

EPA-450/3-76-018-e

April 1977

**DETERMINING
INPUT VARIABLES
FOR CALCULATION
OF IMPACT OF NEW
SOURCE PERFORMANCE
STANDARDS: WORKSHEETS
FOR METALLURGICAL
INDUSTRIES**

U.S. ENVIRONMENTAL PROTECTION AGENCY
Office of Air and Waste Management
Office of Air Quality Planning and Standards
Research Triangle Park, North Carolina 27711

**DETERMINING INPUT VARIABLES
FOR CALCULATION OF IMPACT OF NEW
SOURCE PERFORMANCE STANDARDS:
WORKSHEETS FOR METALLURGICAL
INDUSTRIES**

by

**The Research Corporation of New England
129 Silas Deane Highway
Weathersfield, Connecticut 06109**

Contract No. 68-02-1382

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Prepared for

**U.S. ENVIRONMENTAL PROTECTION AGENCY
Office of Air and Waste Management
Office of Air Quality Planning and Standards
Research Triangle Park, North Carolina 27711**

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Publication No. EPA-450/3-76-018-e

FOREWORD

During 1974, two studies were initiated that ultimately resulted in the establishment of priorities for developing and promulgating New Source Performance Standards (NSPS). The procedures used to determine these priorities produced a great deal of information that is believed to be useful in the industries involved and, accordingly, is being published in this series of reports (EPA-450/3-76-017, EPA-450/3-76-018, EPA-450/3-76-019, and EPA-450/3-76-020). This information is organized as follows:

EPA-450/3-76-017 discusses (1) the mathematical model (Model IV) used to determine NSPS impacts over a 10-year period; (2) the methods used to attain input variables; and (3) the summary tables which are the heart of this study. Included in the summary tables are data related to (1) emission, growth, and replacement rates; (2) present and future production and capacity; (3) nationwide emissions; and (4) NSPS impact. These tables include information on 13 pollutants and nearly 200 stationary source categories.

EPA-450/3-3-76-018-a, -b, -c, -d, -e, and -f are the calculation sheets, showing how the input variables reported in EPA-450/3-76-017 were derived. All information sources, assumptions, and calculations are documented and explained. The appropriate worksheets are arranged alphabetically in the following volumes:

018-a - Stationary Combustion Sources

018-b - Chemical Processing Industries

018-c - Food and Agricultural Industries

018-d - Mineral Products Industries

018-e - Metallurgical Industries

018-f - Miscellaneous Sources (Evaporation Losses, Petroleum

Industry, Wood Products Industry, and Assembly Plants

The 018-a -f series is of interest only to those concerned with the detailed calculations used to determine the Model IV input variables.

EPA-450/3-76-019-a provides additional results and information produced during the priority study. Its major purpose is to describe the computer program used to rank all the Model IV input and output variables by pollutant (these rankings are reported in 019-b and -c). In addition, it contains (1) summaries of the control systems considered "best" for each source, (2) equipment retirement ages, and (3) emission trends for each source category.

EPA-450/3-76-019-b and c present the computer-generated ranked data for each pollutant. Ranking is from highest to lowest for each of the 21 variables, e.g., A (nationwide capacity) and E_u (uncontrolled emission rate). Volume 019-b contains ranked data for particulate, nitrogen oxide (NO_x), and sulfur oxide (SO_x) sources. In Volume 019-c, the remaining pollutant sources are ranked: hydrocarbons, carbon monoxide (CO), fluorides, hazardous material, acid mist, lead, ammonia, sulfides, chlorine, and trace metals.

EPA-450/3-76-020, the final document in this series, takes the objective impact values from EPA-450/3-76-017, adds subjective judgements, and uses these combined criteria to produce a priority list for NSPS development. The report then calculates nationwide emission trends over the next 15 years for each criteria pollutant (particulate, SO_x , NO_x , hydrocarbons, and CO) based on a series of scenarios (e.g., no NSPS, 20 NSPS per year, etc.)

In summary, documents EPA-450/3-76-017 and 020 present the results of this study. Each stands alone, but they also complement each other, with 020 building on the results of 017. The remaining documents (018-a -f and 019-a -c) present additional and/or more detailed information derived from the impact and priority studies.

**WORKSHEET
FOR
METALLURGICAL INDUSTRIES**

Project Number - 32391 New Source Performance Standards

Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 9-6-74

Source Primary Aluminum Smelters

electrolytic reduction of alumina (from bauxite)
electrolyte is Sodium Aluminum Fluoride : (cryolite)

largest source of pollutants " the potline Ref O35 p 4

K: Table 2.7 pg 2-25 in Ref O35 gives production-capacity data from which utilization factors may be obtained. The average K is evaluated for the period 1948 to 1969.

$$\frac{\sum K}{\text{No yrs}} = K_{\text{avg}} = \frac{20.14}{22} = .915$$

We will use $K = .92$ as representative of the normal utilization factor for the primary aluminum industry smelters.

$$\therefore K = .92$$

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Computation Sheet For Industrial Factors

Calculations Done By W. Marone Date 9-6-74

Source Primary Aluminum Smelters

P_c : Ref 035 pg 2-2

1965 - 1968 capacity increase average 8.4% /yr
1968 - 1969 " " " 5.5% /yr

Weighting the growth $\frac{3 \times 8.4 + 5.5}{4} = 7.7\% /yr$

As a conservative approximation will use
 $P_c = .075$ compound as applicable for 1975 - 1985

$$P_c = .075 \text{ compound}$$

A: Using the 1969 Capacity value of 3,888,000 TONS from Table 2.7 of Ref 035 and the growth of 7.5% ($P_c = .075$) we estimate the 1975 Prod Capacity $n = 6 \text{ yrs}$

$$A = 3,888,000 (1 + .075)^6$$

$$\therefore A = 6.00 \times 10^6 \text{ TON/yr Aluminum}$$

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 9-6-74
 Source Primary Aluminum

P_B :

Ref 037 pg 34 Sec 33.2

Asset guideline life 14 yrs

$$P_B \text{ est} = 2 \times \text{IRS} = 28 \text{ yrs}$$

100% Depreciation in 28 yrs

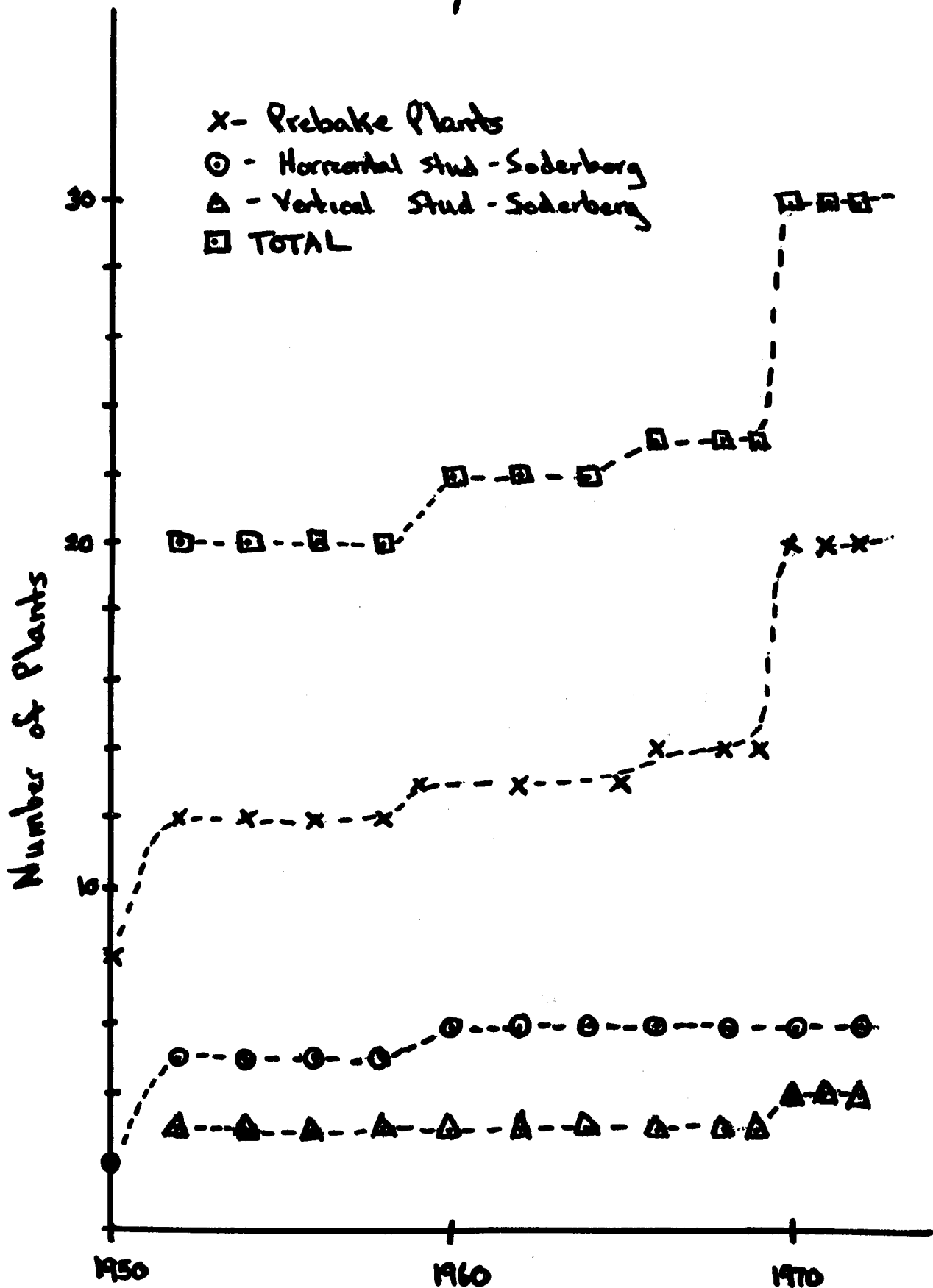
$$\frac{100\%}{28 \text{ yr}} = 3.6\%/\text{yr}$$

We will use 3.5% as an estimate of the
 obsolescence rate.

$$\therefore P_B = .035 \text{ simple}$$

Primary Aluminum

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 9-6-74
 Source Primary Aluminum Smelting

E_u : Particulates

Ref 035 pg 4

Uncontrolled emission factor representative of all cell types (Prebaked & Soderberg)

$$E = 46 \text{ lb / 1000 lb Al produced}$$

$$\therefore E_u = 2 \times 46 = 92 \text{ lb/ton Al}$$

This particulate emission is stated to be composed of a solid fluoride portion

$$9 \text{ lb Solid F / 1000 lb Al} \rightarrow 18 \text{ lb/ton Al}$$

$$\text{Solid Fluoride Content} = \frac{18}{92} = .195 \text{ or } 19.5\%$$

We choose on uncontrolled emission factor to be based on total solids (including Solid F)

$$\therefore E_u = 92 \text{ lb/ton Al}$$

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Computation Sheet For Emission Factors

Calculations Done By W.J. Marzzone Date 9-6-74

Source Primary Aluminum Smelting

Particulate

E_N : Refer to Ref 085 Table 5.2a pg 5-14

highest reported Pt removal eff - 98%
 which includes solid fluoride

We use this for E_N determination
 of particulate emission

$$E_N = .02 \times E_u = .02 \times 92 \frac{\text{lb}}{\text{ton}} \\ = 1.8 \text{ lb/ton}$$

FLUID BED DRY SCRUBBER
 COATED FILTER DRY SCRUBBER
 INJECTED ALUMINA DRY SCRUBBER

$\therefore E_N = 1.8 \text{ lb/ton aluminum}$

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Computation Sheet For Emission Factors

Calculations Done By Z.W. Marrone Date 9-6-74
Source Primary Aluminum Smelting

Hydrocarbons :

Ref 035 pg 4

Hydrocarbon emissions are attributable to Soderberg cells as opposed to the pre-baked potline cells.

pg 3-1b of the above source indicate that the larger number of installations are of the pre-backed version, and moreover, expansion and construction in the Soderberg cells has not been significant nor do we expect this trend to change. Operating problems and control requirements encourage the use of pre-baked potline.

We conclude that the emissions from this industry would not come under NSPS

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 9-6-74
 Source Primary Aluminum Smelter

Sulfur Oxides

E_u : Ref 035 pg 4 and pg 4-8

$$E_u = 30 \text{ lb / 1000 lb Al}$$

or
 $2 \times 30 = 60 \text{ lb / TON Al}$

∴ $E_u = 60 \text{ lb / TON Al}$

Sulfur in carbon anodes is responsible for SO_x emissions
 Petroleum coke of 2.5 - 5.0 % S content is basis of
 carbon anode production.

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Computation Sheet For Emission Factors

Calculations Done By W. Morrone Date 9-6-74

Source Primary Aluminum Smelter

E_N : Sulfur Oxides

Ref 035 pg 10-17 states that "removal of SO_2 from aluminum plants present a difficult problem, because of low concentration in the gas stream."

For the sake of this study until SO_x control becomes evident, we assume $E_N = 0$ which represents maximum impact $(T_S - T_N)$

$$\therefore E_N = 0$$

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Computation Sheet For Emission Factors

Calculations Done By W.J. Monahan Date 9-6-74

Source Primary Aluminum Smelter

E_u :

Fluorides

Ref 035 pg 4

Uncontrolled Emissions

Solid F

9 lb/1000 lb Al

Gaseous F (as HF)

14 lb/1000 lb Al

Total F

23 lb/1000 lb Al

or 46 lb/ton Al

The uncontrolled emission will be considered to represent the total F emission of which

61% is gaseous

39% is solids

Note also that the solid F was included in the particulate emission and represented 19.5% of it.

$$\therefore E_u = 46 \text{ lb/ton Al}$$

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Computation Sheet For Emission Factors

Calculations Done By 23. Maxxone Date 9-6-74
Source Primary Aluminum Smelter

E_N : Fluorides

Ref 160 pg 1-5

NSPS for new sources $2 \text{ lb/TON}_{\text{Al}}$ (total F)

$$\therefore E_N = 2 \text{ lb/TON}_{\text{Al}} \text{ (Total F)}$$

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Computation Sheet For Emission Factors

Calculations Done By V.F. Russo Date 10/18/74

Source Primary Aluminum Smelters

Particulates:

Ref 035, pg 2-2 states that

2 lbs of Bauxite yields 1 lb of Alumina (Al_2O_3)

Using 100% efficiency of Electrolytic Cells

1 lb of Al_2O_3 yields $\frac{3(27)}{102}$ lbs of Al

or

2 lbs of Bauxite yields 0.52 lbs of Al.

1 lb of Bauxite yields 0.26 lbs of Al

Ref. 035 pg 2-22 gives plant locations and capacity for 1970. We will find average (output) capacity for plants in state, get (average) process weight, multi. this output process weight by 3.84 to get it in terms of input process weight and then get the allowable emissions for this input process weight.

i.e. 3.84 lbs Bauxite yields 1 lbs. of Al.

Ref. 093 pg 250 lists actual figure as 3.82 lbs Bauxite \Rightarrow 1 lbs Al. This procedure yields an E_S which assumes that the 1975 production capacity is distributed in the stated states as the 1970 capacity with the same relative capacities. Also, this procedure assumes the size of new plants is the same in 1975 as the average plants capacity in 1970.

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Computation Sheet For Emission Factors

Calculations Done By V.F. Russo Date 10/18/74
 Source Primary Aluminum Smelters

<u>State</u>	<u>N_P</u>	<u>Output 1970 Cap. (10³T/YR)</u>	<u>Input Process Wgt (lbs/HR)*</u>	<u>Plant Allowable Emissions (lbs/HR)**</u>
✓ Tenn.	2	340	149,000	34
✓ N.C.	1	100	88,000	43.4
✓ Ind.	1	225	197,250	46
✓ N.Y.	2	253	110,900	51
✓ Texas	3	561	163,950	90
✓ Mont.	1	175	153,400	47
✓ Ky.	1	90	78,890	41.5
✓ Oreg.	2	228	99,950	44.5
✓ Ia.	1	260	237,950	51
W. Va.	1	163	142,900	48
✓ Ohio	1	240	210,400	50
✓ Ark.	2	185	81,100	31
✓ Md.	1	85	74,500	41.9 ***
✓ Ala.	1	221	193,750	33 ++
Wash.	6	1016	148,500	15 lbs/Ton AL *+

* For average plant capacity by

$$\frac{\text{Output Cap}(\text{lbs/HR}) \times 3.84}{N_P \quad 8760(\text{HRS/YR})}$$

** Ref. 084 & 148

++ For Class I counties

*+ State has specific req. for Al Smelters.

*** For Eastern, Southern, Central, Cumberland - Keyser Regions

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Computation Sheet For Emission Factors

Calculations Done By U.F. Russo Date 10/18/74
 Source Primary Alum. Smelters

Thus

<u>State</u>	<u>A_i</u>	$E_s^o \left(\frac{\text{lbs part.}}{\text{TON AL}} \right)^*$	$A_i E_s^o \left(\frac{\text{lbs part.}}{\text{TON AL}} \right)$
Tenn.	.08	1.752	.140
N.C.	.02	3.802	.076
Ind.	.05	1.791	.090
N.Y.	.06	3.532	.212
Texas	.14	4.216	.590
Mont.	.04	2.353	.094
Ky	.02	4.040	.080
Oreg.	.06	3.419	.205
La.	.06	1.718	.103
W.Va.	.04	2.580	.103
Ohio	.06	1.825	.109
Ark	.04	2.936	.117
Md.	.02	4.318	.086
Ala.	.05	1.308	.065
Wash	.25	15	3.750

*Calculated using $\frac{N_p^i \left(\text{Allow. Emiss } \frac{\text{lbs}}{\text{HR}} \right) 8760 \left(\frac{\text{HR}}{\text{YR}} \right)}{\text{Output Prod. Cap. } \left(\frac{\text{T}}{\text{YR}} \right)}$

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Computation Sheet For Emission Factors

Calculations Done By U.F. Russo Date 10/18/74
Source Primary Alum. Smelters

$$E_s = \sum A_i E_s^o$$

$$E_s = 5.820 \quad \text{lbs Part./Ton Al.}$$

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Computation Sheet For Emission Factors

Calculations Done By V.F. Russo Date 10/21/74
 Source Primary Alum. Smelters

SO₂ Emissions

Only two states have SO₂ regulation applicable for primary aluminum smelters (Ref. 084 and 148)

$$\text{Ind: } E_p (\text{# SO}_2/\text{hr}) = 19.5 P (\text{Tons/HR})^{0.67}$$

Max. emission concentration 2000 ppm

For Indiana, as previously shown,

$$\text{Input Process Wgt (lbs/hr)} = 197,250 \text{ or}$$

$$\text{Process Wgt} = 98.625 \frac{\text{Tons}}{\text{HR}}$$

Thus, the allowable SO₂ emissions are

$$E_p (\text{# SO}_2/\text{hr}) = 19.5 (98.625)^{0.67} \\ \approx 423$$

$$\text{For Ind. } E_p = 423 \frac{\text{# SO}_2}{\text{hr}}$$

or

$$E_s^o = \frac{423 \frac{\text{# SO}_2}{\text{hr}} (8760 \frac{\text{HR}}{\text{YR}})}{225 \times 10^3 \frac{\text{Tons AL}}{\text{YR}}}$$

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Computation Sheet For Emission Factors

Calculations Done By V. F. Russo Date 10/21/74
Source Primary Alum. Smelters

$$E_s^o = 16.5 \frac{\text{lbs SO}_2}{\text{TON AL}}$$

For ha, neglecting Bake cells and considering only
potline effluents.

~~12.5 per cent of total effluents (gums)~~

J Am. Air Poll. Control. Assoc. (Fe. 1973) 23, #2 pg 101
gives

SO₂ emissions as 90 - 300 mg/m³

Taking average of 195 mg/m³

The potline effluents (in ppm) becomes (assuming emission at
standard conditions)

$$10^6 \left(\frac{\text{Moles SO}_2}{\text{Moles Gas}} \right) = \frac{195 \text{ gms}}{\frac{64 \text{ gms/mole} \times 10^6}{\frac{1000 \text{ liters}}{22.4 \text{ liters/mole}}} = 68.25 \text{ ppm}$$

Thus, emissions are below standard.

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Computation Sheet For Emission Factors

Calculations Done By V.F. Russo Date 10/21/74
Source Primary Alum. Smelting

$$\text{or } E_s = (\text{Emission for Ind.}) A_i + (1.00 - A_i) E_u$$

$$\therefore E_s = 16.5 (.090) + (0.91) 60$$

$$E_s = 1.485 + 54.6$$

$$E_s = 56.09$$

165 SO₂
Ton AL.

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Computation Sheet For Emission Factors

Calculations Done By 2d Marrone Date 10-31-74
Source Primary Aluminum Smelter

E_{III}: Fluorides

Ref 160 pg 1-12

Recommended Standards (for State) of Existing
Primary Aluminum Plants

<u>Cell Type</u>	<u>Standard lb/ton Al</u>
Vertical Stud Soderberg	3
Horizontal Stud Soderberg	3
Centerwork Prebaked	4
Sidework Prebaked	2
	<hr/>
Aug	3 lb/ton Al

We assume this average typical off emission limitations to be set on all existing primary aluminum plants.

$$\therefore E_{III} = 3 \text{ lb/ton Al}$$

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Computation Sheet For Emission Factors

Calculations Done By J. Marrone Date 2-7-75

Source Primary Aluminum Smelter

E_S : Fluoride

Of the states having aluminum production capacity only Oregon and Washington specify fluoride regulations applicable to Aluminum (Primary) Production. Montana's Fluoride regulation excludes primary aluminum plants. Washington gives ambient and forage quality standards and doesn't specify emission levels. Oregon gives emission rates for new and-existing plants in terms of total fluoride.

Since fluoride emissions from this industry are in the form of both gaseous and solid constituents then, at least a part of it (solid) will be "de facto" controlled by particulate regulation in those states with no fluoride regulation.

In developing a synthesized value of E_S for this pollutant we make the following assumptions

① Oregon Reg for New : 4.0 lb F / TOW alum
 Existing : 2.5 lb F / TOW alum

② Washington ambient air quality will be obtained by average controlled plant which is assumed to be at 85% level on Total Fluoride

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Computation Sheet For Emission Factors

Calculations Done By D. Marrone Date 2-7-75
 Source Primary Aluminum Smelter

E_g : Fluoride
 cont'd

- ② Solid Fluoride controlled in remaining states by PWR curve (19.5% of Pt E_u is solid Fluoride)
- ④ Gaseous Fluoride is uncontrolled in all states except Oregon and Wash.
- ⑤ E_g will be calculated by weighting allowable emissions by capacity occurrence.
- ⑥ New and Existing E_g will be determined to reflect Oregon's new and existing regulation

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-7-75

Source Primary Aluminum Smelting

<u>State</u>	<u>A_i</u>	<u>k Allowable Emission lb/ton Al₂O₃</u>	<u>Total</u>	<u>A_i x TOTAL</u>	<u>lb/ton Al₂O₃</u>
		<u>Solid</u>	<u>Gaseous</u>	<u>Exist</u>	<u>New</u>
Tenn	.08	.34	28	28.34	2.267
NC	.02	.74	28	28.74	.575
Ind	.05	.35	28	28.35	1.418
NY	.06	.69	28	28.69	1.721
Texas	.14	.82	28	28.82	4.035
Mont	.04	.46	28	28.46	1.138
Ky	.02	.79	28	28.79	.576
Oreg	.06	—	—	10 New, 2.5 Exists	.150
La	.06	.34	28	28.34	1.700
W Va	.04	.50	28	28.50	1.140
Ohio	.06	.36	28	28.36	1.702
Ark	.04	.57	28	28.57	1.143
Md	.02	.84	28	28.84	.577
Ala	.05	.26	28	28.26	1.413
Wash.	.25	—	—	6.9	1.725

$$E_s' = \sum = \frac{19.013}{18.923}$$

Diff. $\approx .09$ lb/ton

Since E_s (exist < new) differ by only .09 lb/ton we will develop only one E_s factor so that $E_s \approx 19$ lb/ton

$$E_s (\text{Fluoride}) = 19 \text{ lb/ton Al produced}$$

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Computation Sheet For Industrial Factors

Calculations Done By W. Marinone Date 9-4-74
 Source Beehive Coke Ovens

Ref 050 "By Product Coke Ovens pg 1

In 1971 the production capacity for beehive coke ovens was about 500,000 TON/yr and it was indicated that no new beehive plants were anticipated to be built in the near future.. This implies that the 1975 Prod Cap. would be equivalent to the 1971 value and that the growth in the industry is nil

A:

$$\therefore A = .5 \times 10^6 \text{ TON/yr coke}$$

P_c:

$$\therefore P_c = 0$$

K:

Assume capacity utilization factor for beehive ovens to be the same as for byproduct ovens $\therefore K=.93$

$$K = .93$$

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Computation Sheet For Industrial Factors

Calculations Done By A. Marrone Date 9-4-74

Source Beehive Coke Ovens

P_B :

The growth in this industry, as indicated above, is not expected to be positive. It may very well have a negative growth rate.

This oven has had a decreasing share of the production and Ref 097 pg 87 identifies this source as an obsolete process. which would be best controlled by replacement with by-product coke plants.

We are assuming that; not only are no new coke plants being built ($P_c=0$) but that existing capacity will not be replaced on obsolescence by other beehive ovens.

This implies that $P_B = 0$

$$\therefore P_B = 0$$

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Computation Sheet For Emission Factors

Calculations Done By W. Monnone Date 9-4-74

Source Beehive Coke Ovens

The coke produced from the coal charged for beehive oven is estimated from the values appear in Ref 04A Table 9-2 pg 120

$$\frac{775,000 \text{ ton coke}}{1,300,000 \text{ ton coal}} \rightarrow \frac{775,000}{1,300,000} = .596 \frac{\text{Ton coke}}{\text{Ton coal}}$$

We obtain uncontrolled emission factors from Ref 075

Table 7.2-1
pg 7.2-2

Particulate

$$E_u = 200 \frac{\text{lb}}{\text{Ton coal}} \times \frac{\text{Ton coal}}{.596 \text{ Ton coke}} \cong 336 \frac{\text{lb}}{\text{Ton coke}}$$

$$\therefore E_u = 336 \text{ lb/Ton coke}$$

Carbon Monoxide

$$E_u = 1 \frac{\text{lb}}{\text{Ton coal}} \times \frac{\text{Ton coal}}{.596 \text{ Ton coke}} \cong 1.7 \text{ lb/Ton coke}$$

$$\therefore E_u = 1.7 \text{ lb/Ton coke}$$

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Computation Sheet For Emission Factors

Calculations Done By A. Marrone Date 9-4-74
 Source Beehive Coke Ovens

Hydrocarbons

$$E_u = 8 \frac{\text{lb}}{\text{TON coal}} \times \frac{\text{coal}}{.5\% \text{ coke}} \approx 13.4 \frac{\text{lb}}{\text{TON coke}}$$

$$\therefore E_u = 13.4 \frac{\text{lb}}{\text{TON COKE}}$$

Ammonia

$$E_u = 2 \frac{\text{lb}}{\text{TON coal}} \times \frac{\text{coal}}{.5\% \text{ coke}} \approx 3.4 \frac{\text{lb}}{\text{TON coke}}$$

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Emission Factors

Calculations Done By W. Mannone Date 5-9-75
 Source Beehive Coke Ovens

Ref 274 pg 32 describes hydrogen sulfide emissions from byproduct coke ovens however, in light of the lack of other data we will approximate that the production of H_2S for beehive oven will be somewhat equivalent

$$\text{Emission} = 6.7 \text{ lb/TON coal charged}$$

$$6.7 \frac{\text{lb}}{\text{TON Coal}} \times \frac{\text{Ton coal}}{.596 \text{ Ton coke}} \approx 11.2 \text{ lb/TON coke}$$

$$E_u(\text{sulfide}) = 11.2 \text{ lb/TON coke}$$

Since this source has values of $P_C = 0$ and $P_B = 0$ it is probable that NSPS would not be set for this industry.

Section 111d may still be imposed on the non-criteria pollutants, therefore we will develop the factors E_S , E_N and E_{111d} for ammonia and sulfide.

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Computation Sheet For Emission Factors

Calculations Done By W. Marzzone Date 5-9-75

Source Beehive Coke Ovens

E_S : Ref 84 & 148

There are no specific regulations for ammonia and sulfides so that we may assume the relationship $E_S = E_N$

$$E_S(\text{ammonia}) = 3.4 \text{ lb/ton coke}$$

$$E_S(\text{sulfide}) = 11.2 \text{ lb/ton coke}$$

As discussed in Ref 097 pg 87 there is no reasonable control method for beehive coke ovens.

"The only practical control method is to replace the ovens with well-designed slot-type ovens with by-product gas recovery systems."

We were unable to obtain specific controls for this source and suspect that little may be done in tune with the economics of this industry.

For max impact assessment we will assume $E_N = 0$ and $E_{\text{III}d} = 0$ for both ammonia and sulfides.

$$E_N(\text{sulfide}) = 0$$

$$E_{\text{III}d}(\text{sulfide}) = 0$$

$$E_N(\text{ammonia}) = 0$$

$$E_{\text{III}d}(\text{ammonia}) = 0$$

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Computation Sheet For Industrial Factors

Calculations Done By 2). Marzzone Date 9-4-74
 Source By Product Coke Ovens

K: Using the data obtained from Ref 050 "By Product Coke ovens" pg 2 and illustrated on the attached graph we are able to calculate utilization factor estimates for the years 1970 through 1977

$$K_{70} = \frac{67.4}{72.2} = .934$$

$$K_{74} = \frac{69.2}{75.0} = .923$$

$$K_{71} = \frac{67.9}{72.2} = .940$$

$$K_{75} = \frac{69.5}{75.5} = .920$$

$$K_{72} = \frac{68.4}{74.2} = .922$$

$$K_{76} = \frac{69.6}{75.5} = .922$$

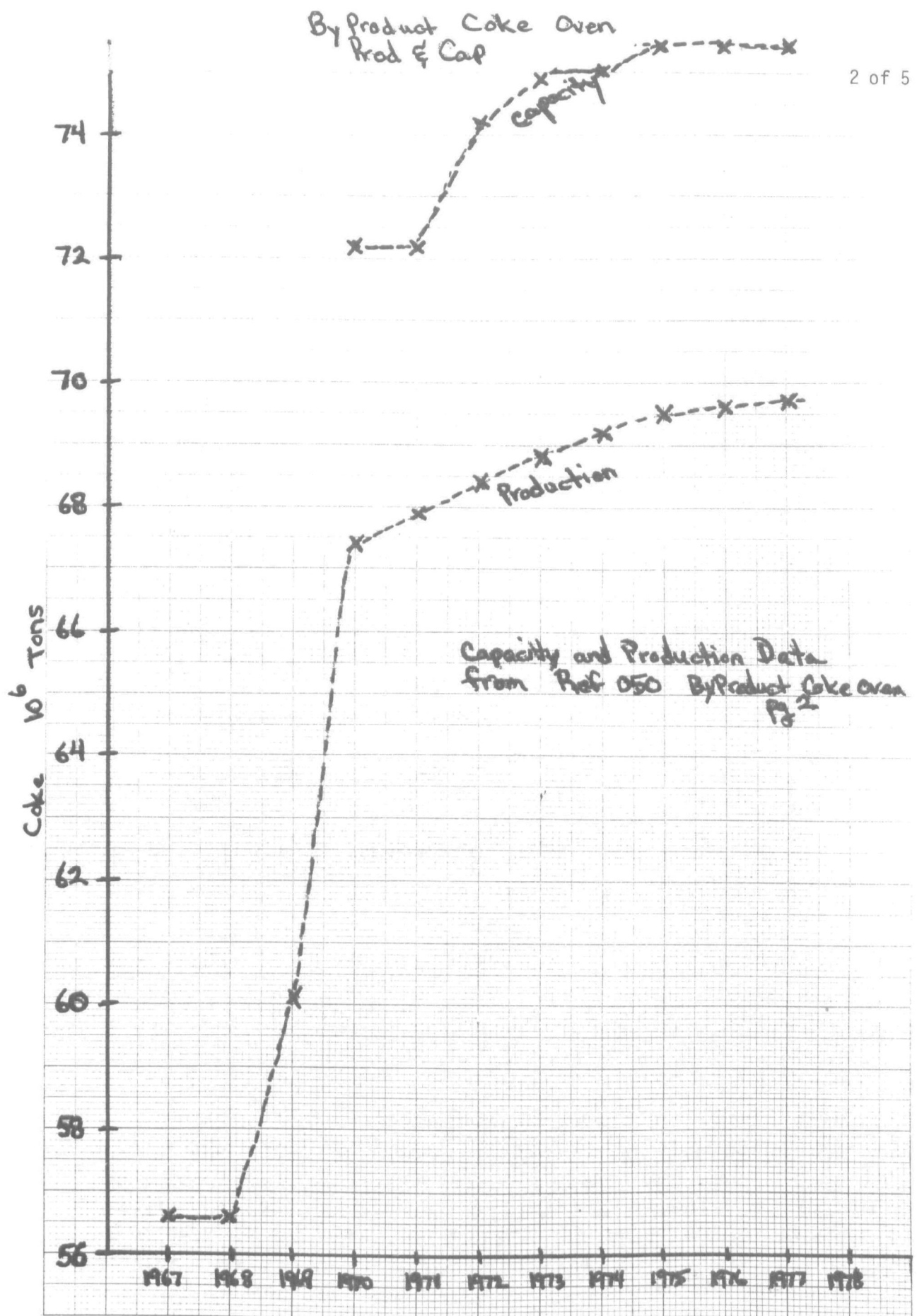
$$K_{73} = \frac{68.8}{74.9} = .919$$

$$K_{77} = \frac{69.7}{75.5} = .923$$

$$K_{\text{Avg}} = \frac{7.475}{8} = .934$$

We will use $K = .93$ and assume it representative of byproduct coke ovens and applicable for 1975-1985

∴ $K = .93$



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Computation Sheet For Industrial Factors

Calculations Done By 2d) Marrone Date 9-4-74
 Source By-Product Coke Oven

A: The 1975 Production capacity is estimate in Ref 050 "Byproduct coke oven pg 2" to be 75.5×10^6 TON/yr

$$\therefore A = 75.5 \times 10^6 \text{ TON/yr}$$

According to Prod. Capacity figures in 1971 for beehive and byproduct coke ovens from the above source;

$$\frac{72.2}{72.2 + .5} \approx .993$$

over 99% of coke production capacity is attributable to byproduct coke ovens.

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Computation Sheet For Industrial Factors

Calculations Done By W. Marone Date 9-4-74

Source By product coke oven

P_C :

The growth of this industry will be directly guided by trends and practices in the steel making plants which consume roughly 90% of the ~~coke~~ coke production.

Looking at the capacity data from Ref 050 pg 2 which is plotted on the attached graph we see that there is a definite leveling off of the production capacity. The estimated level is identical ($75.5 \times 10^6 \text{ T/yr}$) for the period 1975-1977

This implies a negligible growth indication for this industry. True growth predictions for coke were not obtained in the literature.

The simple growth rate from 1970 to 1975 obtained from the data in Ref 050 12 was found to be;

$$\frac{75.5 - 72.2}{75.5} \times 100 \approx 4.37\% \text{ or } .87\%/\text{yr}$$

We make the assumption that a simple growth rate of 1% is representative for the by product coke ovens especially in light of little growth 1975-1977. Steel consumption which should guide coke demand is estimated by Ref 116 p13 is predicted to grow a 3%/yr. It is uncertain whether or not this value includes imports. In all it appears that the 1% growth is adequate.

$$\therefore P_C = .01 \text{ simple)$$

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Computation Sheet For Industrial Factors

Calculations Done By 22 Marrone Date 9-4-74

Source By Product Coke Ovens

P_B :

Ref 031 pg 34 Section 33.0

Asset guideline period 18 yrs

P_B est $\rightarrow 2 \times IRS = 36$ yrs

$\frac{100\%}{36\text{yr}} \cong 2.8\%/\text{yr}$ simple 100% Depreciation in 36 years.

$$\therefore P_B = .028 \text{ simple}$$

Plant locations in Ref 050

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 9-4-74

Source By Product Coke Ovens

The coke produced from the coal feed is estimated from data in Ref 049 Table 9-2 pg 120

$$\frac{63,700,000 \text{ TON coke}}{90,000,000 \text{ ton coal}} \rightarrow \frac{63.7}{90.0} \approx .708 \frac{\text{Ton coke}}{\text{Ton coal}}$$

E₄:

Ref 075 Table 7.2-1

emissions given in terms of lb/TON coal charged we divide these value by $\approx .71$ to obtain the emission factor in terms of lb/ton coke produced.

lb/ton COKE

<u>Operation</u>	<u>Pt</u>	<u>SO_x</u>	<u>CO</u>	<u>HC</u>	<u>NO_x</u>	<u>NH₃</u>
Unloading	.56	—	—	—	—	—
Charging	2.11	.03	.85	3.52	.04	.03
Coking Cycle	.14	—	.85	2.11	.014	.085
Discharging	.85	—	.01	.28	—	.14
Quenching	1.27	—	—	—	—	—
Underfiring	—	5.63	—	—	—	—
<u>Total</u>	<u>4.93</u>	<u>5.66</u>	<u>1.71</u>	<u>5.91</u>	<u>.054</u>	<u>.255</u>

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 9-4-74
 Source By product Coke Ovens

E_u : cont'd

	E_u lb./ton coke produced
Particulate	4.9
SO _x	5.7
CO	1.7
HC	5.9
NO _x	.054
NH ₃	.26

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 10-3-74

Source By product Coke Oven

E_N:

Ref 050 Byproduct Coke ovens pg 6

"There are no well controlled plants either in the U.S. or elsewhere."

"The major effort in the US at this time is to use good operating and maintenance techniques until a control system has been proved"

Efforts are being investigated to control the three most significant sources of pollutants ;

- ① charging
- ② pushing
- ③ quenching

Alternatives range from improved oven design, to add-on control systems and process modification.

The following are some of the controls proposed or being investigated :

- ① control of coal feed flow during charging to permit exhaust pick up for byproduct system
- ② operating at proper oven temp for a proper coking time may eliminate most emissions during pushing
- ③ hooding of hot coke car during pushing
(Ford has a system which is reported to be working well)
- ④ scrubbers may be applied to larry-car during charging
- ⑤ baffle quench towers may reduce Pt. emissions during the coke quench operation

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 10-3-74
 Source By product Coke Oven

E_N : cont'd

- ⑥ dry quenching (now done in Europe) is a relatively new system which may be one solution.
- ⑦ The by-product recovery plant can remove most of the cyanides, phenols, sulfur, tars, ammonia, and organic compounds.

The impact of this industry will certainly depend on the level of control that may be achieved. A high value for $T_S - T_N$ would be indicative of a sizeable control on this industry.

We choose at this point to treat E_N for this industry in two ways; (1) assume levels of control for applicable operations and pollutants and synthesis values of E_N ; (2) set all pollutants $E_N = 0$ whereby we would be flagging this industry ($\text{Max } T_S - T_N$) as a source of emission with no demonstrated or state-of-the-art control technology. The value of $T_S - T_N$ will determine the relative importance of control developments to this industry.

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Computation Sheet For Emission Factors

Calculations Done By W. Marzzone Date 10-3-74
 Source By Product Coke Oven

E_N cont'd

(1) Application of Controls

Assumptions made:

- ① control of pt emission during unloading is within normal collection technology - baghouse system 99% eff
- ② charging control schemes will ultimately achieve 90% eff for particulates.
- ③ discharging (pushing) controls will ultimately achieve 90% eff. for particulates. HC and NH₃ will be picked up by byproduct recovery with 99% control off.
- ④ Quenching will be controlled to the extent of 90% with respect to Particulates
- ⑤ The emissions emanating from the coking cycle will be handled by the by product recovery plant to the extent of 99% for ~~HC NH₃~~ pollutants. This appears to be within current technology!

Unloading

b/tot coke produced

$$\text{Pt E} = .01 \times .56 \approx .006$$

Charging

$$\text{Pt E} = .1 \times 2.11 \approx .21$$

$$\text{SOx E} = E_u = .03$$

$$\text{CO E} = E_u = .85$$

$$\text{HC E} = .01 \times 3.52 = .035$$

$$\text{NO}_x \quad E = E_u = .04$$

$$\text{NH}_3 \quad E = .01 \times .03 = .0003$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marzzone Date 10-3-74
 Source By product Coke oven

E_N : cont'd

Coking Cycle

$$\text{Pt} \quad E = .01 \times .14 \approx .001$$

$$\text{CO} \quad E = E_u = .85$$

$$\text{HC} \quad E = .01 \times 2.11 = .021$$

$$\text{NO}_x \quad E = E_u = .014$$

$$\text{NH}_3 \quad E = .01 \times .085 = .0009$$

Discharging

$$\text{Pt} \quad E = .1 \times .85 = .085$$

$$\text{CO} \quad E = E_u = .01$$

$$\text{HC} \quad E = .01 \times .28 \approx .003$$

$$\text{NH}_3 \quad E = .01 \times .14 \approx .001$$

Quenching

$$\text{Pt} \quad E = .1 \times 1.27 \approx .13$$

Underfiring

$$\text{SO}_x \quad E = E_u = 5.63$$

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Computation Sheet For Emission Factors

Calculations Done By _____ Date _____
 Source _____

E_N "Application of controls" Cont'd

Summary

<u>Operation</u>	<u>PT</u>	<u>SO_x</u>	<u>CO</u>	<u>HC</u>	<u>NO_x</u>	<u>NH₃</u>
Unloading	.006	—	—	—	—	—
Charging	.21	.03	.85	.035	.04	.0003
Coking Cycle	.001	—	.85	.021	.014	.0009
Discharging	.085	—	.01	.003	—	.001
Quenching	.13	—	—	—	—	—
Underfiring	—	5.63	—	—	—	—
Totals =	.43	5.63	1.7	.06	.054	.002

∴

E_N lb/ton coke

Particulate	.43
SO _x	5.63
CO	1.7
HC	.06
NO _x	.054
NH ₃	.002

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 10-3-74
 Source By Product Coke Ovens

E_N cont'd

(2) Maximum T_s-T_N impact

	<u>E_N lb/TON coke</u>
PT	0
SO _x	0
CO	0
HC	0
NO _x	0
NH ₃	0

6-11-75 WAM

For the purposes of this report the impact value T_s-T_N or E_N will be based on actual values of E_N and E_{NLD} not $E_N = E_{NLD} = 0$.

except that there clearly is nothing demonstrated in the way of NO_x or CO control so we will keep these at maximum impact determination $E_N = 0$

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Computation Sheet For Emission Factors

Calculations Done By W. Marone Date 10-15-74

Source By-Product Coke Ovens

E_u : Hydrogen Sulfide

E_u Ref 153 pg 51

$$E_u = 6.7 \text{ lb H}_2\text{S / TON coal charged}$$

applying .71 $\frac{\text{Ton coke}}{\text{Ton coal}}$ as before

$$E_u = \frac{6.7}{.71} \cong 9.4 \text{ lb / TON coke}$$

E_N :

Cooling & scrubbing have been cited Ref 153 pg 51
 50% control

$$E = S \times E_u = .5 \times 9.4 \text{ lb / TON coke}$$

$$E_N = 4.7 \text{ lb / TON coke}$$

maximum impact $E_N = 0$

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Computation Sheet For Emission Factors

Calculations Done By 2d. Manzone Date 6-11-75

Source By product Coke Oven

E_{11d} : NH_3 & H_2S

E_S

NH_3

It is assumed that recovery plant control on NH_3 may be applied to existing source so that

$$E_{11d} = E_N = .002 \text{ lb/Ton coke}$$

H_2S

Control by cooling and scrubbing of H_2S is assumed feasible for existing plant to the same extent as a new plant.

$$E_{11d} = E_N = 4.7 \text{ lb/Ton coke}$$

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Computation Sheet For Emission Factors

Calculations Done By 2) Marrone Date 10-18-74

Source By product Coke Ovens

E_s : Particulates

Ref 050 "Byproduct coke plants" pg 1

1971 Production Capacity	72.2×10^6 T/hr	
" Plants	69	
" No. of States	20	West, Midwest, North Central, South, East, and Northeast.
" No of Ovens	14,895	

Using this information along with .71 lb coke/lb coal we may calculate the process input weight rate.

We assume 8400 hr/yr of operation ($24\text{ hr} \times 350\frac{1}{2}\text{ yr}$)

$$\frac{72.2 \times 10^6 \text{ Ton coke}}{14,895 \text{ ovens}} \times \frac{2000 \text{ lb coke}}{\text{Ton coke}} \times \frac{\text{lb coal}}{.71 \text{ lb coke}} \times \frac{\text{yr}}{8400 \text{ hr}} \cong 1,625 \frac{\text{lb coal}}{\text{hr}}$$

process weight rate
per oven

Since 20 of the 50 states are represented and the makeup spans the different regions of the country and population/size categories we will use the "50-state Process weight curve to estimate the allowable emissions.

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 10-18-74

Source By product Coke Ovens

E_u E_s : particulate cont'd

Allowable Emission = $3.55 \frac{\text{lb pt}}{\text{hr}}$ per oven

$$E_s = \frac{3.55 \frac{\text{lb pt}}{\text{hr/ton}} \times \frac{8400 \text{ hr}}{\text{yr}}}{\frac{72.2 \times 10^6 \text{ TON COKE}}{\text{yr}}} \approx 6.2 \frac{\text{lb pt}}{\text{TON COKE}}$$

Since the value of $E_s > E_u$ we use the value of E_u in impact calculations $E_s = E_u = 4.9 \frac{\text{lb}}{\text{TON}}$

$$\therefore E_s = 4.9 \frac{\text{lb}}{\text{TON COKE}}$$

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Computation Sheet For Emission Factors

Calculations Done By 2). Marrone Date 10-18-74

Source By product Coke Oven

cont'd.

There are ^{no} state regulations (Ref. 084 & 148) which are applicable to Byproduct Coke ovens for SO_x , CO, HC, NO_x , NH_3 , and H_2S .

We therefore assume $E_s = E_u$

$E_s \text{ } \text{SO}_x$	$=$	5.7 lb /TON COKE
$E_s \text{ CO}$	$=$	1.7 lb /TON COKE
$E_s \text{ HC}$	$=$	5.9 lb /TON COKE
$E_s \text{ NO}_x$	$=$	$.054 \text{ lb /TON COKE}$
$E_s \text{ NH}_3$	$=$	$.26 \text{ lb /TON COKE}$
$E_s \text{ H}_2\text{S}$	$=$	9.4 lb /TON COKE

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 12-2-74

Source Primary Copper

E

Basically, there are three operations at Copper Smelting plants

- (A) Roasting
- (B) Smelting
- (C) Converting

Ref 029 p 3.1-1

In 1973 8 of the 15 primary copper smelters do not perform roasting whereas the remaining seven perform all three process steps.

Roasting occurs in multiple-hearth roasters and the newer fluid-bed roasters. pg 3.1-15 $\frac{4}{3}$ multiple hearth
 $\frac{1}{3}$ fluid bed

All but one copper plant use reverberatory furnaces
 (the other uses an electric furnace)

Pierce-Smith converters are the only type converter currently used domestically.

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Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 12-2-74
 Source Primary Copper

K:

Ref 030 pg 6.1-8

during recession of 1971 the copper smelters operated at 82% capacity compared with 90% during the prosperous period of 1970

We assume a utilization factor of .85 within this range and representative of the copper smelting industry during the period 1975 to 1985

$$\therefore K = .85$$

P_c : Ref 030 p 6.1-14

Use assumption of 3% growth rate for copper smelting

\therefore

$$P_c = .03 \text{ compound}$$

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Computation Sheet For Industrial Factors

Calculations Done By 2d. Marzone Date 12-2-74
 Source Primary Copper

P_B : Ref 037 Class 33.2

Asset Guideline period 14 yrs

P_B est $\rightarrow 2 \times \text{IRS} = 28 \text{ yrs}$

100% depreciation $\approx 3.6\%/\text{yr}$ simple
 28 yrs

∴

$$P_B = .036 \text{ simple}$$

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Computation Sheet For Industrial Factors

Calculations Done By D. Marrone Date 12-2-74
 Source Primary Copper

A : Ref 095 pg 233.4531

$$P_{60} = 1,765,100 \text{ TONS/yr}$$

$$P_{75} = P_{60} (1 + P_c)^5 \quad P_c = .03 \text{ compound}$$

$$P_{75} = 1.7651 \times 10^6 (1.03)^5$$

$$P_{75} \approx 2.05 \times 10^6 \text{ TON/yr}$$

and using $K = .85$ we obtain the production capacity (A) in 1975

$$A = \frac{P_{75}}{K} = \frac{2.05 \times 10^6}{.85} \text{ TON/yr}$$

$$\therefore A = 2.41 \times 10^6 \text{ TON/yr} \quad \text{copper produced primary}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-2-74

Source Primary Copper

Particulate

E_u : (Roasting, Smelting, and Converting)

Ref 075 Table 7.3-1

4 units of concentrated ore are required to produce 1 unit weight of copper metal (Note a) Emission factor is stated to be expressed in terms of "units per unit weight of concentrated ore produced".

This at first appears confusing however referring to Ref 049 Table 13-1 pg 262 we obtain emission factors for the above operations in terms of copper produced

Roasting: 168 lb/TON copper

Smelting: 206 lb/TON copper

Converting: 235 lb/TON copper

by using the ratio of 4:1 we convert the data in Ref 075 to lb/TON of copper which agree well with that from 049 except in the case of the smelting furnace.

Roasting: $4 \times 45 = 180$ lb/TON copper

Smelting: $4 \times 20 = 80$ lb/TON copper

Converting: $4 \times 60 = 240$ lb/TON copper

The data base for Ref 075 includes Ref 049 and is believed to be more reliable in this instance so that we will use the uncontrolled emission factors presented in it (corrected to copper production basis).

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-2-74
 Source Primary Copper

Particulate cont'd

E_u :

With the assumption of about $\frac{7}{15} = 47\%$ of production being roasted, and that the total emission to be the sum of roasting, smelting, and converting we may calculate,

$$E = .47 \left(180 \frac{\text{lb}}{\text{TON}} \right) + 80 \frac{\text{lb}}{\text{TON}} + 240 \frac{\text{lb}}{\text{TON}}$$

$$E = 84.6 + 80 + 240$$

$$\therefore E \approx 405 \frac{\text{lb}}{\text{TON}} \text{ copper produced}$$

$$\therefore E_u = 405 \frac{\text{lb}}{\text{TON}} \text{ copper produced}$$

E_N :

Ref 075 Table 7.3-1 Note c states that 99.7% eff has been reported for reducing particulate emissions

$$\therefore E_N = .003 \times E_u = .003 \times 405$$

$$E_N \approx 1.22 \frac{\text{lb}}{\text{TON}}$$

ELECTROSTATIC PRECIPITATORS

$$\therefore E_N = 1.22 \frac{\text{lb}}{\text{TON}} \text{ copper produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-3-74
 Source Primary Copper

Particulates cont'd

E_s :

Ref 029 Table 5-1 gives location of copper smelting plants and daily copper production capacities

We assimilate this data to obtain a breakdown of fractional capacity (total) in each state

<u>State</u>	<u>Copper Capacity T/D</u>	<u>Ai</u>
Michigan	220	.046
Montana	500	.106
Nevada	185	.039
N. Mexico	234	.050
Tennessee	50	.011
Texas	260	.055
Utah	750	.159
Washington	300	.063
Arizona	2228	.471
		$\Sigma = \underline{1.000} \checkmark$

Typical Plant Size

4727 T/D Total for industry of 15 plants

$$\frac{4727}{15} = 315 \text{ T/D copper produced}$$

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Computation Sheet For Emission Factors

Calculations Done By 2). Marrone Date 12-3-74
 Source Primary Copper

Particulate cont'd

E_s :

Referring to Table 5.1 in O29 and using the total SCFM along with the copper production (T/D) for plants 1 to 5 which appear typical in that they all have roasting, smelting, and converting we calculate SCFM: T/D ratios.

Plant	SCFM / T/D
1	1500
2	1323
3	2769
4	2011
5	1523

$$\text{Avg} = 1825 \frac{\text{SCFM}}{\text{T/D}}$$

and for a typical plant the discharge rate is $315 \frac{\text{ft}^3}{\text{min}} \times 1825 \frac{\text{SCFM}}{\text{T/D}}$

$$\text{SCFM} = 574,875$$

Note

Proposed Standards in Ref O29 pg 11-1 and 1.1-2 specifies only opacity limits and not particulate emissions.

The process weight rate is assumed to be $315 \frac{\text{ft}^3}{\text{min}} \times 4 \frac{\text{ton cu ft}}{\text{ton cu}}$

$$1260 \frac{\text{Ton input}}{\text{Day}} \times \frac{\text{day}}{24 \text{hr}} \times \frac{2000 \text{ lb}}{\text{Ton}}$$

$$\text{Process wt} = 105,000 \text{ lb/hr}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-3-74
 Source Primary Copper

Particulate cont'd

E_s :

based on a process weight rate (input) of 105,000 lb/hr and plant size of 315 T/D copper we determine allowable emission based on state process weight rate curves Ref. 084 § 148

Utah specifies 85% control $E_{sc} = 60.8 \text{ lb/TON}$
 Washington specifies no limit $E_s = E_u = 405 \text{ lb/TON}$

Texas 5.07 lb/TON

N. Mexico 3.05 lb/TON

Tenn 2.44 lb/TON

Mich
 Mont
 Nev
 Arizona } 3.43 lb/TON

$$E_s = \sum E_{si} \times A_i$$

$$= .159(60.8) + .063(405) + .055(5.07) + .05(3.05) + .011(2.44) \\ + .662(3.43)$$

$$= 9.67 + 25.52 + .28 + .15 + .03 + 2.27$$

$$E_s \approx 37.9 \text{ lb/TON copper}$$

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-3-74
 Source Primary Copper

Sulfur Oxide

E_u : Ref 075 Table 7.3-1

Using the SO_x emissions for roasting, smelting and converting and the 4:1 ratio for concentrate to copper we may calculate an emission factor based on copper produced. (47% of are roasted does not come into play since all S. is assume to be emitted)

$$E = 4 \times \sum E_p + E_s + E_c$$

$$E = 4 (60 + 320 + 870) = 4 \times 1250 = 5000 \text{ lb/ton copper}$$

∴

$$E_u = 5000 \text{ lb/ton copper}$$

E_N : Ref 075 Table 7.3-1 Note ① States that sulfuric acid recovery and/or lime-dust scrubbing may achieve about 90% reduction.

Ref 030 Table 6-12 shows cost data for various control scheme on copper plants. An efficiency of 99.5% is given for electric & flash smelting plant whereas 98.5% is given for roaster/reverb and reverberatory smelting plants.

We use an efficiency of 99.5% as attainable control tech in 1975

$$E_N = .005 \times E_u = .005 \times 5000$$

$$\therefore E_N = 25 \text{ lb/ton copper}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 4-29-75

Source Primary Copper

Sulfur Oxides cont'd

E_s : Ref 084 § 148 limitation of SO_2 emissions from existing state regs.
 fractional capacity occurrence by state (A_t) is as appears
 under Particulate E_s factor calculations.

From Ref 030 pg 6.1-22

Model Smelter 1000 T/D input ore
 27% Cu
 32% S

We will use this same sulfur cone for determining the sulfur input
 for a 315 Ton copper /day plant with a 4:1 ore to copper ratio

$$315 \frac{\text{Ton}}{\text{d}} \text{ Cu} \times 4 \frac{\text{Ton ore}}{\text{Ton Cu}} = 1260 \frac{\text{Ton}}{\text{d}} \text{ ore}$$

$$.32 \times 1260 \approx 403 \frac{\text{lb}}{\text{d}} \text{ sulfur input}$$

$$403 \frac{\text{lb}}{\text{d}} \times \frac{1}{24 \text{ hr}} \times \frac{2000 \frac{\text{lb}}{\text{ton}}}{\text{Ton}} = 33,583 \frac{\text{lb}}{\text{hr}} \text{ sulfur input}$$

for typical plant

Of the 9 states with copper smelting only Nevada, New Mexico, and Arizona have regulations governing SO_2 emissions from Cu Non-Ferrous smelters. specifically.

Arizona	$lb SO_2/hr = .1 \times \frac{lb}{hr} S / \text{hr}$
Nevada	" " = .2 x "
New Mexico	" " = .8 x " (exist) .2 x " (new)

For other states $E_s = E_u = 5000 lb/\text{ton}$

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Computation Sheet For Emission Factors

Calculations Done By J. Maxone Date 4-29-75
 Source Primary Copper

SO_x/ cont'd

E_s : Allowable emissions

	lb SO _x /hr	lb/TON Cu	ai
Arizona	3358	256	.471
New Mexico	26,866 Ext 6717 New	2047 Ext 512 New	.050
Nevada	6717	512	.039
Remaining 6 states	—	5000	.44

$$\begin{aligned}
 E_s(\text{exist}) &= .471(256) + .05(2047) + .039(2047) + .44(5000) \\
 &= 120.6 + 102.4 + 79.8 + 2200
 \end{aligned}$$

$$E_s(\text{exist}) \approx 2503 \text{ lb/TON copper}$$

$$\begin{aligned}
 E_s(\text{new}) &= .471(256) + .05(2047) + .039(512) + .44(5000) \\
 &= 120.6 + 102.4 + 20 + 2200
 \end{aligned}$$

$$E_s(\text{new}) = 2443 \text{ lb/TON copper}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-4-74
 Source Primary Copper

Summary of Emission Factors

lb/TON COPPER

Pollutant	E_u	E_N	E_s (existing)	E_s (new)
Particulate	405	1.22	37.9	37.9
SO_x	5000	25	2503	2443

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Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 12-23-74

Source Ferroalloy - Electric Submerged Arc Furnace

Ref 239, 240,

Emissions from this industry are generally particulate matter and CO. The level depends ~~highly~~ on the type of alloy, the composition of the alloy, and furnace operation.

We will attempt to evaluate this industry overall by considering the total Ferroalloy production and develop emission estimates from "weighted" values ~~of~~ of production and reported emission factors.

K:

Ref 239 pg 1x-28

Tabulation of parameters for a model furnace is presented on the basis of 90% utilization of full capacity which we assume is representative of this industry.

$$\therefore K = .90$$

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Computation Sheet For Industrial Factors

Calculations Done By W. Marone Date 12-23-74

Source Ferroalloy - Electric Submerged Arc Furnace

$A_3 P_C$:

Ref 239 pg 1-1

Identifies the alloy product groups that are associated with the electric submerged arc furnace, which is identified as the largest source of emissions from this industry.

Our study will include the following:

Silvery Iron (15-22% FeSi)
 50% Ferrosilicon
 65-90% Ferrosilicon
 Silicon Metal
 Calcium Silicon
 High Carbon - Ferromanganese
 Silicomanganese
 Ferromanganese silicon
 High Carbon - Ferrochrome
 Ferrochrome-silicon

<u>1971 Production #</u>
94,801
377,403
109,951
88,888
10,309
266,376
150,383
44,918
113,664
66,685
<hr/>
1,323,278

*Ref 239 pg 1-12 Table II-4

$$P_{71} \approx 1.323 \times 10^6 \text{ TONS/yr}$$

From Ref 239 pg II-1 states than for the ten years before 1972 the production grew at an average rate of 1.5%/yr.

We assume that this overall trend will continue and be applicable to 1971-1985 such that $P_C = .015$ compound

$$P_{75} = P_{71} (1 + .015)^4 = 1.323 \times 10^6 (1.015)^4$$

$$P_{75} \approx 1.40 \times 10^6 \text{ TONS/yr}$$

$\therefore P_C = .015$ compound

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Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 12-23-74

Source Ferroalloy

A: cont'd

With $P_{75} = 1.40 \times 10^6 \text{ ton/yr}$ and a value of $K = .90$
 we may calculate the production capacity in 1975

$$A = \frac{P_{75}}{K} = \frac{1.40 \times 10^6}{.9} \approx 1.56 \times 10^6 \text{ ton/yr}$$

$$\therefore A = 1.56 \times 10^6 \frac{\text{ton}}{\text{yr}}$$

ferro-alloy produced

P_B :

Ref 037 pg 34 Class 33.1

Asset Guidance period = 18 yrs

$P_g \text{ est} \Rightarrow 2 \times \text{IRS} = 36 \text{ yrs}$

$$\frac{100\%}{36 \text{ yrs}} \approx 2.8\%/\text{yr simple}$$

$$\therefore P_B = .028 \text{ simple}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-23-74
 Source Ferroalloy - Electric Submerged Arc Furnace

Particulate

E_u : Uncontrolled emission factors are presented in Table III-4 pg 239 VI-19
 These factors are synthesized by "weighting" by the fractional production obtained for 1971

<u>Alloy</u>	<u>AI 1971</u> <u>Fractional Production</u>	<u>E</u> <u>lb/ton alloy</u>	<u>AI x E</u>
Slvery Iron	.072	116	8.352
50% Ferrosilicon	.285	446	127.110
65-90% Ferrosilicon	.083	673	55.859
Silicon Metal	.067	1200	80.400
Calcium Silicon	.008	1343	10.744
HC Ferromanganese	.201	335	67.335
Silicomanganese	.114	219	24.966
Ferromanganese Silicon	.034	315	10.710
HC Ferrochrome	.086	335	28.880
Ferrochrom-Silicon	.050	831	41.550
<hr/>			
$\Sigma = 1000 /$			

$$\begin{aligned} \Sigma &\approx 455.8 \\ &= 456 \text{ lb/ton} \\ &\text{alloy} \end{aligned}$$

$\therefore E_u = 456 \text{ lb/ton ferro alloy produced}$

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Computation Sheet For Emission Factors

Calculations Done By W.J. Mannone Date 12-23-74

Source Ferroalloy

Particulate cont'd

E_N : Ref 239 pg II-5 to II-8

Discusses control technology for electric arc furnaces in Ferroalloy industry

① 8-9% off by Wet Scrubber (HE) on open furnace for siliconmanganese HE Ferrochromium and ferrosilicon-silicon

② 99.9% by HE Venturi Scrubber on covered furnace for ferromanganese siliconmanganese and 50% ferrosilicon

③ 75-95% overall for semiclosed furnaces

④ overall controls for closed furnaces are generally better than 99%

⑤ cloth filters on open furnaces achieve 99% for ferrochromium-silicon siliconmanganese, silicon metal

We will assume that 99% is achievable in determining
 E_N

$$E_N = .01 \times E_u = .01 \times 456$$

$$E_N = 4.56 \text{ lb/ton}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-23-74
 Source Ferroalloy

Particulate

E_p Ref 239
 pg VIII - 5

size range 7,000 - 50,000 kw furnaces

since we have no info on size distribution we assume that the typical existing Ferroalloy furnace may be represented by the simple average

$$\frac{50+7}{2} = 28.5 \times 10^3 \text{ kw}$$

28.5 megawatts Avg Furnace

Ref 239 Table VIII - 2 gives $\frac{\text{lb charge}}{\text{hr}}$ which we will "weight" according to alloy production (A_i)

<u>Alloy</u>	<u>A_i</u>	<u>(F)</u> <u>lb charge/MW-HR</u>	<u>$A_i \times F$</u>
Silvery Iron	.072	1380	99.36
50% Ferrosilicon	.285	1000	285.00
65-90% Ferrosilicon	.083	1020	84.66
Silicon Metal	.067	700	46.90
Calcium Silicon	.008	660	5.28
HC Ferrromanganese	.201	2500	502.50
Silicomanganese	.114	1410	160.74
Ferrromanganese silicon	.034	1590	54.06
HC Ferrochrome	.086	1900	163.40
Ferrochrom-Silicon	.050	920	46.00

Total \approx 1450 $\frac{\text{lb charge}}{\text{MW HR}}$

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Computation Sheet For Emission Factors

Calculations Done By W. Marone Date 12-23-74
 Source Ferroalloy

Particulate cont'd

E_s :

$$\text{Typical Process Rate} = 1450 \frac{\text{lb charge}}{\text{hr HR}} \times 28.5 \text{ wt\%} = 41,325 \frac{\text{lb}}{\text{hr input}}$$

There are no specific state reg's for Ferroalloy however we apply Gen Process weight rate curve Ref 084 & 148

Ref 239 Table W-2 gives a distribution of Ferroalloy furnaces by state which we assume will be representative for 1975-1988

	No Furn(Ferroalloy)	A_i	<u>Allow Factor lb/hr</u>	$A_i \times E_A$
Ohio	52	.359	31	11.129
Ala	12	.083	23.5	1.951
Penn	0			
Tenn	14	.097	23.5	2.280
Oregon	8	.055	29	1.595
W. Va	25	.172	29	4.988
N.Y.	6	.041	27.5	1.128
Kent.	11	.076	31	2.356
Iowa	5	.034	31	1.054
Wash	6	.041	**	.041 $\times E_u'$
Tex	3	.021	60.6	1.273
SC.	2	.014	31.2	.437
N.J.	1	.007	30	.210
TOTAL	195			

$$\sum * 28.4 + .041 E_u' \frac{\text{lb}}{\text{hr}}$$

** No reported Process Curve assume $E_{SL}' = E_u'$ in (lb/hr)

$$E_g' (\text{lb/hr}) = (28.4 + .041 E_u') \text{lb/hr}$$

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Computation Sheet For Emission Factors

Calculations Done By J. Marrone Date 12-24-74
 Source Ferroalloy

Particulates

E_s :

We must now calculate the typical process output (alloy product) rate. Ref 239 Table III-2 gives the $\frac{\text{lb Prod}}{\text{MW-HR}}$ which may be combined with the previously calculated value of $A_i \times F$ ($\frac{\text{lb charge}}{\text{MW-HR}}$) to determine a "weight" value for ;

$\frac{\text{lb Product}}{\text{MW-HR}}$

$$\text{and } \frac{\text{lb Prod}}{\text{MW-HR}} \times \text{MW} = \frac{\text{lb Prod}}{\text{hr}} \text{ (typical)}$$

<u>Alloy</u>	<u>$A_i \times F$</u>	<u>$(A_i \times F) \times \frac{\text{lb prod/MW-HR}}{\text{lb charge/MW-HR}}$</u>
Silver Iron	99.36	55.44
50% F.S.	285.00	114.00
65-70% FS	84.66	18.84
Silicon Metal	46.90	9.65
Calcium-Silicon	5.28	1.36
HC FM	502.50	167.63
SM	160.74	51.76
FMS:	54.06	12.58
HC FC	163.40	40.94
FCS:	46.00	13.50
<hr/>		
	$\Sigma = 485.7$	$\frac{\text{lb prod}}{\text{MW-HR}}$

$$\text{Typical Prod Rate} = 485.7 \frac{\text{lb Prod}}{\text{MW-HR}} \times 28.5 \text{ MW} = 13,842 \frac{\text{lb}}{\text{hr}}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Mannone Date 12-24-74

Source Ferroalloy

Particulate cont'd

$$E_s: \text{Product Rate} = \frac{\text{Avg Product Rate}}{\text{Ton/Hr}} = \frac{13,842 \text{ lb}}{\text{hr}} \equiv 6.92 \frac{\text{Ton}}{\text{hr}}$$

$$\text{Uncontrolled Emission Rate} = E_{uc} \times 6.92 \frac{\text{Ton}}{\text{hr}} = 456 \frac{\text{lb}}{\text{Ton}} \times 6.92 \frac{\text{Ton}}{\text{hr}}$$

$$E'_{uc} = 3155 \text{ lb/hr}$$

Continuing

$$E'_s \frac{\text{lb}}{\text{hr}} = 28.4 + .041(3155)$$

$$E'_s \approx 158 \text{ lb/hr}$$

With avg product rate at $6.92 \frac{\text{Ton}}{\text{hr}}$ we may calculate the value of E_s in terms of $\frac{\text{lb}}{\text{Ton}}$.

$$E_s = \frac{158 \frac{\text{lb}}{\text{hr}}}{6.92 \frac{\text{Ton}}{\text{hr}}} = 22.8 \frac{\text{lb}}{\text{TON}}$$

Ferro
alloy
produced

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-24-74

Source Ferroalloy

CO

E_u :

Ref 221 pg 29 and pg 30 Table IX

For open furnaces (pg 22 75% of Ferroalloy production) the following values are reported

50% FeSi	133	lb/ton prod
75% FeSi	160	"
90% FeSi	182	"
Si Metal	—	
Si Mn	—	
FeMn	101	
FeCr	104	

We will use the avg from this for E_u estimation

$$E_u = \frac{680}{5} = 136 \text{ lb/TON}$$

∴ $E_u = 136 \text{ lb/TON}$ Ferroalloy produced

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Computation Sheet For Emission Factors

Calculations Done By W.J. Marzzone Date 12-24-74
 Source Ferroalloy

CO/ cont'd

E_N : Ref 239 pg VIII-5 to VIII-8

indicates that CO for open furnaces are often limited by oxidation to CO_2 at the charge surface in the furnace. A large gas volume results which make control for CO difficult (high wt)

Flaring does not appear feasible nor burning for fuel unless the conc is sufficient to support unassisted combustion.

We make the assumption that although unattractive incineration may be employed and 99% eff should be achieved

$$\therefore E_N = .01 \times 136 \cong 1.4 \text{ lb/TON}$$

$$E_N = 1.4 \text{ lb /TON} \quad \text{Ferroalloy produced}$$

E_s : Ref 084 & 148

There are no existing state regulations which may be applied to this source $E_s = E_N$

$$\therefore E_s = 136 \text{ lb /TON} \quad \text{Ferroalloy}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marzzone Date 6-11-75

Source Ferroalloy - Electric Submerged Arc Furnace

Lead

Ref 277 Vol II appendix B

indicates that the range in lead content for particulate emissions from silicomanganese and ferromanganese furnaces is .05-.1 percent.

We assume the average percent (.075%) to be suitable and justified to apply to ferroalloy industry in general.

The emission factors E_u , E_N , E_s will be determined by applying the .075% factor to the corresponding values for particulate which assumes that lead control will be the same as for particles.

$$E_u(\text{lead}) = .00075 \times 456 \text{ lb/ton} = .34 \text{ lb/ton}$$

ferroalloy

$$E_N(\text{lead}) = .00075 \times 4.56 \text{ lb/ton} = .0034 \text{ lb/ton}$$

ferroalloy

$$E_s(\text{lead}) = .00075 \times 22.8 \text{ lb/ton} = .017 \text{ lb/ton}$$

ferroalloy

We assume that existing plants may achieve the same degree of control as new plants so that $E_{IIIId} = E_N$

$$E_{IIIId}(\text{lead}) = .0034 \text{ lb/ton}$$

ferroalloy

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Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 1-6-75

Source Iron & Steel Industry

This study will include the following subcategories which identify specific operations or production methods within the Integrated Iron and Steel Industry

(1) Blast Furnace (Pig Iron Production)

(2) Basic Oxygen Furnace (BOF)

(3) Open Hearth Furnace (OHF)

(4) Electric Arc Furnace (EAF)

(5) Sintering

(6) Scarfing

Steel Production
From Pig Iron & Scrap

By-Product Coke ovens, which are a part of the Integrated Iron and Steel Industry, have been done separately and appears under the heading "By-product Coke Ovens" by W. Marrone.

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Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 1-6-75

Source Iron & Steel Industry

K:

Ref 212 pg III-1 and pg IV-1 points out that steel making capacity has been difficult for this industry to define and quote. Presently, it is no longer reporting "official" capacity figures.

Ref 234 p 3-59 gives a production in 1968 for 80% at 90% of capacity

Ref 212 Appendix C pg C-66 discusses the lag between production and capacity and actually predicts that production will closely approach capacity. Our approx in 1968 $\frac{\text{Prod} = 48 \times 10^6}{\text{Cap} = 52.5 \times 10^6}$
From graph $K_{68} = \frac{48}{52.5} = .914$

In addition, to discussing the difficulty with defining capacity the above ref (212 pg III-1) indicates that this industry may and has often operated at levels above nominal capacity however this can not be expected to occur for long periods.

With lack of definitive info or predictions we must assume a value of K which may be considered representative of the industry. This value of K is to be applied to the whole industry

$$\boxed{K_{\text{All}} = .90}$$

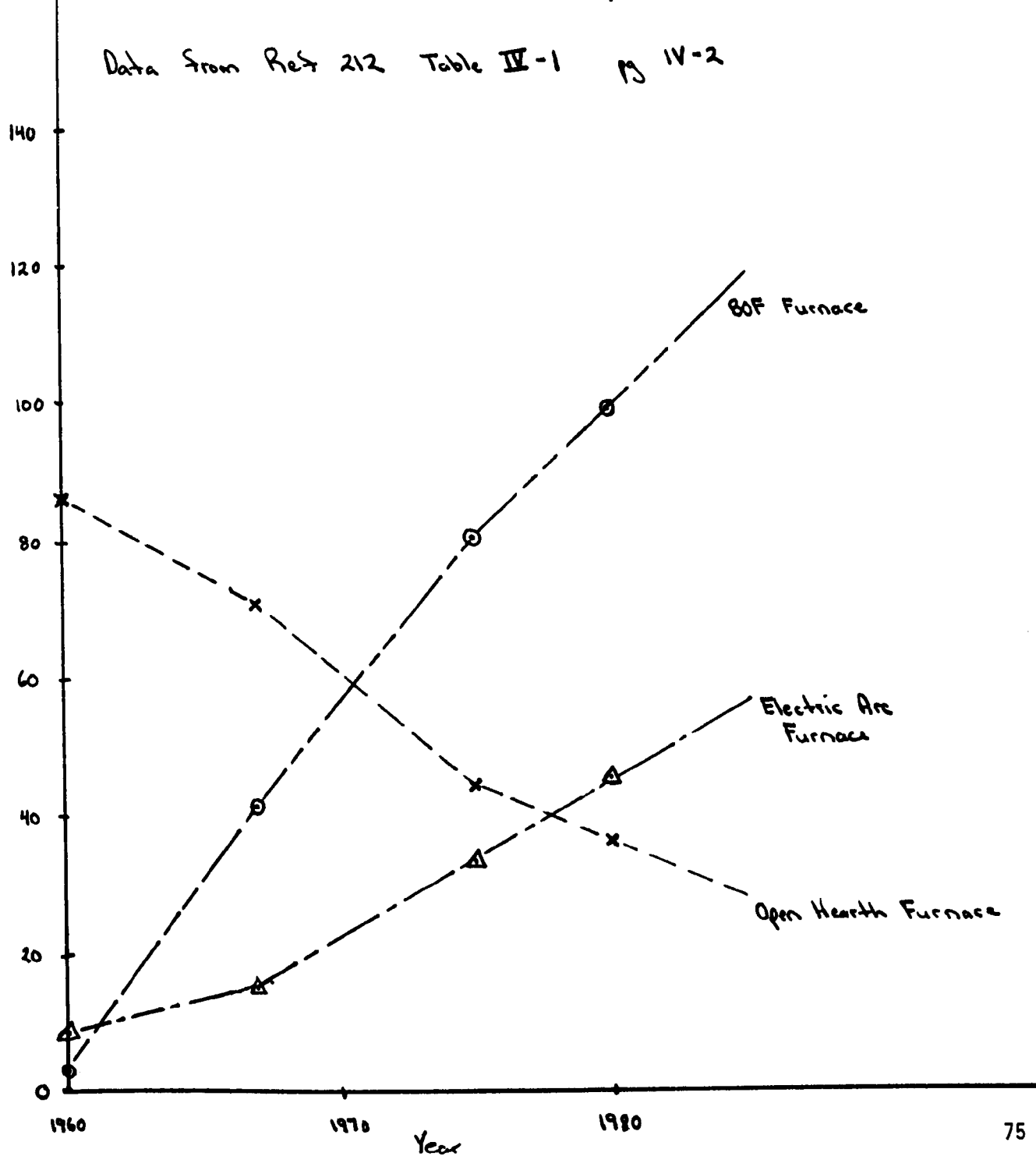
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Source Iron & Steel Industry



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Computation Sheet For Industrial Factors

Calculations Done By W. Marzzone Date 1-6-75

Source Iron & Steel Industry

A:

Steel Furnaces Production

From the graph on pg 3 of the "Industrial Factors" we may obtain the 1975 production of Raw Steel by each of the three major steel producing furnaces.

$$\text{BOF } P_{75} = 80 \times 10^6 \text{ TON/yr}$$

$$\text{EAF } P_{75} = 33 \times 10^6 \text{ TON/yr}$$

$$\text{OxF } P_{75} = 44 \times 10^6 \text{ TON/yr}$$

applying $K= .9$ to these values we may estimate the production capacity (A) in 1975 by: $A = \frac{P_{75}}{K}$

$$A_{\text{BOF}} = 88.9 \times 10^6 \frac{\text{TON}}{\text{YR}} \text{ raw steel produced}$$

$$A_{\text{EAF}} = 36.7 \times 10^6 \frac{\text{TON}}{\text{YR}} \text{ raw steel produced}$$

$$A_{\text{OxF}} = 48.9 \times 10^6 \frac{\text{TON}}{\text{YR}} \text{ raw steel produced}$$

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Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 1-6-75

Source Iron & Steel Industry

A:

Blast Furnace

Pig Iron Production is developed from Ref 212 Table IV-5
 pg IV-6

1975 apparent Pig Iron consumption for raw steel 106.8×10^6 TON
 with a ratio of .68 TON pig iron / TON raw steel

$$P_{75} (\text{Pig Iron}) = 106.8 \times 10^6 \text{ TON/yr}$$

$$\text{applying } K = .9 \quad \therefore A = \frac{106.8 \times 10^6}{.90} = 118.7 \times 10^6 \text{ TON/yr}$$

This value is checked with Total Steel (Raw) Capacity
 is seen to be the value of .68 Pig Iron / Raw Steel holds
 since all this data comes from the same source.

$$\frac{\text{Pig Iron Cap}}{\text{Raw Steel Cap}} = \frac{118.7 \times 10^6}{(88.9 + 36.7 + 48.9) \times 10^6} = \frac{118.7 \times 10^6}{174.5 \times 10^6} \approx .680 \checkmark$$

$$\therefore A (\text{Blast Furnace}) = 118.7 \times 10^6 \text{ TON/yr}$$

pig iron produced

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 1-6-75

Source Iron & Steel Industry

A: cont'd

Surfacing : Ref 047 pg 47

In generating an emission factor it assumes that all steel produced is scurfed [process for removing surface irregularities from semi-finished steel shapes].

Ref 212 pg C-116

presents a plot of % of steel scurfed for the period 1955-1968 with a projection that it will level off at 50%.

We believe this reference to present more definitive data and we make the assumption that surfacing production will be 50% of total steel production

$$A_{\text{scurf}} = .5 \times \sum A_{\text{Furnaces}}$$

$$A_{\text{surfacing}} = .5 \times (88.9 + 36.7 + 48.9) \times 10^6 \frac{\text{TON}}{\text{YR}} \text{ raw steel scurfed}$$

$$\therefore A \approx 87.3 \times 10^6 \frac{\text{TON}}{\text{YR}} \text{ raw steel scurfed}$$

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Computation Sheet For Industrial Factors

Calculations Done By W. Marone Date 1-7-75

Source Iron & Steel Industry

A : cont'd

The most recent production figure for sinter production was for 1967 (Ref 212 Table III-34)

It was shown also in the same reference (Figure C-1) that Sinter was being replaced by pellets in the feed charge of blast furnaces. It is not known how the relative demand for each of these materials stands presently. Our best estimate of sinter production will be developed by assuming that the ratio of sinter produced in 1967 to 1967 pig iron production is valid for 1975-1985. It must be kept in mind however, that the sinter/pellet competition in pig iron production is a major factor in the ultimate production (demand) for these two materials. One interesting factor to also keep in mind is that sintering was basically used to collect fine materials and while not demanded solely by the pig iron production there is a probability that this practice may continue.

Ref 212

1967 Sinter Production	51,068,111 TONS	} Table III-34
1967 Pig Iron Production	86,984,000 TONS	

$$\frac{\text{Sinter}}{\text{Pig Iron}} = \frac{51,068,111}{86,984,000} = .587$$

$$A_{\text{Sinter}} = .587 \times A_{\text{Pig Iron}} = .587 \times 118.7 \times 10^6 \text{ TON/Yr}$$

$$(A_{\text{Sinter}} = 69.6 \times 10^6 \text{ TON/YR} \text{ sinter produced}) 79$$

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 1-7-75

Source Iron and Steel Industry

P_c :

The growth for each of the steel producing furnaces and the whole steel industry is obtained from the data plotted in the accompanying graph (pg 3 of Industrial Factors)

We assume simple growth during period 1975 - 1985

	<u>Production</u> 10^6 TONS	
	<u>1975</u>	<u>1980</u>
BOF	80.0	99.0
EAF	33.0	45.0
OHF	44.0	36.0
TOTAL	<u>157</u>	<u>180</u>

$$\text{For simple growth } P_c = \frac{(1980 - 1975)/5}{1975}$$

$$\text{BOF } P_c = \frac{(99 - 80)/5}{80} = .0475 \text{ simple}$$

$$\text{EAF } P_c = \frac{(45 - 33)/5}{33} = .0727 \text{ simple}$$

$$\text{OHF } P_c = \frac{(36 - 44)/5}{44} = -.0363 \text{ simple}$$

$$\text{Total } P_c = \frac{(180 - 157)/5}{157} = .0292 \text{ simple}$$

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Computation Sheet For Industrial Factors

Calculations Done By 2). Marone Date 1-7-75

Source Iron and Steel Industry

P_c : cont'd

$$\therefore P_c(\text{BOF}) = .048 \text{ simple}$$

$$P_c(\text{EAF}) = .073 \text{ simple}$$

$$P_c(\text{OHF}) = -.036 \text{ simple}$$

Since scarring is related to total raw steel production then we may represent the growth in scarring to the simple growth in total raw steel developed above.

$$P_c(\text{scarring}) = .029 \text{ simple}$$

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Computation Sheet For Industrial Factors

Calculations Done By W. Marone Date 1-7-75

Source Iron and Steel Industry

P_c : cont'd

Blast Furnace production was developed (pg 5 "Ind Factors") from a relationship to total raw steel production. We may therefore relate the growth in Blast Furnace production to the simple growth in raw steel developed above.

$$\therefore P_c(\text{Blast Furnace}) = .029 \text{ simple}$$

From pg 7 of "Industrial Factors" we see that the value of A for sintering was developed directly from a factorization of pig iron production (Blast Furnace) which implies that the growth in sinter production will mirror the blast furnace production growth

$$\therefore P_c(\text{sintering}) = .029 \text{ simple}$$

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Computation Sheet For Industrial Factors

Calculations Done By W. Manrone Date 1-7-75

Source Iron and Steel Industry

P_B :

OPEN HEARTH FURNACES

Ref 212 pg V-10 confirms the negative growth rate estimated for this industry and even suggest that by 1990 → open hearth furnace production will ~~exist~~ exist.

We suspect that since BOF and EAF are replacing the Open Hearth Furnaces and its growth is negative that

P_B for this category is zero

$$\therefore P_B (\text{OHF}) = 0.0$$

All Other Iron & Steel Categories

Ref 037 pg 34 Class 33.1

Asset guideline Period = 18 yrs

$P_B \text{ est} \rightarrow 2 \times \text{IRS} = 36 \text{ yrs}$

$\frac{100\%}{36 \text{ yrs}} = 2.77\%/\text{yr simple}$

$$\therefore P_B \left(\begin{array}{l} \text{BOF} \\ \text{EAF} \\ \text{Blow Furnace} \\ \text{Sintering} \\ \text{Scaling} \end{array} \right) = .028 \text{ simple}$$

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Computation Sheet For Industrial Factors

Calculations Done By W. Mannone Date 1-7-75

Source Iron and Steel Industry

Summary of Industrial Factors

Category	K	P _c	P _e	A
Blast Furnace	.90	.029 s	.028 s	$118.7 \times 10^6 \frac{\text{ton}}{\text{yr}}$ pig iron produced
Basic Oxygen Furnace	.90	.048 s	.028 s	$88.9 \times 10^6 \frac{\text{ton}}{\text{yr}}$ raw steel produced
Electric Arc Furnace	.90	.073 s	.028 s	$36.7 \times 10^6 \frac{\text{ton}}{\text{yr}}$ raw steel produced
Open Hearth Furnace	.90	-.036 s	0.0	$48.9 \times 10^6 \frac{\text{ton}}{\text{yr}}$ raw steel produced
Sintering	.90	.029 s	.028 s	$69.6 \times 10^6 \frac{\text{ton}}{\text{yr}}$ sinter produced
Scarfing	.90	.029 s	.028 s	$87.3 \times 10^6 \frac{\text{ton}}{\text{yr}}$ raw steel scarfed

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Emission Factors

Calculations Done By 2). Mannone Date 1-7-75

Source Iron and Steel Industry

E_u :

Blast Furnace

(Particulates): Ref 075 Table 7.5-1 $E_u = 150 \text{ lb/TON}$ pig iron produced assumed

$$\therefore E_u(\text{P}) = 150 \text{ lb/TON} \quad \text{pig iron produced}$$

Ref 075
Table 7.5-1

(CO): $E_u(\text{CO}) = 1750 \text{ lb/TON}$ pig iron produced

See next page

As will be discussed under E_u for CO from pg 12 of "Emission Factors"
 the exhaust stream containing CO is generally used
 within the plant to recover energy (from heating value)
 Actual emissions are negligible. Therefore the actual
 emission (E') would be very low. The value above
 corresponds to no energy recovery.

(Trace Metals):

Ref 232 p18 $\approx 1.2\%$ of part emission is magnesium

$$.012 \times 150 \frac{\text{lb}}{\text{TON}} = 1.8 \frac{\text{lb}}{\text{TON}} \quad \text{pig iron produced}$$

Ref 230 Table 6-1 .8 lb Manganese/TON pig iron produced

Ref 230 Table 9-1 .03 lb Vanadium/TON pig iron produced

$$\text{Total} = 1.8 + .8 + .03 = 2.63 \text{ lb/TON pig iron}$$

\therefore Trace metals

$$E_u = 2.63 \text{ lb/TON} \quad \text{pig iron produced}$$

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Computation Sheet For Emissions
Industrial Factors

(Revised 1-28-75)

Calculations Done By W. Marrone Date 1-7-75

Source Iron and Steel Industry

Blast Furnace
 E_u for CO:

The "uncontrolled" emission factor of 1750 lb/Ton as obtained above is identified as the emissions from a totally uncontrolled blast furnace operation.

Ref 075 pg 7.5-1 and Ref 212 pg C-38 to C-39 indicate that most of the top gas from blast furnaces is used for heating blast stoves and as a fuel in other operations within the steel plant.

Ref 212 pg C-47 also states that the generation of the recyclable fuel in iron smelting is one feature which contributes to the economy of the blast furnace operation.

Ref 075 pg 7.5-1 Conditions called "slips" ~~can~~ still cause instantaneous emission of CO from a controlled furnace employing energy recovery. This occurrence is becoming minimum due to improved furnace operation and material handling.

We make the judgement that all plants will employ energy recovery so that CO emissions will be negligible during normal production. We feel that emissions during "slips" may still occur sufficiently in existing plants.

We assume that emissions from this situation will only be about .1% of the uncontrolled.

$$E_u = .001 \times 1750 = 1.75 \text{ lb/TON} \quad \text{pg iron produced}$$

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Computation Sheet For Emission Factors

Calculations Done By 2). Marrowe Date 1-7-75

Source Iron and Steel Industry

E_u : cont'd

Basic Oxygen Furnace:

Particulate

Ref 075 Table 7.5-1

$$E_u = 51 \text{ lb/TON raw steel produced}$$

CO

Ref 075 Table 7.5-1

$$E_u = 139 \text{ lb/TON raw steel produced}$$

Fluoride

Ref 075 Table 7.5-1

occurs as CaF_2 solid

$$E'_u = .2 \text{ lb } \frac{\text{CaF}_2}{\text{TON raw steel}}$$

$$\begin{aligned} \text{MW F} &= 19 \\ \text{MW CaF}_2 &= 78 \end{aligned}$$

in terms of F

$$E_u = .2 \times \frac{19}{78} = .0487 \approx .049 \text{ lb/TON}$$

raw

$$\therefore E_u = .049 \text{ lb/TON raw steel produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 1-7-75

Source Iron and Steel Industry

E_u : cont'd

Basic Oxygen Furnace

Trace Metals

Ref 230 p 6-6 Table 6-1	Manganese	1 lb/ton raw steel
Ref 230 p 9-6 Table 9-1	Vanadium	.009 lb/ton raw steel
Ref 232 p 18	Magnesium	.5% of emission (Pt) is magnesium

$$E' = .005 \times 51 = .255 \text{ lb}_{\text{raw steel}}$$

$$\text{Total} = 1 + .009 + .255 \approx 1.26 \text{ lb}_{\text{raw steel}}$$

$$\therefore E_u = 1.26 \text{ lb/ton raw steel produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marzzone Date 1-7-75

Source Iron and Steel Industry

E_u : cont'd

Electric Arc Furnace

Particulate Ref 047 pg 96 used 10 lb/TON for uncontrolled electric arc furnace

Ref 075 Table 7.5-1

$$\begin{array}{ll} E' \text{ for oxygen lance} & 9.2 \text{ lb/TON} \\ E' \text{ for oxygen lance} & 11.0 \text{ lb/TON} \end{array} \quad \left. \begin{array}{l} \text{avg} \approx 10 \text{ lb/TON} \\ \text{ } \end{array} \right.$$

Ref 250 pg 9 reports a figure of up to 30 lb/ton for carbon steel and 15 lb/ton for alloy steels from electric arc furnace operations. These emissions are implied to be from total sources; not just the furnace exhaust. Other prominent sources are (1) charging, (2) tapping, (3) oxygen blowing

We believe the latter source of info to be more representative

$$\therefore E_u = 15-30 \text{ lb/ton}$$

From Ref 212 pg III-16 Table III-19

$$\begin{array}{rcl} \text{Raw Carbon Steel Prod 1167} & = & 113,190 & 88.97\% \\ \text{Raw Alloy Steel Prod 1187} & = & 14,023 & 11.03\% \\ & & \hline & 127,213 \end{array}$$

weighting emission factors by this breakdown we may synthesize a E_u value

$$E = .1103(15) + .8897(30) \approx 1.65 + 26.69 \approx 28.3$$

$$\therefore E_u = 28.3 \text{ lb/TON} \quad \text{raw steel produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 1-7-75

Source Iron and Steel Industry

E_u : cont'd

Electric Arc Furnace

Trace Metals

Ref 232 pg 18

Magnesium

5% of emissions (P+) " magnesium

$$.05 \times 28.3 = 1.42 \text{ lb/TON}$$

Ref 230 pg 6-6

Manganese

Oxygen lance .3 lb/TON
 No lance .2 lb/TON } avg .25 lb/TON

$$\text{Total} = 1.67 \text{ lb/TON raw steel}$$

$$E_u = 1.67 \text{ lb/TON raw steel produced}$$

CO

Ref OTS Table 7.5-1

$$E_u = 18 \text{ lb/TON raw steel produced}$$

Fluorides

Oxygenated and no-oxygenated

gaseous HF .012 lb/TON

MW F = 19

MW HF = 20

solid CaF_2 .238 lb/TON

MW CaF_2 = 78

$$E' = .012 \times \frac{19}{20} = .0114 \text{ lb/TON F as gas}$$

$$E'' = .238 \times \frac{19}{78} = .0580 \text{ lb/TON F as solid}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Mannone Date 1-7-75
 Source Iron and Steel Industry

E_u : cont'd

Electric Arc Furnaces

Fluorides

$$\text{Total Fluorides as "F"} = .0114 + .0580 = .0694 \text{ lb/Ton}$$

$$E_u = .069 \text{ lb/Ton} \text{ raw steel produced}$$

~ 83.5% of this is due to solid emission
 and 16.5% due to gaseous HF emission.

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Computation Sheet For Emission Factors

Calculations Done By W. Marone Date 1-7-75

Source Iron and Steel Industry

E_u : cont'd

Open Hearth Furnace

Particulate Ref 047 pg 96 gives a weighted emission factor between oxygen lanced and non-oxygenlanced open hearth production 17 lb/ton raw steel

based on 8 lb/ton without lancing

21 lb/ton with lancing

and some unspecified % production breakdown of lancing practices.

we determine a breakdown

$$\text{by } 8x + 21y = 17$$

$$\text{and } x + y = 1$$

$$\therefore xy = 1 - y$$

$$8x + 21 - 21y = 17$$

$$13x = 3 \quad x = .23 \quad (23\%)$$

$$\rightarrow y = .77 \quad (77\%)$$

without lancing

$$\therefore E_u = 17 \text{ lb/ton raw steel produced}$$

Trace Metals

Ref 230 pg 6-6

Manganese

.1 lb/ton oxygen lance

.6 lb/ton without oxygen lance

weighting these by the above determined lance/no lance ratio.

$$E = .23(.6) + .77(.1) \approx .138 + .077$$

$$E = .215 \text{ lb/ton}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marvone Date 1-7-75
 Source Iron and Steel Industry

E_u : cont'd

Open Hearth

Trace Metals

Ref 230 pg 9-6 Vanadium .006 lb/tow no oxygen lance
 .01 lb/tow oxygen lance

weighting again

$$E = .23(.006) + .73(.01) = .00138 + .0073$$

$$E = .00868 \text{ lb/tow raw steel}$$

Ref 232 N 18 Magnesium ,3% of Pt emission is magnesium

$$E = .003 \times 17 \frac{\text{lb}}{\text{tow}} = .051 \frac{\text{lb}}{\text{tow}}_{\text{raw steel}}$$

$$\text{Total trace} = .215 + .0087 + .051 \approx .275 \frac{\text{lb}}{\text{tow}}$$

$$E_u = .275 \frac{\text{lb}}{\text{tow}}_{\text{raw steel produced}}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Morezone Date 1-7-75
 Source Iron and Steel Industry

E_u : cont'd

Open Hearth Furnace

Fluorides

Ref 075 Table 7.5-1

$$E' = .1 \text{ lb/TW} \text{ as HF (gaseous)}$$

$$\text{MW F} = 19$$

$$\text{MW CaF}_2 = 78$$

$$\text{MW HF} = 20$$

$$E'' = .03 \text{ lb/TW as CaF}_2 (\text{solid})$$

$$E' = .1 \frac{\text{lb}}{\text{TW}} \times \frac{19}{20} = .095 \text{ lb/TW as F in gaseous form}$$

$$E'' = .03 \frac{\text{lb}}{\text{TW}} \times \frac{19}{78} = .0073 \text{ lb/TW as F in solid}$$

$$E_u \text{ Total F} \approx .102 \text{ lb/TW raw steel}$$

with 7.1% of F from solid

92.9% of F from gas

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 1-7-75

Source Iron and Steel Industry

E_u : cont'd

Sintering

Particulate

Ref. 075 Table 7.5-1

$$E = \text{windbox} + \text{discharge} = 20 + 22 = 42 \text{ lb/ton sinter}$$

$$\therefore E_u = 42 \text{ lb/ton sinter produced}$$

SO_x

Ref 075 pg 7.5-5 note ⑨

3 lb SO₂ / ton sinter at windbox

$$\therefore E_u = 3 \text{ lb/ton sinter produced}$$

CO

Ref 075 pg 7.5.4

$$E_u = 44 \text{ lb/ton sinter produced}$$

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Computation Sheet For Emission Factors

Calculations Done By 2) Marrone Date 1-7-75
Source Iron and Steel Industry

E_u : cont'd

Scarfing

Particulates

Ref 047 pg 97

$$E_u = 3 \text{ lb/TON steel scarfed}$$

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Computation Sheet For Emission Factors

Calculations Done By J. Marzone Date 1-8-75

Source Iron and Steel Industry

E_N : Blast Furnaces
 Ref 075 pg 7.5-1

discusses control strategy for this source. CO generated and emitted is utilized as fuel within the Steel Mill so that we may expect negligible CO emissions due to this practice. Prior to CO oxidation, the exhaust gases must be cleaned of particulate matter. Ref 212 pg V-8 indicates $\approx 30\%$ to heat stoves 70% to rest of plant for heating.

Particulate
 Particulate control via primary cyclone or settling chamber + wet scrubber + venturi or electrostatic precipitation

Table 7.5-1

$$E_N(Pt) = 1.5 \text{ lb/TON pig iron produced}$$

CO

We assume that all the exhaust gas will be utilized in combustion to recover energy or would be flared so that essentially zero CO emissions would result. Emission due to "slips" will also be negligible

$$\therefore E_N(CO) = 0.0 \text{ lb/TON pig iron produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 1-8-75

Source Iron and Steel Industry

E_N : cont'd

Blast Furnace

Trace Metals

We make the assumption that trace metals, being particulate matter in the blast furnace exhaust, will be controlled to the same degree as overall particulate removal

$$ESS = \frac{150 - 1.5}{150} \times 100 = 99\%$$

$$E_N^{(T.M.)} = .01 \times E_N^{(T.M.)} = .01 \times 2.63 = .0263 \frac{\text{lb}}{\text{TON}}$$

$$\therefore E_N^{(\text{Trace Metals})} = .026 \frac{\text{lb}}{\text{TON}} \text{ pig iron produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Manrone Date 1-8-75

Source Iron and Steel Industry

E_N : cont'd

Basic Oxygen Furnace

CO Ref 075 pg 7.5-3

Carbon monoxide may be reduced to 3 lb/ton or less by combustion in the hood, direct flaring, or other ignition means. We will use $E = 3 \text{ lb/ton}$ to represent E_N .

$$\therefore E_N^{(CO)} = 3 \text{ lb/ton raw steel produced}$$

Particulates

Ref 083 indicates that most if not all BOF are (Ref 075 also) equipped with either venturi scrubber or electrostatic precipitators.

Ref 083 pg 563 Design Efficiency for two sizes:
 99.7% & 99.8%

we will use 99.7% in developing E_N

$$E_N = .003 \times E_u = .003 \times 51 \text{ lb/ton} \approx .15 \text{ lb/ton}$$

$$\therefore E_N^{(PT)} = .15 \text{ lb/ton raw steel produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marthone Date 1-8-75

Source Iron and Steel Industry

E_N : cont'd Basic Oxygen Furnace

Trace Metal

We will assume that trace metals will be controlled to the same degree as overall particulate

$$E_N(TM) = .003 \times E_u(TM) = .003 \times 1.26 \frac{\text{lb}}{\text{TON}} = .00378$$

$$\therefore E_N(\text{trace metal}) = .0038 \frac{\text{lb}}{\text{TON}} \begin{matrix} \text{raw} \\ \text{steel produced} \end{matrix}$$

Fluoride

Ref 075 Table 7.5-1

$$E_N' = .002 \frac{\text{lb CaF}_2}{\text{TON steel}}$$

$$\begin{matrix} \text{MW F} = 19 \\ \text{MW CaF}_2 = 78 \end{matrix}$$

This represents an efficiency of $\frac{.3 - .002}{.3} = 99\%$

We will use however the efficiency of 99.7% which was obtained for particulate

$$E_N(F) = .003 \times E_u(F) = .003 \times .049 \approx .00015 \frac{\text{lb}}{\text{TON}}$$

$$\therefore E_N = .00015 \frac{\text{lb}}{\text{TON}} \begin{matrix} \text{raw steel} \\ \text{produced} \end{matrix}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Madrone Date 1-8-75

Source Iron and Steel Industry

E_N : cont'd

Electric Arc Furnace

Particulates

Ref 250 pg 44 Table IV-1

achievable control efficiency for Carbon and Alloy steel plants
 with specified control systems

Carbon Steel - DSE & BE Closed roof 98.8%

Alloy Steel BE & closed roof 94%

Weighting these efficiencies by the ratio of carbon & alloy steel
 determined previously (pg 4 Emission Factors)

= 89% Carbon

= 11% Alloy

$$\text{Weighted eff} = .89(98.8) + .11(94)$$

$$= 87.93 + 10.34 = 98.27$$

we will use 98.3% eff $\Rightarrow E_N = .017 \times E_u = .017 \times 28.3 \frac{\text{lb}}{\text{TON}}$
 $E_N = .48 \frac{\text{lb}}{\text{TON}}$

$$E_N^{(P)} = .48 \frac{\text{lb}}{\text{TON}} \text{ raw steel produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 1-8-75

Source Iron and Steel Industry

E_N : cont'd

Electric Arc Furnace

Trace Metals

attainable efficiency for trace metals is assumed to be the same as for overall particulate

$$E_N^{(TM)} = .017 \times E_u^{(TM)} = .017 \times 1.67 \text{ lb/ton}$$

$$\therefore E_N^{(\text{Trace Metal})} = .028 \text{ lb/ton} \quad \text{raw steel produced}$$

CO

Ref 250 Table IV-1 pg 44

attainable CO reduction for Carbon steel 84%
 " " " " alloy steel 0%

and by weighting by Carbon/Alloy factors.

$$\begin{aligned} \text{Weighted off} &= .89(84) + .11(0) \\ &= 74.76 + 0 \approx 75\% \end{aligned}$$

$$E_N^{(CO)} = .25 \times E_u^{(CO)} = .25 \times 18 \text{ lb/ton} = 4.5 \text{ lb/ton}$$

$$\therefore E_N^{(CO)} = 4.5 \text{ lb/ton} \quad \text{raw steel produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Maronne Date 1-8-75
 Source Iron and Steel Industry

E_N : cont'd

Electric Arc Furnace

Fluorides

Ref 075 Table 7.5-1

with venturi scrubber $E'(HF \text{ gas}) = .0018 \text{ lb}^{\text{HF}}/\text{TON}$

$E'(CaF_2 \text{ sol. d}) = .011 \text{ lb CaF}_2/\text{TON}$

MW F = 19

MW HF = 20

MW CaF₂ = 78

$$E'(\text{gas}) = .0018 \times \frac{19}{20} = .0017 \text{ lb F/TON}$$

$$E'(\text{sol. d}) = .011 \times \frac{19}{78} = .0027 \text{ lb F/TON}$$

$$\overline{E'_\text{TOTAL} .0044 \text{ lb/TON}}$$

$$E_N^{(F)} = .0044 \text{ lb/TON} \text{ raw steel produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marzzone Date 5-8-75

Source Iron and Steel Industry

E_N : cont'd Open Hearth Furnace

Since $P_C < 0$ and $P_B = 0$ we would not expect NSPS to be applied to this industry for criteria pollutants. We will however supply a value for E_N for particulates.

Pics 234 pg 3-83 to 3-87 give mass balance diagrams for typical open hearth controls.

pg 3-86 gives 95% Pt (Total) VENTURE with air break
 and 90% Gaseous HF

pg 3-87 gives 95% Pt VENTURE with air break
 and 0% Gaseous HF with all HF given converted to solid

it is reasonable to expect 95% control of particulate, total fluoride, and trace metal (treated as the same degree of collection ~~as~~ particulate)

$$E_N = .05 \times E_u$$

$$E_N(\text{Pt}) = .05 \times 17 = .85 \text{ lb/TON raw steel}$$

$$E_N(\text{TM}) = .05 \times .275 = .014 \text{ lb/TON raw steel}$$

$$E_N(\text{F}) = .05 \times 102 = .005 \text{ lb/TON raw steel}$$

$$E_N(\text{Pt}) = .85 \text{ lb/TON}$$

$$E_N(\text{TM}) = .014 \text{ lb/TON} \quad \left. \right\} \text{raw steel}$$

$$E_N(\text{F}) = .005 \text{ lb/TON}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Mannone Date 1-8-75

Source Iron and Steel Industry

E_N : cont'd

Sintering

Particulate

Ref 075 Table 7.5-1

using cyclone + elec pre on discharge and cyclone + wet scrub on windbox the emission factor may be reduced to $.04 + .11 = .15 \text{ lb/TON}$

$$\therefore E_N = .15 \text{ lb/TON sinter produced}$$

CO & SO_x

Controls for SO_x & CO were not identified in the literature. It is not likely that these pollutants are presently being controlled from this source. We make the assumption that max impact (hypothetical) would be attained by using $E_N = 0$

$$\therefore E_N (\text{SO}_x) = 0.0$$

$$E_N (\text{CO}) = 0.0$$

assumed
for
max hypoth
impact

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 1-8-75

Source Iron and Steel Industry

E_N : cont'd

Scarfing

Particulate

Ref 047 pg 222 efficiency stated = 90%

by settling chamber, high energy scrubber, or electrostatic precipitator.

While no other info was obtain we don't see any reason why this type of control system should be able to achieve 99.0%.

$$E_N = .01 \times 3 \text{ lb/TON}$$

$$E_N = .03 \text{ lb/TON} \text{ steel scuffed}$$

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Calculations Done By W. Massone Date 1-8-75

Source Iron and Steel Industry

E_{III}d :

It is assumed that existing plants are able to apply control system so as to achieve the same degree of control as specified by E_N for Trace metals and Fluorides. [all categories]

Blast Furnace : $E_{III}d \text{ (Trace Metal)} = .026 \text{ lb/Ton pig iron produced}$

BOF : $E_{III}d \text{ (Trace Metal)} = .0038 \text{ lb/Ton raw steel produced}$
 $E_{III}d \text{ (Fluoride)} = .00015 \text{ lb/Ton raw steel produced}$

EAF : $E_{III}d \text{ (Trace metal)} = .028 \text{ lb/Ton raw steel produced}$
 $E_{III}d \text{ (Fluoride)} = .0044 \text{ lb/Ton raw steel produced}$

Open Hearth : $E_{III}d \text{ (trace metal)} = .014 \text{ lb/Ton raw steel}$
 $E_{III}d \text{ (Fluoride)} = .005 \text{ lb/Ton raw steel}$

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Computation Sheet For Emission Factors

Calculations Done By W. Marwone Date 1-8-75
 Source Iron and Steel Industry

E_s : Ref 148 § 084

SO_x emissions relating to sintering do not exist
 so that we may assign $E_s = E_u$ for this pollutant.

$$\text{Sintering} \quad E_s(\text{SO}_x) = 3 \text{ lb/TON sinter produced}$$

Similarly Carbon Monoxide regulation do not exist which may
 be applied to Iron & Steel plant processes so that $E_s = E_u$

$$\text{Blast Furnace: } E_s(\text{CO}) = 1.750 \text{ lb/TON pig iron produced}$$

$$\text{BOF : } E_s(\text{CO}) = 139 \text{ lb/TON raw steel produced}$$

$$\text{EAF : } E_s(\text{CO}) = 18 \text{ lb/TON raw steel produced}$$

$$\text{Sintering : } E_s(\text{CO}) = 44 \text{ lb/TON sinter produced}$$

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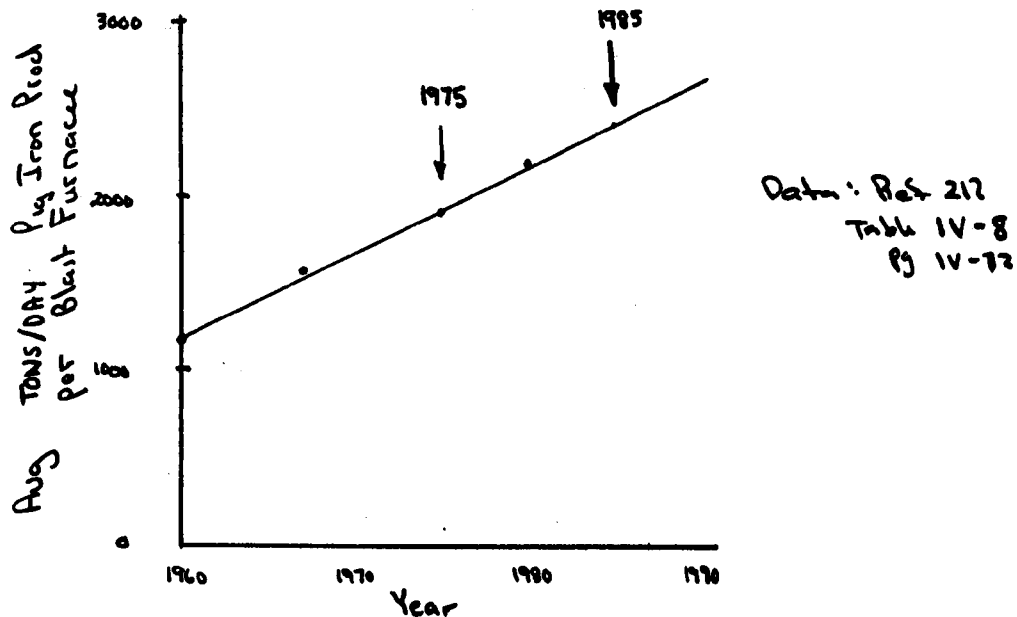
Computation Sheet For Emission Factors

Calculations Done By A. Marrone Date 1-9-75
 Source Iron and Steel Industry

E_s : cont'd

Particulate

Blast Furnace :



According to reference 212 pg IV-12 the average blast furnace pig iron production is increasing due to the replacement of smaller units with larger units in a way such that the total number of units decreases.

We assume the value of $\approx 2400 \frac{\text{Tons}}{\text{day}}$ in 1985 as the representative blast furnace size. and $24 \frac{\text{hr}}{\text{day}}$ schedule

$$\text{Avg Pig Iron Prod per Furnace} = 2400 \frac{\text{Tons}}{\text{day}} \cdot \frac{\text{day}}{24 \text{hr}} = 100 \frac{\text{Tons}}{\text{hr}}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Mannone Date 1-9-75
 Source Iron and Steel Industry

E_g : cont'd Particulate

Blair Furnace Ref 075 pg 7.5-1

To produce 1 unit of pig iron requires:

- (1) 1.55 units iron bearing charge
- (2) .55 " coke
- (3) .20 " limestone
- (4) 2.3 " air

excluding the weight of air the total solids input rate is

$$(1.55 + .55 + .20) / 1.0 = 2.3 / 1.0 = 2.3 \text{ TONS Input/TON} \begin{matrix} \text{pig} \\ \text{iron} \\ \text{produced} \end{matrix}$$

Therefore for an average furnace with 100 $\frac{\text{TON}}{\text{hr}}$ of Pig iron production implies a process weight rate of $2.3 \times 100 = 230 \frac{\text{TON}}{\text{hr}}$ input

$$\text{PWR} = 230 \frac{\text{TON}}{\text{hr}} \text{ or } 460,000 \text{ lb/hr}$$

$$\text{Ref 236 pg VI-11} \quad \text{Avg ACF/TON Pig Iron} = \frac{110+150}{2} = 130,000 \frac{\text{ACF}}{\text{TON}}$$

$$\text{using the average stack temp } 350 + 540 = 445^\circ\text{F}$$

$$130,000 \frac{\text{ACF}}{\text{TON}} \times \frac{704460}{445+460} = 130,000 \times \frac{530}{905} \approx 76,100 \frac{\text{SCF}}{\text{TON}} \begin{matrix} \text{pig} \\ \text{iron} \end{matrix}$$

$$\text{Use density SCF} = .075 \frac{\text{lb}}{\text{SCF}} \Rightarrow \approx 5700 \frac{\text{lb Slue gas}}{\text{TON Pig iron}}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marzzone Date 1-9-75

Source Iron and Steel Industry

E_g : cont'd Particulates

Blast Furnace

From Ref 212 pg III-21 Table III-24

Pig Iron Production by State (1967)

<u>State</u>	<u>Production</u> 10^6 TONS
NY	6.172
Penn	20.542
Maryl, W Va, Kent, Tenn, Tex	10.824
Ala	4.290
Ohio	14.485
Ind	12.167
Ill	6.309
Mich, Minn	7.439
Colo, Utah, Calif.	4.756

For the production data applied to several states we refer to Table III-23 at the above reference and ratio the production for all states by the number of furnaces on blast in the same year.

	<u>No Furn</u>	<u>% at Fur</u>	<u>Production Applied to State</u> (From 10.824×10^6 TON)
Maryland	10	55.6	6.02
W Va	4	22.2	2.40
Kent	2	11.1	1.20
Tenn	0	0	—
Tex	2	11.1	1.20
	18	100% ✓	$\Sigma = 10.82 \checkmark$

Mich	9	90.0	6.70	(From 7.439×10^6 ton)
Minn	1	10.0	.74	
	10		$\Sigma = 7.44 \checkmark$	

Colo	4	36.4	1.73	(From 4.756×10^6 ton)
Utah	3	27.2	1.29	
Calif	4	36.4	1.73	$\Sigma = 4.75 \checkmark$

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Computation Sheet For Emission Factors

Calculations Done By W. Massone Date 1-9-75
 Source Iron and Steel Industry

E_s : cont'd Particulate

Blast Furnace

The amended state distribution is as follows: also presented are the allowable state emissions from Ref 148 & 084 and the weighted allowable based on applying the fraction state capacity (A_i)

<u>State</u>	<u>Prod 10⁶ TONS</u>	<u>A_i</u>	<u>$E_{Allowable}$ lb/TON_{prod}</u>	<u>$A_i \times E_{allow}$</u>
New York	6.11	.071	.63	.0447
Penn	20.54	.236	.52	.1227
Mary	6.02	.069	.60	.0414
W Va	2.40	.028	.45	.0126
Kent	1.20	.014	.60	.0084
Tenn	—	—	—	—
Tex	1.20	.014	1.20	.0168
Ala	4.29	.049	.60	.0294
Ohio	14.49	.167	.60	.1002
Ind	12.17	.140	.60	.0840
Ill	6.31	.073	.45	.0329
Mich	6.70	.077	.58	.0447
Minn	.74	.009	.60	.0054
Colo	1.73	.020	.41	.0082
Utah	1.29	.015	22.50 +	.3375
Calif	1.73	.020	.40 ++	.0080
	$\Sigma 86.98$	$\Sigma = 1.002$ OK		$\Sigma = .897$

Rounding error

$$E_s = \sum E_{allow} \times A_i$$

$$E_s = .897 \text{ lb/ton}$$

+ 85% control required
 # Ref 156 pg 37

.9 < 1.5

$E_s < E_N$

which implies that state Reg are more stringent than apparent technology. We assume $E_s = E_N$ as the best that states can do

NOTE

∴ $E_s \cong .90 \text{ lb/ton}$ pig iron produced
 1.50

see next page

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 1-9-75

Source Iron and Steel Industry

E_g : cont'd Particulates

Blast Furnace:

We see that the value of E_g (.90 lb/ton) is lower than E_N (1.5 lb/ton) which implies that existing state regulations are more stringent than available control technology for this source may attain.

The efficiency of 99% which was used to obtain E_N would have to be improved to $\frac{150 - .9}{150} \times 100 = 99.4\%$

Existing control consist of a cyclone, wet scrubber, and electrostatic precipitator in series and unless control designs are improved this source may find it difficult to meet existing state regulations.

We have taken the position that states could not enforce a level lower than present technology and therefore $E_g = E_N$ (for particulates)

$$E_g = 1.5 \text{ lb/ton} \text{ pig iron produced}$$

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Source Iron and Steel Industry

E_g : cont'd
Particulate

Sintering: since most sintering is done at the same location and related directly to Blast Furnace operation we assume that the distribution of pig iron production by state will approximate the distribution of sinter. \therefore Use A_i from Blast Furnace

Ref 212 pg V-5 Plant Cap range 2000 - 6000 $\frac{\text{TON}}{\text{day}}$ sinter

Taking the avg = 4000 $\frac{\text{TON}}{\text{day}}$ and an using $24 \frac{\text{hr}}{\text{day}}$

$$\text{PWR} = 4000 \frac{\text{TON}}{\text{day}} \times \frac{\text{day}}{24 \text{hr}} = 167 \frac{\text{TON}}{\text{hr}} \text{ or } 330,000 \frac{\text{lb}}{\text{hr}}$$

Ref 236 p VI-13 1.7 - 2.3 scfm / lb/hr exhaust rate for
 avg 2.0 scfm/lb/hr sintering rate

therefore the exhaust from a typical
 167 $\frac{\text{TON}}{\text{hr}}$ sintering unit is $2 \frac{\text{scfm}}{\text{lb/hr}} \times 330,000 \frac{\text{lb}}{\text{hr}} = 660,000 \text{ scfm}$

Allowable emission are obtained from Ref 144 & 084 using the above information

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 1-9-75

Source Iron and Steel Industry

E_s : cont'd
Particulate

Sintering:

Based on
 Ref 148 & D84

<u>State</u>	<u>A_i</u>	<u>E_{Si} lb/ton sinter</u>	<u>$A_i \times E_{Si}$</u>
New York	.071	.359	.0255
Penn	.236	.677+	.1598
Maryl	.069	.339	.0234
W Va	.028	.245	.0069
Kent	.014	.335	.0047
Tex	.014	.659	.0092
Ala	.049	.335	.0164
Ohio	.167	.335	.0559
Ind	.140	.335	.0469
Ill	.073	.228	.0166
Mich	.077	.335	.0258
Minn	.009	.335	.0030
Colo	.020	.233	.0047
Utah	.015	6.300++	.0945
Calif	.020	.240	.0048

$$\Sigma = .4981$$

+ using gr/sch limitation

++ using 85% control required

$$E_s = \sum E_{Si} \times A_i$$

∴

$$E_s \cong .5 \text{ lb/TON sinter produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marone Date 1-9-75
 Source Iron and Steel Industry

E_s : cont'd

Particulate

BOF

Ref 044 pg 50 typical BOF furnace size 250 TON

This unit is characterized as producing
 $470 \frac{\text{TON}}{\text{hr}}$ raw steel and a average
 exhaust flow of 375,000 scfm

New Source Standard for BOF from Ref 044 pg 49

.022 gr/scf

$$\frac{375,000 \frac{\text{scf}}{\text{min}} \times .022 \frac{\text{gr}}{\text{scf}} \times \frac{60 \text{ min}}{\text{hr}} \times \frac{1\text{b}}{7000 \text{ gr}}}{470 \frac{\text{TON}}{\text{hr}}} = .15 \frac{\text{lb}}{\text{TON raw steel}}$$

Ref 212 pg IV-4 gives the 1980 distribution of raw steel production which we will use for a distribution of BOF production in determining E_s .

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Computation Sheet For Emission Factors

Calculations Done By W. Mannone Date 1-9-75
 Source Iron and Steel Industry

E_s : cont'd Particulate

BOF

	<u>Prod 10⁶ TONS</u>
Chicago, Ill	37.8
Pitt, Pa	31.1
N.E. Coast	24.5 ← (assume mainly Pa let 212 Fig III-4)
Youngstown, Pa	12.5
Detroit, Mich	16.9
Western	11.7 ← (Assume 50% Calif., 15% Utah, 20% Colo, 15% Ariz)
Southern	11.7 ← (assume 28% Ala 8% each of
Clev. Ohio	10.3
Buffalo, N.Y.	10.3
Cinn, Ohio	8.9
St. Louis, Mo	4.3
	Fig III-10
	Va Fla NC Miss Tenn Tex SC Okla Geo

Using the above we may redistribute the steel production by state as follows:

<u>State</u>	<u>Prod</u>	<u>A_i</u>
Ala	3.3	.0188
Ill	37.8	.2154
Pa	68.1	.3980
Mich	16.9	.0963
Calif	5.9	.0336
Utah	1.8	.0103
Colo	2.3	.0131
Ariz	1.8	.0103
Va	.9	.0051
NC	.9	.0051
Tenn	.9	.0051
SC	.9	.0051
Geo	.9	.0051
Fla	.9	.0051
Miss	.9	.0051
Tex	.9	.0051
Okla	.9	.0051
Ohio	19.2	.1094
NY	10.3	.0587
	<hr/>	
	175.5	

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Calculations Done By W. Marrone Date 1-9-75

Source Iron and Steel Industry

E_s : cont'd BoF

assuming that the BoF will be distributed in the same proportion as raw steel production we will develop a value for E_s by weighting each state E_{sc} by the value of A_i

E_{sc} is obtained from Ref 148 and 084 using a process weight rate of 470 TON/hr or 940,000 lb/hr - call it 1000,000 lb/hr

<u>State</u>	<u>Allowable lb/hr</u>	<u>Allow x Ai</u>
Ala	46.8	.880
Ill	69.0	14.863
Pa	47.4	18.391
Mich	69.0	6.645
Calif	40.0	1.344
Utah	3600.0 +	37.080
Colo	46.7	.612
Ariz	69.0	.711
Va	69.0	.352
NC	69.0	.352
Tenn	69.0	.352
SC	69.0	.352
Geo	262.0	1.336
Fla	46.7	.238
Miss	264.0	1.346
Tex	150.0	.765
Okla	69.0	.352
Ohio	69.0	7.549
NY	71.1	4.168

$$E_s' = \sum = 97.69 \text{ lb/hr}$$

+ 85% control required

$$E_s = \frac{97.69 \text{ lb/hr}}{470 \text{ TON/hr}} \approx .208 \text{ lb/TON raw steel produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marone Date 1-9-75

Source Iron and Steel Industry

E_s & cont'd

Particulate

BOF

The value of E_s determined above is representative of existing BOF facilities whereas the NPS standard developed is for all new facilities.

$$E_s^{pt} (\text{Existing}) = .208 \text{ lb/ton raw steel produced}$$

$$E_s^{pt} (\text{New}) = .15 \text{ lb/ton raw steel produced}$$

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Source Iron and Steel Industry

E_s : cont'd Particulate

[E.A.F]

On the following page a graph was developed from data in ref 250 to permit an estimation of the average furnace size.

We estimate 40 Ton of Cap as representative of the average existing plant

New plants according to the same ref have been on the order of 300 Ton cap

In order to evaluate this industry both for existing and new plants we will use the following info

- ① Existing Plant 40 Ton cap
- ② New Plant 300 Ton Cap
- ③ Furnace cycle 5 hrs (within range 3.5-7 Ref pg 49)
- ④ 4500 scfm/Ton cap exhaust flow (avg value p 49)

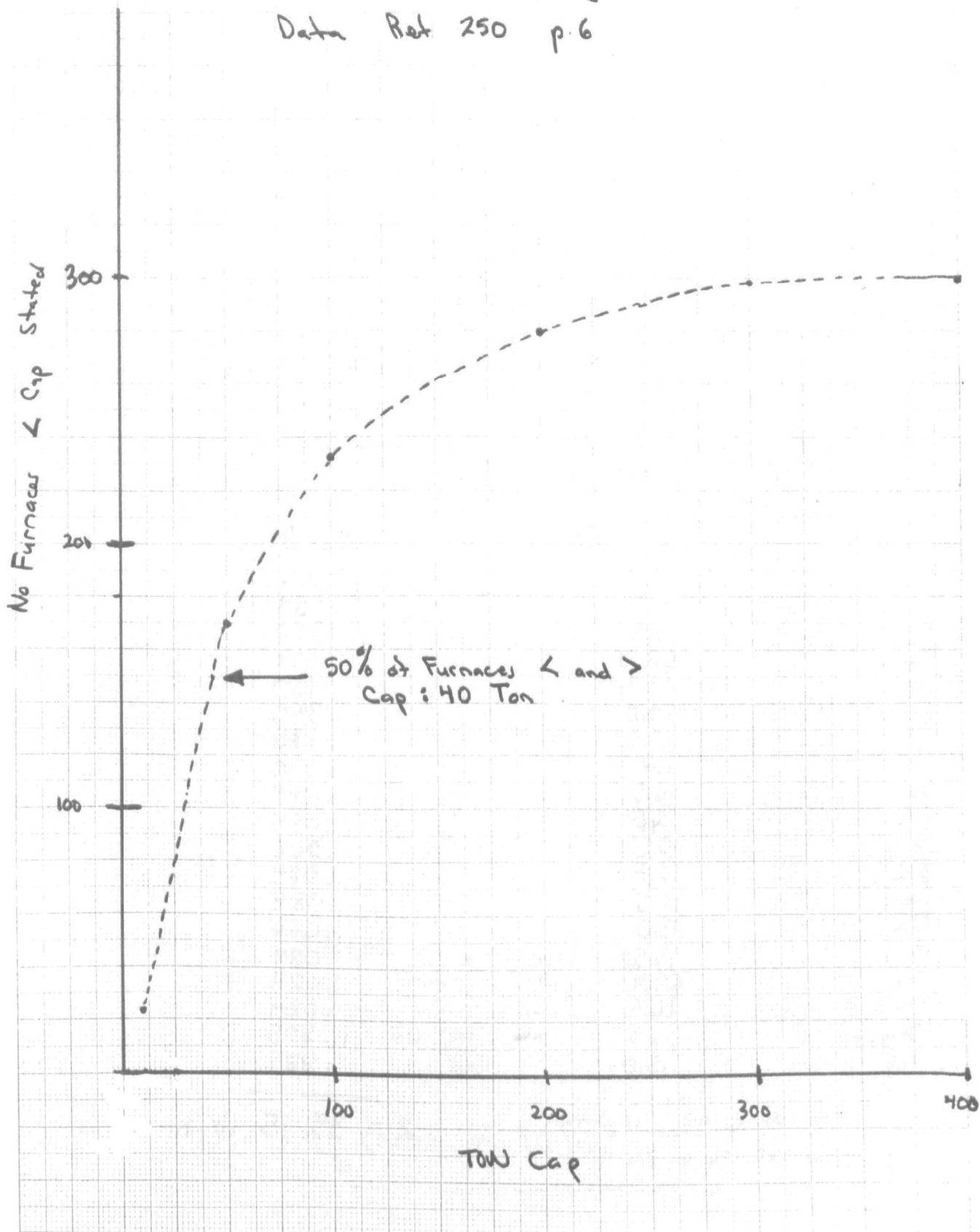
$$\text{PWR existing} = \frac{40}{5} = 8 \frac{\text{Ton}}{\text{hr}} \text{ or } 16,000 \text{ lb/hr}$$

$$\text{PWR now} = \frac{300}{5} = 60 \frac{\text{Ton}}{\text{hr}} \text{ or } 120,000 \text{ lb/hr}$$

EAF

Determination of Avg EAF

Data Rat 250 p.6



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Calculations Done By W.J. Marrone Date 1-9-75

Source Iron and Steel Industry

E_s : cont'd Particulate

EAF.

Assuming the EAF production to be distributed in the same manner as raw steel production then we may use the A_i value presented on pg 32 of the "Emission Factors"

Exit reg are developed from Ref 148 and 084 using the above information

<u>State</u>	<u>Allow lb/hr</u>	<u>$A_i \times Allow$</u>
Ala	13.0	.244
Ill	16.5	3.554
Pa	30.8 +	11.950
Mich	162.0	15.601
Calif	13.7 ++	.460
Utah	34.0	.350
Colo	13.0	.170
Ariz	16.5	.170
Va	16.5	.084
NC	16.5	.084
Tenn	13.0	.066
SC	16.5	.084
Geo	16.5	.084
Fla	13.0	.066
Miss	16.5	.084
Tex	24.4	.124
Oklas	16.5	.084
Ohio	16.5	1.805
NY	15.0	.881

+ using gr/sct regulation

++ Ref 156 pg 37

$$\sum = 35.95 \text{ lb/hr}$$

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Computation Sheet For Emission Factors

Calculations Done By 2). Marone Date 1-9-75
 source Iron and Steel Industry

E_s : cont'd Particulate

EAF

$$E_s' = \frac{35.95 \text{ lb/hr}}{8 \text{ TON/hr}} = 4.49 \text{ lb/TON}$$

We use this factor to be E_s (existing) for EAF

$$E_s \text{ (existing)} = 4.49 \text{ lb/TON raw steel produced}$$

For new sources at 300 TON cap $\frac{60 \text{ TON}}{\text{hr}}$ and
 NSPS of .0052 gr/sec [Ref 250 pg 13]

$$\text{Using } 4500 \frac{\text{scfm}}{\text{TON cap}} \times 300 \text{ TON} = 1,350,000 \text{ scfm}$$

$$\frac{.0052 \frac{\text{gr}}{\text{sec}} \times 1,350,000 \frac{\text{scf}}{\text{min}} \times \frac{60 \text{ min}}{\text{hr}} \times \frac{16}{7000} \frac{\text{lb}}{\text{gr}}}{60 \frac{\text{TON}}{\text{hr}}} = 1.00 \frac{\text{lb}}{\text{TON}}$$

$$\therefore E_s \text{ (new)} = 1.0 \text{ lb/TON raw steel produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marzzone Date 5-9-75

Source Iron and Steel Industry

E_s : cont'd Particulate

Open Hearth Furnace

Ref 212 pg II-10 suggest a range of typical plant size for open hearth furnaces 100-200 TON Head

Ref 234 pg 3-83 describes a typical 200 TON Head furnace which we will use to develop E_s assuming it to be close to the average.

200 Ton input

Gases N_2 180 TONS

155 TON steel output

O_2 5 TONS

CO_2 75 TONS

260 TONS

$$\frac{260 \times 2000}{155} = 3355 \frac{\text{lb gas}}{\text{TON raw steel}}$$

It will be assumed that the distribution (A_i) of raw steel production appearing on pg 32 of "Emission Factors" may be used to represent O.H. furnace distribution

Ref 084 & 148

only Mich and Pa (of the states with O.H. furnace prod) have regulations specific to this source

Mich .15 lb/1000 lb gas

Pa .81 lb/ton steel [from allowable $\frac{\text{lb/hr}}{\text{hr}} = .76 (F \times w)^{.42}$]

where $F = 20 \text{ lb/ton}$

$w = 20 \frac{\text{Ton input}}{\text{hr}}$

$hr = 10 \text{ hrs}$

$$\frac{\text{lb/hr}}{\text{hr}} \times \text{hr} = \frac{\text{lb}}{\text{ton steel}}$$

All other states will be assumed to control particulate by application of the process weight rate table.

$$\frac{200 \text{ TON}}{10 \text{ hr}} = 20 \frac{\text{TON}}{\text{hr}} \text{ or } 40,000 \frac{\text{lb}}{\text{hr}} \text{ process weight rate}$$

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Calculations Done By W. Marzzone Date 5-9-75

Source Iron and Steel Industry

E_j: cont'd Particulate

Open Hearth Furnace

<u>State</u>	<u>A_i</u>	<u>Allowable lb/Ton</u>	<u>Allow × A_i</u>
Ala	.0188	1.45	.0273
Ill	.2154	.84	.1809
Pa	.3880	.81	.3143
Mich	.0963	.50	.0482 +
Calif	.0336	1.93	.0615 ++
Utah	.0103	2.55	.0263 *
Colo	.0131	1.48	.0194
Ariz	.0103	1.93	.0199
Va	.0051	1.93	.0098
NC	.0051	1.93	.0098
Tenn	.0051	1.48	.0075
SC	.0051	1.93	.0098
Geo	.0051	1.93	.0098
Fla	.0051	1.45	.0074
Miss	.0051	1.93	.0098
Tex	.0051	3.87	.0197
Oklahoma	.0051	1.93	.0098
Ohio	.1094	1.93	.2111
NY	.0587	1.74	.1021

$$\sum = 1.1044 \text{ lb/ton steel}$$

+ based on allowable .15 lb/1000 lb gas.
 and 3355 lb gas/ton steel

** based on 85% control required

++ use allowable table for LARPCD per 156 pg 37

$$\therefore E_S = 1.1 \text{ lb/ton raw steel}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 1-10-75
 Source Iron and Steel Industry

E_s : cont'd Particulate Scarfing

Information regarding a representative size scarfing operation could not be obtained from the literature nor was there sufficient information to directly determine the size.

We will make a rough estimate by making the following assumptions:

1975	(1) BOF	avg size	470 TON/HR	pg 31 "Emission Factors"
	(2) EAF	" "	8 TON/HR	pg 35 "Emission Factors"
	(3) OHF	" "	15 TON/HR	pg 10 Ref 212
	(4)	$\frac{88.9}{174.5} \approx .51$	or 51% of cap by BOF	
	(5)	$\frac{36.7}{174.5} \approx .21$	or 21% of cap by EAF	
	(6)	$\frac{48.9}{174.5} \approx .28$	or 28% of cap by OHF	

weighting the representative sizes by the capacities we may obtain an estimate of "typical" furnace size.

$$\begin{aligned} \text{Avg Size} &= (.51 \times 470) + (.21 \times 8) + (.28 \times 15) \\ &= 239.7 + 1.68 + 4.2 \end{aligned}$$

$$\text{Avg Size} = 245 \text{ TON/HR}$$

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Computation Sheet For Emission Factors

Calculations Done By 2). Mannone Date 1-10-75

Source Iron and Steel Industry

E_s : cont'd Particulate

Scarfing

Using 245 TON/HR and assuming that an average plant will have 3 such furnaces and also that 50% (see pg 6 "Industrial Factors") of raw steel produced is scuffed.

$$.5 \times (3 \times 245) \approx 368 \frac{\text{TON}}{\text{HR}} \text{ for scarfing operation}$$

$$\text{PWR} = 368 \frac{\text{TON}}{\text{HR}} \text{ or } 736,000 \text{ lb/HR}$$

Scarfing will be assumed to occur in the same states as previously identified for steel production and distributed according to the value of A_i (pg 32 "Factor") used in developing E_s factors for the BOF + EAF

Allowable State emission for this process weight rate will be obtained from Part 148 + 084 by applying the Particulate Process Weight Rate Tables. State allowable levels will be weighted by A_i to determine an overall E_s

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Computation Sheet For Emission Factors

Calculations Done By W. Marzzone Date 1-10-75

Source Iron and Steel Industry

E_s : cont'd

Particulates Scarfing

<u>State</u>	<u>Allowable lb/hr</u>	<u>$A_i \times A_{allow}$</u>
Ala	65.0	1.222
Ill	57.0	12.278
Pa	32.0	12.416
Mich	65.0	6.260
Calif	40.0	1.344
Utah	166.0+	1.710
Colo	44.5	.583
Ariz	65.0	.670
Va	65.0	.332
NC	65.0	.332
Tenn	46.0	.235
SC	65.0	.332
Geo	210.0	1.071
Fla	45.5	.232
Miss	210.0	1.071
Tex	137.0	.699
Okla	65.0	.332
Ohio	65.0	.7.111
NY	68.0	<u>3.992</u>
$\Sigma = 50.9$		lb/hr

+ 15% control required

$$E'_s = 50.9 \text{ lb/hr}$$

$$E_s = \frac{50.9 \text{ lb/hr}}{368 \text{ Ton/hr}} = .138 \text{ lb/TON}$$

$\therefore E_s = .138 \text{ lb/TON}$ steel scarfed

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 5-9-75

Source Iron and Steel Industry

E_s cont'd Ref 148 & 084

Fluoride emission regulations do not exist for Iron & Steel plants and strictly speaking the value of E_s should be assumed equal to E_u . The fluoride emissions occur from the steelmaking furnaces. [BOF, EAF, DHF]

Both gaseous and solid fluoride compounds are emitted

For BOF all solid CaF_2 emission 100% of F in solid

For EAF 83.5% of F in solid form
 16.5% of F in gaseous form

by default particulate emission regulations will be, in effect stipulating a fluoride regulation; at least for the solid portion of the total fluoride emission. The gaseous fluoride is assumed to be permitted uncontrolled.

Ref 234 in pg 3-81 & 3-83 all F emitted from OH furnace
 " in gaseous HF form

$$\therefore E_s = E_u = .102 \text{ lb/tow steel}$$

Open Hearth Furnace

$$E_s(F) = .102 \text{ lb/tow raw steel}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marzzone Date 1-10-75

Source Iron and Steel Industry

E_s : cont'd

Fluoride

BOF

From pg 2 "Emission Factors"

$$E_u(F) = .049 \text{ lb/ton}$$

$$E_u(Pt) = 51 \text{ lb/ton}$$

Fluoride allowable E_s will be determined by ratiosing the $E_u(F)$ to $E_u(Pt)$ and applying this to $E_s(Pt)$ to obtain $E_s(F)$

$$E_s(F)(exit) = \frac{.049}{51} \times E_s^{Pt}(exit) = \frac{.049}{51} \times .208 \approx .0002 \text{ lb/ton}$$

$$E_s(F)_{New} = \frac{.049}{51} \times E_s^{Pt}_{New} = \frac{.049}{51} \times .15 \approx .00015 \text{ lb/ton}$$

assumed = to E_u
 instead of .000144

EAF

$$E_u(F) = .069$$

$$E_u(Pt) = 28.3$$

83.5% is solid and will be applied under E_s
 16.5% is gas and is consider uncontrollable

$$E_s = .165 (.069) + \frac{.069 (.835)}{28.3} [E_s^{Pt}]_{exit}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marzone Date 1-10-75
 Source Iron and Steel Industry

E_s cont'd

Fluoride

EAF

$$E_s(F) \text{ new} = .165(.069) + \frac{.069(.835)}{28.3} [1.0]$$

$$E_s(F) \approx .0114 + .0020$$

$$E_s(F) \text{ new} \approx .0134 \text{ lb/ton raw steel produced}$$

$$E_s(F)_{\text{exist}} = .165(.069) + \frac{.069(.835)}{28.3} [4.49]$$

$$= .0114 + .0091$$

$$E_s(F)_{\text{exist}} \approx .021 \text{ lb/ton raw steel produced}$$

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Computation Sheet For Emission Factors

Calculations Done By 2. Morone Date 1-10-75
Source Iron and Steel Industry

E_s : cont'd

Trace Metals

Trace metals will be treated similarly as fluoride since particulate reg will be default limit the allowable trace metal emissions

Blast Furnace

$$E_s(\text{Trace Met}) = \frac{E_u(\text{TM})}{E_u(\text{PT})} \times E_s(\text{PT})$$

$$E_s(\text{TM}) = \frac{2.63}{150} \times .90 \approx .016 \text{ lb/ton} \text{ pig iron produced}$$

This value E_s is less than E_N and may be discussed similarly as was the $E_s < E_N$ for particulate "Emission Factor" pg 28

$$E_s(\text{Trace Metal}) = .016 \text{ lb/ton} \text{ pig iron produced}$$

Note $E_s < E_N$ and as discussed on pg 27 & 28 of Emission Factor we set $E_s = E_N$

$$E_s(\text{TM}) = .026 \text{ lb/ton} \text{ pig iron produced}$$

BOF

$$E_s(\text{TM})_{\text{exist}} = \frac{E_u(\text{TM})}{E_u(\text{PT})} \times E_s(\text{PT})_{\text{exist}}$$

$$E_s(\text{TM})_{\text{exist}} = \frac{1.26}{51} \times .208 = .0051 \text{ lb/ton} \text{ raw steel produced}$$

$$E_s(\text{TM})_{\text{New}} = \frac{1.26}{51} \times E_s(\text{PT})_{\text{New}}$$

$$E_s(\text{TM})_{\text{New}} = \frac{1.26}{51} \times .15 = .0037 \text{ lb/ton}$$

This is assumed equivalent to E_N since it differs only slightly

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 1-10-75

Source Iron and Steel Industry

E_s cont'd

Trace Metals

BOF

$$E_s(\text{Trac Met})_{\text{New}} = .0038 \text{ lb/tow raw steel produced}$$

EAF

$$E_s(\text{TM})_{\text{exist}} = \frac{E_u(\text{TM})}{E_u(\text{PT})} \times E_s(\text{PT})_{\text{exist}}$$

$$E_s(\text{TM})_{\text{exist}} = \frac{1.67}{28.3} \times 4.49 \approx .265 \text{ lb/tow raw steel produced}$$

$$E_s(\text{TM})_{\text{now}} = \frac{1.67}{28.3} \times E_s(\text{PT})_{\text{now}}$$

$$E_s(\text{TM})_{\text{now}} = \frac{1.67}{28.3} \times 1.00 \approx .059 \text{ lb/tow raw steel produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marone Date 5-9-75

Source Iron & Steel Industry

E_s cont'd

Trace Metals

OHF

$$E_s(\text{TM}) = \frac{E_u(\text{TM})}{E_u(\text{P4})} \times E_s(\text{P4}) = \frac{.275}{17} \times 1.1 \cong .018 \frac{\text{lb}}{\text{ton}}$$

$$E_s(\text{TM}) = .018 \frac{\text{lb}}{\text{ton raw steel}}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Mannone Date 1-8-75

Source Iron and Steel Industry

Summary of Emission Factors

Category	Units	Pollutant					<u>Comments</u>
			V	E _N	E _N	E _{Ind}	
Blast Furnace	lb./TON pig iron produced	Pt	150	1.5			1.5
		CO	1.750	0.0			1.750
		TH met	2.63	.026	.026	.026	
Basic Oxygen Furnace	lb./TON raw steel produced	Pt	51	.15			.208 Exist .15 New
		CO	139	3			139
		TH met	1.26	.0038	.0038	.0038	.0051 Exist .0038 New
		F	.049	.00015	.00015	.00015	.0002 Exist .00015 New
Electric Arc Furnace	lb./TON raw steel produced	Pt	28.3	.48			4.49 Exist 1.00 New
		CO	18	4.5			18
		TH met	1.67	.028	.028	.028	.265 Exist .059 New
		F	.069	.0044	.0044	.0044	.021 Exist .0134 New
Open Hearth Furnace	lb./TON raw steel produced	Pt	17	.85			1.1
		TH	.275	.014	.014	.014	.018
		F	.102	.005	.005	.005	.102
Sintering	lb./TON sinter produced	Pt	42	.15			.50
		SO ₂	3.0	0.0 *			3.0
		CO	44	0.0 *			44
Scouring	lb./TON steel scoured	Pt	3.0	.03			.138

* set equal to zero for max hypothetical
emission NSPS impact evaluation

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Industrial Factors

Calculations Done By J. Marrone Date 12-2-74

Source Primary Lead

P_c : Ref 030 pg 6.3-5

"The overall growth rate in lead consumption for the United States is estimated at 2-3% per year between 1971 and 1975. It is doubtful that this rate will change appreciably after 1975."

This statement clearly indicated a growth in lead consumption i.e. production however,

pg 6.3-5 also states

"It is doubtful that any new smelters will be constructed in the United States in the near future since the industry as a whole is currently under utilized with regard to pig lead production capacity and the domestic consumption of lead appears to be relatively stable. Whereas, it is true that some pig lead is imported, it is doubtful that domestic lead industry would add additional capacity in order to try to capture the market share that is currently going to foreign markets."

We interpret this to mean that production capacity will not be growing, yet that production will which is equivalent to saying that growth will be taken up by a increasing value of K.

With this in mind we assume the following:

$$\textcircled{1} \ A \text{ in } 1975 = A \text{ 1985}$$

$$\textcircled{2} \ P_c = 0$$

$$\textcircled{3} \ K \text{ increasing from 1975 to 1985}$$

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Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 12-2-74

Source Primary Lead

P_c :

Since capacity growth in the near future appears nil
 and we assume this covers the period 1975-1985
 we have;

$$P_c = 0$$

A : Ref 030 Table 6-23 pg 6.3-2

Capacity 1971 = 765,000 TONS/yr

Assuming no new capacity between 1971 and 1975
 then

Capacity 1975 = 765,000 TON/yr Lead produced

∴

$$A = .765 \times 10^6 \text{ TON/yr Lead produced}$$

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Computation Sheet For Industrial Factors

Calculations Done By J. Marrone Date 12-2-74
 Source Primary Head

K: Ref 030 pg 6.3-1

Capacity utilization in 1971 was 85% for the industry as a whole (four producers) however, two of the four producers were operating at 99% utilization of capacity.

We make the assumption that the existing value of K (1975) is .85 and that by the year 1985 the value will increase to .99.

Note

This is a loose assumption especially since new capacity may at any time be developed, which would realize a growth in capacity ($P_c > 0$) and a decrease in K to some level significantly less than .99.

$$\therefore K_{\text{existing 1975}} = .85$$

$$K_{\text{new 1985}} = .99$$

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Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 12-2-74
 Source Primary Head

P_B : Ref 037 Section Class 33.2

Asset guideline period 14 yrs

P_B est \rightarrow 2xIRS = 28 yrs

100% depreciation \approx 3.6%/yr simple
 28 yrs

$$P_B = .036 \text{ simple}$$

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Emission Factors

Calculations Done By W. Marzzone Date 12-3-74
 Source Primary Lead

Particulate

E_u : Ref 049 Table 13-1

Particulate emissions from

- ① Sintering 520 lb/Ton Lead
- ② Blast Furnace 250 lb/Ton Lead
- ③ Cross Reverb Furnace 20 lb/Ton Lead

$$E_u = \Sigma E = 520 + 250 + 20 = 790 \text{ lb/Ton Lead}$$

$$\therefore E_u = 790 \text{ lb/Ton Lead}$$

E_N : Ref 075 Table 7.6-1 Note ⑥ gives collection efficiency for
 electrostatic precipitator 96%
 baghouse system 99.5%

Ref 029 Table 5.4 Plant # 6 uses a baghouse-spray system
 on the sintering, blast furnace & reverb furnace
 with 99.8% for sinter and 99.9% for blast & reverb
 as collection efficiency.

We assume a collection efficiency of 99.85%
 as attainable for lead smelters.

$$E_N = .0015 \times E_u = .0015 \times 790 \cong 1.19 \text{ lb/Ton Lead}$$

$$E_N = 1.19 \text{ lb/Ton Lead}$$

Project Number - 32391 New Source Performance Standards

Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-3-74

Source Primary Lead

Particulates cont'd

E_S :

Ref 029 Table 5-4 pg 5.3-3

Provides a distribution of capacity of lead smelting by state from which fractional capacity occurring in each state may be determined

<u>State</u>	<u>Capacity T/day Lead</u>	<u>A_i</u>
Idaho	350	.199
Montana	105	.060
Missouri	1152	.656
Texas	150	.085
		<u>$\Sigma = 1.000$</u> ✓

6 plants \approx 1757 Total T/D Lead

Average Plant size \approx 293 T/D Lead

A rough estimate of SCFM/T/D is obtained from the data in the Table (5.4) with the exclusion of Plant 4 since part of its scfm is dilution and ventilation air.

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Computation Sheet For Emission Factors

Calculations Done By J. Marrone Date 12-4-74
 Source Primary lead

Particulate cont'd

E_s :

Plant	SCFM/T/0
①	990
②	690
③	647
④	1602
⑤	3066

Aug = 1399

For a 293 T/0 lead smelting plant

$$\text{SCFM} = 293 \text{ T/0} \times 1399 \frac{\text{SCFM}}{\text{T/0}} = 409,907 \text{ SCFM}$$

A proposed limit of .022 gr/scf (Ref 029 pg 1.1-2) is used to calculate a value for E_s (new) particulate.

$$E = \frac{.022 \frac{\text{gr}}{\text{scf}} \times 409,907 \frac{\text{scf}}{\text{min}} \times 60 \frac{\text{min}}{\text{hr}} \times \frac{16}{7000} \frac{\text{gr}}{\text{lb}} \times \frac{24}{\text{day}}}{293 \text{ T/day}} = 6.33 \text{ lb/TON Lead}$$

$E_s(\text{new}) = \cancel{6.33} \text{ lb/TON Lead}$

3.06 lb/TON Lead

Note since $E_s(\text{existing}) < E_s(\text{new})$ the $E_s(\text{new}) = E_s(\text{existing})$

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Computation Sheet For Emission Factors

Calculations Done By J. Marrone Date 12-4-74
 Source Primary Lead

Particulates cont'd

E_s :

For existing plants the emission limit will be determined by applying the generalized process weight rate table for the four states having lead smelting capacity

Ref 075 Table 7.6-1 2 Ton ore \rightarrow 1 TON Lead

$$\begin{aligned} \text{Average Process Weight Rate} &= 293 \frac{\text{lb}}{\text{ton}} \times \frac{\text{day}}{24 \text{ hr}} \times 2 \frac{\text{ton}}{\text{lead}} \times \frac{2000 \frac{\text{lb}}{\text{ton}}}{\text{ton}} \\ &= 48,833 \text{ lb/hr input} \end{aligned}$$

Ref 084 § 148

<u>State</u>	<u>E_{si} lb/TON Lead</u>	<u>$A_i \times E_{si}$</u>
Idaho	2.87	.57
Montana	2.87	.17
Missouri	2.87	1.88
Texas	5.12	.44

$$E_s(\text{existing}) = \sum E_{si} \times A_i = 3.06 \text{ lb/TON lead}$$

$$\therefore E_s(\text{existing}) = 3.06 \text{ lb/TON lead}$$

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Computation Sheet For Emission Factors

Calculations Done By 2). Marone Date 4-29-75

Source Primary lead

$\text{SO}_x /$

E_u : Ref 015 Table 7.6-1 and Notes ① and ②

$$E = 660 \text{ lb/ton of ore conc}$$

2 ton ore \Rightarrow 1 ton metal lead

$$\therefore E_u = 660 \frac{\text{lb}}{\text{ton ore}} \times \frac{2 \text{ ton ore}}{1 \text{ ton lead}} = 1320 \text{ lb/ton lead}$$

$$\therefore E_u = 1320 \text{ lb/ton lead}$$

E_N : Ref 030 pg 7.5-17

best level of control achievable (98.5%) based on

- (1) utilization of recirculating sinter machine
- (2) scrubbing (DMA!) to weak SO_2 stream of blast furnace
- (3) DA acid plant to strong stream of sinter machine

$$E_N = .015 \times E_u = .015 \times 1320 = 19.8 \text{ lb/ton lead}$$

$$\therefore E_N = 19.8 \text{ lb/ton lead}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marzzone Date 4-29-75
 Source Primary Lead

~~SO_x~~ / cont'd

E_s :

Ref 148 § 084

None of the four states have existing regulation pertaining to SO_x emissions from non-ferrous smelters specifically.

In this case $E_s = E_u = 1320 \text{ lb/ton lead}$.

$$\therefore E_s = 1320 \text{ lb}^{\text{SO}_x}/\text{TON lead}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-3-74

Source Primary Lead

Lead

E_u : Ref 136 "Primary Lead Smelters"

$$E_u = 37 \text{ lb/ton Lead}$$

This is a second-hand value based on data from one test which we hope would be confirmed when more data is available.

E_N :

We will assume that the collection efficiency of lead will be identical as for the total particulate emission (99.8%)

$$\therefore E_N = .0015 \times E_u = .0015 \times 37 \text{ lb/ton Lead}$$

$$\therefore E_N = .056 \text{ lb/ton Lead}$$

E_{IIIId} : It is assumed that the fabric filters may be retrofitted on existing plants to achieve the same efficiency as for new plants

$$\therefore E_{IIIId} = E_N = .056 \text{ lb/ton Lead}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-3-75
 Source Primary Lead

~~head~~ / cont'd

E_s : Ref 48 § 084

Specific regulation for lead do not exist
 however we assume the lead may be
 controlled "de facto" via the particulate

PWR tables

$$E_s = \frac{37 \text{ lb Pb}}{790 \text{ lb Pt}} \times 3.06 = .14 \text{ lb/TON}$$

E_s
Lead

Lead

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-4-74
 Source Primary Lead

Summary of Emission Factors

lb/TON Lead

Pollutant	E _u	E _N	E _{ind}	E _s (existing)	E _s (new)
Particulate	790	1.19	/	3.06	3.06
SO _X	1320	19.8	/	1320	1320
Lead	37	.056	.056	.14	.14

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Computation Sheet For Industrial Factors

Calculations Done By 2) Marrone Date 12-2-74

Source Primary Zinc

P_c : Ref 030 pg 6.2-4

growth rate of 3.5% /yr cited for domestic consumption
 which we assume will coincide with growth in
 zinc capacity

$$\therefore P_c = .035 \text{ compound}$$

P_B : Ref 037 Class 33.2

Asset guideline period 14 yrs

$$P_B \text{ est} \rightarrow 2 \times \text{IRS} = 28 \text{ yrs}$$

$$\frac{100\% \text{ depreciation}}{28 \text{ yrs}} \approx 3.6\% / \text{yr simple}$$

$$\therefore P_B = .036 \text{ simple}$$

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Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 12-2-74
 Source Primary Zinc

K:

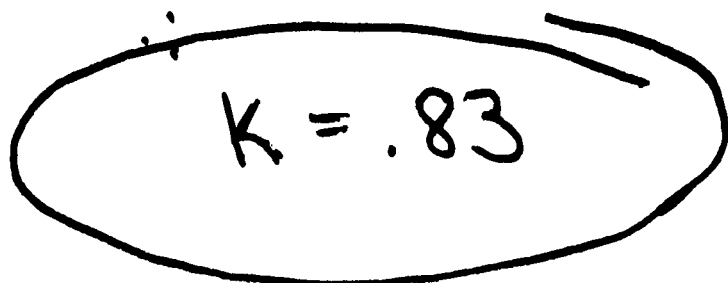
Ref 095 pg 233. 6833 under Note(3)

Capacity Primary Zinc Slab 1969 was 1,225,700 Tons/yr

Production Primary Zinc Slab 1969 was 1,020,900 Ton/yr

$$K = \frac{1,020,900}{1,225,700} \approx .83$$

We assume this value representative of primary zinc refineries



$$K = .83$$

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Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 12-2-74
 Source Primary Zinc

A :

Ref 030 pg 6.2-5

1975 total domestic demand = 1,450,000 Ton/yr

Slab zinc imports = 25% of Total

If we assume consumption = production then we calculate domestic production in 1975 as;

$$P_{75} = 1.45 \times 10^6 \frac{\text{Ton}}{\text{yr}} \times .75 \approx 1.088 \times 10^6 \frac{\text{Ton}}{\text{yr}}$$

$$\text{and employing } K = .83 \rightarrow A = \frac{P_{75}}{K} = \frac{1.088 \times 10^6}{.83}$$

$$A = 1.31 \times 10^6 \frac{\text{Ton}}{\text{yr}} \text{ zinc produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-3-74

Source Primary Zinc

Particulates

E_u : This factor will be developed by combining emissions from roasting, sintering, and distillation

Ref 049 Table 13-1 pg 262

90% of production is roasted with 75% in fluid-bed-suspension roasters
 15% in Multiple-hearth

60% of production is sintered

60% of production is distilled

Emission factors as presented in the Table will be weighted according to fraction of production for each operation

$$E = .75(2000 \frac{\text{lb}}{\text{TON zinc}}) + .15(333 \frac{\text{lb}}{\text{TON zinc}}) + .10(0) + .6(180) + .6(12) + .4(0) + .4(0)$$

$$E = 1500 + 49.95 + 0 + 108 + 7.2 + 0 + 0$$

$$E \approx 1665 \text{ lb/TON zinc produced}$$

∴

$$E_u = 1665 \text{ lb/TON zinc produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-3-74
 Source Primary Zinc

Particulate cont'd.

E_N : Ref 075 p7.7-1 cites the use of electrostatic precipitators and baghouse as acceptable devices for controlling zinc smelting plants

We assume that an efficiency of 99.85% (comparable to lead smelter) is applicable to zinc smelter

$$E_N = .0015 \times E_u = .0015 \times 1665 \approx 2.50 \text{ lb/TON}_{\text{Zinc}}$$

Note we see that the calculate value of E_s (new) on pg.— is less than E_N

$$E_s(\text{new}) = 2.25 \text{ lb/TON}$$

$$E_N \text{ calc.} = 2.50$$

This would at first imply that the NSPS proposed are stringent for this industry in requiring a control efficiency of about 99.9%. However, Ref 030 p7.5-21 and 7.5-26 indicates that the applicability of a baghouse system to zinc smelter should achieve 99.9% or better efficiency

$$\text{We therefore assume } E_N = .001 \times E_u \approx 1.67 \text{ lb/TON}$$

$$E_N = 1.67 \text{ lb/TON}$$

It is important to see that the existing and proposed regulations are closely controlling this industry.

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-3-74
 Source Primary Zinc

Particulars cont'd.

E_s : Ref 029 Table 5-3 gives the location and zinc smelting production (T/D) by state for the seven zinc producing smelters.

<u>State</u>	<u>Zinc Capacity T/D</u>	<u>A_i</u>
Texas	435	.207
Oklahoma	415	.197
Idaho	300	.143
Pa.	955	.453
		<u>$\sum = 1.000 \checkmark$</u>

Typical Plant Size

2105 T/D Total for industry of 7 plants

$$\frac{2105}{7} \approx 300 \text{ T/D zinc produced}$$

From the same table we calculate the SCFM/T/D zinc for the representative plants ④, ⑥, ⑦, ⑧

- ④ - 325 SCFM/T/D
- ⑥ - 764 SCFM/T/D
- ⑦ - 422 SCFM/T/D
- ⑧ - 480 SCFM/T/D

$$\left. \begin{array}{l} \\ \\ \\ \end{array} \right\} \text{Avg} \approx 498 \text{ SCFM/T/D zinc}$$

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-3-74

Source Primary Zinc

Particulates cont'd

E_s : For a 300 Tons zinc plant :

$$\text{SCFM} = 300 \frac{\text{T}/\text{hr}}{\text{hr}} \times 498 \frac{\text{SCFM}}{\text{T}/\text{hr}} = 149,400 \text{ SCFM}$$

E_s (new)

A proposed limit (Ref 029 p1.1-2) of .022 $\frac{\text{gr}}{\text{scf}}$ is used to calculate E_s (new) particulates.

$$E = \frac{.022 \frac{\text{gr}}{\text{scf}} \times 149,400 \frac{\text{scf}}{\text{min.}} \times 60 \frac{\text{min}}{\text{hr}} \times \frac{1\text{b}}{7000\text{gr}} \times \frac{24 \text{ hr}}{\text{day}}}{300 \frac{\text{Tons zinc}}{\text{day}}}$$

$$E_s(\text{new}) = 2.25 \frac{\text{lb}}{\text{ton zinc}}$$

E_s (existing)

For existing plants the emission limit will be set by using the process weight rate curve for the affected states.

$$\begin{aligned} \text{Process Weight Rate} &= 300 \frac{\text{Tons zinc}}{\text{hr}} \times \frac{2\text{hr}}{24\text{hr}} \times \frac{2 \frac{\text{Tons zinc}}{\text{Ton zinc}} \times 2000 \frac{\text{lb}}{\text{Ton}}}{\text{Ton}} \\ &= 50,000 \frac{\text{lb}}{\text{hr}} \text{ input} \end{aligned}$$

Ref 075
Table 7.7-1

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Calculations Done By W. Marrone Date 12-3-74

Source Primary Zinc

Particulate cont'd.

E_S :		E_{Si} lb/TON zinc	$A_i \times E_{Si}$
Ref 084	Texas	5.04	1.04
5/148	Oklahoma	2.84	.56
	Idaho	2.84	.41
	Penn.	4.09	1.85

$$E_S(\text{existing}) = \sum E_{Si} \times A_i \\ = 3.86 \text{ lb/TON zinc}$$

∴ $E_S(\text{existing}) = 3.86 \text{ lb/TON zinc}$

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Computation Sheet For Emission Factors

Calculations Done By Z.J. Marrone Date 4-29-75

Source Primary Zinc

SO_x / E_u : Ref. 075 Table 7.7-1 Note @

$$E = 1100 \text{ lb/TON zinc} \text{ and } 2 \frac{\text{Ton zinc}}{\text{Ton zinc}}$$

$$E = 1100 \frac{\text{lb}}{\text{TON zinc}} \times 2 \frac{\text{Ton zinc}}{\text{Ton zinc}}$$

$$E = 2200 \text{ lb SO}_x / \text{TON zinc produced}$$

$$\therefore E_u = 2200 \text{ lb SO}_x / \text{TON zinc}$$

E_N : Ref 030 pg 7.5-13 to 7.5-15

We assume application of DA acid plant to roaster and similar control to sinter machine obtaining a control efficiency of 98.5%.

$$E_N = .015 \times E_u = .015 \times 2200 = 33 \text{ lb / TON zinc}$$

$$\therefore E_N = 33 \text{ lb / TON zinc}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marzzone Date 4-29-75
 Source Primary Zinc

SO_x / cont'd.

Ref 084 / 148

E_s : Of the 4 states with zinc smelting capacity only Oklahoma has specific reg's. for SO₂ emissions (new plant) only ($A_i = .197$)

For other states ($A_i = 1.0 - .197 = .803$) $E_{S_i} = E_u = 2200 \frac{\text{lb}}{\text{TON}}$

Ref 030, 6.2-25

30% Sulfur in zinc ore

For a 300 T/D zinc production plant

$$.3 \times 300 \frac{\text{Ton zinc}}{\text{day}} \times \frac{2 \frac{\text{Ton ore}}{\text{Ton zinc}}}{\text{Ton zinc}} = 180 \frac{\text{Ton S}}{\text{day}}$$

$$180 \frac{\text{Ton S}}{\text{day}} \times \frac{1}{24 \text{ hr}} \times 2000 \frac{\text{lb}}{\text{Ton}} = 15,000 \frac{\text{lb}}{\text{hr}} \text{ S input}$$

$$E_s^{\text{new}} \text{ only} \quad 16/\text{hr SO}_2 = .564 (16/\text{hr S})^{.85} \\ = .564 (15,000)^{.85} \approx 2000 \frac{\text{lb}}{\text{hr}} \text{ SO}_2$$

$$E_s'' \text{ (new)} = \frac{2000 \frac{\text{lb}}{\text{hr}}}{300 \frac{\text{Ton}}{\text{day}} \times \frac{1}{24 \text{ hr}}} = 160 \frac{\text{lb}}{\text{TON}}$$

$$E_s \text{ (new)} = .197(160) + .803(2200) \\ = 31.5 + 1766.6 \approx 1798 \frac{\text{lb}}{\text{TON}}$$

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Calculations Done By W.J. Marzzone Date 4-29-75
 Source Primary Zinc.

SO_x / cont'd.

E_s :

since there are no exist reg. $E_s(\text{exist}) = E_u = 2200 \text{ lb/ton}$
 and $E_s(\text{new}) = 1798 \text{ lb/ton}$

$$E_s(\text{exist}) = 2200 \text{ lb/ton zinc}$$

$$E_s(\text{new}) = 1798 \text{ lb/ton zinc}$$

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Computation Sheet For Emission Factors

Calculations Done By J. Manone Date 12-4-74
 Source Primary Zinc

Summary of Emission Factors

lb/ton zinc

Pollutant	E_u	E_M	$E_s(\text{exist})$	$E_s(\text{new})$
Particulate	1665	1.67	3.86	2.25
SO _x	2200	33	2200	1798

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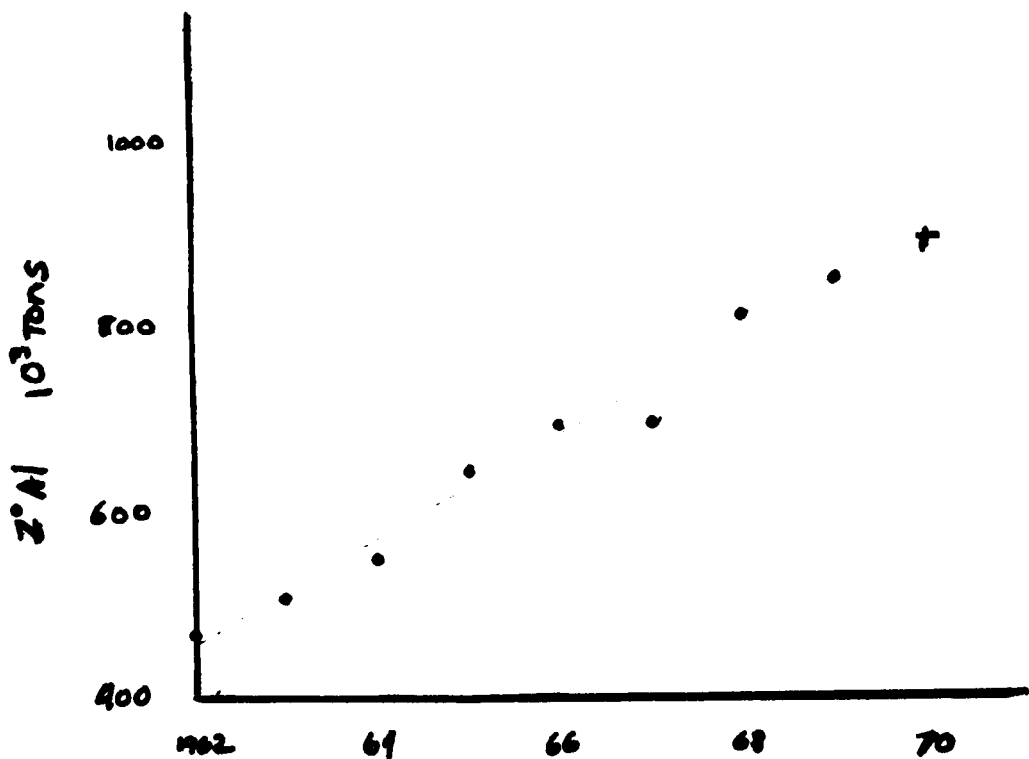
Computation Sheet For Industrial Factors

Calculations Done By HOPPER Date 1/8/75

Source SECONDARY Aluminum

Ref (95) gives Z^o Al. production data from 1930 - 1969

The 1969 value is 856.2×10^3 tons



REF (132) p1 gives the 1970 production of $(781,000) \times (1.15) = 898.2 \times 10^3$
 This is compatible with Ref (95)

The avg. (simple) growth between 1962 & 1970 is

$$\frac{898.2 \times 10^3 - 461.8 \times 10^3}{8} = 54.55 \times 10^3 \frac{\text{tons}}{\text{yr.}}$$

$$1975 \text{ Production} = 898.2 \times 10^3 + 5(54.55 \times 10^3) = 1.171 \times 10^6 \text{ tons}$$

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Computation Sheet For Industrial Factors

Calculations Done By Hopper Date 1/8/75

Source 2° Al

$$P_c = \frac{54.55 \times 10^3}{1.171 \times 10^6} = 4.66\%$$

$$P_c = 0.0466$$

simple ← Assume applicable
to sweating &
reverb. furnaces

Of the total 2° Al produced, about 81% of the mat'l comes from new scrap; the remaining 19% comes from old scrap. (Rot 132 p1)

We will assume that 50% of all old scrap and 10% of all new scrap is processed through sweat furnaces. This is based on the following points:

(1) Old scrap is relatively "dirty" and can be mixtures of other elements

(2) New scrap is relatively clean as it consists of turnings, borings, clippings etc.

$$\begin{aligned} \text{Answer}_{\text{Furn.}} &= \left[(10)(.81)(1.171 \times 10^6) + (50)(.19)(1.171 \times 10^6) \right] \div K \\ &= [9.485 \times 10^4 + 1.112 \times 10^5] = K(2.061 \times 10^5) \div K \end{aligned}$$

2° Al sweat furnaces & 2° Pb sweat furnaces are similar in nature & in many cases are interchangeable (Rot 132 p113 ff)
 We will use a K = to that for 2° Pb (See "INDUSTRIAL Factors Secondary Lead")

$$K = 0.68$$

SWEAT
FURNACE

$$\text{Answer} = \frac{2.061 \times 10^5}{0.68} = 0.303 \times 10^6 \text{ tons Al}$$

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Computation Sheet For Industrial Factors

Calculations Done By Hopper Date 1/9/75

Source 2° Aluminum

$$A_{\text{SWEAT FURNACE}} = 0.303 \times 10^6 \text{ Tons Al}$$

We will assume that all 2° Al has been processed through a reverberatory furnace.

No information could be found for K specifically for this operation. We will use the capacity utilization value for "ADVANCED PROCESS NON-FERROUS METALS". The avg K for 1965-1975 is 82%.

REF 14 P 48-TAB 1 & P 50 TAB 2

$$K_{\text{REVERB FURN.}} = 0.82$$

$$A_{\text{REVERB}} = 1.171 \times 10^6 / 0.82 = 1.428 \times 10^6$$

$$A_{\text{REVERB}} = 1.428 \times 10^6 \text{ Tons Al}$$

We will assume the absolute rate to be equal for both furnace types and this will be a function of twice the IRS depreciation allowance for the mfg. of non-ferrous metals. Ref 3 DP 34

$$P_B = \frac{1}{(2)(14 \text{ yrs})} = 0.036$$

$$P_B = 0.036 \quad \text{simple}$$

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Computation Sheet For Emission Factors

Calculations Done By HOPPER Date 1/7/75

Source Secondary Aluminum

SWEAT FURNACES

From Ref (89) p 116

$$E_{Sp} = 33 \frac{\text{lb}}{\text{TON AL}}$$

A/B's followed by a bighouse is recommended as the control equip't (Ref (49) p 438) Ref (49) TNB 19-12 p 439 indicates an $\eta = 94.1\%$. using such a system

$$E_{Np} = (1 - .94)(33) = 1.98$$

$$E_{Np} = 1.98 \frac{\text{lb}}{\text{TON AL}}$$

Ref (89) p 116 indicates the avg. capacity of a sweat furnace is 1000 lb/HR of Al. For a feed/product ratio of

$$\frac{1.0}{0.6} = 1.67 \quad (\text{Same page, same Ref})$$

the typical PWR = $1670 \frac{\text{lb}}{\text{HR}} = 0.835 \text{ TPH}$

Ref (132) p 1 states that in 1970, there were 75 aluminum smelters in the U.S. Since we do not have the specific geographical distribution of capacity, we will use the generalized PWR curve.

For a PWR = 1670 lb/hr , the allowable rate is $3.68 \frac{\text{lb}}{\text{hr}}$

$$E_{Sp} = \frac{3.68 \frac{\text{lb}}{\text{hr}}}{1000 \frac{\text{lb AL}}{\text{hr}}} \times 2000 \frac{\text{lb}}{\text{TON}} = 7.36 \frac{\text{lb}}{\text{TON}}$$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 1/19/75

Source 2° Alum.

$$E_{SP} = 7.36 \frac{LB}{TON} \text{ Alum.}$$

REVERBATORY FURNACES - PARTICULATES

Ref 214 p 3-115, 116 states that particulate emissions from REVERB furnaces are 2-4 LB/ton (ASSUME 3) during the smelting operation & 600 LB per ton Cl₂ during demagging.

Ref 132 TAB 1 STATES

300 LB/HR of PART. per 600 LB/HR of Cl₂ for a 75,000# capacity FURNACE.

$$\frac{300(2000)}{600} = 1000 \frac{LB}{TON Cl_2} \quad \text{This value is in agreement.}$$

On a ^{LB}/ton Al basis:

Ref 132 p 5 states that the avg batch cycle time is ~24 hrs. for a unit rate @ 75,000# Al

$$\frac{75,000}{(24)(2000)} = 1.563 \frac{Tons Al}{HR}$$

$$E_{UP} = 3.0 + \frac{300}{1.563} = 195 \frac{LB}{TON}$$

$$E_{UP} = 195 \frac{LB}{TON} \text{ Alum.}$$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 1/10/75

Source 2^o Aluminum

Ref 132 TAB 4 The use of a wet (fluid bed) scrubber can reduce emissions of particulate to 1.5 lb/hr

$$E_{NP} = \frac{1.5}{1.563} = 0.96$$

$$\eta = 99.5\%$$

$$E_{NP} = 0.96 \text{ lb/ton Alum.}$$

To determine Es, the following assumptions are made

75,000 lb/MELT of Al (Ref 132 TAB 4)

$$\text{FEED/PRODUCT} = \frac{1 \text{ # SCRAP}}{0.9 \text{ # Al}}$$

$$\text{WRFUX} = (0.333)(\text{WT SCRAP}) \text{ Ref } (119) \text{ p 7.8.45, 46}$$

21 hrs/batch Ref 132 p5

$$\frac{(75,000)(1.333)}{(21)(0.90)} = 1630 \text{ lb/hr}$$

Again, we will use the gen'l PWR curve since the geographical distribution of capacity is not known.

Allow $\approx 7.5 \text{ lb/hr}$
 for 1.563 TPH of Al

$$E_{SP} = 7.5 / 1.563 = 4.80 \text{ lb/ton}$$

$$E_{SP} = 4.80 \text{ lb/ton Al}$$

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Computation Sheet For Emission Factors

Calculations Done By Hoppe Date 1/10/75

Source 2° Aluminum.

Ref (132) TAB 3 states that Cl₂ emissions range between 0.1 & 0.3
 LB Cl₂ / LB Cl₂ feed

Assuming the avg = 0.2

$$\frac{0.2 \frac{\text{LB Cl}_2}{\text{LB Cl}_2 \text{ feed}} \times 600 \frac{\text{LB Cl}_2 \text{ feed}}{\text{hr}}}{1.563 \frac{\text{Tons Al}}{\text{hr}}} = 76.8 \frac{\text{lb}}{\text{ton Al}}$$

$$EU_{Cl_2 \text{ raw}} = 76.8 \frac{\text{lb}}{\text{ton Alum.}}$$

Ref (132) TAB 4 indicates that a wet scrubber will reduce chlorine emissions to "fence". We will ∴ assume 99.9%

$$EN = (1 - .999)(76.8) = 0.077$$

$$EN_{Cl_2 \text{ fence}} = 0.077 \frac{\text{lb}}{\text{ton Al}}$$

We will assume that a wet scrubber can be retrofit to existing installations as a gen'l rule. ∴ E_{ref} = EN

$$E_{III(d)} = 0.077 \frac{\text{lb}}{\text{ton Al}}$$

Since there are no reg's for Cl₂, E_S = EU.

$$ES_{Cl_2 \text{ raw}} = 76.8 \frac{\text{lb}}{\text{ton Alum.}}$$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 1/10/75

Source 2^o Aluminum

Acid Mist (HCl)

Ref(132) TAB + STATES hydrogen chloride emissions of 237 lb/ton for a 75,000 lb furnace (1.563 TPH Al)

$$EU_{HCl} = \frac{237}{1.563} = 151.6$$

$$EU_{HCl} = 151.6 \text{ lb/ton Al}$$

Same Ref + TAB

The wet scrubber can reduce the HCl emission to "trace"
 We will assume an $R = 99.9\%$

$$EN_{HCl} = 151.6 (1 - .999) = 0.152 \text{ lb/ton Al}$$

$$EN_{HCl} = 0.152 \text{ lb/ton Al}$$

As was the case with Cl₂

$$E_{III(d)} = 0.152 \text{ lb/ton Al}$$

Since there are no regs for acid mist, $E_S = EU$

$$ES_{HCl} = 151.6 \text{ lb/ton Al}$$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 1/13/75

Source 2° Aluminum

Sweat Furnace

Pær.

	<u>E_U</u>	<u>E_N</u>	<u>E_S</u>
	33	1.98	7.36

Reverb Furnace

	<u>E_U</u>	<u>E_N</u>	<u>E_S</u>	<u>E_{IIIId}</u>
Pt	195	.96	4.80	—
Ch	76.8	.077	76.8	.077
Acid Mist	151.6	.152	151.6	.152

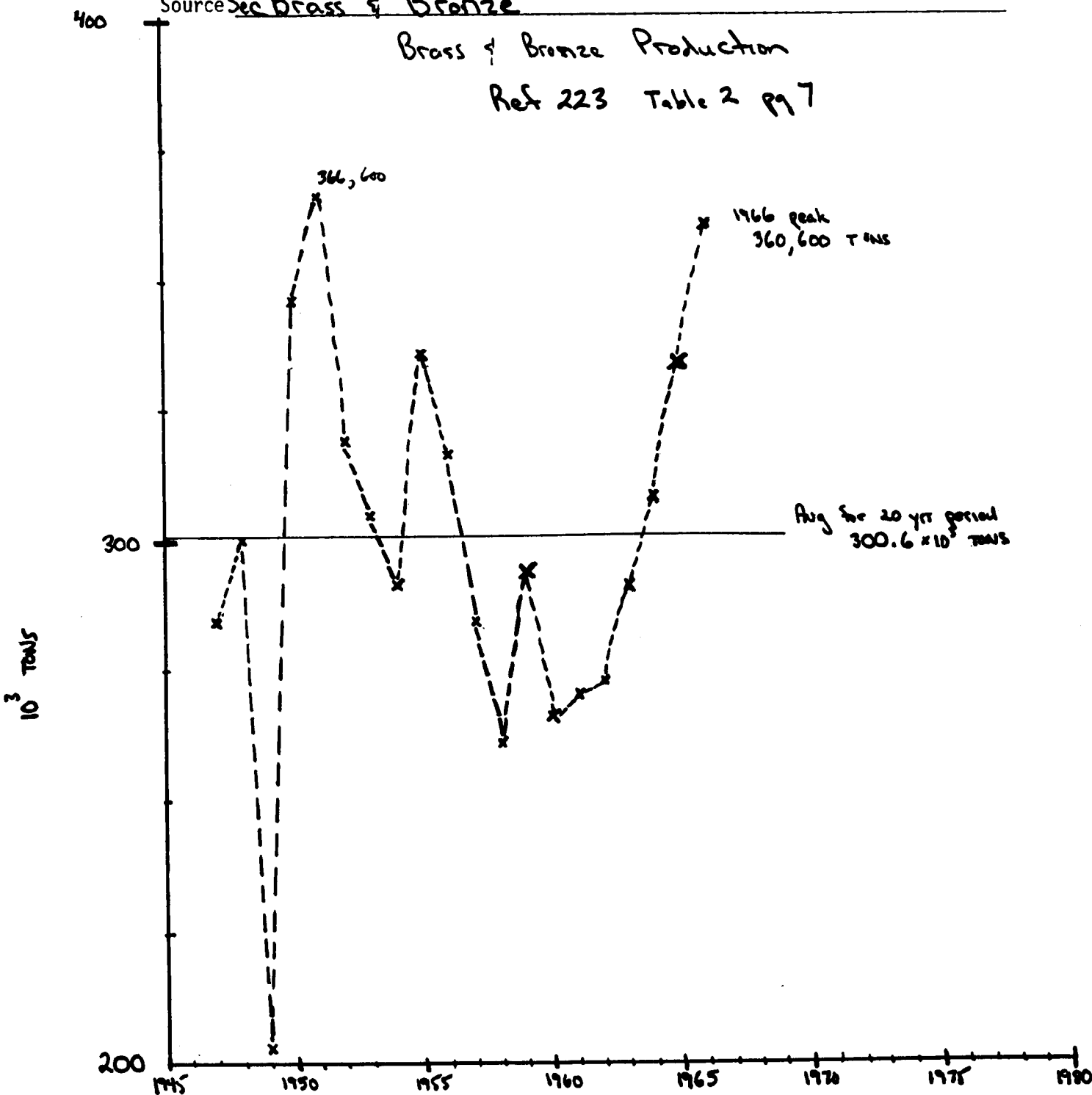
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Project Number - 32391 New Source Performance Standards

Computation Sheet For Industrial Factors

Calculations Done By W. Morrone Date 12-26-74

Source Sec Brass & Bronze



Project Number - 32391 New Source Performance Standards

Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 12-26-74

Source See Brass & Bronze

P_c :

The preceding plot of brass and bronze production indicates a fluctuating production history for this industry during the period 1947-1966

Ref 044 pg 48 confirms that 1965-1966 was a peak and that production has declined since then. It goes on to say the excess capacity exists within the industry and few, if any, new plants are expected to be constructed in the next few years.

We make the assumption that existing capacity will be sufficient to meet brass & bronze demands. and the new capacity will not be constructed during 1975-1985

$$\therefore P_c = 0$$

K: Information for the determination of K was not found in the literature. Since it has been mentioned that overcapacity exists we make a best estimate by comparing the average production rate of $\approx 300 \times 10^3$ Tons/yr with some likely maximum

est Peak = 375,000 Tons/yr

Avg = 300,000 Tons/yr

$$K = \frac{300,000}{375,000} = .8$$

$$\therefore K = .80$$

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Computation Sheet For Industrial Factors

Calculations Done By W. Marzzone Date 12-26-74

Source See Brass & Bronze

A:

Since we have assumed $P_c = 0$ and used an average production and estimate peak for determination of K then the value of A (1975 production capacity) is represented by the peak production level of 375,000 TON/yr used above:

$$\therefore A = .375 \times 10^6 \text{ TON/yr}$$

P_B : Ref 037 pg 34 Class 33.2

Asset Guideline period = 14 yrs

P_B est $\rightarrow 2 \times \text{IRS} = 28 \text{ yrs}$

$$\frac{100\%}{28 \text{ yrs}} \approx 3.6\%/\text{yr} \text{ simple}$$

$$\therefore P_B = .036 \text{ simple}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-26-74
 Source Sec Brass & Bronze

Postulates

Ref 223 pg 1-2

Emissions from this industry are mainly attributable to refining furnaces (reverb, rotary, electric, crucible [blast])

Ref 044 pg 45 states

(1) 95% of production is from reverberatory furnaces which has highest emission level

(2) with the exception of blast (crucible) furnace, which has emission comparable to reverberatory, all other furnaces have lower emissions levels

E_u :

Ref 075 Table 7.9-1

$$E = 70 \text{ lb/TOW metal charged}$$

Ref 223 pg 2

avg emission factor = 60-80 lb/TOW ingot
 for reverberatory furnace

We will use 70 lb/TOW metal produced

$$E_u = 70 \text{ lb/TOW metal produced}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-26-74

Source Sec Brass & Bronze

Particulate cont'd

E_N :

Ref 075 pg 7.9-2

The use of a baghouse, which is the predominant control device used in this industry, may achieve collection efficiencies of 95-99.6 %

We will use a value of 99.6% as attainable in developing E_N

$$E_N = .004 \times E_u = .004 \times 70 \text{ lb/ton}$$

$$\therefore E_N = .28 \text{ lb/ton}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marone Date 12-26-74

Source Sec Brass + Bronze

Particulate

E_s :

New Source Performance Standards have been proposed for this industry so that a value of E_s (new) will be determined.

Ref 044 pg 46 states that the proposed standard will require emissions to be limited to 1.0 - 1.5 lb/hr from a 25 T/O Furnace

Note

assuming 24 hr

Ref 271 pg 9312 states that the proposed standard is the same as the proposed

$$\frac{1.0 \text{ lb}}{25 \text{ T} \div 24 \text{ hr}} = .96 \text{ lb/TON}$$

> avg 1.2 lb/TON

$$\frac{1.5 \text{ lb}}{25 \text{ T} \div 24 \text{ hr}} = 1.44 \text{ lb/TON}$$

Based on the avg this represents $\approx 98\%$ off

$$E_s(\text{new}) = 1.2 \text{ lb/TON}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-26-74
 Source See Brass & Bronze

Particulate cont'd

Ref 044 pg 45-46 discusses regulation in terms of
 a typical 25 ton/day furnace

assuming a production schedule of 24 hr/day we
 obtain $\frac{25 \times 2000}{24 \text{ hr}} = 2083 \text{ lb/hr} \approx 2000 \frac{\text{lb}}{\text{hr}}$ or $1 \frac{\text{ton}}{\text{hr}}$
 as a process input weight rate.

A breakdown of brass & bronze production by state was not obtained however by referring to Ref 223 & 124 we were able to see that several states were identified as major contributors to brass and bronze production.

We develop an E_g (existing) factor by making the following assumptions:

- (1) the states identified are New York, Penn, Ohio, Alabama, Illinois, Kansas, Virginia, Delaware.
- (2) allowable emission for each state based on process weight of 2000 lb/hr will be given equal weighting so that $E_g \text{ avg} = \frac{\sum E_g}{N}$ where $N = 8$.
- (3) allowable emissions for any other states not covered will be considered to be equal to this avg

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Computation Sheet For Emission Factors

Calculations Done By Z. Manone Date 12-26-74

Source See Brass & Bronze

<u>State</u>	<u>lb/hr</u> <u>Allowable Emission</u>
NY	3.65
Penn	2.25*
Ohio	4.10
Alabama	3.60
Illinois	4.10
Kansas	4.10
Virginia	4.10
Delaware	1.20*

* based on 9g/sec regulation
 compared to regulation in R-044
 for new source

.022 g/sec " proposed
 $\approx 1.2 \text{ lb/hr}$

$$\text{Avg} = 3.39 \text{ lb/hr}$$

$$E_s = \frac{3.39 \text{ lb/hr}}{2083 \frac{\text{lb}}{\text{hr}} \times \frac{\text{Ton metal}}{2000 \text{ lb}}} = 3.25 \text{ lb/TON}$$

metal produced

$$E_s (\text{existing}) = 3.25 \text{ lb/TON}$$

metal produced

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Source Sec Brass & Bronze

Lead and Zinc

Ref 044 p45 identifies most of the particulate emissions to be lead oxides and zinc oxides

45 to 77% Zn Oxides
 1 to 13% Pb Oxides in emissions

We will use the average of these percentages in determining the emission factors. $E_{N\text{Pb}} = .065 \times E_{u\text{Pb}}$
 $E_{N\text{Zn}} = .61 \times E_{u\text{Zn}}$

It is assumed that (1) control devices will be equally effective on these constituents as for total particulate so that $E_{N\text{Pb}} = .004 \times E_{u\text{Pb}}$
 $E_{N\text{Zn}} = .004 \times E_{u\text{Zn}}$

(2) Since controls that are installed for particulate source are controlling Pb & Zn simultaneously we must assume that the allowable emissions are a function of the E_s for particulate and proportional to the Zn and Pb content.

$$E_s(\text{new}) \text{ Pb} = .065 \times E_s(\text{new})$$

$$E_s(\text{new}) \text{ Zn} = .61 \times E_s(\text{new})$$

$$E_s(\text{existing}) \text{ Pb} = .065 \times E_s(\text{existing})$$

$$E_s(\text{existing}) \text{ Zn} = .61 \times E_s(\text{existing})$$

(3) Existing plants are assumed to be able to apply baghouses to revert to surfaces and achieve same control efficiency as new plant $\therefore E_{\text{old}} = E_N(\text{Pb or Zn})$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 12-26-74

Source See Brass & Bronze

head of zinc Cont'd

E_u :

$$E_u \text{ Pb} = .065 \times 70 \approx 4.6 \text{ lb/ton}$$

$$E_u \text{ Zn} = .61 \times 70 = 42.7 \text{ lb/ton}$$

$$\therefore E_u(\text{Pb}) = 4.6 \text{ lb/ton metal produced}$$

$$E_u(\text{Zn}) = 42.7 \text{ lb/ton metal produced}$$

E_N :

$$E_N = .004 \times E_u$$

$$E_{\text{III}d} = E_N$$

$$E_N(\text{Pb}) = .004 \times 4.6 \frac{\text{lb}}{\text{ton}} = .018$$

$$E_N(\text{Zn}) = .004 \times 42.7 \frac{\text{lb}}{\text{ton}} = .17$$

$$\therefore E_{\text{III}d} = E_N(\text{Pb}) = .018 \text{ lb/ton metal produced}$$

$$E_{\text{III}d} = E_N(\text{Zn}) = .17 \text{ lb/ton metal produced}$$

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Emission Factors

Calculations Done By W. Monroe Date 12-26-74

Source See Brass & Bronze

Lead and Zinc cont'd

$$E_s(\text{new}) \text{ Pb} = .065 \times 1.2 \text{ lb/tow} = .078 \text{ lb/tow metal prod}$$

$$E_s(\text{new}) \text{ Zn} = .61 \times 1.2 \text{ lb/tow} = .73 \text{ lb/tow metal prod}$$

$$E_s(\text{existing}) \text{ Pb} = .065 \times 3.25 \text{ lb/tow} = .21 \text{ lb/tow metal produced}$$

$$E_s(\text{existing}) \text{ Zn} = .61 \times 3.25 \text{ lb/tow} = 1.98 \text{ lb/tow metal produced}$$

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Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 1-28-75

Source Core Ovens - Iron Foundry

K:

We will use the value of $K = .93$ obtained for Cast Iron Foundry as representative of core ovens.

$$K = .93$$

P_c : The growth in core production is assumed a direct effect of cast iron growth. From "Cast Iron Foundry" by W. Marrone - plot of Cast Iron production and total production year "a growth rate (simple) of:

$$P_c = \frac{25.2 - 21.4}{5} = .035 \text{ simple}$$

$$P_c = .035 \text{ simple}$$

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Computation Sheet For Industrial Factors

Calculations Done By W. Monroe Date 1-28-71

Source Core Oven - Iron Foundry

A :

A will be assumed equivalent to the sum of cupola and electric cast iron production capacity.

$$A_{\text{cupola}} = 14.9 \times 10^6$$

$$A_{\text{elec}} = \frac{8.1 \times 10^6}{23.0 \times 10^6 \text{ ton/yr}}$$

$$\therefore A = 23 \times 10^6 \frac{\text{ton}}{\text{yr}} \text{ metal casted}$$

P_B :

Use P_B obtained for electric furnace in Cast Iron which was derived from IRS Tables Ref 037

$$P_B = .028 \text{ simple}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Morrone Date 1-28-75

Source Core Ovens - Iron Foundry

Hydrocarbons

E_u : Ref 047 pg 175 gives figures for 1968 of 18,000,000 ton metal cast and 10,500,000 ton of sand used. This gives us a factor of:

$$\frac{10.5 \times 10^6}{18 \times 10^6} = .583 \frac{\text{TON sand}}{\text{TON iron cast}}$$

From Ref 046 p 313 for typical core mixture of:

930 lb sand
 7.5 lb core oil. { 45% linseed oil
 28% gum resin
 27% kerosene

9.0 lb cereal binder

3.0 lb kerosene

38 lb water

HC Emissions are assumed to be 45% + 27% of core oil and 3 lb kerosene or $.72(7.5 \text{ lb}) + 3.0 = 5.4 + 3.0 = 8.4 \text{ lb HC}$

$$\frac{8.4 \text{ lb HC}}{930 \text{ lb sand}} \times \frac{2000 \text{ lb}}{\text{TON}} \approx 18 \frac{\text{lb HC}}{\text{TON sand}}$$

Then

$$E_u = .583 \frac{\text{TON sand}}{\text{TON iron cast}} \times 18 \frac{\text{lb HC}}{\text{TON sand}} \approx 10.5 \frac{\text{lb}}{\text{TON iron cast}}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 1-28-75
 Source Core Ovens - Iron Foundry

Hydrocarbons cont'd

E_N :

Emission reduction from Core ovens may be achieved through modification of core oil mixture however this may not always be feasible.

Ref 046 pg 314 - 315 indicates that afterburners are generally applied to this source for control of organic material.

Without any definitive test data to define an efficiency we will assume that at least 90% can be achieved for this source $E_N = .10 \times E_u$

$$E_N = 1.05 \text{ lb / TON} \quad \begin{matrix} \text{iron} \\ \text{cast} \end{matrix}$$

E_S :

State Regulations for hydrocarbons' are not applicable to core ovens so that we may state that $E_S = E_u$

Ref 084 § 148

$$E_S = 10.5 \text{ lb / TON} \quad \begin{matrix} \text{iron} \\ \text{cast} \end{matrix}$$

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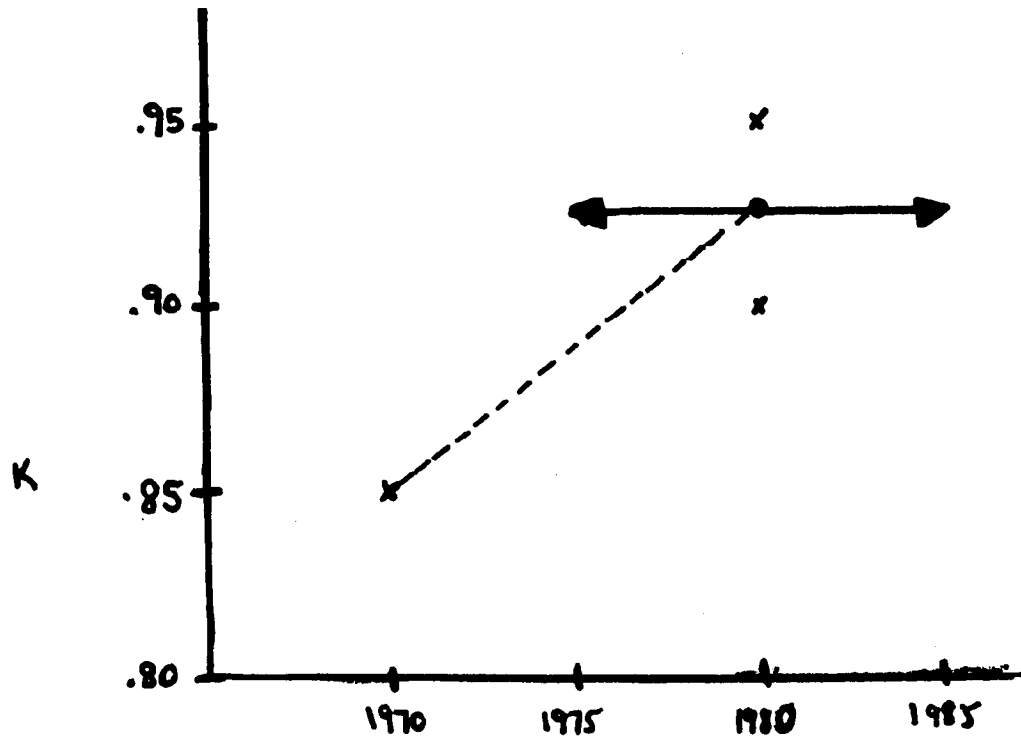
Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 8-19-74

Source Cast Iron Foundry (Furnaces)

K: Reference O7 pg 51 states that the 1970 production of gray iron casting was 17,923,000 TONS or about 85% of capacity. The estimated 1980 Prod/Cap ratios is expected to be 90 - 95%. ^{some ref} pg 51

We plot this information for better visualization.



We will take the simple average of 90 and 95% in 1980 and use this value to represent the typical K applicable over the 1975-1985 period

$$\frac{90+95}{2} = 92.5 \Rightarrow \text{assume } K = .93$$

$$K = .93$$

Project Number - 32391 New Source Performance Standards

Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 8-19-74

Source Cast Iron Foundry (Furnaces)

K: cont'd

The gray iron production is assumed to include the production of ductile and malleable iron.

Cast iron, which is the general term used for casting manuf. having the characteristics of one of the above irons, is performed by melting metal scrap, rerun material, flux, etc. in a furnace. The three most commonly employed are;

- (a) cupola
- (b) electric arc
- (c) induction.

We will assume the utilization factor ($K = .93$) is the same industry wide.

Since the emission factors vary widely between furnaces and the growth rates are different for each, we are compelled to treat each separately, and employ individual values for A , P_e and P_B .

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Computation Sheet For Industrial Factors

Calculations Done By W. Marrone Date 8-19-74
 Source Cast Iron Foundry (Furnaces)

A:

Ref 050 "Gray-Iron Foundries" pg 1 states that in 1970 90% of the metal poured for gray iron was done in cupolas.

Ref 067 pg II-11 vol I states that in 1969 85% of all ^{iron} metal melted was performed in cupolas. It goes on to project that only 50% of the total will be melted in cupolas; the balance being taken up by electric arc and induction. Reverberatory is referred to be a small source due to it small % of the production total.

We will assume that the 85% figure for 1969 applies to 1970 & is representative to the ~~cast iron~~ industry. It is believed that the 90% figure from Ref 050 represents gray iron only and not the ductile or malleable components combined (Total cast iron)

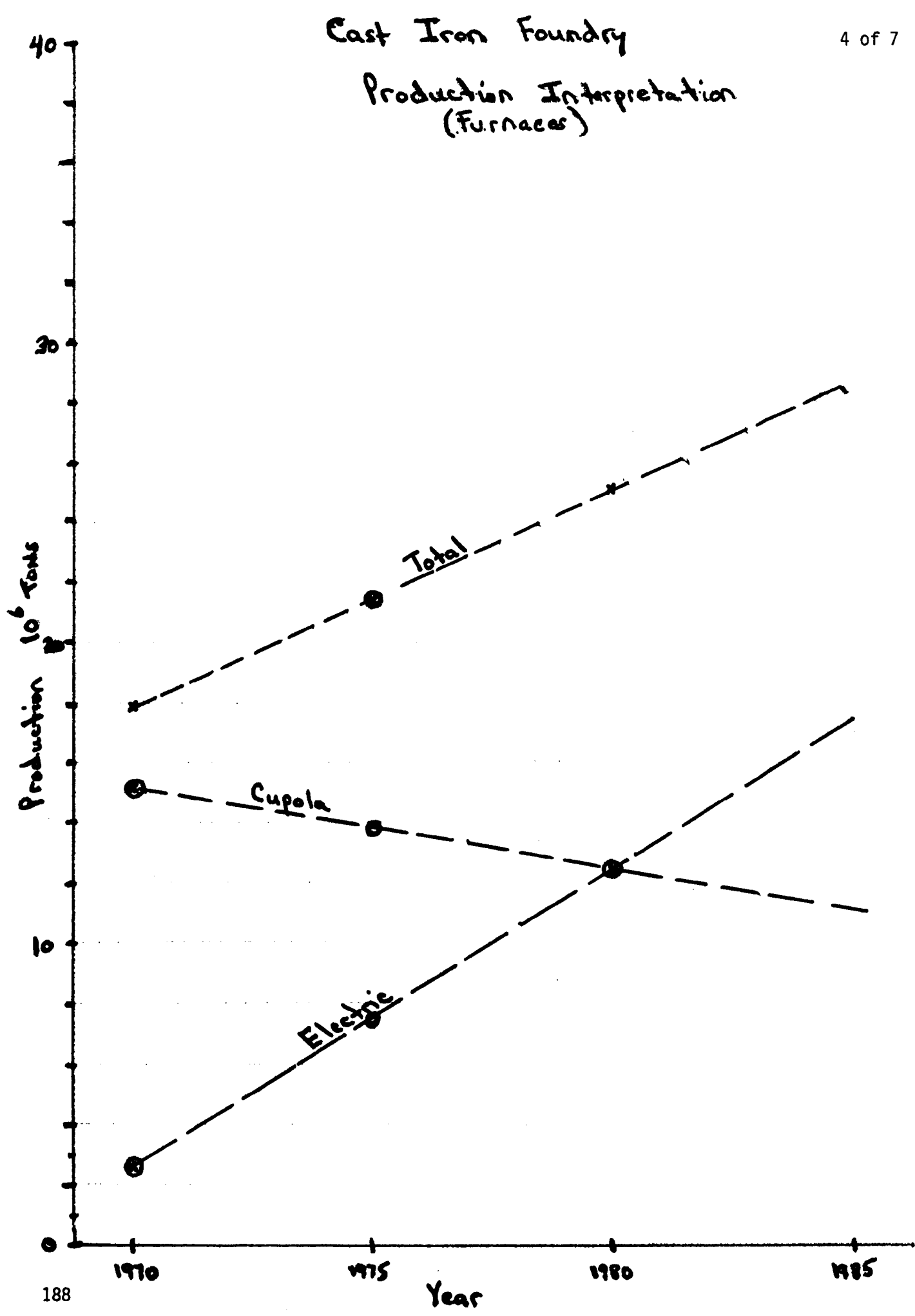
The attached Graph plots the total production information for the period 1970 - 1980 and the cupola and electric furnace prod. values.

The Total production is based on Ref 007 Table 8.1 and the 85% (1970) cupola component and the 50% (1980) cupola component.

Cast Iron Foundry

4 of 7

Production Interpretation (furnaces)



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Computation Sheet For Industrial Factors

Calculations Done By William Marone Date 8-19-74
 Source Cast Iron Foundry (Furnaces)

A: cont'd

This industry will be divided into the cupola
 and electric furnace processes.

Referring to the graph we prepared, we may obtain an estimate for the 1975 production figures of cupolas and electric furnaces. They were arrived at by straight-line interpolation between the data points plotted from information found in the literature as referenced previously.

$$1975 \text{ Production Total} = 21.4 \times 10^6 \text{ Tons/yr}$$

$$1975 \text{ Cupola Production} = 13.9 \times 10^6 \text{ Tons/yr}$$

$$1975 \text{ Electric Production} = 7.5 \times 10^6 \text{ Tons/yr}$$

Applying the value of $K = .93$ we calculate the production capacities in 1975;

$$K = .93$$

$$A_{\text{cupola}} = \frac{13.9}{.93} \times 10^6$$

$$A_{\text{cupola}} = 14.9 \times 10^6 \text{ Tons/yr}$$

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Computation Sheet For Industrial Factors

Calculations Done By William Monroe Date 8-19-74

Source Cast Iron Foundry (Furnaces)

A: cont'd

$$A_{\text{electric}} = \frac{7.5}{.93} \times 10^6$$

$$A_{\text{electric}} = 8.1 \times 10^6 \text{ Ton/yr}$$

$P_c \approx$

We will take the growth of each of these to be represented by the simple growth above base year 1975. Use Production curves on attached graph.

$$P_c \text{ Cupola} = \frac{1980 - 1975}{1975} \div 5 = -.018 \quad \text{simple}$$

$$P_c = \frac{12.6 - 13.9}{13.9} \div 5 = -.018$$

$$P_c \text{ Electric} = \frac{1980 - 1975}{1975} \div 5 = .136 \quad \text{simple}$$

$$P_c = \frac{12.6 - 7.5}{7.5} \div 5 = .136$$

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Computation Sheet For Industrial Factors

Calculations Done By William Marone Date 8-19-74

Source Cast Iron Foundry (Furnaces)

P_B : cupola As indicated in Ref 067 pg II-11 cupolas are being replaced industry-wide by electric furnaces. This trend is expected to continue undiminished, which would imply that replacement of existing equipment by ^{cupolas} is not expected. We therefore assume that

$$P_{B \text{ cupola}} = 0$$

NOTE

Since the P_C for cupola is a negative value and the P_B was assumed to be 0, this particular source does not come under NSPS

P_B :
electric

Ref 037 pg 34 Section 33.1

Asset guideline period 18 yrs

P_B est $\rightarrow 2 \times \text{IRS} = 36 \text{ yrs}$

$$\frac{100\%}{36 \text{ yrs}} \cong 2.8\%/\text{yr simple}$$

$$\therefore P_{B \text{ electric}} = .028 \text{ simple}$$

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Computation Sheet For Emission Factors

Calculations Done By W. Marrone Date 8-20-74
 Source Cast Iron Foundry (Furnaces)

We have concluded that there is a decline in the cupola capacity (negative growth) and no replacement of obsolete capacity ($P_B = 0$) so that Cupola Furnaces would not be considered under NSPS.

Emission factors will be developed for the Electric furnaces.

Particulate

E_u : Ref 075 Table 7.10-1 pg 7.10-1

Electric Arc: $E_u = 4.5 \text{ lb/TON metal}$

Ref 145 pg 216

Electric Arc $5-10 \text{ lb/TON metal}$

We will use ~~an~~ average value for an emission factor from electric furnaces

$$E_u = \frac{\frac{5+10}{2} + 1.5}{2} = 4.5 \text{ lb/TON}$$

$E_u = 4.5 \text{ lb/TON metal}$

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Computation Sheet For Emission Factors

Calculations Done By William Marrone Date 8-19-74
 Source Cast Iron Foundry (Furnaces)

E_N : Ref 050 "Gray Iron Foundry" pg 4-5

Control for Electric Arc Furnace are mostly
 baghouses 95-99% eff

No control information for Electric Induction
 was found.

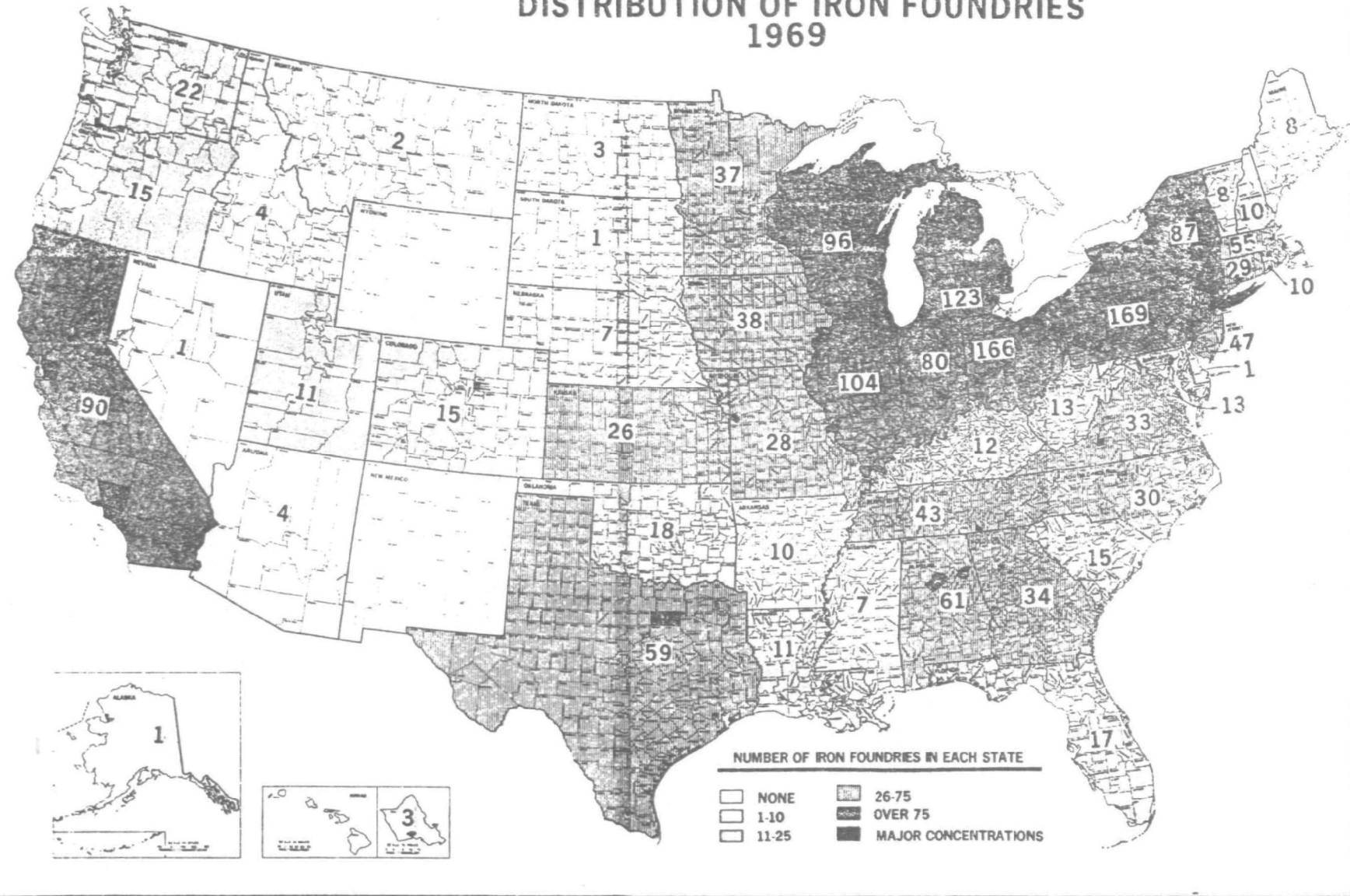
We assume an attainable efficiency for both
 types of electric furnaces to be 99% which
 will be applied to the whole electric furnace
 portion of the Cast Iron Foundry industry

$$E_N = .01 \times E_u = .01 \times 4.5 = .045 \text{ lb/ton}$$

$$E_N = .045 \text{ lb/ton metal}$$

Ref 068

DISTRIBUTION OF IRON FOUNDRIES 1969



SOURCE: PENTON PUBLISHING CO.

A

B

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Computation Sheet For Emission Factors

Calculations Done By R. J. Londergan Date 00-24-74
 Source Cast Iron Foundry

Particulates

Using the information presented on p.4 of the production factors calculation, electric furnace production in 1970 was 2.7×10^6 TONS. From Ref. 007, p.52, we see that there were a total of 1040 electric furnaces in operation in 1970. The process weight rate (average) is therefore

$$\frac{2.7 \times 10^6 \text{ TON/YR}}{1040 \text{ furnaces}} \times \frac{1 \text{ YR}}{365 \text{ day}} \times \frac{1 \text{ day}}{16 \text{ hr.}} \times \frac{2000 \text{ LB}}{\text{TON}} \approx 900 \text{ LB/HR}$$

assuming a 16-hour/day average utilization. No geographic distribution for electric furnaces is available, so we utilize the 50-state average allowable emissions curves, as discussed in Vol # 1. The resulting allowable emissions rate is 2.4 LB/HR, corresponding to

$$E_S = (2.4 \text{ LB/HR}) \times \frac{1}{(.45 \text{ TON/HR})} = 5.3 \text{ LB PT.}$$

value is larger than

TON IRON

This ~~value exceeds~~ E_u ; therefore we have

$$E_S = E_{cl} = 4.5 \text{ LB/TON}$$

) since E_S cannot in fact be larger than E_u .

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Computation Sheet For Industrial Factors

Calculations Done By Hornee Date 9/11/74
 Source Secondary Copper

1970 Production = 1257×10^3 tons Ref (95) p 233.4530

Growth 1960 - 1970 Ref (95) p 233.4530

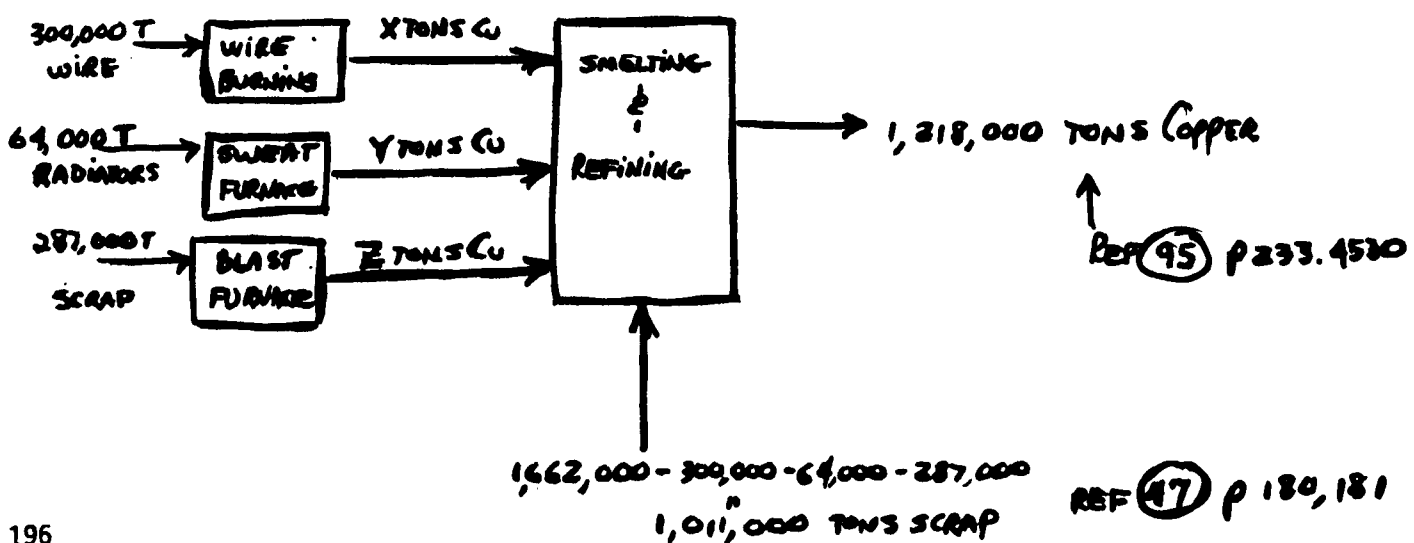
$$P_c = \sqrt[19]{\frac{1257}{871.4}} - 1.0 = 3.74\%$$

$P_c = 3.74\% \text{ compound}$

Emissions are the result of

- ① Materials Preparation Ref (49) p 406 TNS 19-1
- ② Smelting & Refining

(49) REF (1) p 406 TNS 19-1 EMISSION FACTORS ARE PRENTED ON THE BASIC OF UNIT WEIGHT OF SCRAP NOT UNIT WEIGHT OF COPPER PRODUCED. THE FOLLOWING FLOW CHART MAY HELP IN REDFINING EMISSION RATES AND DETERMINING BREAKDOWNS IN "A".
 1968 VALUES



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Computation Sheet For Industrial Factors

Calculations Done By Homer Date 9/12/74
 Source Secondary Copper

Ref (1) STATES VARIOUS % COMBUSTIBLES FOR WIRE. THE AVG P 181 IS ABOUT 26% OR 74% COPPER

$$\left(300,000 \text{ TONS WIRE} \right) \left(0.74 \frac{\text{TONS COPPER}}{\text{TON WIRE}} \right) = X \text{ TONS CU} \approx 222,000 \text{ TONS CU}$$

Ref (89) p 116 STATES 60% AS THE RECOVERABLE PERCENT OF ALUMINUM IN SCRAP USED IN SWEATING FURNACES. WE WILL ASSUME THIS % FOR COPPER USED IN SWEATING FURNACES.

$$\left(64,000 \text{ TONS RAD'S} \right) \left(0.6 \frac{\text{TONS CU}}{\text{TON RAD}} \right) = Y \text{ TONS CU} = 38,400 \text{ TONS CU}$$

Ref (46) p 302 STATES 70% AS THE RECOVERABLE PERCENT OF LEAD FROM SCRAP USED IN BLAST FURNACES. WE WILL ASSUME THIS % FOR COPPER USED IN BLAST FURNACES.

$$\left(287,000 \text{ TONS SCRAP} \right) \left(0.7 \frac{\text{TONS CU}}{\text{TON SCRAP}} \right) = 200,900 \text{ TONS CU}$$

$$\Sigma \text{Copper} = 461,300 \text{ TONS 1968}$$

1975 "Production"
 COPPER OUTPUT
 FROM M&L'S
 HANDLING

$$= \left(0.4613 \times 10^6 \text{ tons} \right) \left(1.0374 \right) = 0.596 \times 10^6 \text{ tons}$$

Assume to be same
 as the whole
 industry

No direct information regarding "K" could be located other than DDC statistics on capacity utilization rates for "ADVANCED PROCESSED NON-FERROUS METALS". THE AVERAGE "K" FOR 1965-1973 IS 82%

$$K = 0.82$$

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Computation Sheet For Industrial Factors

Calculations Done By Hopper Date 7/12/74
 Source Screaming Comee

$$A_{\text{mores}} = \frac{0.596 \times 10^6 \text{ TONS}}{0.82} = 0.727 \times 10^6$$

$$A_{\text{comes}} = 0.727 \times 10^6 \text{ TONS}$$

$$A_{\text{comes}} = \frac{(1.257 \times 10^6)(1.0374)^5}{0.82} = 1.842 \times 10^6 \text{ TONS}$$

$$A_{\text{comes}} = 1.842 \times 10^6 \text{ TONS}$$

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Computation Sheet For Industrial Factors

Calculations Done By ZJ. Marnone Date 9-26-74

Source Secondary Copper

P_B : Ref 037 pg 34 section 33.2

Asset guideline life 14 yrs.

P_B est = $2 \times \text{IRS}$ 28 yrs.

$$\frac{100\%}{28 \text{ yrs.}} \approx 3.5\% \text{ /yr}$$

simple

$$\therefore P_B = .035 \text{ simple}$$

We assume that the value of P_B is applicable to both the material handling and smelting/refining operations of this industry.

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Computation Sheet For Emission Factors

Calculations Done By Homer Date 9/12/74
 Source SOURCE OF COPPER

AS MENTIONED ON P1 OF THE CALCULATION SHEET FOR INDUSTRIAL FACTORS, EMISSION FACTORS ARE PRESENTED ON THE BASIS OF UNIT WEIGHT OF SCRAP, NOT UNIT WEIGHT OF COPPER PRODUCED.

Ref (4) p406 TAB 19-1

A. MATERIALS Preparation

- ① WIRE BURNING - 275 #/ton scrap
- ② SWEEPING FURNACES - 15 #/ton scrap
- ③ BLAST FURNACES - 50 #/ton scrap

From p2 of industrial factor calc sheet

$$0.74 \frac{\text{TONS CU}}{\text{TON WIRE}} \Rightarrow \frac{275}{0.74} = 372 \frac{\text{#}}{\text{TON CU}}$$

$$0.6 \frac{\text{TONS CU}}{\text{TON RAD'S}} \Rightarrow \frac{15}{0.6} = 25 \frac{\text{#}}{\text{TON CU}} (\text{Sweeping})$$

$$0.7 \frac{\text{TONS CU}}{\text{TON SCRAP}} \Rightarrow \frac{50}{0.7} = 71.4 \frac{\text{#}}{\text{TON CU}} (\text{Blast})$$

Also from p2

222,000	TONS CU FROM WIRE	-	18 %
38,400	TONS CU FROM SWEEPING FURNACES	-	8 %
200,900	TONS CU FROM BLAST FURNACES	-	14 %
<u>461,300</u>	TONS TOTAL		

$$\bar{E}_{U_{\text{HANDLING}}} = 372(0.18) + 25(0.08) + 71.4(0.14)$$

$$= 179 + 2 + 31.4$$

$$E_{U_{\text{HANDLING}}} = 212 \frac{\text{#}}{\text{TON COPPER}}$$

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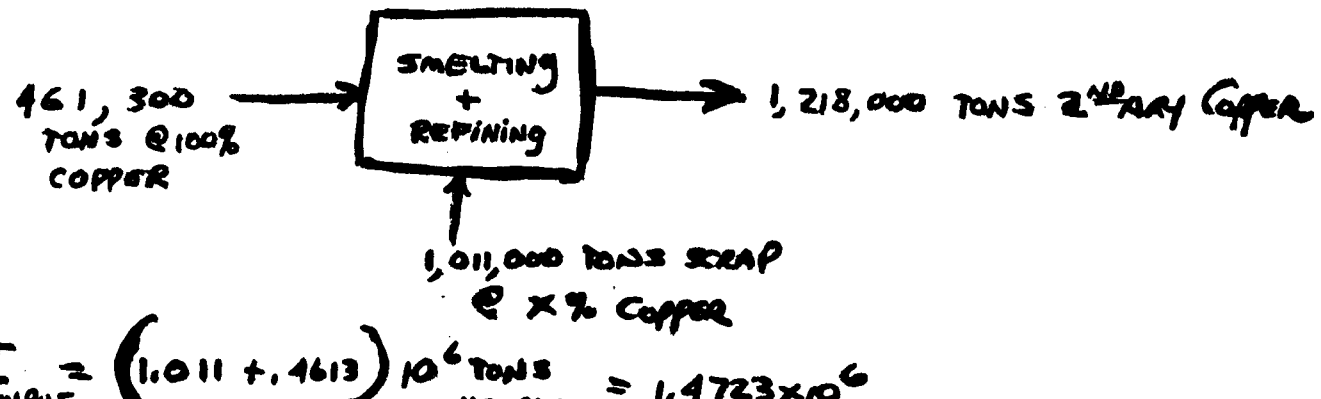
Computation Sheet For Emission Factors

Calculations Done By Homer Date 9/12/74

Source Secondary Copper

Ref (49) p 406 TAB 19-1 gives

$$\text{EF}_{\text{SMELT.}} = 70 \text{ #/ton SCRAP}$$



$$\frac{\text{TONS COPPER}}{\text{TON SCRAP INPUT}} = \frac{1.218 \times 10^6}{1.4723 \times 10^6} = 0.827$$

$$\text{EF}_{\text{SMELT.}} = 70 / 0.827 = 84.6 \text{ #/ton Cu}$$

$$\text{EF}_{\text{SMELTING + REFINING}} = 84.6 \text{ #/ton Copper}$$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 9/12/74
 Source Secondary Copper

REF (49) p 915 STATES AN efficiency of 99.6% has
 been achieved with A baghouse - the only generally
 ACCEPTED COLLECTION DEVICE.

$$E_{NP} = EU (1 - .996) = EU (.004)$$

$$E_{NP_{\text{smelting}}} = 0.848 \text{ #/ton copper}$$

$$E_{NP_{\text{refining}}} = 0.338 \text{ #/ton copper}$$

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Computation Sheet For Emission Factors

Calculations Done By V. P. Russo Date 10/21/74

Source Secondary Copper
Particulates:

Ref 155 states for 1967 that 42 plants are in the
 2nd day Copper Industry. Assuming each plant
 performs all of the functions (i.e. wire burning, sweating,
 blast furnaces, smelting and refining).

Wire Burning

300,000 T Wire yields 222,000 Tons Cu (See previous analysis)

$$\text{For single Pl. Process Wgt. Rate} = \frac{300,000 \text{ Tons/YR}}{(42) 8760 \text{ HRS/YR}} \frac{\text{lbs/ton}}{\text{HR}} \\ \approx 1630 \text{ lbs/HR}$$

Allowable Emissions (50 State Numerical Average) = 3.60 $\frac{\text{lbs}}{\text{HR}}$ Particulate

$$\text{Copper Production per hour} = \frac{222,000 \text{ Tons} \times 2000 \text{ lbs/ton}}{(42) 8760} \frac{\text{TONS}}{\text{HR}}$$

$$\approx 1206 \text{ lbs/HR} = 0.603 \frac{\text{Tons}}{\text{HR}}$$

$$\text{For wireburn } E_S^o = \frac{3.60 \text{ lbs/HR}}{0.603 \frac{\text{Tons}}{\text{HR}}} = 5.97 \text{ lbs Part/Ton Cu}$$

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Emission Factors

Calculations Done By U. F. Russo Date 10/21/24

Source Secondary Copper

Radiators

64,000 TONS yields 38,400 TONS Cu

$$\text{For single plant Process Wgt. Rate} = \frac{64,000 \text{ TON/YR}}{(42) (8760)} \frac{2000 \text{ lbs/TON}}{\text{also } .174 \text{ TONS/HR}}$$

$$\approx 350 \text{ lbs/HR}$$

$$\text{Allowable Emissions} = 1.25 \text{ lbs Part./HR}$$

(50 State Average)

$$\text{Copper Production} = \frac{38,400 \text{ TONS YR}}{(42) (8760)} \frac{2000 \text{ lbs/TON}}{=.104 \text{ TONS/HR}}$$

$$= 208.74 \text{ lbs/HR} = .104 \text{ TONS/HR}$$

$$E_s^o = \frac{1.25 \text{ lbs Part./HR}}{.104 \text{ TON Cu/HR}} = 12.02 \text{ lbs Part./TON Cu.}$$

Blast Furnace

287,000 TONS Yields 200,900 TONS Cu

$$\text{Single Plant Process Wgt. Rate} = 1560 \text{ lbs/HR}$$

$$\text{Allowable Emissions} = 3.5 \text{ lbs Part./HR}$$

(50 State Average)

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Computation Sheet For Emission Factors

Calculations Done By V. Russo Date 10/21/74
 Source Secondary Copper

$$\text{Copper Production} = \frac{200,900 \text{ Ton Cu/Yr}}{(42) 8760 \text{ HR/YR}} \\ = 0.546 \text{ TONS/hr}$$

$$E_S^0 = \frac{3.5 \text{ lbs/HR}}{.546 \text{ TONS/hr}} = 6.41 \text{ lbs Part/Ton Cu}$$

For handling

$$E_S^{\text{Handling}} = 5.97(48) + 12.02(28) + 6.41(44) \\ = 2.86 + 0.96 + 2.82$$

$$E_S^{\text{Handling}} = 6.64 \text{ lbs Part/Ton Cu.}$$

SMELTING & REFINING

$$1.4723 \times 10^6 \text{ TONS yields } 1.218 \times 10^6 \text{ TONS } 2^{\text{nd}} \text{ary Cu}$$

For Single Plant Process Wgt. Rate = 8000 lbs/HR

$$\text{Allowable Emissions} = 11.5 \text{ lbs/HR} \\ (\text{50 State Average})$$

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Computation Sheet For Emission Factors

Calculations Done By V. F. Russo Date 10/21/74
 Source Secondary Copper

$$\text{Cu Production} = \frac{1.218 \times 10^6 \text{ TONS Cu}}{(42)(8760)}$$

$$= 3.31 \frac{\text{TONS}}{\text{HR}}$$

Smelting & Refining

$$E_S = \frac{11.5 \frac{\text{LBS}}{\text{HR}}}{3.31 \frac{\text{TONS}}{\text{HR}}} = 3.47 \frac{\text{Lbs. Paint}}{\text{TON Cu}}$$

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Industrial Factors

Calculations Done By Hopper Date 9/6/74
 Source Secondary Lead

PRODUCTION IN 1971 = 555×10^3 TONS REF (95)
p 233, 4830

REF (41) p 41 STATES

$$P_c = 3.2\%$$

Assume capo

REF (41) p 41 STATES

2 new plants/year

1-2 plants will be modified

Assume that the capacity of a modified plant equals
 THE CAPACITY OF A NEW PLANT. Then

$$P_B = \frac{1.5}{2} (3.2\%) = 2.4\%$$

$$P_B = 2.4\%$$

Assume simple

REF (41) p 39 STATES typical plant capacity = 50 TPD
 " " p 41 STATES 45 operating plants in 1971

Assuming 365 days/yr open

$$(45 \text{ plants}) \left(50 \frac{\text{TONS}}{\text{DAY/PLANT}} \right) \left(365 \frac{\text{DAYS}}{\text{YR}} \right) = \text{Capacity} = 821.3 \times 10^3 \text{ tons}$$

$$K = \frac{\text{Prod}}{\text{Cap}} = \frac{555 \times 10^3}{821.3 \times 10^3} = 0.68$$

$$K = 0.68$$

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Computation Sheet For Industrial Factors

Calculations Done By Horne Date 9/6/74

Source SECONDARY LEAD

$$A = \frac{(555 \times 10^3)}{(0.68)} \times (1.032)^4 = 0.926 \times 10^6 \text{ TONS}$$

$$A_{\text{TOTAL}} = 0.926 \times 10^6 \text{ TONS}$$

SECONDARY LEAD is produced by one of three processes

- (a) POT FURNACES
- (b) BLAST FURNACES (CUPOLAS)
- (c) REVERBATORIUM FURNACES

It appears as though emissions differ for each, so well will break up "A" accordingly & assume K, P_c & P_s to hold for all three cases.

Ref (49) states (p406 TAB 19-1) the following breakdown, in terms of scrapfeed, to each process

POT	53,000	—	7%
BLAST	119,000	—	16%
REVERB.	551,000	—	77%
TOTAL	726,000		

Assuming that this breakdown holds for output as well as input.

$$A_{\text{POT}} = 0.065 \times 10^6 \text{ TONS}$$

$$A_{\text{BLAST}} = 0.118 \times 10^6 \text{ TONS}$$

$$A_{\text{REVERB}} = 0.713 \times 10^6 \text{ TONS}$$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 9/16/74
 Source Secondary Lead

PARTICULATES

BLAST FURNACES

REF (49) p 406 TAB 19-1

$$E_U = 190 \frac{\#}{TON SCRAP}$$

We need this value in terms of lead product, not scrap

REF (49) p 302 STATES

$$\frac{\# LEAD}{\# CHARGE} = 0.70$$

$$190 \frac{\#}{TONSCRAP} \times \frac{1 TON SCRAP}{0.7 TON LEAD} = 271$$

$$E_{UP}_{BLAST} = 271 \frac{\#}{TON LEAD}$$

TOTAL PARTICULATE

REF (44) p 37 STATES THAT THE % OF LEAD IN THE EMISSION IS 6.3%

$$E_{UPB}_{BLAST} = 17.1 \frac{\#}{TON LEAD}$$

REVERBERATORY FURNACES

REF (49) p 406 TAB 19-1

$$E_U = 130 \frac{\#}{TONSCRAP}$$

$$\frac{\# LEAD}{\# CHARGE} = 0.47$$

REF (49) p 300 STATES

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Computation Sheet For Emission Factors

Calculations Done By _____ Date 9/9/74

Source Sintering Lead

$$EV_{P_{REVER}} = \frac{130}{0.47} = 277$$

$$EV_{P_{REVER}} = 277 \frac{\#}{TONLEAD}$$

TOTAL PARTICULATE

FOR 1 LEAD CONCENTRATION OF 63 %

$$EV_{P_{REVER}} = 175 \frac{\#}{TONLEAD}$$

POT

REF (19) pg 06 TAB 19-1

$$EV = 0.8 \frac{\#}{TONSCRAP}$$

$$EV_{POT} = \frac{0.8}{0.59} = 1.36$$

$$EV_{POT} = 1.36 \frac{\#}{TONLEAD}$$

FOR 63% LEAD

$$EV_{P_{POT}} = 0.857 \frac{\#}{TONLEAD}$$

NO % LEAD RELATED
 -SMALL COLLODE
 FOUND IN THE UT -
 EXTRATE.
 ASSUME AVG OF
 OTHER TWO METHODS
 THIS APPEARS
 REASONABLE SINCE
 POT-FURNACES
 FURNISH ONLY 7%
 OF THE TOTAL LEAD

$$\frac{0.7 + .47}{2} = .59$$

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Computation Sheet For Emission Factors

Calculations Done By _____ Date 9/9/74
 Source Secondary Lead

Sulfur Dioxide

Reactorary

Ref (49) p 427 TAB 19-10 STATES

- (A) 0.104% SO₂ for 2500#/hr (scav) plant
- (B) 10,400 SCFM

$$\frac{\text{# SO}_2}{\text{TON LEAD}} = \frac{(0.00104 \frac{\text{FT}^3 \text{SO}_2}{\text{FT}^3 \text{HR}})(10,400 \frac{\text{FT}^3 \text{HR}}{\text{HR}})(530\%) \left(\frac{64 \frac{\text{# SO}_2}{\text{# mole}}}{\text{TON LEAD}}\right) \left(60 \frac{\text{min}}{\text{HR}}\right)}{(192^\circ\text{R})(359 \frac{\text{FT}^3 \text{SO}_2}{\text{# mole}})(2500 \frac{\text{# LEAD}}{\text{HR}})(0.47) \left(\frac{\text{TON LEAD}}{2000 \text{# LEAD}}\right)}$$

$$= 212 \frac{\text{# SO}_2}{\text{TON LEAD}}$$

$$E_{V_{SO_2}} = 212 \frac{\text{#}}{\text{TON LEAD}}$$

Blast Ref (49) p 427 TAB 19-10 STATES

- (A) 0.03% SO₂ for 2670#/hr (exp) plant
- (B) 13,000 SCFM

$$\frac{\text{# SO}_2}{\text{TON LEAD}} = \frac{(.0003)(13000)(530)(64)(60)(2000)}{(492)(359)(2670)(.7)} = 48$$

$$E_{V_{SO_2}} = 48 \frac{\text{#}}{\text{TON LEAD}}$$

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Computation Sheet For Emission Factors

Calculations Done By HOPPER Date 9/9/74
 Source Secondary Lead

No mention of SO_x emissions from pot furnaces could be found in the literature. Therefore, for the present, we will assume

$$E_{V_{SO_2}} = 0 \text{ POT}$$

Carbon Monoxide

REF (41) p 37 STATES

"AN AFTERBURNER IS NOT NEEDED IN THE REVERBERATORY furnace since the excess air and temperature are sufficient to incinerate carbon monoxide and hydrocarbons."

SEE PAGES 14 & 15 OF THESE CALC. FOR ADDITIONAL CO CALCULATIONS

$$E_{V_{CO}} = E_{A/I} = 0 \text{ BLAST}$$

REF (49) "THE QUANTITY OF AIR CONTAMINANTS ... POT FURNACES ... IS MUCH LESS THAN THAT FROM REVERBERATORY OR BLAST FURNACES" p 24 (E_{V_{CO}} = 0) PT

MANY REFERENCES STATE CO EMISSIONS TO BE A PROBLEM IN BLAST FURNACES. "None," HOWEVER, TELL WHAT ONE MIGHT EXPECT IN TERMS OF EMISSION LEVELS.

REF (89) p 97 refers to an A/B AS A CONTROL METHOD. (Many references make the same claims.) Since an A/B IS HARDLY 100% EFFICIENT IN CONVERTING CO → CO₂, WE WILL ASSUME

$$E_{V_{CO/A/B}} = 0$$

$$E_{V_{BLAST}} = ?$$

UNTIL BETTER EV INFO CAN BE DETERMINED

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 12/27/74

Source 2° LEAD

SEE PAGES 15-18 of these calc's for factors regarding NO_x.

9/9/74

Ref (1) p 37 indicates a control efficiency of 97% on an average emission of 35 #/TON_{LEAD} for blast and reverberatory furnaces using a baghouse or high energy scrubber

$$E_N = 35 (1 - .97) = 1.05 \text{ LB/TON}$$

$$E_{NP_BLAST} = 1.05 \text{ #/TON}_{LEAD}$$

TOTAL
PARTICULATE

$$E_{N_BLAST} = 1.05 (0.63) = 0.662$$

$$E_{N_BLAST} = 0.662 \text{ #/TON}_{LEAD}$$

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Computation Sheet For Emission Factors

Calculations Done By HOPPER Date 9/9/74
 Source 2° LEAD

Reverb

$$EN_{REVERB} = 1.05 \frac{\#}{\text{ton LEAD}} \quad \leftarrow \text{TOTAL PARTICULATE}$$

$$EN_{REVERB} = 0.662 \frac{\#}{\text{ton LEAD}}$$

POT

Applying the same efficiency to pot furnaces as
 blast & reverb,

$$EN_{POT} = 1.36(1-.97) = .041$$

$$EN_{POT} = 0.041 \frac{\#}{\text{ton LEAD}} \quad \leftarrow \text{TOTAL PARTICULATE}$$

$$EN_{POT} = .041(.63) = 0.026$$

$$EN_{POT} = 0.026 \frac{\#}{\text{ton LEAD}}$$

SOX Ref(75) p7.1-2 TAB 7.11-1

$$\eta_{SOX_BLAST} = \frac{90 - .8}{90} = 99\% \text{ with NaOH scrubber}$$

214 $EN_{SOX_BLAST} = 48(1-.99) = 0.48 \frac{\#}{\text{ton}}$

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Computation Sheet For Emission Factors

Calculations Done By Hansen Date 9/10/74
 Source SECONDARY LEAD

$$E_{N_{SO_x}} = 0.18 \frac{\#}{ton-LEAD}$$

Assumed same efficiency for reverse furnace SO_2 removal system.

$$E_{N_{SO_x}} = 2.12 \frac{\#}{ton-LEAD}$$

Summary

	<u>EU</u>		<u>EN</u>	
	<u>SL.</u>	<u>REVERSE</u>	<u>SL.</u>	<u>REVERSE</u>
TOTAL PART.	271	277	1.36	1.05
LEAD PART.	171	175	0.857	0.66
SO_x	48	212	[0]	0.48
CO	[?]	0	[0]	[0]
NO_x	[0]	[0]	[0]	[0]

[] NEED DATA TO PROVE OR MORE DEFINITIVE NUMBERS

SEE P 19 FOR A FINAL SUMMARY. DO NOT USE THE ABOVE VALUES.

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 12/27/74
 Source 2° LEAD

It is felt that either a baghouse or high energy scrubber can be retrofit to existing installations. Therefore,

$$E_{III(d)} = E_N$$

$$E_{III(d)} = 0.662 \text{ lb/ton LEAD}$$

Pb
Blast

$$E_{III(d)} = 0.662 \text{ lb/ton LEAD}$$

Pb
Reverb

$$E_{III(d)} = 0.026 \text{ lb/ton LEAD}$$

Pb
Pot

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 12/5/74

Source Secondary Lead

PARTICULATE ES

FOR 1971, Ref(49) p406 gives the following:

Pot 53,000 tons (77%) }
 Blast 119,000 tons (16%) } of scrap
 Reverb 554,000 tons (77%) }

Ref(44) states 45 plants in 1971

Assuming that 7% of the plants use pot furnaces,
 16% use blast furnaces and 77% use reverberatory furnaces,
 the typical hourly production rates are

$$\text{Pot} = \frac{53,000}{(.07)(45)(8760)} = 1.9207 \text{ TPH of scrap}$$

$$\text{Blast} = \frac{119,000}{(.16)(45)(8760)} = 1.8867 \text{ TPH of scrap}$$

$$\text{Reverb} = \frac{554,000}{(.77)(45)(8760)} = 1.8252 \text{ TPH of scrap}$$

CAPACITY

Converting to #/HR, using Refs (84) & (48) (to get the generalized PWR curve)

$$\text{Pot} - 5649 \text{ pph} - 8.6 \text{ #/HR allowable}$$

$$\text{Blast} - 5599 \text{ pph} - 8.4 \text{ " " "$$

$$\text{Reverb} - 5368 \text{ pph} - 8.1 \text{ " " "$$

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Computation Sheet For Emission Factors

Calculations Done By HOPPER Date 12/5/74

Source Secondary LEAD

E_s for Particulates

For BLAST furnaces $\frac{\# \text{LEAD}}{\# \text{charge}} = 0.7$

$$\text{So } \frac{\% \text{ of } 2^{\circ} \text{ LEAD}}{\text{PRODUCT PER PLANT}} = 5649 (.7) = 3954 \%_{\text{HR}} = 1.977 \text{ TPH}$$

 For Reverb furnaces, $\frac{\# \text{LEAD}/\# \text{CHARGE}}{} = .47$

$$= 5549 (.47) = 2608 \%_{\text{HR}} = 1.304 \text{ TPH}$$

 For POT furnaces, $\frac{\# \text{LEAD}/\# \text{CHARGE}}{} = .59$

$$= 5368 (.59) = 3167 = 1.584 \text{ TPH}$$

$$E_{S_{\text{BLAST}}} = \frac{8.6}{1.977} = 4.35 \frac{\# \text{TON}}{\text{LEAD}} \Rightarrow E_{S_{\text{BLAST}} \text{ EXIST}}$$

$$E_{S_{\text{Reverb}}} = \frac{8.4}{1.304} = 6.44 \frac{\# \text{TON}}{\text{LEAD}} \Rightarrow E_{S_{\text{REVERB}} \text{ EXIST}}$$

$$E_{S_{\text{POT}}} = \frac{8.1}{1.584} = 5.11 \frac{\# \text{TON}}{\text{LEAD}} \Rightarrow E_{S_{\text{POT}} \text{ EXIST}}$$

} FOR EXISTING INSTALLATIONS

For new installations, for blast & reverb furnaces, the regulation is 0.022 g/c/dscf (Ref 44 p37).

* Since this is greater than E_u , we will set $E_u = E_s$

$$\therefore (E_{S_{\text{POT}}} = 1.36 \frac{\# \text{TON}}{})$$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 12/5/74
Source 2° LEAD

Ref (44) p 37 indicates that this regulation would correspond to an emission rate of 2.6 #/hr for a 50 TPD plant

$$E_{SP_{new}} = \frac{2.6(24)}{50} = 1.25$$

$$E_{SP_{new}} = 1.25 \text{ #/ton LEAD}$$

Rgs for Pb furnaces are based on opacity only. State control would satisfy the reg'n
 $\therefore E_{S_{new}} = E_{S_{exist}}$
Accuracy Env is below E_S

There are no specific rgs for Pb emissions. However, de facto control of Pb is accomplished through control of particulate emissions. Therefore, since the emission of particulate matter is 63% of Es for Pb would be 63% of the Es for particulates.

$$E_{S_{PART}} = 2.74 \text{ #/ton Pb}$$

$$E_{S_{PART}} = 4.06 \text{ #/ton Pb}$$

$$E_{S_{POT}} = 0.86 \text{ #/ton Pb}$$

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Computation Sheet For Emission Factors

Calculations Done By HOPPER Date 12/5/74

Source 2° LEAD

$$E_s^s_{\text{BLAST}} = 0.788 \frac{\#}{\text{ton Pb}}$$

Pb
New

$$E_s^s_{\text{REVRB}} = 0.788 \frac{\#}{\text{ton Pb}}$$

Pb
New

$$E_s^s_{\text{PT}} = 0.86 \frac{\#}{\text{ton Pb}}$$

Pb
New

E_s summary for Part's & Lead

	<u>PARTICULATE</u>		<u>LEAD</u>	
	<u>EXISTING</u>	<u>NEW</u>	<u>EXISTING</u>	<u>NEW</u>
BLAST	4.35	1.25	2.74	0.79
REVRB	6.44	1.25	4.06	0.79
PT	1.36	1.36	0.86	0.86

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Computation Sheet For Emission Factors

Calculations Done By Hopper

Date 12/13/74

Source 2° LEAD

SO₂ E_s

THREE STATES have SO₂ reg's. which would be applicable to
2° Pb plants (Ref 84 + 148)

ILL. 2000 ppm

IND. 19.5 (P)^{.67}

N.J. 2000 ppm

For Reverb furnaces, the uncontrolled SO₂ emissions are 1040 ppm
(See p 3 of these calcs) ∴ E_s = E_u in ILL & N.J.

For Blast furnaces, the uncontrolled SO₂ emissions are 300 ppm
∴ E_s = E_u in ILL & N.J.

TYPICAL PWR for all furnaces ≈ 1.90 TPH (see p 8)

$$\text{Allow IND} = \frac{19.5(1.90)^{.67}}{(1.90)(.47)} = \frac{(1.534)(19.5)}{(1.90)(.47)} = 33.5 \frac{\text{#SO}_2}{\text{Ton Pb}} (\text{Reverb})$$

$$\begin{aligned} &\text{REVERB} \\ &= \frac{(19.5)(1.90)^{.67}}{(1.90)(.70)} = 22.4 \frac{\text{#SO}_2}{\text{Ton Pb}} (\text{Blast}) \\ &\uparrow \\ &\text{BLAST} \end{aligned}$$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 12/13/74
 Source 2° LEAD

Based on capacity apportioned according to value added from
 Ref 155 for the entire 2° metals industry (sic 331), 5.5%
 of the industry is located in Indiana

So

$$E_{SO_2 \text{ REVERB}} = (.055)(33.5) + 2/2(1-.055) = 200.34 + 1.84$$

$$E_{SO_2 \text{ REVERB}} = 202 \frac{\#}{ton \text{ LEAD}}$$

$$E_{SO_2 \text{ BLAST}} = (.055)(22.4) + 48(1-.055) = 45.36 + 123$$

$$E_{SO_2 \text{ BLAST}} = 46.6 \frac{\#}{ton \text{ LEAD}}$$

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Computation Sheet For Emission Factors

Calculations Done By Happer Date 12/26/74
 Source 2° LEAD

CO

(a) BLAST FURNACES

From Ref (236) p vi-59 "LEAD BLAST FURNACES OR CUPOLAS ARE VERY SIMILAR TO THE CUPOLAS UTILIZED IN THE GRAY IRON FOUNDRY INDUSTRY." WE WILL ASSUME CO EMISSIONS TO BE THE SAME AS FOR GRAY IRON BLAST FURNACES.

REF (75) p 7.10-1 TAB 7.10-1

$$E_{U_{CO}} = 145 \text{ #/TON METAL CHARGED}$$

From pg of these calc's

$$\left(\frac{\# \text{ LEAD}}{\# \text{ CHARGE}} \right)_{\text{BLAST}} = 0.7$$

$$E_{U_{CO}} = \frac{145}{0.7} = 207$$

$$E_{U_{CO}} = 207 \text{ #/TON LEAD}$$

REF (55) p 7.10-1

A WELL - DESIGNED AFTERBURNER CAN CONTROL EMISSIONS TO 9 LB/TON METAL CHARGED

$$9/0.7 = E_{N_{CO}} = 12.86$$

$$E_{N_{CO}} = 12.9 \text{ #/TON LEAD}$$

Since there are no reg's for CO, $E_S = E_U$

$$E_{S_{CO}} = 207 \text{ #/TON LEAD}$$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 12/27/74
 Source 2° LEAD

from p4 of these calc's

$$E_{U_{CO}} = E_{N_{CO}} = 0$$

REVERB SOURCE B

ALSO

$$E_{U_{CO}} = E_{N_{CO}} = 0$$

POT POF

NO_x

(a) BLAST FURNACE

Employing the same assumption of the gray iron blast furnace parallel,

From Ref ⑦ p 18

$$E_{U_{NO_x}} = 0.11 \text{ #/ton}_\text{NET}$$

BLAST

From p4 of these calc's $\frac{\#LENO}{\#CHARGE} = 0.7$

$$E_{U_{NO_x}} = 0.11/0.7 = 0.157$$

BLAST

$$E_{U_{NO_x}} = 0.157 \text{ #/ton}_\text{LENO}$$

No efforts have been made to control NO_x emissions. As we have done in THE PAST, EN will be set EQUAL TO 0.0 TO PROVIDE A mechanism of determining THE NEED FOR CONTROL EQUIPMENT & THE RELATIVE PRIORITY TO OTHER SOURCES FOR WHICH CONTROL HAS ALSO NOT BEEN DEMONSTRATED.

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Computation Sheet For Emission Factors

Calculations Done By _____ Date 12/27/74

Source 2° LEAD

$$E_{N_{NO_x}} = 0$$

From Ref (84), the gen'l reg't for NO_x limitation for solid fuels (coke in this case) is 0.7 #/10⁶ BTU

The heat energy reg'd per ton of lead was not available in the literature directly. However, a material balance for a "THEORETICAL GRAY IRON CUPOLA" is presented on p576 of REF (77). We will use this to det'n the heat reg'ts (approx) to convert the reg into proper units.

268 LB/HR of coke for 2000 LB/HR of metal output

Ref (101) p 9.5 TAB 9-5 Aug of heating value ≈ 14,000 BTU/LB

$$\frac{268 \text{ LB/HR} \times 14,000 \text{ BTU/LB}}{2000 \text{ LB LEAD / HR}} = 1876 \frac{\text{BTU}}{\text{LB LEAD}}$$

$$0.7 \frac{\# \text{ NO}_x}{10^6 \text{ BTU}} \times \frac{1876 \text{ BTU}}{\text{LB LEAD}} \times \frac{2000 \text{ LB LEAD}}{1 \text{ TON LEAD}} = 263 \frac{\text{LB}}{\text{TON LEAD}}$$

Since this is > E_U, E_S = E_U

$$E_{S_{NO_x}} = 0.157 \frac{\# \text{ TON LEAD}}{\text{BLAST}}$$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 12/27/74

Source 2° LEAD

(b) Reverbatory Furnace

Using the grey iron industry analogy

$$\text{REF } \textcircled{7} \text{ p. 18 } E_U = 2.5 \frac{\text{LB}}{\text{TON MELT}}$$

From p 1 of these calc's

$$\frac{\# \text{LEAD}}{\# \text{charge}} = 0.17$$

$$E_U = 2.5 / 0.17 = 5.32$$

$$E_{U_{NOx}} = 5.32 \frac{\text{LB}}{\text{TON LEAD}}$$

REVERB

As we did for blast furnaces, $E_N = 0.0$

$$E_{N_{NOx}} = 0.0$$

REVERB

Since no direct info could be found regarding heat input req'ts for reverb. furnaces. We will assume the same as blast furnaces,
 $1876 \cdot \text{BTU/LB LEAD}$

We will also assume the fuels used are gas & oil & that they are used in a 50/50 proportion.

From Ref 81 the Reg for gaseous fuel is $0.2 \# / 10^6 \text{ BTU}$
 & for liquid fuel is $0.3 \# / 10^6$. The avg is $0.25 \frac{\text{LB}}{10^6 \text{ BTU}}$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 12/27/74
 Source 2^o LEAD

$$(0.25 \frac{\#}{10^6 \text{ BTU}}) (1876 \frac{\text{BTU}}{\#}) (2000) = 0.938 \frac{\#}{\text{TON LEAD}}$$

$$E_{S_{NOx}} = 0.938 \frac{\text{LB}}{\text{TON LEAD}}$$

Reverb

POT FURNACES

As noted on p 4 of these calc's, "The quantity of air contaminants.... pot furnaces.... is much less than that from reverberatory or blast furnaces" We will assume, therefore, that $E_U = E_N = 0$

$$E_{U_{NOx}}_{POT} = E_{N_{NOx}}_{POT} = 0$$

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Emission factors

Calculations Done By HOPER Date 12/27/74

Source 2° LEAD

SUMMARY

PART	<u>EU</u>		<u>EN</u>		<u>ES</u>		<u>ENV(d)</u>	
	BLAST	REVERSE POT	BLAST	REVERSE POT	BLAST	REVERSE POT	BLAST	REVERSE POT
PART	271	277 1.36	1.05	1.05 .041	4.35*	6.41* 1.36*	—	—
NO _X	.157	5.32 Negl.	0.0	0.0 Negl.	.157	.938 —	—	—
SO _X	.48	212 Negl.	.48	2.12 Negl.	46.6	202 —	—	—
CO	207	Negl. Negl.	12.9	Negl. Negl.	207	—	—	—
Pb	171	175 .857	.66	.66 .026	274*	4.06*.86	.66	.66 .026

* Existing. For New, ↓

PART	<u>BLAST</u>	<u>REVERSE</u>	<u>POT</u>
	1.25	1.25	1.36
Pb	.79	.79	.86

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Computation Sheet For Industrial Factors

Calculations Done By Hopper Date 9/10/74

Source Secondary Magnesium

Very little information could be found regarding the secondary magnesium industry.

Ref (95) 233.5020 A states secondary production in 1971

$$\approx 13.42 \times 10^3 \text{ tons}$$

"Growth" has been very erratic

For the purpose of this study, we will assume a straight line growth using data (ref (95)) from 1964-1971

$$\frac{13.42 - 11.79}{(7)} = 0.233 \times 10^3 \text{ tons/yr}$$

$$\frac{13.42 + (0.233 \times 10^3)(1)}{10^3} = 1975 \text{ PROD} = 14.35 \times 10^3 \text{ tons}$$

$$\frac{0.233}{14.35} = 1.6\%$$

$$P_C = 1.6\% \quad (\text{simple-MAX})$$

No direct information regarding "K" could be located OTHER THAN DOC statistics on capacity utilization rates FOR "ADVANCED PROCESS NON-FERROUS METALS." THE AVG K FOR 1965-1973 IS 82%

$$K = 0.82$$

REF (144)

P 48,50

TAB 1 TAB 2

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Industrial Factors

Calculations Done By HOPPER Date 9/10/74
 Source SECONDARY magnesium

$$A = \frac{14.35 \times 10^3}{0.82} = 17.5 \times 10^3$$

$$A = 17.5 \times 10^3 \text{ TONS/yr}$$

" .0175 $\times 10^6$ TON/yr

WAM 9-27-74

P_B: Ref 037 pg 34 Section 33.2

Asset guideline period = 14 yrs
 P_B est \rightarrow 2xIRS = 28 yrs

$$\frac{100\%}{28 \text{ yrs}} \approx 3.5\%/\text{yr simple}$$

$$\therefore P_B = .035 \text{ simple}$$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 9/10/74

Source Secondary Magnesium

Very little information could be found regarding the secondary magnesium industry.

REF (75) P 7.12-1 TAB 7.12-1 gives

$$E_{Mg} = 4 \frac{\#}{\text{TON METAL}} \text{ PROCESSED}$$

Since no info could be found on the % recovery of Mg from scrap, we will assume the same as 100%.
 (See calc'n sheet for secondary Mg)

$$\frac{\# Mg}{\text{TON METAL PROCESSED}} = 0.59$$

$$E_{Mg} = 4 / .59$$

$$E_{Mg} = 6.78 \frac{\#}{\text{TON Mg}}$$

$$E_{Mg} = 0.4 / .59$$

SAME REF

$$E_{Mg} = 0.68 \frac{\#}{\text{TON Mg}}$$

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Computation Sheet For Industrial Factors

Calculations Done By Hopper Date 12/26/74

Source 2^o Mg

Ref (222) pg-29 indicates a typical capacity of about 1 ton
 We will assume this to 2000 of feed scrap

Ref (236) pvi-61 indicates a typical cycle is about 8 hours

$$\therefore PWR = \frac{2000 \text{ LBS}}{8 \text{ hrs}} = 250 \text{ LBS/HR}$$

Since we do not have the specific geographical distribution of plants, we will use the gen'l PWR curve generated from Refs (34) + (48)

$$\text{Allowable} = 0.9 \text{ LBS/HR}$$

$$\text{For a } \cancel{\text{feed}} \text{ product} = \frac{1}{.59} = 1.695$$

$$\frac{250}{1.695} = 147.5 \text{ LBS/HR} = 0.0738 \text{ TONS/HR Mg}$$

$$ES_{P_{Mg}} = \frac{0.9}{0.0738} = 12.2 \%/\text{TON Mg}$$

Since this is > Eu, Es = Eu

$$ES_p = 6.78 \%/\text{TON Mg}$$

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Computation Sheet For Industrial Factors

Calculations Done By Hoppe Date 12/26/74

Source 2^o Mg.

Since no emissions factors could be found for NO_x, CO, SO_x & HC, we will generate them based on the following assumptions:

(a) Emission factors for NO_x + SO_x + CO for boilers (AP-42)

(b) Heat req'ts based on a typical process weight, consisting of pure Mg. being raised in temperature from 70°F to the melting point through melting & up to the pouring temp.

(c) A heat transfer efficiency = 85%

Although the process is 8 hours in length, the furnace will be operating only 6 hours (Ref. 236 p VI-61)

$$Q = \left[(333 \frac{\text{lb}/\text{hr}}{\text{solid}}) \left(C_{P_{Mg}} \frac{\text{Btu}}{\text{lb} \cdot \text{hr}} \right) (T_{MELT} - 70^\circ\text{F}) + (333)(\lambda_w) + (333)(C_{P_{air}}) \frac{(T_{pour} - T_{air})}{(\text{lb}_g)} + (\text{SCFM})(60)(\rho_{air})(C_{P_{air}}) \right] \div (0.85)$$

$$T_{MELT} = 1200^\circ\text{F} \quad (\text{Ref. } 236 \text{ p VI-61})$$

$$T_{POUR} = 1500^\circ\text{F} \quad (\text{Ref. } 232 \text{ p 6-29})$$

$$\rho_{air} = 0.075$$

$$\text{SCFM} = 1100 + 800 / 2 = 950 \text{ SCFM} \quad (\text{Ref. } 236 \text{ p VI-61})$$

From Ref. 101 p 3-118 TAB 3-173

$$C_{P_{Mg}} = \left[5.20 + 0.00133T - \frac{67800}{T^2} \right] \div \text{MW} \quad (24.3)$$

$$C_{P_{Mg}} = 7.4 \div \text{MW}$$

$$\bar{T}_{Ave.} = \frac{(70 + 1200)}{1.8} - 32 + 273 = 608^\circ\text{K}$$

$$\bar{T}_{Ave.} = \frac{(1200 + 1500) \div 2 - 32}{1.8} + 273 = 1005^\circ\text{K}$$

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Computation Sheet For Industrial Factors

Calculations Done By Apper Date 12/26/74

Source 2° Mg.

$$C_{Pmg} = \left[6.20 + .00133(608) - \frac{67800}{608^2} \right] \div 24.3 = 0.2809 \text{ BTU/HR}$$

$.8086$ $.1834$

$$C_{Pmg} = 7.4 \div 24.3 = .3045 \text{ BTU/HR}$$

Ref (101) p 3-108 TAB 3-169

$$\lambda = \frac{(2160)}{24.3} \times 1.8 = 160 \text{ BTU/HR}$$

$$Q = \left[(333)(.2809)(1200-70) + (333)(160) + (333)(.3045)(1500-1200) + (750)(60)(.075)(.14) \right] \div .85$$

$$= \left[1.057 \times 10^5 + 5.328 \times 10^4 + 3.042 \times 10^4 + 1.026 \times 10^3 \right] \div .85$$

$$Q = 2.24 \times 10^5 \text{ BTU/HR}$$

Ref (236) p VI-61 "Natural gas is usually the heat fuel for the pots."

From Ref (75) p 1-4-2 TAB 1-4-1

$$NO_x \text{ EMISSIONS} = 120 \text{ lb/10^6 FT}^3$$

$$EV_{NO_x} = \frac{(2.24 \times 10^5)(120 \times 10^{-6})(2000)}{(1050)(147.5)} = 0.347 \text{ lb NO}_x / \text{ton Mg.}$$

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Emission Factors

Calculations Done By Hoppe Date 12/26/74

Source 2° Mg

$$E_{U_{NO_x}} = 0.347 \text{ #/TON}_Mg$$

As we have done in the past, E_N will be set = 0.0 to provide a mechanism of determining the need for control equip't & the relative priority to other sources for which control has also not been demonstrated.

$$E_{N_{NO_x}} = 0.0$$

From Ref 64, the gen'l reg't for NO_x limitation for gass is $0.2 \text{ #}/10^6 \text{ BTU}$

$$\frac{(0.2)(2.24 \times 10^5)(2000)}{147.5} = 0.607 \text{ #/TON}_Mg$$

Since this is greater than E_U , $E_U = E_S$

$$E_{S_{NO_x}} = 0.347 \text{ #/TON}_Mg$$

CO

Ref 75 P 14-2 TAB 1.4-1

$$CO_{EMISSION} = 20 \text{ #}/10^6 \text{ FT}^3$$

$$E_{U_{CO}} = \frac{(2.24 \times 10^5)(20 \times 10^{-6})(2000)}{(1050)(147.5)} = 0.0579$$

$$E_{U_{CO}} = 0.0579 \text{ #/TON}_Mg$$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 12/26/74
 Source 2⁰ Mg

Since we also have HC emissions, the use of thermal incineration could be an effective control measure. Assuming an $\eta = 99.9\%$.

$$E_{NCO} = (1 - .999)(.0579) \approx 0$$

$$E_{NCO} = 0$$

Since there are no Regs for CO, $E_S = E_U$

$$E_{S_{\infty}} = 0.0579 \text{ #/ton}$$

SO_X

Ref ⑦5 PLT-2 TAB 1.4-1

$$SO_X \text{ emission} = 0.6 \text{#/10}^6 ft^3$$

$$E_{U_{SOX}} = \left(\frac{.6}{20}\right)(.0579) = .0017$$

$$E_{U_{SOX}} = 0.0017 \text{ #/ton}$$

As we did for NO_X, we will set $E_N = 0.0$. Also, it appears that the emissions are too low to be of any consequence.

$$E_{N_{SOX}} = 0.0$$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 12/26/74
 Source 2° Mg

Converting EU to

$$\frac{(.0017)(147.5)}{(2000)(2.24 \times 10^5)} = .0006 \text{ #/ton hr}$$

↑ Since this is far less than any State Reg for SO₂, E_S = EU

$$E_{S_{SO_2}} = 0.0017 \text{ #/ton hr}$$

H/C The scrap used in melting furnaces contains organic mat'l which results in the emissions of hydrocarbons during the first part of the operation. No mention of emissions could be found in the literature, however for the purpose of evaluating the emission impact of controls, we will make the following unqualified assumption.

$$\frac{\# \text{ organic mat'l}}{1000 \# \text{ FURNACE FEED}}$$

Since the FEED/Product Ratio = 1/.59 = 1.695

$$EU_{HC} = (1.695)(2) = 3.39 \text{ #/ton}$$

$$EU_{HC} = 3.39 \text{ #/ton}$$

Since we have chosen thermal incineration for CO emissions with an assumed η = 99.9%

$$EN_{CO} = (1-.999)(3.39) = 0.0034$$

$$EN_{CO} = 0.0034 \text{ #/ton}$$

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Computation Sheet For Industrial Factors

Calculations Done By Hoppe Date 12/26/74

Source 2^o Mgr.

Since there are no reg's for H/C $E_S = E_U$

$$E_{S_{HC}} = 3.39 \text{ #/ton mg}$$

Summary

	<u>E_U</u>	<u>E_N</u>	<u>E_S</u>
P _r	.78	.68	.78
NO _x	.347	0.0	.347
SO _x	.0017	0.0	.0017
CO	.058	0	.058
HC	3.39	.0094	3.39

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Industrial Factors

Calculations Done By K.Tower Date 1/22/75
 Source Steel Foundries

Steel foundries produce steel castings by melting steel metal and pouring it into molds. The melting of steel for castings is accomplished primarily in one of three types of furnaces: direct electric arc, electric induction and open hearth. Specific information regarding the fractional utilization of these furnaces could not be found. We assumed K to be equal to that for the Iron + Steel Industry → 0.90 (See "Industrial Factors, Iron + Steel Industry , p. 12.)

$$K = 0.90$$

The growth rate for the Steel foundry industry was assumed to be that for direct electric arc furnaces of the Iron + Steel industry (See "Industrial factors, Iron + Steel Industry, p/12) → .073. Specific information concerning the growth rate for electric induction furnaces could not be found. In addition the open hearth furnace from the Iron + Steel Industry has a negative growth rate, so we assumed its negative growth would be absorbed by the other furnaces.

$$P_c = .073$$

simple

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Industrial Factors

Calculations Done By K. Tower Date 1/22/75

Source Steel Foundries

Ref 037, p.34 section 33.0 states that the asset guideline period for the manufacture of primary metals (including assets used in the smelting of ferrous metals) is 18 years.

Assuming twice the IRS allowable

$$P_B = \frac{1}{(2)(18)} = .028$$

$$\boxed{P_B = .028}$$

Ref 236, p. VI-63 indicates there are 400 steel foundries operating in the U.S with plant sizes ranging from 25 to 240 tons/day operating 24 hrs/day, 7 days/wk. for large plants and 8 hrs/day for small plants. We assume the average steel foundry is the average of 25 and 240 tons/day or 132.5 tons/day. Since this reference was dated October, 1971 we assume the data represents 1970 values.

$$\text{Prod}_{1970} = 132.5 \frac{\text{TONS}}{\text{DAY}} \times 365 \frac{\text{days}}{\text{yr}} \times 400 \text{plants} = 19.3 \times 10^6 \frac{\text{TONS}}{\text{YR}}$$

$$A = \frac{\text{Prod}_{1970}}{K} \left(1 + (1975-1970) P_C \right)$$

$$= \frac{19.3 \times 10^6 \frac{\text{TONS}}{\text{YR}}}{0.90} \left(1 + 5(.073) \right) = 29.3 \times 10^6 \frac{\text{TONS}}{\text{YR}}$$

$$\boxed{A = 29.3 \times 10^6 \frac{\text{TONS}}{\text{YR}}}$$

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Emission Factors

Calculations Done By K. Tower Date 1/22/75
 Source Steel foundries

Ref 222^{p-11, table 2-3} gives the following table for emission factors of Steel Foundries.

Type of Process	Uncontrolled Emissions ($\frac{\text{lb}}{\text{ton processed}}$)	
	Particulate	NO_x
Electric Arc Furnace	13	0.2
Open Hearth	11	0.013
Induction Electric	0.1	—

Ref. 222, p. 6-11 also states the following:

Electric induction furnaces are generally small in capacity, usually $\frac{1}{4}$ to 1 ton in size. Since open hearth furnaces have a negative growth rate as previously referenced we assume that by 1985 there will be no open hearth furnaces. We also assume that electric induction furnaces contribute only 20% to the total capacity of Steel foundries.

$$E_{u_{P_t}} = (.80)(13 \frac{\text{lbs}}{\text{ton}}) + (.20)(0.1 \frac{\text{lbs}}{\text{ton}}) = 10.4 + .02 = 10.42 \frac{\text{lbs}}{\text{ton}}$$

$$E_{u_{P_t}} = 10.4 \frac{\text{lbs}}{\text{ton}}$$

$$E_{u_{NO_x}} = 0.2 \frac{\text{lbs}}{\text{ton}}$$

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Computation Sheet For Emission Factors

Calculations Done By K. Tower Date 1/22/75
 Source Steel Foundries

Since the majority of Steel Foundries utilize either the electric arc furnace or the electric induction furnace and no combustion of fuel is involved we assume there are no hydrocarbon or SO_x emissions.

Ref 222, p. 6-14 indicates that a 99% control efficiency can be expected of particulate emissions with the use of a baghouse.

$$E_{N_{P_f}} = (.01) \times (10.4 \frac{\text{lbs}}{\text{TON}}) = .104 \frac{\text{lbs}}{\text{TON}}$$

$$E_{N_{P_f}} = .104 \frac{\text{lbs}}{\text{TON}}$$

Since^{there} is no available technology for the control of NO_x emissions we assume a 100% control to determine maximum potential impact from control technology

$$E_{N_{NO_x}} = 0.00 \frac{\text{lbs}}{\text{TON}}$$

E_s

The average process weight rate for a steel foundry is as follows:

$$\frac{(132.5 \frac{\text{TONS}}{\text{DAY}})(2000 \frac{\text{lbs}}{\text{TON}})}{24 \frac{\text{HRS}}{\text{DAY}}} = 1.1 \times 10^4 \frac{\text{lbs}}{\text{hr}}$$

We assume the average steel foundry operates 24 hrs/day.

From the general process weight curve Ref 084 & Ref 143 the allowable emissions from an average steel foundry is $12.2 \frac{\text{lbs}}{\text{hr}}$.

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Computation Sheet For Emission Factors

Calculations Done By K.Tower Date 1/23/75
 Source Steel foundries

Since no specific information could be found regarding the geographical distribution of steel foundries in the U.S. we assume they are equally distributed among all the states.

$$E_{S_{Pr}} = \frac{(12.2 \frac{hr}{day})(24 \frac{hrs}{day})}{132.5 \frac{TON}{DAY}} = 2.2 \frac{lbs}{TON}$$

$$E_{S_{Pr}} = 2.2 \frac{lbs}{TON}$$

Since we assume that all steel foundries will use either an electric arc furnace or an electric induction furnace by 1985 they will ^{not} come under any NO_x regulations. Therefore $E_{S_{NO_x}} = E_{u_{NO_x}}$.

$$E_{S_{NO_x}} = 0.2 \frac{lbs}{TON}$$

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Computation Sheet For Emission Factors

Calculations Done By K. Tower Date 1/23/75
 Source Steel Foundries

Summary of Emission Factors

	E_4	E_N	E_S
Particulate	10.4 lb_s/ton	.104 lb_s/ton	2.2 lb_s/ton
NO_x	0.2 lb_s/ton	0.00 lb_s/ton	0.2 lb_s/ton
SO_x	—	—	—
Hydrocarbon	—	—	—

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Computation Sheet For Industrial Factors

Calculations Done By Harrer Date 9/10/74
 Source Secondary Zinc

Ref (95) 233,6832 Prod 1969 = 376.4×10^3 TONS
 GROWTH between 1960-1969

$$P_c = \sqrt[9]{\frac{376.4}{265.8}} - 1.0 = 3.94\%$$

$P_c = 3.9\%$ compound

$$A_{prod} = 376.4 \times 10^3 (1.039)^6$$

$$A = 473.5 \times 10^3 \text{ tons}$$

No information could be found for K specifically FOR THIS INDUSTRY. WE WILL USE THE CAPACITY UTILIZATION VALUE FOR "ADVANCED PROCESS NON-FERROUS METALS." THE AVERAGE K FOR 1965-1973 WAS 82%

$$K = 0.82$$

$$A = 473.5 \times 10^3 / 0.82$$

$$A = 577.4 \times 10^3 \text{ tons}$$

Ref (144)
 P48-TM81
 P50-TM82

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Computation Sheet For Industrial Factors

Calculations Done By _____ Date _____

Source _____

AS A RESULT OF THE APPARENTLY DIFFERENT EMISSION FACTORS FROM THE TWO BASIC ZINC PRODUCING PROCESSES (SWEATING & DISTILLATION), "A" WILL BE DIVIDED ACCORDING TO THE RATIO OF ZINC PRODUCED BY EACH PROCESS.

Ref (4) p106

IN 1968 233,000 TONS ZN RECOVERED IN DISTILLATION FURNACES

Ref (95) 233,6832

IN 1968 354,700 TONS OF SECONDARY ZN

$$\therefore \text{1968 ZINC BY SWEATING} = 354,700 - 233,000 = 121,700$$

% BY DISTILL = 66%

% BY SWEAT = 34%

$$A_{\text{sweat}} = 196.3 \times 10^3 \text{ TONS ZN}$$

$$A_{\text{distill}} = 381.1 \times 10^3 \text{ TONS ZN}$$

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Computation Sheet For Industrial Factors

Calculations Done By 2J. Marrone Date 9-27-74

Source Secondary Zinc

P_B :

We will assume that the obsolescence rate will be identical for sweating and distillation surfaces and that this factor may be estimated from IRS Tables

Ref 037 pg 34 Section 33.2

Asset guideline period 14 yrs

$$P_B \text{ est} = 2 \times \text{IRS} = 28 \text{ yrs}$$

$$\frac{100\%}{28 \text{ yrs}} \cong 3.5\%/\text{yr simple}$$

∴

$$P_B = .035 \text{ simple}$$

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Computation Sheet For Emission Factors

Calculations Done By Homer Date 2/10/74
 Source Secondary Zinc

Ref (75) P 7.14-2 TAB 7.14-1 Precursors

Smelting Process

Metallic scrap - 11 #/ton FROM KETTLE SMELT FURNACE
 - 13 #/ton FROM REVERSED SMELT FURNACE
 AVG = 12 #/ton

Assume the average = 12#/ton

Residual Scrap

25#/ton	- KETTLE
32#/ton	- REVERB
AVG = 28.5#/ton	

Assume the average = 28.5#/ton

Ref (49) p 106 TAB 19-1 gives 1968 breakdown

METALLIC	=	52,000 TONS	=	30%
RESIDUAL	=	119,000 TONS	=	70%
				171,000 TONS

$$E_{UP_{SMELT}} = (.3)(12) + .7(28.5) = 3.6 + 19.95 = 23.55$$

$$E_{UP_{SMELT}} = 23.6 \frac{\#}{TON ZN}$$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 9/10/74
 Source Secondary Zinc

Distillation Furnaces

REF (75) P 7.14-2 TAB 7.14-1

Refiner - 47#/ton

Horizontal Muffle - 45#/ton AVG = 46#/ton

$$EN_{\text{distill}} = 16 \text{#/ton Zn}$$

Ref (49) p 135 gives $\eta = 96.3\%$ for baghouse on a smelting furnace

$$EN_{\text{smelt}} = 236 (1-.963) = 0.872$$

$$EN_{\text{smelt}} = 0.872 \text{#/ton Zn}$$

Assuming that comparable efficiencies can be obtained on distillation furnaces

$$EN_{\text{distill}} = 16(1-.963) = 1.70 \text{#/ton Zn}$$

$$EN_{\text{distill}} = 1.70 \text{#/ton Zn}$$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 9/10/74
Source Secondary Zinc

No information could be found on NO_x emissions.

No emission data could be found for CO & HC; however,
REF 49 p434 STATES CONTROL OF THESE EMISSIONS CAN BE
ACHIEVED WITH AFTERBURNING OR INCINERATION FOR "COMBUSTIBLE
MATTER DISCHARGED FROM A LOW-TEMPERATURE SWEATING
FURNACE".

As a general rule, this will result in a controlled emission
essentially EQUAL TO ZERO

$$\therefore EN_{CO} = 0 \\ SWATT$$

$$EN_{HC} = 0 \\ SWATT$$

HOWEVER

$$EU_{CO} = ? \\ SWATT$$

$$EU_{HC} = ? \\ SWATT$$

IT CAN BE INFERRED FROM REF 49 p431 THAT THERE
ARE NO CO EMISSIONS FROM DISTILLATION FURNACES DUE TO HIGH
TEMPS & THUS THE OXIDATION OF CO TO CO₂

$$\therefore EU_{CO} = EN_{CO} = 0 \\ DISTILL$$

THE SCRAP USED IN SWEATING FURNACES CONTAINS ORGANIC
MATERIALS WHICH RESULTS IN THE EMISSION OF HYDROCARBONS.
HOWEVER, THE ZINC CHARGED TO DISTILLATION FURNACES IS
"RELATIVELY PURE" BY COMPARISON. WE WILL, THEREFORE,
ASSUME

$$EN_{HC} = EU_{HC} = 0 \\ DISTILL$$

Project Number - 32391 New Source Performance Standards

Computation Sheet For Emission Factors

Calculations Done By Hoppe Date 9/11/74
Source Secondary Zinc

Summary:

	<u>E_N</u>		<u>E_U</u>	
	<u>DISTILL</u>	<u>SWEAT</u>	<u>DISTILL</u>	<u>SWEAT</u>
Pr	1.70	0.872	16.0	23.6
CO	0	[0]	0	?
HC	0	[0]	0	?
NO _x				

SEE p 12 of those Cars for an
UPDATED Summary

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Computation Sheet For Emission Factors

Calculations Done By UF RUSSO Date 10/29/74
 Source Secondary Zinc

Ref. 157 gives the number of establishments in the Zinc Industry as 17 plants in 1967.*
 Assuming the #plants are in same ratio as production i.e. assume same average capacity

Sweating

$$\begin{aligned} \text{Output Process Wgt Rate} &= \frac{A_{\text{sweating}} (\text{lbs/yr})}{.34 (1 + P_c)^8 N_p (8760 \text{ hrs/year})} \\ &= \frac{(196.3 \times 10^3)(2000)}{.34 (1 + 0.039)^8 (8760) N_p} \\ &\approx \frac{97066}{17} \text{ lbs/hr} \\ &\approx 5700 \text{ lbs/hr} \end{aligned}$$

Assuming the same input wgt./output wgt. relationship as for Secondary copper sweating

$$\begin{aligned} \text{Input Process Wgt Rate} &= \left(\frac{38,400}{64,000} \right) (5700) \text{ lbs/hr} \\ &= 9515 \text{ lbs/hr} \end{aligned}$$

* Plants in 2nd non-ferrous industry's were apportioned according to 1967 production

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Computation Sheet For Emission Factors

Calculations Done By KF. Russo Date 10/29/74
 Source Secondary Zinc

Using the 50 State Average Process wgt. Rate,
 the allowable Emissions are

$$\text{Allowable Emissions} = 11.0 \text{ lbs Part/HR}$$

Thus.

$$E_S^{\text{Secondary}} = \frac{11.0 \text{ lbs Part/HR}}{5700 \text{ lbs/HR}} \times 2000 \text{ lbs/Ton}$$

$$E_S^{\text{Secondary}} = 3.86$$

lbs Part
Ton 2nd day Zinc

Distillation Furnaces.

Assuming input wgt/output wgt. relationship
 is same as for copper blast furnaces.

$$\text{Input Process wgt.Rate} = 15,830 \text{ lbs/HR}$$

Using 50 state average:

$$\text{Allowable Emissions} = 15.0 \text{ lbs Part/HR}$$

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Computation Sheet For Emission Factors

Calculations Done By Russo Date 10/29/74

Source Zinc

Thru

$$E_S^{\text{Dist.}} = \frac{15.0 \text{ lbs Part/Hr}}{11,080 \text{ lbs/Hr}} \times \frac{2000 \text{ lbs/Ton}}{1 \text{ Ton}}$$

$$E_S^{\text{Dist.}} = 2.71 \text{ lbs Part/Ton 2nd any Zinc}$$

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Project Number - 32391 New Source Performance Standards

Computation Sheet For Emission Factors

Calculations Done By Hopper Date 10/24/74
 Source 2° Zinc

Since no emission factors could be located for gaseous emissions,
 we will generate them based on the following assumptions:

- (a) Emission factors for NO_x , CO for boilers (12.42)
- (b) Heat ex'ts based on a typical PWR consisting
 of pure Zn being raised in temperature from 70°F
 to the melting point, through melting & up to the
 pouring temperature
- (c) A heat transfer efficiency = 85%

Ref (157) gives the No. of 2° Zn establishments = 17 (1967)

$$1969 = 1967 (1 + 0.039)^2$$

$$\frac{376.4 \times 10^3}{(1.039)^2} = 1967 \text{ Pwrs.} = 348.7 \times 10^3 \text{ Pwrs}$$

$$\frac{348.7 \times 10^3}{17} = 20,510 \frac{\text{tons}}{\text{yr. plant}}$$

From Ref (236) pvi-66, of single operation lists from 10-12 hrs
 Assume ∴, # hours operation per year = $8760/2 = 4380$ hrs.

$$\frac{20,510 \text{ tons yr (2000)}}{4380 \text{ hrs}} = 9365 \text{ #/hr} \quad \text{Ref (236) same page}$$

falls of $\pm 10,000$ to/me plant; so, this value appears reasonable.

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 Source 2^o ZINC

From Ref ④ p 406 TAB 19-1 gives 1968 scrap input
 @ 171,000 tons (swelling)

From PQ of these calculations 1968 production (swelling)
 = 121,700 tons

$$\frac{\text{Feed}}{\text{Product}} = \frac{171,000}{121,700} = 1.405$$

$$\text{INPUT} = (1.405)(9365) = 13,157 \text{ lb/hr}$$

$$\begin{aligned} \text{Heat Reg}^t &= (13,157 \text{ lb/hr}) \left(C_{P_{\text{air}}} \frac{870}{18^{\circ}\text{F}} \right) (T_{\text{out}} - 70^{\circ}\text{F}) + (13,157) (\lambda_{\text{Zn}}) \\ &\quad + (13,157) \left(C_{P_{\text{Zn}}} \right) (T_{\text{out}} - T_{\text{out}}) \\ &\quad + (\text{SCFM})(60) \left(\rho_{\text{air}} \right) \left(C_{P_{\text{air}}} \right) \div 0.85 \end{aligned}$$

From Ref ②36 p II-66 SCFM = 20-30 per lb/hr (Assume 25)

$$(13,157)(25) \approx 329,000 \text{ SCFM}$$

$$C_{P_{\text{air}}} = 0.24 \text{ BTU/lb.°F}$$

$$\rho = 0.075 \text{ lb/ft}^3$$

$$\left. \begin{array}{l} T_{\text{out}} \approx 800^{\circ}\text{F} \\ T_{\text{out}} \approx 100^{\circ}\text{F} \end{array} \right\} \text{Ref } ②36 \text{ p II-66}$$

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Calculations Done By Hopper Date 12/26/74

Source Z° Zinc

Ref(101) P 3-122 TAB 3-173

$$\rho_{\text{Zn}} = 5.25 + .00270 \bar{T}^{\circ}\text{K} \div \text{mw} \quad (65.4)$$

$$\rho_{\text{Zn}} = 7.59 + .00055 \bar{T}^{\circ}\text{K} \div \text{mw}$$

$$\frac{T_{\text{avg}}}{S} = \frac{(70^{\circ} + 800^{\circ}\text{F}) - 32}{1.8} + 273 = 738^{\circ}\text{K}$$

$$\frac{T_{\text{avg}}}{L} = \frac{(800 + 1100) - 32}{1.8} + 273 = 1311^{\circ}\text{K}$$

$$\rho_{\text{Zn}} = .1107 \text{ BRU}/\#^{\circ}\text{F}$$

$$\lambda = 1595 \frac{\text{cal}}{\text{mole}}$$

Ref(101)
 P 3-109
 TAB 3-169

$$\rho_{\text{Zn}} = .1271 \text{ BRU}/\#^{\circ}\text{F}$$

$$\pi = \frac{1595}{65.4} \times 1.8 = 43.9 \text{ BRU}/\#$$

$$Q = \left[(13,157)(.1107)(800 - 70) + (13,157)(43.9) + (13,157)(.1271)(1100 - 800) \right. \\ \left. + (929,000)(60)(.015)(.24) \right] \div .85$$

$$= \left[1,063 \times 10^6 + .578 \times 10^6 + .502 \times 10^6 + .355 \right] \div .85$$

$$Q = 2.91 \times 10^6 \text{ BRU}/\text{HR}$$

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Calculations Done By Hoppe Date 12/26/74
 Source 2° Zinc

Ref (244) p II-6 indicates primary fuels are natural gas
 to oil & "the fuel used most is natural gas".

We will assume 100% to simplify the calc's..

NO_x

From Ref (75) p 1.4-2 TAD 1.4-1

$$\text{NO}_x \text{ emissions} = 120 \frac{\text{MM}}{10^6 \text{ FT}^3}$$

$$\frac{2.99 \times 10^6 \frac{\text{BTU}}{\text{HR}}}{1050 \frac{\text{BTU}}{\text{FT}^3}} = 2800 \frac{\text{FT}^3}{\text{HR}}$$

Ref (75)
 P 1.4-1

$$(2800 \frac{\text{FT}^3}{\text{HR}}) \left(20 \times 10^6 \frac{\#}{\text{FT}^3} \right) = .336 \frac{\#}{\text{HR}}$$

$$E_{\text{NO}_x} = \frac{(0.336)(2000)}{(9365)} = 0.0718 \frac{\#\text{NO}_x}{\text{TON ZN}}$$

$$E_{\text{NO}_x} = 0.0718 \frac{\#\text{NO}_x}{\text{TON ZN}}$$

Assume the same for distillation

$$E_{\text{NO}_x} = 0.0718 \frac{\#\text{NO}_x}{\text{TON ZN}}$$

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 Source 2^o Zinc

As we have done in the past, E_N will be set = 0.0 to provide a mechanism of determining THE NEED FOR CONTROL EQUIP'T + the relative priority to OTHER SOURCES FOR WHICH CONTROL HAS ALSO NOT BEEN DEMONSTRATED

$$E_{N_{NO_x}} = 0.0 \\ \text{SUGAR}$$

$$E_{N_{NO_x}} = 0.0 \\ \text{DISTILL.}$$

From Ref ⑧4 the gen'l reg't for NO_x limitation for gas is $0.2 \text{#/10}^6 \text{ BTU}$

$$\frac{0.2 \text{#/10}^6 \text{ BTU} \times 2.94 \times 10^6 \frac{\text{BTU}}{\text{HR}} \times 2000}{9365} = 0.126 \text{ #/TONZN}$$

9365

Since this is greater than E_U , $E_U = E_S$

$$E_{S_{NO_x}} = 0.0718 \text{ #/tonZN} \\ \text{SUGAR}$$

$$E_{S_{NO_x}} = 0.0718 \text{ #/tonZN} \\ \text{DISTILL.}$$

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Computation Sheet For Emission Factors

Calculations Done By Hopper Date 12/26/74

Source 20% Zinc

CO

Ref(75) p 1.4-2 TAB 1.4-1

CO Emission = $20 \text{ #}/10^6 \text{ ft}^3$

$$E_{U_{CO}} = \frac{(2800)(20 \times 10^{-6})(2000)}{9365} = 0.012$$

$$E_{N_{CO}} = 0.012 \text{ #/ton ZN}$$

p 8 of these calc's discusses the use of incineration to control combustible matter from a sweating furnace

Assuming AN R = 99.9%

$$E_{N_{CO}} = 0.012(1 - .999) \approx 0.0$$

$$E_{N_{CO}} = 0$$

Since there are no reg's. for CO, $E_U = E_S$

$$E_{S_{CO}} = 0.012 \text{ #/ton ZN}$$

H/C

The scrap used in sweating furnaces contains organic mat's which results in the emission of hydrocarbons. No mention of emissions could be found in the literature.

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Computation Sheet For Emission Factors

Calculations Done By Hoyer Date 12/26/74
 Source 2# Zinc

For the purpose of evaluating the emission impact, we will make the following unqualified assumption

$$1\text{# HYDROCARBON} / 1000 \text{# FEED MAT'L} = 2\text{# TON FEED}$$

Since $\text{t/feed} / \text{product ratio} = 1.405$

$$E_{UHC} = \frac{2\text{#/TON}_{feed}}{\text{SWAT}} \times 1.405 \frac{\text{TONFEED}}{\text{TON ZN}} = 2.81$$

$$E_{UHC} = 2.81 \text{#/TON}_{ZN}$$

Since we are using incineration with an assumed $\eta = 99.9\%$

$$E_{UHC} = 2.81 \left(1 - .999\right) = 0.0028$$

$$E_{NHC} = 0.0028 \text{#/TON}_{ZN}$$

Since there are no H/C Regs $E_S = EU$

$$E_{SHC} = 2.81 \text{#/TON}_{ZN}$$

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Computation Sheet For Emission Factors

Calculations Done By Hoppe Date 12/26/74

Source 2°ZN

Summary

	<u>Eu</u>		<u>En</u>		<u>Es</u>	
	<u>distill.</u>	<u>swat</u>	<u>distill</u>	<u>swat</u>	<u>distill</u>	<u>swat</u>
Pt	46.0	23.6	1.70	.872	2.71	3.86
CO	—	.012	—	0	—	.012
HC	—	2.81	—	.0028		2.81
NOx	.0718	.0718	0.0	0.0	.0718	.0718

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(Please read Instructions on the reverse before completing)

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16. ABSTRACT The purpose of this document is to present the results of a study to determine the impact of new source performance standards on nationwide emissions. The work presented covers 14 potential pollutants from approximately 200 source categories for the year 1985. The results are being used by EPA as input to the development of an overall standard setting strategy. The report contains information regarding controlled and uncontrolled emission factors, State emission limitations, industrial capacity, utilization, growth and retirement rates. The results of this study have been published as three volumes which encompass ten separate documents. This document contains Appendix 4E of Volume II - Calculation Sheets for the Metallurgical Industry.			
17. KEY WORDS AND DOCUMENT ANALYSIS			
a. DESCRIPTORS	b. IDENTIFIERS/OPEN ENDED TERMS	c. COSATI Field/Group	
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