of Ljungstrom air heaters has evaluated the feasibility of employing the air heater process environment in conjunction with limestone injection as a proactive sulfuric acid control strategy (Hamel, 2003, and Bowes, 2006).

Air heater surfaces follow a pattern of alternately heating and cooling as the heat exchange elements move from the relatively hot flue gas to the cooler combustion air. As reported during field tests of commercial equipment (Saranuc, 1999), this temperature profile introduces a strong gradient in sulfuric acid concentration across the exit plane of the air heater. The cyclic thermal conditions as described by Hamel (2003) reveal that a significant portion of the air heater basket surface metal is exposed to flue gas temperature below the sulfuric acid dewpoint. This phenomenon may promote sulfuric acid deposition on the heat exchanger metal surface, compared to a tube-type heat exchanger in which the metal temperature is always above the sulfuric acid dewpoint.

Figure 4-2 presents data obtained from a survey of power plant operators and general literature sources describing SO_3/H_2SO_4 removal (and the corresponding F2 factors) across commercial, Ljungstrom-type air heaters. These data are presented as a function of SO_2 content of the flue gas for the host unit. Most data shown are for low sulfur eastern bituminous coal, for which air heater H_2SO_4 removal varies significantly between 0% and 69%, averaging 50% as indicated by the horizontal line. Two high sulfur (>2.5%) eastern cases are shown, along with limited data for PRB coal.

The data point showing 0% removal at approximately 600 ppm SO₂ is believed suspect, as an identical companion unit firing the same coal exhibited sulfuric acid capture of 38%. This measurement was not included in the F2 calculation.

Figure 4-2 presents two points reported by Hamel (2003) based on a low sulfur eastern bituminous-fired unit where SO₃ was "spiked" into the flue gas to elevate the concentration entering the air heater to 80 and 122 ppm. These two points are plotted separately on Figure 4-2 versus an estimated flue gas SO₂ content that could generate such values (using the average SO₂ oxidation rate for high sulfur, eastern bituminous coal in Figure 4-1. It should be noted these values measured for the "spiked" flue gas significantly exceed those measured for the two high sulfur eastern coal cases.

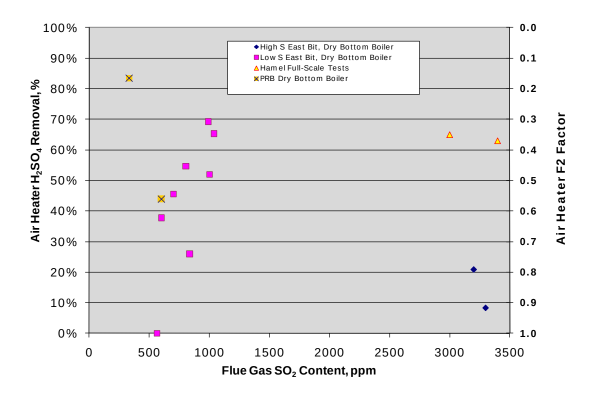


Figure 4-2 Removal of Sulfuric Acid by Ljungstrom-Type Air Heaters

The F2 factor for the air heater, calculated as [1 – Percent Removal], is estimated excluding the "spiked" SO₃ and suspect 0% removal measurement. The F2 factors for the air heater for low sulfur eastern bituminous, medium-high sulfur eastern bituminous, and PRB coals are shown in Table 4-4. Data are not currently available for other coal types. For those coals, as well as fuel oils, the model user could consider adopting the F2 factor for PRB if the ash is strongly alkaline and the low sulfur, eastern bituminous value if the ash is acidic.

Table 4-3	
Summary of F2 Factors for Air Heater Removal of Sulfuric Acid	

Boiler Type	Fuel	F2	Standard Deviation	Comment
All Boilers	Low S Eastern Bit	0.50	0.15	Average of measurements at 7 units.
All Boilers	Med-High S Eastern Bit (S >2.5%)	0.85	n/a	Based on two data points.
All Boilers	PRB	0.36	n/a	Based on two data points.

The standard deviation of the reported measurements for low sulfur eastern bituminous coal is about one third of the average measurement. Inadequate data prevent calculating a standard deviation for F2 factors for the other fuels.

Particulate and SO3 Control Processes

This section discusses the available data on removal of SO₃ by ESPs, FGDs, and alkali injection.

ESPs and **Baghouses**

An ESP provides extended residence time at relatively low temperatures, allowing contact between sulfuric acid and fly ash particles, as well as having collecting plates that can retain sulfuric acid particles. These features contribute to the removal of sulfuric acid. The ESP is the flue gas contacting device with perhaps the longest residence time; for large units, usually 10 seconds and in some cases, up to 15 seconds. Given the low flue gas velocities of 2 to 4 actual feet per second (0.6 - 1.2 meters per second), and the opportunity for heat loss at or near the walls, sulfuric acid condensation can be significant.

Figure 4-3 plots sulfuric acid removal by the ESP and the corresponding F2 factors as a function of the SO₂ content of the flue gas for the host unit, from a survey of plant operators. Most data shown are for a cold–side ESP and low sulfur eastern bituminous coal; three data points are shown for high sulfur coal (>2.5%). Also shown is a single data point for a hot-side ESP and four data points for low sulfur eastern bituminous coal. Similar to the case for the air heater, the 0% sulfuric acid removal at approximately 800 ppm SO₂ is suspect, as a companion unit at the same site firing the identical coal had 50% sulfuric acid removal. Accordingly, this "zero" removal datum, although shown in Figure 4-3, is not used in the analysis.

The F2 factor for the cold-side ESP, calculated as [1- Percent Removal], is estimated using all data except the 0% removal point. The F2 factors for ESP capture for low sulfur eastern bituminous, medium-high sulfur eastern bituminous, and PRB coals are listed in Table 4-5. Data are not currently available for other coal types. Western subbituminous coals (e.g., non-PRB) could consider adopting the F2 factor for PRB if the ash is strongly alkaline; western coals with acidic ash (e.g., bituminous) may consider adopting the low sulfur, eastern bituminous value. This guidance also applies to for fuel oils or any other coal that is not listed in the table.

Data describing the reduced H₂SO₄ penetration for one hot-side ESP (ESPh) is based on only a single test series. It should be noted that although this F2 factor is used for all coals, the measurement is from a power plant burning a low-sulfur, eastern bituminous coal. Comparison of F2 factors for a cold-side ESP burning low and high sulfur eastern bituminous coals show a

significant difference between the two coal types. As a result, total H₂SO₄ releases for hot-side ESP configurations may have a low bias; however, there is insufficient data to recommend alternate values.

Wet ESP's operate in much the same manner as a traditional ESP; however, due to the lower operating temperature and saturated environment, higher power levels are achievable, resulting in higher collection efficiency for sulfuric acid mists. Data describing sulfuric acid removal for a Wet ESP is based on two sets of paired measurements, both of which demonstrated 88% removal. One data point was from a high sulfur, eastern bituminous, coal-fired boiler and the other from a petroleum-coke fired boiler. Flue gas SO₂ was not reported for these tests; therefore, the data points are not included in Figure 4-3. Until further data are available, EPRI recommends using a wet ESP F2 factor of 0.12 for all fuels.

Baghouses show high H₂SO₄ removal based on very limited data. The average of two data points is 90% removal.

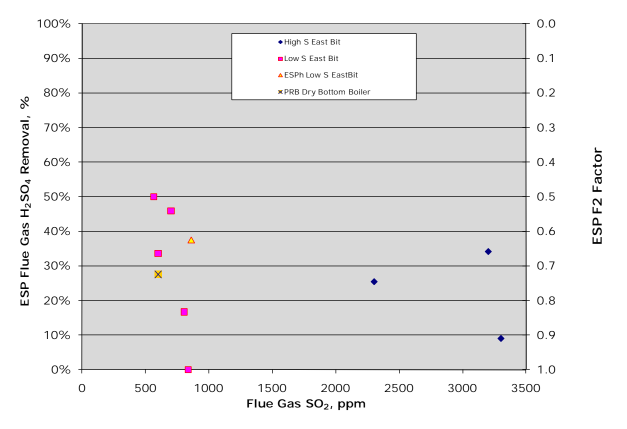


Figure 4-3 Removal of Sulfuric Acid by Cold-Side ESPs and One Hot-Side ESP

Equipment Type	Coal Type	F2 Factor	Standard Deviation	Comment or Observation
Cold-side ESP	Low S Eastern Bit	0.63	0.15	Average of measurements at 4 units.
Cold-side ESP	High S Eastern Bit (>2.5%)	0.77	0.13	Average of measurements at 3 units.
Cold-side ESP	Subbituminous (PRB)	0.72	n/a	Based on one measurement at one unit.
Hot-side ESP	All	0.63	n/a	Based on one measurement at one unit.
Wet ESP	All	0.12	n/a	Average of measurements at two units
Baghouse	Subbituminous coal	0.10	n/a	Two data points.

 Table 4-4

 Summary of F2 Factors for Particulate Control Devices (ESP, Baghouse)

n/a - not applicable – too few points to calculate.

The standard deviation of the reported measurements for low sulfur eastern bituminous coal is about one fourth of the average measurement, while the standard deviation of the measurements for the high sulfur eastern bituminous coal is about one sixth of the average. Inadequate data prevent calculating a standard deviation for F2 factors for the other fuels.

FGD

FGD process equipment rapidly cools or quenches flue gas, condensing a significant portion of the sulfuric acid into submicron droplets that can escape the process environment, confounding capture. Buckley (2002) notes that for condensation to occur, sulfuric acid generally must be supersaturated. However, fly ash particles can provide a nucleus for condensation of sulfuric acid, even at conditions that are not thermodynamically supersaturated. Buckley also projects sulfuric acid condensation on surfaces where equipment walls are lower in temperature than the flue gas (common in commercial equipment). However, the thin laminar boundary layer at the wall limits mass transfer, and for FGD equipment this mechanism provides no appreciable removal. Ironically, it is the high saturation conditions in this laminar layer near surfaces that are key to producing fine sulfuric acid mist.

Srivastava (2004) suggests that the condensed submicron droplets, once formed, are sufficiently small so that they follow the flow streamlines and avoid contact with the remaining wetted walls, liquid sheets, and droplets in the flow path. Although some degree of sulfuric acid removal is observed in FGD equipment, the amount is highly variable and depends on the design of the system. Buckley (2002) estimates FGD equipment removes 40-70% of the sulfuric acid, and Srivastava an average of 50%.

Figure 4-4 summarizes data reported in the public domain and from utility-specific tests describing FGD removal (and corresponding F2 factors) of sulfuric acid. The percent of sulfuric acid removal is depicted as a function of FGD inlet SO₂ concentration. Removals range from 14% to 78%, with an average of about 50%. The shaded portion of Figure 4-4 represents the upper and lower bounds of the data reported by Buckley (2002), while the solid horizontal line reflects the average reported by Srivastava (2004).

Table 4-6 summarizes the F2 factors for FGD process equipment, including results for a wet spray tower burning bituminous coal, and also one burning a blend of PRB/lignite. Given the limited data, the latter is assumed to serve as the basis of an F2-factor for 100% firing of PRB or lignite. Data for a wet venturi -type combined particulate/FGD process is also available, but only for a bituminous coal. Similar to the case for a wet spray tower, the lack of data merits assuming sulfuric acid removal for all coals is described by that measured for bituminous coal.

F2 factors for the use of magnesium-based additives in oil-fired boilers are also included in Table 4-6. These additives are used to control furnace slagging caused by the vanadium in the oil or to control sulfuric acid emissions or both. The fuel oil vanadium can also catalyze SO_2 to SO_3 oxidation, but the additive, when added to the oil, tends to effectively bind up the vanadium, partially reducing its catalytic effect. Addition of magnesium-based additives in the fuel oil tend to be less effective in controlling the emissions of sulfuric acid than the same additive sprayed into the furnace downstream of the flame zone. The factors for Mag-Ox addition are derived from a single data point; as such, it is recommended to use measured facility data if available

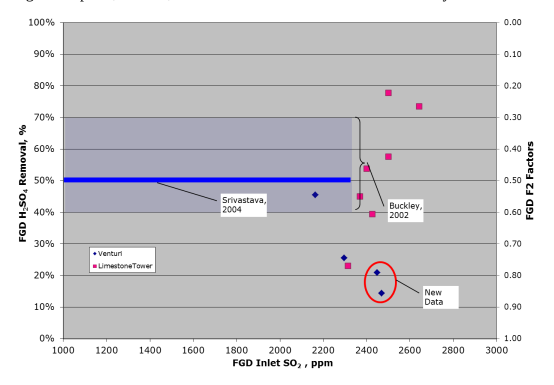


Figure 4-4 Removal of Sulfuric Acid by Flue Gas Desulfurization Equipment: Various FGD Designs, Coals

Table 4-5Summary of F2 Factors for Wet and Dry FGD Equipment and Additives

FGD Type	Coal Type	F2 Factor	Standard Deviation	Comment or Observation
Wet: Spray Tower	E. Bituminous	0.47	0.17	Seven data points.
Wet: Spray Tower	PRB or Lignite	0.40	n/a	Two data points.
Wet: Venturi Tower	All coals	0.73	0.13	Four data points from three units, bituminous coal only; limited data merit assigning same factor for other coals.
Dry FGD and baghouse	All coals	0.01	n/a	Two data points.
Mg-Ox mixed w/fuel oil	All fuel	0.50	n/a	One data point.
Mg-Ox into furnace	All fuel	0.25	n/a	One data point.

5 ESTIMATING GUIDELINE: MULTIPLE FUEL OR BLENDED FUEL BOILERS

Estimating sulfuric acid production of steam boilers firing multiple or blended fuels – such as natural gas, fuel oil, and perhaps coal – uses the same approach as for single fuels. This will consist of determining the contribution of each source (combustion, SCR or SNCR NOx control, and FGC) for each different fuel, and adjusting for loss or removal. It is assumed for multiple fuel boilers that the contribution of each fuel can be separated and treated individually. The same approach is to be used for boilers that fire a blended coal. This assumption is particularly important when considering blends of PRB with low or high sulfur eastern bituminous coal, as this approach accounts for the role of alkalinity in PRB ash in mitigating SO₃.

This approach of treating blends of coal as separate fuels in proportion to the mass burn rate is believed to be conservative with regard to SO₃ emissions when considering PRB coal, in that it will likely project H₂SO₄ emissions higher than actual. The ability of PRB coal to neutralize SO₃ by an amount that is greater than a direct proportion of the coal blend is due to the extremely high content of alkaline material, particularly CaO. However, data to quantify this impact with any reasonable degree of confidence are not available at present. Accordingly, the conservative approach is to presume an impact in direct proportion to the coal blend until a database can be developed.

The sequence of calculations is performed for the first component of the blend, including calculations for SCR and FGC, if applicable. The sequence is repeated for each component. When complete, the total manufacture of sulfuric acid is calculated by adding all of the manufacture totals for all fuels from all processes. Likewise, the release is summed over all processes and fuels.

As an example, if a unit with an SCR and FGC burns mostly coal, but uses natural gas in a NOx reburn process and also disposes of used oil by combustion in the furnace, then the following sequence of calculations would be required:

- 1. Coal fuel
 - a. Combustion manufacture
 - b. Combustion release
 - c. SCR manufacture
 - d. SCR release
 - e. FGC manufacture
 - f. FGC release
- 2. Natural gas
 - a. Combustion manufacture
 - b. Combustion release
 - c. SCR manufacture

- d. SCR release
- e. FGC manufacture
- f. FGC release
- 3. Used oil
 - a. Combustion manufacture
 - b. Combustion release
 - c. SCR manufacture
 - d. SCR release
 - e. FGC manufacture
 - f. FGC release
- 4. Sum manufacture and releases
 - a. Manufacture

= 1a + 1c + 1e + 2a + 2c + 2e + 3a + 3c + 3e

b. Release

= 1b + 1d + 1f + 2b + 2d + 2f + 3b + 3d + 3f

All of the manufactured results would be summed together and the releases summed also to give the final result. Example 8 in Appendix A details this calculation procedure.

6 ESTIMATING GUIDELINE: COMBUSTION TURBINES

Natural gas-fired sources typically have negligible content of sulfur in the fuel, thus sulfuric acid production is negligible. Facilities that burn only natural gas are not required to participate in the TRI reporting program. However, sources that are co-located with coal units will need to be included in the estimates for total sulfuric acid release for the site. The calculation described in this section can also be used to estimate sulfuric acid emissions for Prevention of Significant Deterioration (PSD) review of new natural gas-fired generation sites.

The calculations for gas-fired units are structured very much the same as for coal- and oil-fired sources. For simple cycle combustion turbines, the only source of sulfuric acid is the sulfur in the fuel. The EPA AP-42 emissions factor suggests a value of 2000 grains of sulfur per million cubic feet of natural gas as a default sulfur content value. This value is equivalent to approximately 3.5 ppm of sulfur in the raw natural gas.

A methodology to estimate manufacture and release of sulfuric acid for simple cycle and combined cycle units is provided below.

Manufacture and Release for Simple Cycle Units

Given the current configuration of simple cycle units, any sulfuric acid *manufactured* is *released*; thus, the estimates of sulfuric acid are the same. This is because there is no equipment located following the simple cycle arrangement that removes sulfuric acid. Even though simple cycle units can be equipped with SCR, and the use of SCR with sulfur-containing fuels will manufacture SO₃, the exit gas temperature is too high to allow condensation of SO₃ or reaction with ammonia slip. Selective non-catalytic reduction is not applied to simple cycle or combined cycle combustion turbines; therefore, no estimation procedure is provided for that configuration.

Accordingly, the equations for formation of sulfuric acid from natural gas (NG) combustion are:

$$EM_{SC} = K \bullet F1 \bullet E2_{NG} Eq. 6-1$$

where,

EMsc = total H₂SO₄ manufactured from combustion, lbs/yr

K = Molecular weight and units conversion constant

= 98.07 / 64.04 • 2000 = 3,063

 $98.07 = Molecular weight of H_2SO_4$

 $64.04 = Molecular weight of SO_2$

2,000 = Conversion from tons per year to pounds per year.

F1 = Fuel Impact Factor for NG

 $E2_{NG}$ = Sulfur dioxide (SO₂) emissions either: (1) recorded by a continuous emissions monitor, tons/yr, or (2) calculated from fuel burn data, tons/yr.

SO₂ emissions can be obtained through a calculation using the heat input of natural gas.

$$E2_{NG} = K_b \bullet B_{NG} \bullet S \qquad \qquad Eq. \ 6-2$$

where,

 $E2_{NG}$ = Total SO₂ production from NG combustion, tons/yr

 K_b = Molecular weight and units conversion constant = 0.0001359

 $B_{NG} = Burn of NG in Tbtu/yr$

S = Sulfur content of natural gas, in grains per million standard cubic feet (Mscf), typically 2000 gr/ 10^6 scf per EPA AP-42.

The derivation of constant K_b is presented in Text Box D.

Text Box D: Derivation of Molecular Weight and Units Conversion Constant, K _b K _b is determined from the following equation.								
$(K_b \bullet B)$	$_{\rm NG} \bullet S) = t$	ons SO ₂ per	yr =					
\mathbf{B}_{NG} TBtu1 scf nat gas1 scf nat 10^{12} Btu1 ton S1 ton S1 ton mole 								
Year	10 ⁶ scf nat gas	1050 Btu	TBtu	7000 gr S	2000 lbs S	32 tons S	1 ton mol S	1 ton mole SO ₂
Grouping terms,								
$= (B_{NG} \bullet S) \bullet 64 / (1050 \bullet 32 \bullet 14) = (B_{NG} \bullet S) \bullet 0.0001359$								
Therefore, the value of K_b is equal to 0.0001359 tons SO ₂ /(TBtu • grains S/million SCF NG).								

The SO₂ emissions can also be calculated from the volume of natural gas burned:

$$E2_{NG} = K_{NG} \bullet N1 \bullet S \qquad \qquad Eq. 6-3$$

where,

 $E2_{NG}$ = total SO₂ production from NG combustion, tons/yr

 K_{NG} = Molecular weight and units conversion constant = 1.427 • 10⁻⁷

N1 = NG burn in million standard cubic feet (Mscf) per year

S = NG sulfur content in grains per million standard cubic feet; use EPA's value of $2000 \text{ gr}/10^6 \text{ scf}$ as default

The derivation of constant K_{NG} is presented in Text Box E.

Text Box E: Derivation of Molecular Weight and Units Conversion Constant, K_{NG}						
$(K_{NG} \bullet N1 \bullet S)$	$S = tons SO_2 per y$	r =				
N1 10 ⁶ scf	N1 10^6 scf S gr S b S 1 ton S 1 ton mol S 1 ton mole SO ₂ 64 tons SO ₂					
Year	Year 10^6 scf nat gas 7000 gr S2000 lbs S32 tons S1 ton mol S1 ton mole SO2					
Grouping terms, $(N1 \bullet S) \bullet 64 / (7000 \bullet 2000 \bullet 32) = (N1 \bullet S) \bullet 1.427 \bullet 10^{-7}$ Therefore, the value of K _{NG} is equal to $1.427 \bullet 10^{-7}$ tons SO ₂ /(grains S)						

Table 6-1 presents the F1 factors for simple cycle units as a function of stack temperature, as sulfuric acid vapor is related to the temperature of the exhaust. As simple cycle combustion turbines (CT) exhaust is usually around 1000°F, and TRI rules require the reporting of sulfuric acid (not of SO₃), the amount manufactured and released depends on stack temperature. Table 6-1 combines the temperature-based SO₃ to H₂SO₄ conversion with the SO₂ to SO₃ conversion to give the Fuel Impact Factor, F1.

Table 6-1 Fuel Impact (F1) Factors for a Simple CT

Stack T, °F	F1
300	0.055
400	0.055
500	0.047
600	0.022
700	0.0055
750	0.0027
800	0.0013
850	0.00071
900	0.00039
950	0.00022
1000	0.00013
1050	0.00008
1100	0.00005
1150	0.00003
1200	0.00002

Combined Cycle Units

Sulfuric Acid Manufactured

For combined cycle units, sources of *manufacture* of sulfuric acid are combustion, oxidation of SO_2 across catalyst used for NOx control and recently, oxidation of SO_2 across catalyst used for CO control.

For combustion, the manufacture of sulfuric acid for combined-cycle units is described by Equation 6-4.

$$EM_{CCcom} = K \bullet F1 \bullet E2_{NG} Eq. 6-4$$

where,

EM_{CCcom} = Total H₂SO₄ manufactured from combustion, lbs/yr

= 98.07/64.04 • 2000 = 3,063

98.07 = Molecular weight of H₂SO₄;

64.04 = Molecular weight of SO₂;

2,000 =Conversion from tons per year to pounds per year.

- F1 = Fuel Impact Factor for NG (See Table 4-1)
- $E2_{NG}$ = Sulfur dioxide (SO₂) emissions either: (1) recorded by a continuous emissions monitor, tons/yr, or (2) calculated from fuel burn data, tons/yr.

Recently, it has been shown that CO catalysts in combined cycle units can oxidize SO_2 to SO_3 in a manner similar to SCRs. Equation 6-5 describes the manufacture of sulfuric acid across the CO catalyst. For the case of combined cycle combustion turbines, the CO catalyst SO_2 oxidation rate (S3) will be approximately 0.1. This value is an estimate based on data from a prominent catalyst vendor. A more accurate value can be obtained from field measurements of SO_2 oxidation, or from the vendor's specification of the CO catalyst.

$$\mathsf{EM}_{\mathsf{CC}_\mathsf{CO}} = \mathsf{K} \bullet \mathsf{S3} \bullet \mathsf{f}_{\mathsf{COops}} \bullet \mathsf{E2} \qquad \qquad \mathsf{Eq. 6-5}$$

where,

 EM_{CC_CO} = Total H₂SO₄ manufactured from the CO catalyst, lbs per year

K = Molecular weight and units conversion constant

$$=$$
 98.07/64.04 • 2000 $=$ 3,063

98.07 =Molecular weight of H_2SO_4

 $64.04 = Molecular weight of SO_2$

2000 = Conversion from tons per year to pounds per year

- S3 = CO catalyst SO₂ oxidation rate (specified as a decimal, typically from 0.1-0.6)
- f_{COops} = Operating factor of CO catalyst system, or the fraction of fuel burn when the flue gas is directed through the CO catalyst. This value should reflect the fraction of hours that the CO catalyst processed flue gas. This value will be site-specific but can be approximated as 0.8 for year-round peaking operation, 0.98 for year-round base-loaded operation, or 0.43 for seasonal operation.

E2 =
$$SO_2$$
 produced, tons per year

Equation 6-6 describes the manufacture of sulfuric acid in an SCR. The amount of sulfuric acid manufactured due to SO₂ oxidation across the CO catalyst (EM_{CC_CO}) must be subtracted from the total amount of SO₂ produced (E2), because the CO catalyst will reduce the amount available for conversion by the SCR catalyst. For the case of combined cycle combustion turbines, the SCR catalyst SO₂ oxidation rate (S2) will be approximately 0.03. A more precise value can be obtained from either field test reports of SO₂ oxidation, or from the vendor's specification of the SCR catalyst or process. The Technology Impact Factor for SCR, F3_{SCR}, will be 1.0.

$$EM_{CCSCR} = [(K \bullet E2) - EM_{CC-CO}] \bullet f_{sops} \bullet S2 \bullet F3_{SCR}$$
 Eq. 6-6

where,

 EM_{CCSCR} = Total H₂SO₄ manufactured from SCR, lbs per year

= 98.07/64.04 • 2000 = 3,063

98.07 = Molecular weight of H₂SO₄;

64.04 = Molecular weight of SO₂;

2,000 = Conversion from tons per year to pounds per year

- E2 = SO_2 produced, tons per year
- EM_{CC_CO} = Total H₂SO₄ manufactured from the CO catalyst, lbs per year
- S2 = SCR catalyst SO₂ oxidation rate (specified as a decimal, typically from 0.001-0.03)
- f_{sops} = Operating factor of SCR system, or the fraction of coal burn when the flue gas is directed through the SCR, whether NH₃ reagent is injected to derive NOx reduction or not. This value should reflect the hours the SCR reactor processed flue gas, which will be site-specific but can be approximated by generally 0.8 for year-round peaking operation, 0.98 for year-round baseloaded operation, or 0.43 for seasonal operation
- $F3_{SCR}$ = Technology Impact Factor, for SCR (equal to 1.0, refer to Table 4-2)

The total sulfuric acid *manufacture* (TSAM) for a combined cycle unit is estimated using the equation below:

Sulfuric Acid Released

The sulfuric acid released by all sources in combined cycle units is estimated as shown in Equation 6-8. This equation takes into account the reducing effect of ammonia slip from the SCR equipment in a similar method as described in Section 4:

$$TSAR_{cc} = [TSAM_{cc} - (K_s \bullet B \bullet f_{sreagent} \bullet S_{NH3})] \bullet F2_{cc} \qquad Eq. 6-8$$

where,

TSAR _{CC}	=	Total H ₂ SO ₄ released, lbs per year
TSAM _{CC}	=	Total H ₂ SO ₄ manufactured from all sources, lbs per year
Ks	=	Conversion factor = 3,799 (see Text Box B)
В	=	Fuel burn in Tbtu/yr
fsreagent	=	Fraction of SCR operation with reagent injection, when residual NH ₃ is produced that will remove SO ₃ .
SNH3	=	NH ₃ slip from SCR, ppmv at 6% O ₂ , wet (averages 0.75 ppmv over catalyst guarantee period. Actual NH ₃ slip data should be used if available)
F2 _{CC}	=	Technology Impact Factor for the combined cycle heat exchanger, equal to a value of 0.5.

Examples 9 and 10 of Appendix A illustrates a gas-fired combined cycle plant calculation.

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A EXAMPLE CALCULATIONS

Example 1: Conventional Coal-Fired Boiler with an ESP

A 500 MW pulverized coal-fired (PC) boiler, equipped with a cold-side electrostatic precipitator, burns an Eastern bituminous coal. The coal used in the reporting year is 1,126,938 tons with a weighted average sulfur concentration of 2.0% and a heating value of 12,000 Btu/lb.

Solution

Manufactured

 $E2 = K1 \bullet K2 \bullet C1 \bullet S1$ $E2 = 0.02 \bullet 0.95 \bullet 1,126,938 \bullet 2.0 = 42,824 \text{ tons SO}_2/\text{yr}$ $EM_{\text{Comb}} = K \bullet F1 \bullet E2$ $F1_{\text{eastbit}} = 1.1163E-6 \bullet \text{SO}_2 + 0.0064876$ $SO_2 = S1 \bullet \frac{K_{F1}}{HV} = 2.0 \bullet \frac{10,003,602}{12,000} = 1667 \text{ ppm}$

 $F1_{eastbit} = 1.1163 E\text{-}6 \bullet + 0.0064876 = 0.0083$

 $EM_{Comb} = 3063 \bullet 0.0083 \bullet 42,824 = 1,088,710 lbs H_2SO_4/yr$

The 25,000 lbs/yr threshold has been exceeded; therefore, a release estimate must be made and the result reported on Form R.

Released

 $TSAR = EM_{Comb} \bullet F2_{APH} \bullet F2_{ESP}$

TSAR = 1,088,710 • 0.50 • 0.63 = 342,944 lbs H₂SO₄/yr

Example 2: SCR Added to Example 1

A 500 MW PC boiler, equipped with a cold-side electrostatic precipitator, burns an Eastern bituminous coal. The plant is equipped with an SCR process that operates during the ozone season only, so that 0.43 of the coal burn occurred with the SCR operating. The remaining time that the SCR is not in service the flue gas is bypassed around the SCR. The SCR catalyst SO₂ oxidation rate specified in the design is 0.75%, and the ammonia slip is estimated to be 0.75 ppmv. The coal used in the reporting year is 1,126,938 tons with a weighted average sulfur concentration of 2.0% and a heating value of 12,000 Btu/lb.

Solution

Manufactured

 $E2 = K1 \bullet K2 \bullet C1 \bullet S1$ E2 = 0.02 • 0.95 • 1,126,938 • 2.0 = 42,824 tons SO₂/yr

Combustion

 $EM_{comb} = K \bullet F1 \bullet E2$

 $F1_{eastbit} = 1.1163E-6 \bullet SO2 + 0.0064877$

$$SO_2 = S1 \bullet \frac{K_{F1}}{HV} = 2.0 \bullet \frac{10,003,602}{12,000} = 1667 \ ppm$$

 $F1_{eastbit} = 1.1163E-6 \bullet + 0.0064877 = 0.0083$

 $EM_{Comb} = 3063 \bullet 0.0083 \bullet 42,824 = 1,088,710 lbs H_2SO_4/yr$

SCR

 $EM_{SCR} = K \bullet S2 \bullet f_s \bullet E2$

 $EM_{SCR} = 3063 \bullet 0.0075 \bullet 0.43 \bullet 42,824 = 423,023 lbs H_2SO_4/yr$

Total

 $TSAM = EM_{comb} + EM_{SCR}$

 $TSAM = 1,088,710 + 423,023 lbs H_2SO_4/yr$

 $TSAM = 1,511,733 lbs H_2SO_4/yr$

The 25,000 lbs/yr threshold has been exceeded; therefore, a release estimate must be made and the result reported on Form R.

Released

Coal Burn

 $\mathbf{B} = \mathbf{C}\mathbf{1} \bullet \mathbf{H}\mathbf{V} \bullet \mathbf{K}_{\mathbf{B}}$

 $B = 1,126,938 \text{ tons/yr} \bullet 2000 \text{ lbs/ton} \bullet 12,000 \text{ Btu/lb} \bullet 1 \text{ Tbtu/}10^{12} \text{ Btu}$

B = 27.05 Tbtu/yr

Ammonia Slip

 $NH3_{SCR} = K_s \bullet B \bullet f_{sreagent} \bullet S_{NH3}$

NH3_{SCR} = 3799 • 27.05 • 0.43 • 0.75 = 33,141 lbs NH₃/yr

Total Releases

 $TSAR = [(EM_{Comb} + EM_{SCR/SNCR}) - (NH3_{SCR})] \bullet F2_{APH} \bullet F2_{ESP}$

 $TSAR = [(1,088,710 + 423,023) - (33,141)] \bullet 0.50 \bullet 0.63$

 $TSAR = 465,756 lbs H_2SO_4/yr$

Example 3: Alkali Injection Added to Example 2

A 500 MW PC boiler, equipped with a cold-side electrostatic precipitator, burns an Eastern bituminous coal. The plant is equipped with an SCR process that operates during the ozone season only, so that 0.43 of the coal burn occurred with the SCR operating. The remaining time that the SCR is not in service the flue gas is bypassed around the SCR. The SCR catalyst SO₂ oxidation rate specified in the design is 0.75%, and the ammonia slip is estimated to be 0.75 ppmv.

The plant employs injection of alkali material (e.g., lime, sodium bicarbonate, sodium bisulfate, etc.) to control SO₃ emissions at the stack. The plant operator has conducted a series of field tests to determine the amount of alkali material that must be injected to limit flue gas SO₃ concentration at the stack. The test contractor has issued a report defining the amount of alkali that should be added to limit the stack SO₃ to 5 ppm, as measured in the flue gas at 6.6% O₂, and 8.8% moisture. The alkali injection system will be operated throughout the entire year, so that no more than 5 ppm of SO₃ at the stated conditions of excess O₂ and moisture exist in the stack.

The coal burn rate in the reporting year is 1,126,938 tons, with a weighted average sulfur concentration of 2.0%, and a heating value of 12,000 Btu/lb.

The calculation of manufactured sulfuric acid is the same as Example 1, and is repeated here:

Solution

Manufactured

 $E2 = K1 \bullet K2 \bullet C1 \bullet S1$

 $E2 = 0.02 \bullet 0.95 \bullet 1,126,938 \bullet 2.0 = 42,824 \text{ tons } SO_2/yr$

Combustion

 $EM_{comb} = K \bullet F1 \bullet E2$

 $F1_{eastbit} = 1.1163E-6 \bullet SO_2 + 0.0064877$

$$SO2=S1 \bullet \frac{K_{F1}}{HV} = 2.0 \bullet \frac{10,003,602}{12,000} = 1667 \text{ ppm}$$

 $F1_{eastbit} = 1.1163E-6 \bullet + 0.0064877 = 0.0083$

 $EM_{Comb} = 3063 \bullet 0.0083 \bullet 42,824 = 1,088,710 lbs H_2SO_4/yr$

SCR

 $EM_{SCR} = K \bullet S2 \bullet f_s \bullet E2$

Total

 $TSAM = EM_{comb} + EM_{SCR}$

TSAM = 1,088,710 + 423,023 lbs H₂SO₄/yr

TSAM = 1,511,733 lbs H₂SO₄/yr

The 25,000 lbs/yr threshold has been exceeded; therefore, a release estimate must be made and the result reported on Form R.

Released

The total sulfuric acid released is based on the maximum value allowed by alkali injection, as measured in the stack, and the volume flow rate of combustion products processed.

The concentration of SO_3 measured at the stack can be converted to a mass emission rate, as described by Equation 4-15

 $ER_{ALKINJ} = K_{alkali} \bullet B \bullet S_{SO3} \bullet F_{alkali} \bullet F2_x$

where,

ERALKIN	11	= Total H ₂ SO ₄ released from unit equipped with alkali injection, lbs per year
Kalkali	=	Constant, equal to 3,799, with units of lbs $H_2SO_4/(Tbtu ppmv SO_3)$ (SO ₃ measured at @ 6% O ₂ and 8.1% H ₂ O)
В	=	27.05 Tbtu/yr, for the specific case of Example 1
S _{SO3}	=	SO_3 content as measured in the stack or particulate collector exit, corrected to a concentration basis of ppmv at 6% O_2 , 8.1% moisture
Falkali	=	fraction of operation with alkali addition to remove SO ₃ .
F2 _x	=	1 (as the SO ₃ measurement is in the stack, there are no control technology components downstream of the measurement)

The conversion factor K_{alkali} is equal to 3,799 (see Text Box B). This constant is derived for the case where flue gas SO₃ is reported in terms of 6% oxygen and wet flue gas at 8.1% H₂O. However, the SO₃ concentration of 5 ppm in the stack is measured at slightly different conditions of excess O₂ and moisture (6.6% oxygen and 8.8% moisture and therefore, a new value of K_{alkali} must be calculated:

$$\begin{split} & K_{alkali} \left(O_2, H_2 O \right) = 3799 \ (6\% \ O_2, 8.1\% \ H_2 O \right) \bullet \left[(100\text{-}8.1)/(100\text{-}new \ H_2 O) \right] \\ & \bullet \left[(20.9 - 6.0)/(20.9 - new \ O_2) \right] \\ & K_{alkali} \left(O_2, H_2 O \right) = 3799 \ (6\% \ O_2, 8.1\% \ H_2 O \right) \bullet \left[(100\text{-}8.1)/(100\text{-}8.8) \right] \\ & \bullet \left[(20.9 - 6.0)/(20.9 - 6.6) \right] \\ & K_{alkali} \ (6.6\%, 8.8\%) = 3799 \bullet \left[1.0077 \right] \bullet \left[1.034 \right] \\ & K_{alkali} \ (6.6\%, 8.8\%) = 3958 \end{split}$$

Total sulfuric acid release is then calculated as follows:

 $ER_{ALKINJ} = [3,958] \bullet [27.05] \bullet [5.0] \bullet [1] \bullet [1]$

 $ER_{ALKINJ} = 535,320 lbs/yr$

 $TSAR = ER_{ALKINJ}$

TSAR = 535,320 lbs/yr

Example 4: 500 MW PRB-fired Boiler with ESP, SCR

A 500 MW PC boiler, equipped with a cold-side electrostatic precipitator, burns a PRB coal. The plant is equipped with an SCR process that operates during the ozone season only, so that 0.43 of the coal burn occurred with the SCR operating. The SCR is bypassed when not in use. The SCR catalyst SO₂ oxidation rate specified in the design is 2%, and the ammonia slip is estimated to be 0.75 ppmv. The coal used in the reporting year is 1,384,552 tons with a weighted average sulfur concentration of 0.40% and a heating value of 8,500 Btu/lb.

Solution

Manufactured

 $E2 = K1 \bullet K2 \bullet C1 \bullet S1$

 $E2 = 0.02 \bullet 0.875 \bullet 1,384,552 \bullet 0.40 = 9,692 \text{ tons } SO_2/yr$

Combustion

 $EM_{comb} = K \bullet F1 \bullet E2$

EM_{comb} = 3063 • 0.0019 • 9,692 = 56,405 lbs H₂SO₄/yr

SCR

```
EM_{SCR} = K \bullet S2 \bullet f_s \bullet f_{3SCR} \bullet E2
```

 $EM_{SCR} = 3063 \bullet 0.020 \bullet 0.43 \bullet 0.17 \bullet 9,692 = 43,402 lbs H_2SO_4/yr$

Total

 $TSAM = EM_{comb} + EM_{SCR}$ $TSAM = 56,404 + 43,402 \text{ lbs } H_2SO_4/yr$ $TSAM = 99,806 \text{ lbs } H_2SO_4/yr$

The 25,000 lbs/yr threshold has been exceeded; therefore, a release estimate must be made and the result reported on Form R.

Released

Coal Burn

 $\mathbf{B} = \mathbf{C1} \bullet \mathbf{HV} \bullet \mathbf{K}_{\mathbf{B}}$

 $B = 1,384,552 \text{ tons/yr} \bullet 2000 \text{ lbs/ton} \bullet 8,500 \text{ Btu/lb} \bullet 1 \text{ Tbtu/}10^{12} \text{ Btu}$

B =23.54 Tbtu/yr

Ammonia Slip

 $NH3_{SCR} = K_s \bullet B \bullet f_{sreagent} \bullet S_{NH3}$

NH3_{SCR} = 3799 • 23.54 • 0.43 • 0.75 = 28,841 lbs NH₃/yr

Total Releases

 $TSAR = [(EM_{Comb} + EM_{SCR/SNCR}) - (NH3_{SCR})] \bullet F2_{APH} \bullet F2_{ESP}$ $TSAR = [(56,404 + 43,402) - (28,841)] \bullet 0.36 \bullet 0.72$ $TSAR = 18,394 \text{ lbs } H_2SO_4/yr$

Example 5: FGC Added to Example 1

A 500 MW PC boiler, equipped with a cold-side electrostatic precipitator, burns an Eastern bituminous coal. The plant is equipped with a FGC process that injects both SO3 and NH3 upstream of the air preheater. The SO3 is injected at 7 ppmv at 6% O2 wet, and the ammonia at 3 ppmv also at 6% O2 wet. The FGC system operates whenever the plant is on, except during startup and shutdown, with an operating factor estimated at 0.9. The coal used in the reporting year is 1,126,938 tons with a weighted average sulfur concentration of 2.0% and a heating value of 12,000 Btu/lb.

Solution

Manufactured

 $E2 = K1 \bullet K2 \bullet C1 \bullet S1$ E2 = 0.02 • 0.95 • 1,126,938 • 2.0 = 42,824 tons SO₂/yr

Combustion

 $\mathbf{EM}_{\mathbf{Comb}} = \mathbf{K} \bullet \mathbf{F1} \bullet \mathbf{E2}$

 $F1_{eastbit} = 1.1163E-6 \bullet SO_2 + 0.0064877$

$$SO_2 = S1 \bullet \frac{K_{F1}}{HV} = 2.0 \bullet \frac{10,003,602}{12,000} = 1667 \ ppm$$

 $F1_{eastbit} = 1.1163E-6 \bullet + 0.0064877 = 0.0083$

 $EM_{Comb} = 3063 \bullet 0.0083 \bullet 42,824 = 1,088,710 \text{ lbs } H_2SO_4/yr$

FGC

B = 1,126,938 tons/yr • 2000 lbs/ton • 12000 Btu/lb • 1 Tbtu/10¹² Btu

B = 27.05 Tbtu/yr

 $EM_{FGC} = K_e \bullet B \bullet f_e \bullet I_s \bullet F3_{FGC}$

 $EM_{FGC} = 3799 \bullet 27.05 \bullet 0.9 \bullet 7 \bullet 1 = 647,407 \text{ lbs } H_2SO_4/yr$

Total

 $TSAM = EM_{Comb} + EM_{FGC}$ TSAM = 1,088,710 + 647,407 lbs H₂SO₄/yr TSAM = 1,736,117 lbs H₂SO₄/yr

The 25,000 lbs/yr threshold has been exceeded; therefore, a release estimate must be made and the result reported on Form R.

Released

Coal Burn

 $B = C1 \bullet HV \bullet K_B$

B = 1,126,938 tons/yr • 2000 lbs/ton • 12000 Btu/lb • 1 Tbtu/10¹² Btu

 $B=27.05 \ Tbtu/yr$

FGC Ammonia Injection

 $NH3_{FGC} = K_e \bullet B \bullet f_e \bullet I_{NH3}$

NH3_{FGC} = 3799 • 27.05 • 0.9 • 3.0 = 277,460 lbs NH₃/yr

Total Releases

 $TSAR = [(EM_{Comb} + EM_{FGC}) - (NH3_{FGC})] \bullet F2_{APH} \bullet F2_{ESP}$

 $TSAR = [(1,088,710 + 647,407) - (277,460)] \bullet 0.50 \bullet 0.63$

TSAR = 459,477 lbs H₂SO₄/yr

Example 6: SCR and FGC Added to Example 1

A 500 MW PC boiler, equipped with a cold-side electrostatic precipitator, burns an Eastern bituminous coal. The plant is equipped with an SCR process that operates during the ozone season only, so that 0.43 of the coal burn occurred with the SCR operating. The SCR is bypassed when not in operation. The SCR catalyst SO₂ oxidation rate specified in the design is 0.75%, and the ammonia slip is estimated to be 0.75 ppmv. The plant is also equipped with a FGC process that injects both SO₃ and NH₃ upstream of the air preheater. The SO₃ is injected at 7 ppmv at 6% O₂ wet, and the ammonia at 3 ppmv also at 6% O₂ wet. The FGC system operates whenever the plant is on, except during startup and shutdown, with an operating factor estimated at 0.9. The coal used in the reporting year is 1,126,938 tons with a weighted average sulfur concentration of 2.0% and a heating value of 12,000 Btu/lb.

Solution

Manufactured

Total (from previous examples)

 $TSAM = EM_{Comb} + EM_{SCR} + EM_{FGC}$

 $TSAM = 1,088,710 + 423,023 + 647,407 \ lbs \ H_2SO_4/yr$

 $TSAM = 2,159,140 lbs H_2SO_4/yr$

The 25,000 lbs/yr threshold has been exceeded; therefore, a release estimate must be made and the result reported on Form R.

Released

Total Releases (from previous examples)

 $TSAR = [(EM_{Comb} + EM_{SCR} + EM_{FGC}) - (NH3_{SCR} + NH3_{FGC})] \bullet F2_{APH} \bullet F2_{ESP}$ $TSAR = [(1,088,710 + 423,023 + 647,407) - (33,141 + 277,460)] \bullet 0.50 \bullet 0.63$ $TSAR = 582,290 \text{ lbs } H_2SO_4/\text{yr}$

Example 7: SCR and Downstream FGC Added to Example 1

A 500 MW PC boiler, equipped with a cold-side electrostatic precipitator, burns an Eastern bituminous coal. The plant is equipped with an SCR process that operates during the ozone season only, so that 0.43 of the coal burn occurred with the SCR operating. The SCR is bypassed when not operating. The SCR catalyst SO₂ oxidation rate specified in the design is 0.75%, and the ammonia slip is estimated to be 0.75 ppmv. The plant is also equipped with a FGC process that injects both SO₃ and NH₃ <u>downstream</u> of the air preheater. The SO₃ is injected at 7 ppmv at 6% O₂ wet, and the ammonia at 3 ppmv also at 6% O₂ wet. The FGC system operates whenever the plant is on, except during startup and shutdown, with an operating factor estimated at 0.9. The coal used in the reporting year is 1,126,938 tons with a weighted average sulfur concentration of 2.0% and a heating value of 12,000 Btu/lb.

Solution

Manufactured

Total (from previous examples)

 $TSAM = EM_{Comb} + EM_{SCR} + EM_{FGC}$

TSAM= 1,088,710 + 423,023 + 647,407 lbs H₂SO₄/yr

 $TSAM = 2,159,140 lbs H_2SO_4/yr$

The 25,000 lbs/yr threshold has been exceeded; therefore, a release estimate must be made and the result reported on Form R.

Released

Total Releases (from previous examples)

 $TSAR = \{ [(EM_{Comb} + EM_{SCR}) - NH3_{SCR}] \bullet F2_{APH} + (EM_{FGC_afterAPH} - NH3_{FGC_afterAPH}) \} \bullet F2_{ESP}$

Check that subtotal from upstream sources is positive:

 $TSAR_{upstream} = [(1,088,710 + 423,023) - 33,141] = 1,478,592 > 0$

 $TSAR = [(1,088,710 + 423,023) - 33,141] \bullet 0.50 + (647,407 - 277,460)] \bullet 0.63$

 $TSAR = 698,823 lbs H_2SO_4/yr$

Example 8: Coal-Fired Boiler with FGC, NG Startup Fuel, and Used Oil Co-Firing

A 500 MW PC boiler, equipped with a cold-side electrostatic precipitator, burns an Eastern bituminous coal as the main fuel. The plant is equipped with an SCR process that operates during the ozone season only, so that 0.43 of the coal burn occurred with the SCR operating. The SCR is bypassed when not in operation. The SCR catalyst SO₂ oxidation rate specified in the design is 0.75%, and the ammonia slip is estimated to be 0.75 ppmv. The plant is also equipped with a FGC process that injects both SO₃ and NH₃ upstream of the air preheater. The SO₃ is injected at 7 ppmv at 6% O₂ wet, and the ammonia at 3 ppmv also at 6% O₂ wet. The FGC system operates whenever the plant is on, except during startup and shutdown, with an operating factor estimated at 0.9.

The coal used in the reporting year is 1,126,938 tons with a weighted average sulfur concentration of 2.0% and a heating value of 12,000 Btu/lb.

Natural gas is used as a startup fuel, with 0.5 Tbtu per year. During startup, neither the SCR nor the FGC system is used. Used oil is also burned, with 483.2 tons burned (0.0185 Tbtu/yr) in the year. Since the used oil is burned when the unit is at full load, it is burned while the SCR and FGC are both operating. The used oil has a sulfur content of 0.1% from analysis.

Solution

Coal fuel calculations

Manufactured from coal

Total (from previous examples)

 $TSAM = EM_{Comb} + EM_{SCR} + EM_{FGC}$

TSAM= 1,088,710 + 423,023 + 647,407 lbs H₂SO₄/yr

TSAM = 2,159,140 lbs H₂SO₄/yr

Released from coal

Total Releases (from previous examples) TSAR_{coal} = 582,290 lbs H₂SO₄/yr

Natural gas fuel calculations

Manufactured from natural gas

Combustion

$$\begin{split} E2_{NG} &= K_b \bullet B_{NG} \bullet S \\ E2_{NG} &= 0.0001359 \bullet 0.5 \ Tbtu/yr \bullet 2000 \ gr/10^6 \ scf \\ E2_{NG} &= 0.136 \ tons \ SO_2/year \\ EM_{Comb} &= K \ \bullet F1 \ \bullet E2_{NG} \end{split}$$

 $EM_{Comb} = 3063 \bullet 0.01 \bullet 0.136$

 $EM_{Comb} = 4.17 lbs H_2SO_4$ manufactured

No sulfuric acid is manufactured in either the SCR or FGC.

Released from natural gas

Total Releases

 $TSAR_{NG} = EM_{Comb} \bullet F2_{APH} \bullet F2_{ESP}$

 $TSAR_{NG} = 4.17 \bullet 0.50 \bullet 0.63$

 $TSAR_{NG} = 1.31 lbs H_2SO_4/yr$

F2 is taken to be equal to 0.50 for the air heater and 0.63 for the ESP.

Used oil fuel calculations

Manufactured from used oil

Combustion

 $E2 = K1 \bullet K2 \bullet C1 \bullet S1$ $E2 = 0.02 \bullet 1.0 \bullet 483.2 \bullet 0.1$ $E2 = 0.966 \text{ tons } SO_2/\text{year}$ $EM_{Comb} = K \bullet F1 \bullet E2$

 $EM_{Comb} = 51.8 lbs H_2SO_4$ manufactured

 $EM_{Comb} = 3063 \bullet 0.0175 \bullet 0.966$

SCR

 $EM_{SCR} = K \bullet S2 \bullet f_s \bullet E2$

 $EM_{SCR} = 3063 \bullet 0.0075 \bullet 1.0 \bullet 0.966$

 $EM_{SCR} = 22.2$ lbs H₂SO₄ manufactured

FGC

 $EM_{FGC} = K_e \bullet B \bullet f_e \bullet I_s \bullet F3_{FGC}$

 $EM_{FGC} = 3799 \bullet 0.0185 \bullet 1.0 \bullet 7.0 \bullet 1$

 $EM_{FGC} = 492 lbs H_2SO_4$ manufactured

Total Manufactured

 $TSAM = EM_{Comb} + EM_{SCR} + EM_{FGC}$

 $TSAM = 51.8 + 22.2 + 492 lbs H_2SO_4/yr$

 $TSAM = 566 lbs H_2SO_4/yr$

Released from used oil

Ammonia Slip

 $NH3_{SCR} = K_s \bullet B \bullet f_{sreagent} \bullet S_{NH3}$

NH3_{SCR} = 3799 • 0.0185 • 1.0 • 0.75 = 52.7 lbs NH₃/yr

FGC Ammonia Injection

 $NH3_{FGC} = K_e \bullet B \bullet f_e \bullet I_{NH3}$

 $NH3_{FGC} = 3799 \bullet 0.0185 \bullet 1.0 \bullet 3.0 = 211 lbs NH_3/yr$

Total Releases

 $TSAR_{oil} = [(EM_{Comb} + EM_{SCR} + EM_{FGC}) - (NH3_{SCR} + NH3_{FGC})] \bullet F2_{APH} \bullet F2_{ESP}$

 $TSAR_{oil} = [(51.8 + 22.2 + 492) - (52.7 + 211)] \bullet 0.5 \bullet 0.63$

 $TSAR_{oil} = 95 lbs H_2SO_4/yr$

F2 is taken to be equal to 0.50 for the air heater and 0.63 for the ESP

Grand totals for all fuels

Manufactured

 $TSAM = TSAM_{coal} + TSAM_{NG} + TSAM_{oil}$

 $TSAM = 2,159,140 + 4.17 + 566 lbs H_2SO_4/yr$

TSAM = 2,159,710 lbs H₂SO₄/yr

Released

 $TSAR = TSAR_{coal} + TSAR_{NG} + TSAR_{oil}$ $TSAR = 582,290 + 1.31 + 95 \text{ lbs } H_2SO_4/\text{yr}$ $TSAR = 582,386 \text{ lbs } H_2SO_4/\text{yr}$

Example 9: Natural Gas Combined Cycle Plant

A natural gas-fired combined-cycle unit uses 12.3 Tbtu/yr. of natural gas, with the standard sulfur content.

Solution

Manufactured

$$\begin{split} E2_{NG} &= K_b \bullet B_{NG} \bullet S \\ E2_{NG} &= 0.0001359 \bullet 12.3 \bullet 2000 \\ E2_{NG} &= 3.343 \text{ tons } SO_2/yr \Box \end{split}$$

Combustion

 $EM_{CC} = K \bullet F1 \bullet E2_{NG}$

 $EM_{CC} = 3063 \bullet 0.0555 \bullet 3.343$

 $EM_{CC} = 568 \text{ lbs } H_2SO_4/yr$

Released

 $TSAR_{NG} = EM_{Comb} \bullet F2_{CC}$

 $TSAR_{NG} = 568 \bullet 0.5$

 $TSAR_{NG} = 284 \ lbs \ H_2SO_4/yr$

 $F2_{CC} = 0.5$ because the low temperature of the back-end tubes of the HRSG act like an air heater.

Example 10: SCR and CO catalyst added to Example 9

A natural gas-fired combined-cycle unit uses 12.3 Tbtu/yr. of natural gas, with the standard sulfur content. The plant is equipped with an SCR process that operates continually (operating factor of 1.0). The SCR catalyst SO₂ oxidation rate specified in the design is 2.0%, and the ammonia slip is estimated to be 0.75 ppmv. The plant is also equipped with a CO catalyst, which also is in continuous operation, and has an estimated SO₂ oxidation rate of 10%.

Solution

```
Manufactured

E2_{NG} = K_b \bullet B_{NG} \bullet S

E2_{NG} = 0.0001359 \bullet 12.3 \bullet 2000

E2_{NG} = 3.343 \text{ tons } SO_2/yr \square

Combustion

EM_{CC} = K \bullet F1 \bullet E2_{NG}

EM_{CC} = 3063 \bullet 0.0555 \bullet 3.343

EM_{CC} = 568 \text{ lbs } H_2SO_4/yr

CO Catalyst

EM_{CC_CO} = K \bullet S3 \bullet f_{COops} \bullet E2

EM_{CC_CO} = 3063 \bullet 0.1 \bullet 1.0 \bullet 3.343

EM_{CC_CO} = 1024 \text{ lbs } H_2SO_4/yr
```

SCR

```
EM_{CCSCR} = [(K \bullet E2) - EM_{CC-CO}] \bullet f_{sops} \bullet S2 \bullet F3_{SCR}EM_{CCSCR} = [(3063 \bullet 3.343) - 1024] \bullet 1.0 \bullet 0.02 \bullet 1.0
```

```
EM_{CCSCR} = 184 lbs H_2SO_4/yr
```

Total

 $TSAM_{CC} = EM_{Comb} + EM_{CC_CO} + EM_{SCR}$

 $TSAM_{CC} = 568 + 1024 + 184$

 $TSAM_{CC} = 1776 lbs H_2SO_4/yr$

Released

```
TSAR_{CC} = [TSAM - (K_s \bullet B \bullet f_{sreagent} \bullet S_{NH3})] \bullet F2_{cc}TSAR_{CC} = [1776 - (3799 \bullet 12.3 \bullet 1.0 \bullet 0.75)] \bullet 0.5TSAR_{CC} = -16,635 = 0 \text{ lbs } H_2SO_4/\text{yr}
```

The total releases are zero because the actual result is a negative value, indicating that the ammonia slip is greater than the sulfuric acid, thereby forming ABS, which is not reportable.

 $F2_{CC} = 0.5$ because the low temperature of the back-end tubes of the HRSG act like an air heater.

B CHRONOLOGY OF CHANGES TO THE EPRI SULFURIC ACID ESTIMATION MODEL

Modification	Description
Stack correction for CEMs SO ₂ measurements	Added equation using wall affects factor (WAF) for CEMs measurements in conjunction with Method 2H.
Total Sulfuric Acid Release Methodology (TSAR)	Changed application location of F2 _{Hotside_ESP} and F2 _{Mag-Ox} such that they apply only to EM _{COMB} .
Partial-Year SCR/SNCR	Added methodology to calculate the Total Sulfuric Acid Release if injection for SCR/SNCR is partial-year.

EPRI, 2018 (EPRI Report 3002012398)

EPRI, 2012 (EPRI Report 1023790)

Modification	Description
SCR Factor (F3 _{SCR})	Changes made to Table 4-2 to clarify proper use of the F3 _{SCR} factor.
Alkali Injection	 Changed description of Falkali such that it is fraction of coal burn instead of fraction of operating year, to be consistent with other operating factors Added alternate approach for calculating releases with alkali injection to be used if the utility does not have measured data. F3_{ALKINJ} factor is introduced.
Total Sulfuric Acid Release Methodology (TSAR)	Reorganized release equations such that sources are summed upstream of the APH and ammonia from SCR/SNCR is applied to reduce releases based on ABS formation, then the F2 factor for the APH is applied. Then, any downstream source of SO ₃ or ammonia from downstream FGC is added, followed by application of all remaining F2 factors.
FGC Factor (F3 _{FGC})	Corrected the $F3_{FGC}$ factor. The $F3_{FGC}$ factor was introduced in the 2001 version of the report, but as actual data has become available to update the F2 factors, the $F3_{FGC}$ factor has become outdated. $F3_{FGC}$ will now account only for alkalinity in PRB coal, and be applied directly to the manufacture equation.

EPRI, 2012 (EPRI Report 1023790), continued

Combined Cycle	Introduced a potential third source of sulfuric acid manufacture and release from oxidation across CO catalysts.
Technology Factors (F2)	Added Wet ESP, all fuels: 0.1. Changed FGD, venturi, all coals from 0.65 to 0.73 (2 new data points).
	Updated and corrected Figure 4-4 to reflect two new data points, change the shaded area to reflect the proper data range, and deleted an inaccurate PRB/lignite data point
	Changed the F2 factor in Table 4-6 for Mg-Ox into furnace from 0.5 to 0.25 to rectify an error introduced in the 2004 version of the report.
Examples	Updated examples to account for new release equation methodology.

EPRI, 2010a (EPRI Report 1020636)

Modification	Description
Fuel Impact Factor (F1)	Combined low and high-sulfur eastern bituminous coals into a single category with a linear curve fit. F1 factor based on fuel SO ₂ content calculated from fuel sulfur content:
	$F1_{ebit} = 1.1163E-6 \bullet SO_2 + 0.0065$
	Changed the following F1 factors based on additional data:
	Subbit/PRB – changed from 0.0018 to 0.0019
	Lignite – changed from 0.0048 to 0.0044
Technology Factors (F2)	Changed factors for APH and ESP due to addition and re- evaluation of data:
	APH, low-sulfur east. bit. – changed from 0.49 to 0.50
	APH, PRB – changed from 0.56 to 0.36
	ESP, low-sulfur east. bit – changed from 0.49 to 0.63
	ESP, PRB – changed from 0.73 to 0.72

Alkali Injection	Added a procedure to determine total releases from partial year injection.
	Incorporated correction to alkali injection conversion factor, Kalkali.
Report Organization	Moved F2 Factors from Section 3 to Section 4.
Examples	Updated examples to account for new F1 and F2 factors.

EPRI, 2010a (EPRI Report 1020636) (continued)

EPRI, 2008 (EPRI Report 1016384)

Modification	Description
Alkali Injection	Added a procedure to estimate a stack release using a measured or specified stack gas SO ₃ concentration, to support estimates on units with alkali injection.
	Errata page published to correct error in alkali injection conversion factor, K _{alkali} .
Combustion Turbines	Clarified procedures for estimating emissions from simple and combined cycle CTs.
Blended Coals	New method calculates SO ₃ manufacture by calculating the SO ₃ emission rate separately for each of the component coals, and proportioning the contribution of SO ₃ from each. The prior method assumed the SO ₃ production was the same as that of the major coal component.
Examples	Provided new example calculations for alkali injection and blended coal.

EPRI, 2007 (Report 1014773) changes from Harriso	on, 2005
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Modification	Description
Formatting	Adapted Southern Company paper (Harrison, 2005) to EPRI format.
Fuel (F1) Factors	Changed the following F1 factors based on additional data:
	Low S E. Bit, Dry Boiler – from 0.0080 to 0.0082
	Subbit. (including PRB), Dry Boiler – from 0.0014 to 0.0018
	Subbit. (including PRB), Cyclone – from 0.003 to 0.0018

Technology (F2) Factors	Added new F2 Factors:
	Air Preheater, PRB fuel - 0.56
	Cold-side ESP, PRB fuel – 0.73
SCR	Added a Technology Impact Factor for SCR, F3 _{SCR} .

EPRI, 2007 (Report 1014773) changes from Harrison, 2005 (continued)

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