METHOD 19-DETERMINATION OF SULFUR DIOXIDE REMOVAL EFFICIENCY AND PARTICULATE MATTER, SULFUR DIOXIDE, AND NITROGEN OXIDES EMISSION RATES

1. Applicability and Principle

- 1.1 Applicability. This method is applicable for (a) determining particulate matter (PM), sulfur dioxide (SO₂), and nitrogen oxides (NO_x) emission rates; (b) determining sulfur removal efficiencies of fuel pretreatment and SO₂ control devices; (c) determining overall reduction of potential SO₂ emissions from steam generating units or other sources as specified in applicable regulations; and (d) determining SO₂ rates based on fuel sampling and analysis procedures.
 - 1.2 Principle.
- 1.2.1 Pollutant emission rates are determined from concentrations of PM, SO_2 , or NO_x , and oxygen (O_2) or carbon dioxide (CO_2) along with F factors (ratios of combustion gas volumes to heat inputs).
- $1.2.2\,$ An overall SO_2 emission reduction efficiency is computed from the efficiency of fuel pretreatment systems (optional) and the efficiency of SO_2 control devices.
- 1.2.3 The sulfur removal efficiency of a fuel pretreatment system is determined by fuel sampling and analysis of the sulfur and heat contents of the fuel before and after the pretreatment system.
- 1.2.4 The ${\rm SO_2}$ removal efficiency of a control device is determined by measuring the ${\rm SO_2}$ rates before and after the control device.
- 1.2.5 The inlet rates to SO_2 control systems and when SO_2 control systems are not used, SO_2 emission rates to the atmosphere may be determined by fuel sampling and analysis (optional).
- 2. Emission Rates of Particulate Matter, Sulfur Dioxide, and Nitrogen Oxides

Select from the following sections the applicable procedure to compute the PM, SO_2 , or NO_x emission rate (E) in ng/J (lb/million Btu). The pollutant concentration must be in ng/scm (lb/scf) and the F factor must be in scm/J (scf/million Btu). If the pollutant concentration (C) is not in the appropriate units, use the following table to make the proper conversion:

Conversion Factors for Concentration

From	 To	Multiply by		
g/scm	ng/scm ng/scm ng/scm ng/scm ng/scm ng/scm lb/scf lb/scf	10°		

An F factor is the ratio of the gas volume of the products of combustion to the heat content of the fuel. The dry F factor (F_d) includes all components of combustion less water, the wet F factor (F_w) includes all components of combustion, and the carbon F factor (F_c) includes only carbon dioxide.

Note: Since F_w factors include water resulting only from the combustion

of hydrogen in the fuel, the procedures using $F_{\rm w}$ factors are not applicable for computing E from steam generating units with wet scrubbers or with other processes that add water (e.g., steam injection).

2.1 Oxygen-Based F Factor, Dry Basis. When measurements are on a dry basis for both O_2 (% O_{2d}) and pollutant (C_d) concentrations, use the following equation:

$$E = C_d F_d \frac{20.9}{(20.9 - 8O_{2d})}$$
 Eq. 19-1

- 2.2 Oxygen-Based F Factor, Wet Basis. When measurements are on a wet basis for both O_2 (O_{2w}) and pollutant (O_{w}) concentrations, use either of the following:
 - 2.2.1 If the moisture fraction of ambient air (B_{wa}) is measured:

$$E = C_w F_w \frac{20.9}{(20.9(1-B_{wa}) - 8O_{2w})}$$
 Eq. 19-2

Instead of actual measurement, B_{wa} may be estimated according to the procedure below. (Note: The estimates are selected to ensure that negative errors will not be larger than -1.5 percent. However, positive errors, or over-estimation of emissions, of as much as 5 percent may be introduced depending upon the geographic location of the facility and the associated range of ambient moisture):

- 2.2.1.1 $$B_{\text{wa}}\text{=}0.027$$. This value may be used at any location at all times.
- 2.2.1.2 B_{wa} =Highest monthly average of B_{wa} that occurred within the previous calendar year at the nearest Weather Service Station. This value shall be determined annually and may be used as an estimate for the entire current calendar year.
- 2.2.1.3 B_{wa}=Highest daily average of Bwa that occurred within a calendar month at the nearest Weather Service Station, calculated from the data from the past 3 years. This value shall be computed for each month and may be used as an estimate for the current respective calendar month.
 - 2.2.2 If the moisture fraction (B_{ws}) of the effluent gas is measured:

$$E = C_w F_d \frac{20.9}{(20.9(1-B_{wg}) - \%O_{2w})}$$
 Eq. 19-3

- 2.3 Oxygen-Based F Factor, Dry/Wet Basis.
- 2.3.1 When the pollutant concentration is measured on a wet basis (\texttt{C}_{w}) and \texttt{O}_{2} concentration is measured on a dry basis $(\$\texttt{O}_{2d})$, use the following equation:

$$E = \frac{C_w F_d}{(1 - B_{ws})} \frac{20.9}{(20.9 - 8O_{2w})}$$
 Eq. 19-4

2.3.2 When the pollutant concentration is measured on a dry basis (C_{d}) and the O_2 concentration is measured on a wet basis $(\text{\%O}_{\text{2w}})\text{,}$ use the following equation:

$$E = C_{d}F_{d} = \frac{20.9}{20.9 - \frac{O_{2w}}{(1 - B_{ws})}}$$
 Eq. 19-5

2.4 Carbon Dioxide-Based F Factor, Dry Basis. When measurements are on a dry basis for both $\rm CO_2$ (%CO $_{\rm 2d})$ and pollutant (C $_{\rm d}$) concentrations, use the following equation:

$$E = C_d F_c \frac{100}{{}^{8}CO_{2d}}$$
 Eq. 19-6

2.5 Carbon Dioxide-Based F Factor, Wet Basis. When measurements are on a wet basis for both $\rm CO_2$ (%CO $_{\rm 2w}$) and pollutant (C $_{\rm w}$) concentrations, use the following equation:

$$E = C_w F_c \frac{100}{{}^8CO_{2w}}$$
 Eq. 19-7

- 2.6 Carbon Dioxide-Based F Factor, Dry/Wet Basis.
- 2.6.1 When the pollutant concentration is measured on a wet basis (C_w) and CO_2 concentration is measured on a dry basis $(%CO_{2d})$, use the following equation:

$$E = \frac{C_w F_c}{(1 - B_{ws})} \frac{100}{%CO_{2d}}$$
 Eq. 19-8

2.6.2 When the pollutant concentration is measured on a dry basis (C_{d}) and CO_2 concentration is measured on a wet basis $(\text{%CO}_{2w})\text{,}$ use the following equation:

$$E = C_d (1 - B_{ws}) F_c \frac{100}{\$CO_{2w}}$$
 Eq. 19-9

- 2.7 Direct-Fired Reheat Fuel Burning. The effect of direct-fired reheat fuel burning (for the purpose of raising the temperature of the exhaust effluent from wet scrubbers to above the moisture dew-point) on emission rates will be less than ±1.0 percent and, therefore, may be ignored.
- $2.8\,$ Combined Cycle-Gas Turbine Systems. For gas turbine-steam generator combined cycle systems, determine the emissions from the steam generating unit or the percent reduction in potential SO_2 emissions as follows:
- 2.8.1 Compute the emission rate from the steam generating unit using the following equation:

$$E_{bo} = E_{co} + (E_{co} - E_{g}) \frac{H_{g}}{H_{b}}$$
 Eq. 19-10

where:

 E_{bo} = pollutant emission rate from the steam generating unit, ng/J (1b/million Btu).

 E_{co} = pollutant emission rate in combined effluent, ng/J

(lb/million Btu).

 $E_{\rm g}$ = pollutant rate from gas turbine, ng/J (lb/million Btu). $H_{\rm b}$ = heat input rate to the steam generating unit from fuels fired in the steam generating unit, J/hr (million Btu/hr). $H_{\rm g}$ = heat input rate to gas turbine from all fuels fired in the

 H_g = heat input rate to gas turbine from all fuels fired in t gas turbine, J/hr (million Btu/hr).

 $2.8.1.1\,$ Use the test methods and procedures section of Subpart GG to obtain E_{co} and $E_g.$ Do not use F_w factors for determining E_g or $E_{co}.$ If an SO_2 control device is used, measure E_{co} after the control device.

2.8.1.2 Suitable methods shall be used to determine the heat input rates to the steam generating units (H_b) and the gas turbine (H_a) .

2.8.2 If a control device is used, compute the percent of potential SO_2 emissions (% $P_{\rm s}$) using the following equations:

$$E_{bi} = E_{ci} - (E_{ci} - E_{g}) \frac{H_{g}}{H_{b}}$$
 Eq. 19-11

$$P_{s} = 100 (1 - \frac{E_{bo}}{E_{bi}})$$
 Eq. 19-12

where:

 ${\tt E}_{\tt bi}$ = pollutant rate from the steam generating unit, ng/J

(lb/million Btu)

 E_{ci} = pollutant rate in combined effluent, ng/J (lb/million Btu).

Use the test methods and procedures section of Subpart GG to obtain E_{ci} and E_{g} . Do not use F_{w} factors for determining E_{g} or E_{ci} .

F Factors

Use an average F factor according to Section 3.1 or determine an applicable F factor according to Section 3.2. If combined fuels are fired, prorate the applicable F factors using the procedure in Section 3.3.

3.1 Average F Factors. Average F factors ($F_{\rm d}$, $F_{\rm w}$, or $F_{\rm c}$) from Table 19-1 may be used.

Table 19-1-F Factors for Various Fuels¹

	F _d		F _w		F_c	
Fuel type	dscm/J	dscf/10 ⁶ Btu	wscm/J 	wscf/10 ⁶ Btu	scm/J 	scf/10 ⁶ Btu
Coal: Anthracite ² 1,970	2.71*10 ⁻⁷	10	 ,100 2.83 ²	 *10 ⁻⁷	10,540	0.530*10 ⁻⁷
Bituminous ² 1,800	2.63*10 ⁻⁷	 9 	 ,780 2.86 ³ 	*10 ⁻⁷	10,640	0.484*10 ⁻⁷
Lignite 1,910	2.65*10 ⁻⁷	9	,860 3.21 ³	*10 ⁻⁷	11,950	0.513*10 ⁻⁷
Oil ³ 1,420	2.47*10 ⁻⁷	9	,190 2.77	*10 ⁻⁷	10,320	0.383*10 ⁻⁷
Gas:		1	1			
Natural 1,040	2.43*10 ⁻⁷	8	,710 2.85	10-7	10,610	0.287*10 ⁻⁷
Propane 1,190	2.34*10 ⁻⁷	8	,710 2.74	*10 ⁻⁷	10,200	0.321*10 ⁻⁷
Butane 1,250	2.34*10 ⁻⁷	8	,710 2.79	*10 ⁻⁷	10,390	0.337*10 ⁻⁷
Wood 1,830	2.48*10 ⁻⁷	9	,240		0.492	2*10 ⁻⁷
Wood Bark. 1,920	2.58*10 ⁻⁷	9	,600		0.516	5*10 ⁻⁷
	2.57*10 ⁻⁷	9	,570		0.488	3*10 ⁻⁷
Solid Waste	, 					

¹ Determined at standard conditions: 20 °C (68 °F) and 760 mm Hg (29.92 in. Hg).

3.2 Determined F Factors. If the fuel burned is not listed in Table 19-1 or if the owner or operator chooses to determine an F factor rather than use the values in Table 19-1, use the procedure below:

3.2.1 Equations. Use the equations below, as appropriate, to compute the F factors:

$$F_{d} = K \frac{[(K_{hd} %H) + (K_{c} %C) + (K_{s} %S) + (K_{n} %N) - (K_{o} %O)]}{GCV_{w}}$$
 Eq. 19-13

$$K = \frac{(K_{hw} %H) + (K_{c} %C) + (K_{s} %S) + (K_{n} %N) - (K_{o} %O) + (K_{w} %H_{2})}{GCV_{w}}$$
Eq. 19-14

 $^{^{2}}$ As classified according to ASTM D388-77.

³ Crude, residual, or distillate.

$$F_{c} = K \frac{(K_{cc} %C)}{GCV}$$
 Eq. 19-15

(Note.-Omit the H_2O term in the equations for F_w if %H and %O include the unavailable hydrogen and oxygen in the form of $\text{H}_2\text{O.})$ where:

 $F_{\rm d}, F_{\rm w}, F_{\rm c}$ = volumes of combustion components per unit of heat content, scm/J (scf/million Btu).

%H, %C, %S, %N, %O, and %H $_2$ O=concentrations of hydrogen, carbon, sulfur, nitrogen, oxygen, and water from an ultimate analysis of fuel, weight percent. GCV = gross calorific value of the fuel consistent with the ultimate analysis, kJ/kg (Btu/lb).

K = conversion factor, 10^{-5} (kJ/J)/(%) [10^{6} Btu/million Btu].

 $\begin{array}{lll} {\rm K_{hd}} & = & 22.7 \; ({\rm scm/kg}))[(3.64 \; ({\rm scf/lb})/(\$)]. \\ {\rm K_c} & = & 9.57 \; ({\rm scm/kg})[(1.53 \; ({\rm scf/lb})/(\$)]. \\ {\rm K_s} & = & 3.54 \; ({\rm scm/kg}) \; [(0.57 \; ({\rm scf/lb})/(\$)]. \\ {\rm K_n} & = & 0.86 \; ({\rm scm/kg} \; [0.14 \; ({\rm scf/lb})/(\$)]. \\ {\rm K_o} & = & 2.85 \; ({\rm scm/kg}) \; [0.46 \; ({\rm scf/lb})/(\$)]. \\ {\rm K_{hw}} & = & 34.74 \; ({\rm scm/kg}) \; [(5.57 \; ({\rm scf/lb})/(\$)]. \\ {\rm K_w} & = & 1.30 \; ({\rm scm/kg}) \; [(0.21 \; ({\rm scf/lb})/(\$)]. \\ {\rm K_{cc}} & = & 2.0 \; ({\rm scm/kg}) \; [(0.321 \; ({\rm scf/lb})/(\$)]. \end{array}$

- 3.2.2 Use applicable sampling procedures in Section 5.2.1 or 5.2.2 to obtain samples for analyses.
- 3.2.3 Use ASTM D3176-74 (incorporated by reference-see § 60.17) for ultimate analysis of the fuel.
- 3.2.4 Use applicable methods in Section 5.2.1 or 5.2.2 to determine the heat content of solid or liquid fuels. For gaseous fuels, use ASTM D1826-77 (IBR-see § 60.17) to determine the heat content.
- 3.3 F Factors for Combination of Fuels. If combinations of fuels are burned, use the following equations, as applicable unless otherwise specified in applicable subpart:

$$F_{d} = \sum_{k=1}^{n} X_{k} F_{dk}$$
 Eq. 19-16

$$F_{w} = \sum_{k=1}^{n} X_{k} F_{wk}$$
 Eq. 19-17

$$F_{c} = \sum_{k=1}^{n} X_{k} F_{ck}$$
 Eq. 19-18

where:

 X_k = fraction of total heat input from each type of fuel k. n = number of fuels being burned in combination.

- 4. Determination of Average Pollutant Rates
 - 4.1 Average Pollutant Rates from Hourly Values. When hourly average

pollutant rates (E_h) , inlet or outlet, are obtained (e.g., CEMS values), compute the average pollutant rate (E_a) for the performance test period (e.g., 30 days) specified in the applicable regulation using the following equation:

$$E_a = \frac{1}{H} \sum_{j=1}^{n} E_{hj}$$
 Eq. 19-19

where:

 E_a = average pollutant rate for the specified performance test period, ng/J (lb/million Btu).

 E_h = hourly average pollutant, ng/J (lb/million Btu).

H = total number of operating hours for which pollutant rates are determined in the performance test period.

4.2 Average Pollutant Rates from Other than Hourly Averages. When pollutant rates are determined from measured values representing longer than 1-hour periods (e.g., daily fuel sampling and analyses or Method 6B values), or when pollutant rates are determined from combinations of 1-hour and longer than 1-hour periods (e.g., CEMS and Method 6B values), compute the average pollutant rate ($\rm E_a$) for the performance test period (e.g., 30 days) specified in the applicable regulation using the following equation:

$$E_{a} = \frac{\sum_{j=1}^{D} (n_{d}E_{d})_{j}}{\sum_{j=1}^{D} n_{dj}}$$
 Eq. 19-20

where:

 $E_{\rm d}$ = average pollutant rate for each sampling period (e.g., 24 hr Method 6B sample or 24-hr fuel sample) or for each fuel lot (e.g., amount of fuel bunkered), ng/J (lb/million Btu).

 $n_{\rm d}$ = number of operating hours of the affected facility within the performance test period for each Ed determined.

D = number of sampling periods during the performance test period.

4.3 Daily Geometric Average Pollutant Rates from Hourly Values. The geometric average pollutant rate $(E_{\rm ga})$ is computed using the following equation:

$$E_{ga} = EXP \left[\frac{1}{n} \sum_{j=1}^{n} ln(E_{hj}) \right]$$
 Eq. 19-20a

where:

 E_{ga} = daily geometric average pollutant rate, ng/J (lbs/million Btu) or ppm corrected to 7 percent O_2 .

 E_{hj} = hourly arithmetic average pollutant rate for hour "j," ng/J (lb/million Btu) or ppm corrected to 7 percent O_2 .

n = total number of hourly averages for which pollutant rates are available within the 24 hr midnight to midnight daily period.

ln = natural log of indicated value.

EXP = the natural logarithmic base (2.718) raised to the value enclosed by brackets.

- Determination of Overall Reduction in Potential Sulfur Dioxide Emission 5.
- Overall Percent Reduction. Compute the overall percent SO, reduction (%R_o) using the following equation:

$$R_{\circ} = 100[1.0 - (1.0 - \frac{R_{f}}{100})(1.0 - \frac{R_{g}}{100})]$$
 Eq. 19-21

where:

%R_f = SO₂ removal efficiency from fuel pretreatment, percent. SO₂ removal efficiency of the control device, percent.

5.2 Pretreatment Removal Efficiency (Optional). Compute the SO, removal efficiency from fuel pretreatment (R_f) for the averaging period (e.g., 90 days) as specified in the applicable regulation using the following equation:

$$R_{f} = 100[1.0 - \frac{\sum_{j=1}^{n} \frac{8S_{pj}}{GCV_{pj}} L_{pj}}{\sum_{j=1}^{n} \frac{8S_{rj}}{GCV_{rj}} L_{rj}}]$$
Eq. 19-22

where:

 $\rm \$S_p, \ \S_r = sulfur content of the product and raw fuel lots, respectively, dry basis weight percent.

 GCV_p , GCV_r = gross calorific value for the product and raw fuel lots, respectively, dry basis, kg/kg (Btu/lb).

 L_p , L_r weight of the product and raw fuel lots, respectively,

metric ton (ton).

number of fuel lots during the averaging period.

Note: In calculating $R_{\rm f},$ include $\rm Note$ and GCV values for all fuel lots that are not pretreated and are used during the averaging period.

- 5.2.1 Solid Fossil (Including Waste) Fuel-Sampling and Analysis. Note: For the purposes of this method, raw fuel (coal or oil) is the fuel delivered to the desulfurization (pretreatment) facility. For oil, the input oil to the oil desulfurization process (e.g., hydrotreatment) is considered to be the raw fuel.
- 5.2.1.1 Sample Increment Collection. Use ASTM D2234-76 (IBR-see § 60.17), Type I, Conditions A, B, or C, and systematic spacing. As used in this method, systematic spacing is intended to include evenly spaced increments in time or increments based on equal weights of coal passing the collection area.

As a minimum, determine the number and weight of increments required per gross sample representing each coal lot according to Table 2 or Paragraph 7.1.5.2 of ASTM D2234-76. Collect one gross sample for each lot of raw coal and one gross sample for each lot of product coal.

5.2.1.2 ASTM Lot Size. For the purpose of Section 5.2 (fuel pretreatment), the lot size of product coal is the weight of product coal from one type of raw coal. The lot size of raw coal is the weight of raw coal used to produce one lot of product coal. Typically, the lot size is the weight of coal processed in a 1-day (24-hour) period. If more than one type of coal is treated and produced in 1 day, then gross samples must be collected and analyzed for each type of coal. A coal lot size equaling the 90-day quarterly fuel quantity for a steam generating unit may be used if representative

sampling can be conducted for each raw coal and product coal.

Note: Alternative definitions of lot sizes may be used, subject to prior approval of the Administrator.

- 5.2.1.3 Gross Sample Analysis. Use ASTM D2013-72 to prepare the sample, ASTM D3177-75 or ASTM D4239-85 to determine sulfur content (%S), ASTM D3173-73 to determine moisture content, and ASTM D2015-77 or ASTM D3286-85 to determine gross calorific value (GCV) (all methods cited IBR-see § 60.17) on a dry basis for each gross sample.
- 5.2.2 Liquid Fossil Fuel-Sampling and Analysis. See Note under Section 5.2.1.
- 5.2.2.1~ Sample Collection. Follow the procedures for continuous sampling in ASTM D270-65 (Reapproved 1975) (IBR-see § 60.17) for each gross sample from each fuel lot.
- 5.2.2.2 Lot Size. For the purpose of Section 5.2 (fuel pretreatment), the lot size of a product oil is the weight of product oil from one pretreatment facility and intended as one shipment (ship load, barge load, etc.). The lot size of raw oil is the weight of each crude liquid fuel type used to produce a lot of product oil.

Note: Alternative definitions of lot sizes may be used, subject to prior approval of the Administrator.

- 5.2.2.3 Sample Analysis. Use ASTM D129-64 (Reapproved 1978), ASTM D1552-83, or ASTM D4057-81 to determine the sulfur content (%S) and ASTM D240-76 (all methods cited IBR-see § 60.17) to determine the GCV of each gross sample. These values may be assumed to be on a dry basis. The owner or operator of an affected facility may elect to determine the GCV by sampling the oil combusted on the first steam generating unit operating day of each calendar month and then using the lowest GCV value of the three GCV values per quarter for the GCV of all oil combusted in that calendar quarter.
- 5.2.3 Use appropriate procedures, subject to the approval of the Administrator, to determine the fraction of total mass input derived from each type of fuel.
- 5.3 Control Device Removal Efficiency. Compute the percent removal efficiency (R_{α}) of the control device using the following equation:

$$R_g = 100[1.0 - \frac{E_{ao}}{E_{ai}}]$$
 Eq. 19-23

where:

 E_{ao} , E_{ai} = average pollutant rate of the control device, outlet and inlet, respectively, for the performance test period, ng/J (lb/million Btu).

- 5.3.1 Use continuous emission monitoring systems or test methods, as appropriate, to determine the outlet SO_2 rates and, if appropriate, the inlet SO_2 rates. The rates may be determined as hourly (E_h) or other sampling period averages (E_d) . Then, compute the average pollutant rates for the performance test period $(E_{ao}$ and $E_{ai})$ using the procedures in Section 4.
- 5.3.2 As an alternative, as-fired fuel sampling and analysis may be used to determine inlet SO_2 rates as follows:
- 5.3.2.1 Compute the average inlet ${\rm SO_2}$ rate $({\rm E_{di}})$ for each sampling period using the following equation:

$$E_{di} = K \frac{%S}{GCV}$$
 Eq. 19-24

where:

 E_{di} = average inlet SO₂ rate for each sampling period d, ng/J

(lb/million Btu)

%S = sulfur content of as-fired fuel lot, dry basis, weight percent.

GCV = gross calorific value of the fuel lot consistent with the sulfur

analysis, kJ/kg (Btu/lb).

K = $2*10^7$ [(kg)(ng)/(%)(J)]{ $2*10^4$ (lb)(Btu/(%))(million Btu)} After calculating E_{di} use the procedures in Section 4.2 to determine the average inlet SO2 rate for the performance test period (E_{ai}).

5.3.2.2 Collect the fuel samples from a location in the fuel handling system that provides a sample representative of the fuel bunkered or consumed during a steam generating unit operating day.

For the purpose of as-fired fuel sampling under Section 5.3.2 or Section 6, the lot size for coal is the weight of coal bunkered or consumed during each steam generating unit operating day. The lot size for oil is the weight of oil supplied to the "day" tank or consumed during each steam generating unit operating day.

For reporting and calculation purposes, the gross sample shall be identified with the calendar day on which sampling began. For steam generating unit operating days when a coal-fired steam generating unit is operated without coal being added to the bunkers, the coal analysis from the previous "as bunkered" coal sample shall be used until coal is bunkered again. For steam generating unit operating days when an oil-fired steam generating unit is operated without oil being added to the oil "day" tank, the oil analysis from the previous day shall be used until the "day" tank is filled again.

Alternative definitions of fuel lot size may be used, subject to prior approval of the Administrator.

5.3.2.3 Use ASTM procedures specified in Section 5.2.1 or 5.2.2 to determine the sulfur contents (%S) and gross calorific values (GCV).

5.4 Daily Geometric Average Percent Reduction from Hourly Values. The geometric average percent reduction ($R_{\rm ga}$) is computed using the following equation:

$$R_{ga} = 100[1-EXP[\frac{1}{n}\sum_{j=1}^{n}ln(\frac{E_{jo}}{E_{ji}})]$$
 Eq. 19-24a

where:

 R_{α} = daily geometric average percent reduction.

 E_{io} , E_{ji} = matched pair hourly arithmetic average pollutant rate,

outlet and inlet, respectively, ng/J (lb/million Btu) or ppm

corrected to 7 percent 02.

n = total number of hourly averages for which paired inlet and

outlet pollutant rates are available within the 24-hr

midnight to midnight daily period.

ln = natural log of indicated value.

EXP = the natural logarithmic base (2.718) raised to the value

enclosed by brackets.

Note: The calculation includes only paired data sets (hourly average) for the inlet and outlet pollutant measurements.

6. Sulfur Retention Credit for Compliance Fuel

If fuel sampling and analysis procedures in Section 5.2.1 are being used to determine average SO_2 emission rates (E_{as}) to the atmosphere from a

coal-fired steam generating unit when there is no SO_2 control device, the following equation may be used to adjust the emission rate for sulfur retention credits (no credits are allowed for oil-fired systems) (Edi) for each sampling period using the following equation:

$$E_{di} = 0.97 K \frac{%S}{GCV}$$
 Eq. 19-25

where:

 E_{di} = average inlet SO_2 rate for each sampling period d, ng/J

(lb/million Btu)

%S = sulfur content of as-fired fuel lot, dry basis, weight percent.

GCV = gross calorific value of the fuel lot consistent with the sulfur

analysis, kJ/kg (Btu/lb).

 $K = 2*10^7 [(kg)(ng)/(%)(J)] {2*10^4 (lb)(Btu/(%))(million Btu)}$

After calculating E_{di} use the procedures in Section 4-2 to determine the average SO_2 emission rate to the atmosphere for the performance test period (E_{ao}) .

- 7. Determination of Compliance When Minimum Data Requirement Is Not Met
- 7.1 Adjusted Emission Rates and Control Device Removal Efficiency. When the minimum data requirement is not met, the Administrator may use the following adjusted emission rates or control device removal efficiencies to determine compliance with the applicable standards.
- 7.1.1 Emission Rate. Compliance with the emission rate standard may be determined by using the lower confidence limit of the emission rate (E_{ao}^*) as follows:

$$E_{ao} * = E_{ao} - t_{0.95} S_o$$
 Eq. 19-26

where:

 S_{\circ} = standard deviation of the hourly average emission rates for each performance test period, ng/J (lb/million Btu).

 $t_{0.95}$ = values shown in Table 19-2 for the indicated number of data points

7.1.2 Control Device Removal Efficiency. Compliance with the overall emission reduction (R_0) may be determined by using the lower confidence limit of the emission rate (E_{ao} *) and the upper confidence limit of the inlet pollutant rate (E_{ai} *) in calculating the control device removal efficiency (R_0) as follows:

$$R_g = 100[1.0 - \frac{E_{ao}*}{E_{ai}*}]$$
 Eq. 19-27

$$E_{ai} * = E_{ai} + t_{0.95}S_{i}$$
 Eq. 19-28

where:

 S_i = standard deviation of the hourly average inlet pollutant rates for each performance test period, ng/J (lb/million Btu).

Table 19-2-Values for $t_{0.95}$

n ¹	 t _{0.95}	n ¹	t _{0.95}	 n¹ 	t _{0.95}	
2 3 4 5 6 7	6.31 2.42 2.35 2.13 2.02 1.94	8 9 10 11 12-16 17-21	1.89 1.86 1.83 1.81 1.77 1.73	22-26 27-31 32-51 59-91 92-151 152 or more	1.71 1.70 1.68 1.67 1.66 1.65	

 1 The values of this table are corrected for n-1 degrees of freedom. Use n equal to the number (H) of hourly average data points.

7.2 Standard Deviation of Hourly Average Pollutant Rates. Compute the standard deviation ($\rm S_e$) of the hourly average pollutant rates using the following equation:

$$S_{e} = \sqrt{\frac{1}{H} - \frac{1}{H_{r}}} \sqrt{\frac{\sum_{j=1}^{H} (E_{hj} - E_{a})^{2}}{(H-1)}}$$
 Eq. 19-29

where:

S = standard deviation of the hourly average pollutant rates for each
 performance test period, ng/J (lb/million Btu).

 ${\rm H_r}$ = total numbers of hours in the performance test period (e.g., 720 hours for 30-day performance test period).

Equation 19-29 may be used to compute the standard deviation for both the outlet (S_{\circ}) and, if applicable, inlet (S_{i}) pollutant rates.