United States Environmental Protection Agency Office of Water Regulations and Standards (WH-552) Industrial Technology Division Washington, DC 20460

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Development FINAL Document for Effluent Limitations Guidelines and Standards for the Nonferrous Metals Manufacturing Point Source Category

Volume V Primary Precious Metals and Mercury Secondary Precious Metals Secondary Silver Secondary Mercury



ORGANIZATION OF THIS DOCUMENT

This development document for the nonferrous metals manufacturing category consists of a general development document which considers the general and overall aspects of the regulation and 31 subcategory specific supplements. These parts are organized into 10 volumes as listed below.

The information in the general document and in the supplements is organized by sections with the same type of information reported in the same section of each part. Hence to find information on any specific aspect of the category one would need only look in the same section of the general document and the specific supplements of interest.

The ten volumes contain contain the following subjects:

- Volume I General Development Document
- Volume II Bauxite Refining Primary Aluminum Smelting Secondary Aluminum Smelting
- Volume III Primary Copper Smelting Primary Electrolytic Copper Refining Secondary Copper Refining Metallurgical Acid Plants
- Volume IV Primary Zinc Primary Lead Secondary Lead Primary Antimony
- Volume V Primary Precious Metals and Mercury Secondary Precious Metals Secondary Silver Secondary Mercury
- Volume VI Primary Tungsten Secondary Tungsten and Cobalt Primary Molybdenum and Rhenium Secondary Molybdenum and Vanadium
- Volume VII Primary Beryllium Primary Nickel and Cobalt Secondary Nickel Secondary Tin
- Volume VIII Primary Columbium and Tantalum Secondary Tantalum Secondary Uranium
- Volume IX Primary and Secondary Titanium Primary Zirconium and Hafnium
- Volume X Primary and Secondary Germanium and Gallium Primary Rare Earth Metals Secondary Indium

DEVELOPMENT DOCUMENT

for

EFFLUENT LIMITATIONS GUIDELINES AND STANDARDS

for the

NONFERROUS METALS MANUFACTURING POINT SOURCE CATEGORY

VOLUME V

Primary Precious Metals and Mercury Secondary Precious Metals Secondary Silver Secondary Mercury

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NONFERROUS METALS MANUFACTURING POINT SOURCE CATEGORY

DEVELOPMENT DOCUMENT SUPPLEMENT

for the

Primary Precious Metals and Mercury Subcategory

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SECTION I

SUMMARY

This document provides the technical basis for promulgating effluent limitations based on best practicable technology (BPT) and best available technology (BAT) for existing direct dischargers, pretreatment standards for new indirect dischargers (PSNS), and standards of performance for new source direct dischargers (NSPS) for plants in the primary precious metals and mercury subcategory.

The primary precious metals and mercury subcategory is comprised of eight plants. Of the eight plants, one discharges directly to rivers, lakes, or streams; none discharge to publicly owned treatment works (POTW); and seven achieve zero discharge of process wastewater.

EPA first studied the primary precious metals and mercury subcategory to determine whether differences in raw materials, final products, manufacturing processes, equipment, age and size of plants, and water usage, required the development of separate effluent limitations and standards for different segments of the subcategory. This involved a detailed analysis of wastewater discharge and treated effluent characteristics, including the sources and volume of water used, the processes used, the sources of pollutants and wastewaters in the plant, and the constituents of wastewaters, including priority pollutants. As a result, nine subdivisions have been identified for this subcategory that warrant separate effluent limitations. These include:

- 1. Smelter wet air pollution control,
- 2. Silver chloride reduction spent solution,
- 3. Electrolytic cells wet air pollution control,
- 4. Electrolyte preparation wet air pollution control,
- 5. Calciner wet air pollution control,
- 6. Calciner quench water,
- 7. Calciner stack gas contact cooling water,
- 8. Condenser blowdown, and
- 9. Mercury cleaning bath water

Several distinct control and treatment technologies (both inplant and end-of-pipe) applicable to the primary precious metals The Agency analyzed and mercury subcategory were identified. both historical and newly generated data on the performance of nonwater quality including their technologies, these impacts and air quality, solid waste generation, environmental and energy requirements. EPA also studied various flow reduction techniques reported in the data collection portfolios (dcp) and plant visits.

Engineering costs were prepared for each of the control and

treatment options considered for the subcategory. These costs were then used by the Agency to estimate the impact of implementing the various options on the subcategory. For each control and treatment option that the Agency found to be most effective and technically feasible in controlling the discharge of pollutants, the number of potential closures, number of employees affected, and impact on price were estimated. These results are reported in a separate document entitled "The Economic Impact Analysis of Effluent Limitations and Standards for the Nonferrous Metals Manufacturing Industry."

After examining the various treatment technologies, the Agency has identified BPT to represent the average of the best existing technology in the nonferrous metals manufacturing industry. Metals removal based on chemical precipitation and sedimentation and ion exchange technology is the basis for the BPT limitations. Oil skimming was selected as the technology basis for oil and grease limitations. To meet the BPT effluent limitations based on this technology, the primary precious metals and mercury subcategory is expected to incur a capital cost of \$42,200 and an annual cost of \$26,800.

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For BAT, the Agency has built upon the BPT technology basis by adding in-process control technologies which include recycle of process water from air pollution control waste streams. Filtration is added as an effluent polishing step to the end-ofpipe treatment scheme. To meet the BAT effluent limitations based on this technology, the primary precious metals and mercury subcategory is estimated to incur a capital cost of \$43,025 and an annual cost of \$27,300.

NSPS are equivalent to BAT. In selecting NSPS, EPA recognizes that new plants have the opportunity to implement the best and most efficient manufacturing processes and treatment technology. As such, the technology basis of BAT has been determined as the best demonstrated technology.

EPA is not promulgating PSES for the primary precious metals and mercury subcategory because there are no indirect dischargers. For PSNS, the Agency selected end-of-pipe treatment and inprocess flow reduction control techniques equivalent to NSPS.

The best conventional technology (BCT) replaces BAT for the control of conventional pollutants. BCT is not being promulgated because the methodology for BCT has not yet been finalized.

The mass limitations and standards for BPT, BAT, NSPS, and PSNS are presented in Section II.

SECTION II

CONCLUSIONS

EPA has divided the primary precious metals and mercury subcategory into nine subdivisions or building blocks for the purpose of effluent limitations and standards. These subdivisions are:

- (a) Smelter wet air pollution control,
- (b) Silver chloride reduction spent solution,
- (c) Electrolytic cells wet air pollution control,
- (d) Electrolyte preparation wet air pollution control,
- (e) Calciner wet air pollution control,
- (f) Calcine quench water,
- (g) Calciner stack gas contact cooling water,
- (h) Condenser blowdown, and
- (i) Mercury cleaning bath water.

BPT is promulgated based on the performance achievable by the application of chemical precipitation and sedimentation technology (lime and settle) and ion exchange as a polishing step technology along with preliminary treatment consisting of oil skimming for selected waste streams. The following BPT effluent limitations are promulgated:

(a) Smelter Wet Air Pollution Control BPT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of gol	d and silver sme	lted
Lead	0.546	0.260
Mercury	0.325	0.130
Silver	0.533	0.221
Zinc	1.898	0.793
Gold	0.130	
Oil and grease	26.000	15.600
Total suspended solids	53.300	25.350
pH		nge of 7.5 to 10.0 all times

(b) <u>Silver</u> Chloride Reduction Spent Solution BPT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
ng/troy ounce of silv	ver reduced in so	olution
lead	0.168	0.080
lercury	0.100	0.040
Silver	0.164	0.068
linc	0.584	0.244
Gold	0.040	
)il and grease	8.000	4.800
otal suspended solids	16.400	7.800
OH	Within the rar	nge of 7.5 to 10.0
		all times
c) Electrolytic Cel	ls Wet Air Pollu	tion Control DDM
e, <u>miccerorycic</u> cei	The Wet All Pollo	tion <u>Control</u> BPT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of go	ld refined electro	olytically
Lead	83.160	39.600
Mercury	49.500	19.800
Silver	81.180	33.660
Zinc	289.100	120.800
Gold	19.800	
Oil and grease	· 3,960.000	2,376.000
Total suspended solids	8,118.000	3,861.000
pH	Within the ran	nge of 7.5 to 10.0 L times
	at ai.	LUIMES

(d) Electrolyte Preparation Wet Air Pollution Control BPT

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
mg/troy ounce of silv	ver in electroly	ce produced
Lead	0.021	0.010
Mercury	0.013	0.005
Silver	0.021	0.009
Zinc	0.073	0.031
Gold	0.005	
Oil and grease	1.000	0.600
Total suspended	2.050	0.975
solids	2.000	0.373
pH		nge of 7.5 to 10.0
· · · · ·	at al.	l times
(e) <u>Calciner</u> <u>Wet</u> <u>Ai</u>	r Pollution Cont	rol. BPT
		rol BPT Maximum for
(e) <u>Calciner Wet Ai</u> Pollutant or Pollutant Property	r Pollution Cont Maximum for Any One Day	
Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
Pollutant or Pollutant Property mg/kg (lb/million lb	Maximum for Any One Day	Maximum for Monthly Average
Pollutant or Pollutant Property mg/kg (lb/million lb Lead	Maximum for Any One Day s) of mercury con 78.200	Maximum for Monthly Average ndensed
Pollutant or Pollutant Property mg/kg (lb/million lb Lead Mercury	Maximum for Any One Day s) of mercury con 78.200 46.550	Maximum for Monthly Average ndensed 37.240 18.620
Pollutant or Pollutant Property mg/kg (lb/million lb Lead Mercury Silver	Maximum for Any One Day s) of mercury con 78.200 46.550 76.320	Maximum for Monthly Average ndensed 37.240 18.620 31.650
Pollutant or Pollutant Property mg/kg (lb/million lb Lead Mercury Silver Zinc	Maximum for Any One Day s) of mercury con 78.200 46.550 76.320 271.900	Maximum for Monthly Average ndensed 37.240 18.620
Pollutant or Pollutant Property mg/kg (lb/million lb Lead Mercury Silver Zinc Gold	Maximum for Any One Day s) of mercury con 78.200 46.550 76.3?0 271.900 18.600	Maximum for Monthly Average ndensed 37.240 18.620 31.650 113.600
Pollutant or Pollutant Property mg/kg (lb/million lb Lead Mercury Silver Zinc Gold Oil and grease Total suspended	Maximum for Any One Day s) of mercury con 78.200 46.550 76.320 271.900	Maximum for Monthly Average ndensed 37.240 18.620 31.650
Pollutant or Pollutant Property mg/kg (lb/million lb Lead Mercury Silver Zinc Gold Oil and grease Total suspended solids	Maximum for Any One Day s) of mercury con 78.200 46.550 76.320 271.900 18.600 3,724.000 7,634.000	Maximum for Monthly Average ndensed 37.240 18.620 31.650 113.600 2.234.000 3,631.000
Pollutant or Pollutant Property mg/kg (lb/million lb Lead Mercury Silver Zinc Gold Oil and grease Total suspended	Maximum for Any One Day s) of mercury con 78.200 46.550 76.320 271.900 18.600 3,724.000 7,634.000 Within the ra	Maximum for Monthly Average ndensed 37.240 18.620 31.650 113.600 2.234.000

(f) <u>Calcine</u> <u>Quench</u> <u>Water</u> BPT

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
mg/kg (lb/million lbs) of mercury co	ndensed
Lead	7.392	3.520
Mercury	4.400	1.760
Silver	7.216	2.992
Zinc	25.700	10.740
Gold	1.760	
Oil and grease	352.000	211.200
Total suspended solids	721.600	343.200
pH	Within the ray	nge of 7.5 to 10.0
E	at al	l times
(g) <u>Calciner</u> Stack <u>G</u>	as Contact Cool	ing <u>Water</u> BPT
(g) <u>Calciner</u> <u>Stack</u> <u>G</u> Pollutant or		· · · · · · · · · · · · · · · · · · ·
	as <u>Contact</u> <u>Cool</u> Maximum for Any One Day	ing Water BPT Maximum for Monthly Average
Pollutant or	Maximum for Any One Day	Maximum for Monthly Average
Pollutant or Pollutant Property	Maximum for Any One Day) of mercury con	Maximum for Monthly Average ndensed
Pollutant or Pollutant Property mg/kg (lb/million lbs	Maximum for Any One Day) of mercury con 1.743	Maximum for Monthly Average
Pollutant or Pollutant Property mg/kg (lb/million lbs Lead	Maximum for Any One Day) of mercury con	Maximum for Monthly Average ndensed 0.830
Pollutant or Pollutant Property mg/kg (lb/million lbs Lead Mercury Silver Zinc	Maximum for Any One Day) of mercury con 1.743 1.038	Maximum for Monthly Average ndensed 0.830 0.415
Pollutant or Pollutant Property mg/kg (lb/million lbs Lead Mercury Silver Zinc Gold	Maximum for Any One Day) of mercury con 1.743 1.038 1.702 6.059 0.415	Maximum for Monthly Average ndensed 0.830 0.415 0.706
Pollutant or Pollutant Property mg/kg (lb/million lbs Lead Mercury Silver Zinc Gold Oil and grease	Maximum for Any One Day) of mercury con 1.743 1.038 1.702 6.059 0.415 83.000	Maximum for Monthly Average ndensed 0.830 0.415 0.706
Pollutant or Pollutant Property mg/kg (lb/million lbs Lead Mercury Silver Zinc Gold	Maximum for Any One Day) of mercury con 1.743 1.038 1.702 6.059 0.415	Maximum for Monthly Average ndensed 0.830 0.415 0.706 2.532

(h) <u>Condenser</u> <u>Blowdown</u> BPT

	Maximum for Any One Day	Maximum for Monthly Average	
mg/kg (lb/million lbs)) of mercury con	Idensed	•
Lead	5.796	2.760	
Mercury	3.350	1.380	
Silver	5.658	2.346	
Zinc	20.150	8.418	,
Gold	1.380		
Oil and grease	276.000	165.600	
Total suspended	565.800	269.100	
solids			
pH and the second secon	ata	nge of 7.5 to 10.0 11 times	
		4	
and an and the second secon The second se	ا و ه را مراجع مراجع المراجع و مراجع مراجع المراجع المراجع المراجع المراجع المراجع المراجع المراجع ا	an a	
(i) Mercury Cleaning	Bath Water BPI		
(i) <u>Mercury</u> <u>Cleaning</u>	<u>Bath</u> <u>Water</u> BPI		ŕ.
(i) <u>Mercury</u> <u>Cleaning</u> Pollutant or	<u>Bath Water</u> BPT Maximum for	Maximum for	· .
	Maximum for	Maximum for	
Pollutant or	Maximum for Any One Day	Maximum for Monthly Average	
Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	
Pollutant or Pollutant Property mg/kg (lb/million lbs)	Maximum for Any One Day) of mercury con	Maximum for Monthly Average Idensed	
Pollutant or Pollutant Property mg/kg (lb/million lbs) Lead	Maximum for Any One Day) of mercury con 0.588	Maximum for Monthly Average Idensed 0.280	
Pollutant or Pollutant Property mg/kg (lb/million lbs) Lead Mercury	Maximum for Any One Day) of mercury con 0.588 0.350 0.574	Maximum for Monthly Average Idensed 0.280 0.140	
Pollutant or Pollutant Property mg/kg (lb/million lbs) Lead Mercury Silver	Maximum for Any One Day of mercury con 0.588 0.350	Maximum for Monthly Average Idensed 0.280 0.140 0.238	· · · · · · · · · · · · · · · · · · ·
Pollutant or Pollutant Property mg/kg (lb/million lbs) Lead Mercury Silver Zinc Gold	Maximum for Any One Day) of mercury con 0.588 0.350 0.574 2.044	Maximum for Monthly Average Idensed 0.280 0.140 0.238	
Pollutant or Pollutant Property mg/kg (lb/million lbs) Lead Mercury Silver Zinc	Maximum for Any One Day of mercury con 0.588 0.350 0.574 2.044 0.140	Maximum for Monthly Average Indensed 0.280 0.140 0.238 0.854 	

BAT is promulgated based on the performance achievable by the application of chemical precipitation, sedimentation, and multimedia filtration (lime, settle, and filter) and ion exchange end-of-pipe polishing treatment technology and inprocess flow reduction methods, along with preliminary treatment consisting of oil skimming for selected waste streams. The following BAT effluent limitations are promulgated:

(a) <u>Smelter Wet Air Pollution Control</u> BAT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of gold	d and silver smel	lted
Lead	0.364	0.169
Mercury	0.195	0.078
Silver	0.377	0.156
Zinc	1.326	0.546
Gold	0.130	

(b) Silver Chloride Reduction Spent Solution BAT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of size	lver reduced in so	olution
Lead	0.112	0.052
Mercury	0.060	0.024
Silver	0.116	0.048
Zinc	0.408	0.168
Gold	0.040	

(c) <u>Electrolytic</u> <u>Cells</u> <u>Wet</u> <u>Air</u> <u>Pollution</u> <u>Control</u> BAT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	
mg/troy ounce of gold	refined electro	olytically	
Lead	5.544	2.574	
Mercury	2.970	1.188	
Silver	5.742	2.376	
Zinc	20.200	8.316	
Gold	1.980	— —	•
	·		······

(d) Electrolyte Preparation Wet Air Pollution Control BAT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Avęrage
mg/troy ounce of s	ilver in electroly	te produced
Lead	0.014	0.0065
Mercury Silver	0.0075 0.015	0.0030 0.0060
Zinc	0.051	0.021
Gold	0.005	— —

(e) <u>Calciner Wet</u> <u>Air</u> <u>Pollution</u> <u>Control</u> BAT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/kg (lb/million lk	os) of mercury con	ndensed
Lead	6.160	2.860
Mercury	3.300	1.320
Silver	6.380	2.640
Zinc	22.440	9.240
Gold	2.200	·

(f) Calcine Quench Water

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/kg (lb/million lbs) of mercury con	ndensed
Lead	4.928	2.288
Mercury Silver	2.640 5.104	1.056 2.112
Zinc Gold	17.950 1.760	7.392

(g) <u>Calciner Stack</u> <u>Gas</u> <u>Contact</u> <u>Cooling</u> <u>Water</u> BAT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/kg (lb/million lbs)	of mercury con	ndensed
Lead Mercury Silver Zinc Gold	1.162 0.623 1.204 4.233 0.415	0.540 0.249 0.498 1.743

(h) Condenser Blowdown

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/kg (lb/million lbs) of mercury con	ndensed
Lead	3.864	1.794
Mercury	2.070	0.828
Silver	4.002	1.656
Zinc	14.080	5.796
Gold	1.380	

SECT - II

SECT - II

(i) Mercury Cleaning Bath Water BAT

Maximum for Any One Day	Maximum for Monthly Average
) of mercury con	ndensed
0.392	0.182
0.210	0.084
0.406	0.168
1.428	0.588
0.140	
	Any One Day) of mercury con 0.392 0.210 0.406 1.428

NSPS are promulgated based on the performance achievable by the application of chemical precipitation, sedimentation, and multimedia filtration (lime, settle, and filter) and ion exchange as a polishing step technology, and inprocess flow reduction control methods, along with preliminary treatment consisting of oil skimming for selected waste streams. The following effluent standards are promulgated for new sources:

(a) <u>Smelter</u> <u>Wet Air</u> <u>Pollution</u> <u>Control</u> NSPS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of go	ld and silver sme	lted
Lead	0.364	0.169
Mercury	0.195	0.078
Silver	0.377	0.156
Zinc	1.326	0.546
Gold	0.130	
Oil and grease	13.000	13.000
Total suspended solids	19.500	15.600
рН		nge of 7.5 to 10.0 all times

(b) Silver Chloride Reduction Spent Solution NSPS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of sil	ver reduced in s	olution
Lead	0.112	0.052
Mercury	0.060	0.024
Silver	0.116	0.048
Zinc	0.408	0.168
Gold	0.000	
Oil and grease	4.000	4.000
Total suspended solids	6.000	4.800
рН		nge of 7.5 to 10.0 all times

(c) <u>Electrolytic</u> <u>Cells</u> <u>Wet</u> <u>Air</u> <u>Pollution</u> <u>Control</u> NSPS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
		olytically
mg/troy ounce of gold	l relined electr	Orycrearry
Lead	5.544	2.574
Mercury	2.970	1.188
Silver	5.742	2.376
Zinc	20.200	8.316
Gold	1.980	
Oil and grease	198.000	198.000
Total suspended solids	297.000	237.600
pH	Within the at	ange of 7.5 to 10.0 all times

(d) Electrolyte Preparation Wet Air Pollution Control NSPS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of si	lver in electrolyte	produced
Lead	0.014	0.0065
Mercury	0.0075	0.0030
Silver	0.015	0.0060
Zinc	0.051	0.021
Gold	0.005	
Oil and grease	0.500	0.500
Total suspended solids	0.750	0.600
PH	Within the rang	e of 7.5 to 10.0
	at all t	imes

(e) <u>Calciner</u> <u>Wet</u> <u>Air</u> <u>Pollution</u> <u>Control</u> NSPS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	· · ·
mg/kg (lb/million 1)	os) of mercury con	ndensed	
Lead	6.160	2.860	
Mercury	3.300	1.320	
Silver	6.380	2.640	
Zinc	22.440	9.240	2
Gold	2.200		
Oil and grease	220.000	220.000	
Total suspended solids	330.000	264.000	
pH		nge of 7.5 to 10.0 all times	

(f) <u>Calciner Quench Water</u> NSPS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/kg (lb/million lb	s) of mercury co	ndensed
Lead	4.928	2.288
Mercury	2.640	1.056
Silver	5.104	2.112
Zinc	17.950	7.392
Gold	1.760	
Oil and grease	176.000	176.000
Total suspended	264.000	211.200
solids		
Нq		ange of 7.5 to 10.0
	, at	all times
(g) <u>Calciner</u> <u>Stack</u>	Gas Contact Cool	ing Water NSPS
	<u>Gas Contact Cool</u> Maximum for	ing Water NSPS Maximum for
(g) <u>Calciner Stack</u> Pollutant or Pollutant Property	· · · · · · · · · · · · · · · · · · ·	Maximum for
Pollutant or	Maximum for Any One Day	Maximum for Monthly Average
Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average ndensed 0.540
Pollutant or Pollutant Property mg/kg (lb/million lb	Maximum for Any One Day os) of mercury co 1.162 0.623	Maximum for Monthly Average ndensed 0.540 0.249
Pollutant or Pollutant Property mg/kg (lb/million lb Lead	Maximum for Any One Day os) of mercury co 1.162 0.623 1.204	Maximum for Monthly Average ndensed 0.540 0.249 0.498
Pollutant or Pollutant Property mg/kg (lb/million lt Lead Mercury	Maximum for Any One Day os) of mercury co 1.162 0.623 1.204 4.233	Maximum for Monthly Average ndensed 0.540 0.249
Pollutant or Pollutant Property mg/kg (lb/million lb Lead Mercury Silver	Maximum for Any One Day os) of mercury co 1.162 0.623 1.204 4.233 0.415	Maximum for Monthly Average ndensed 0.540 0.249 0.498 1.743
Pollutant or Pollutant Property mg/kg (lb/million lb Lead Mercury Silver Zinc Gold Oil and grease	Maximum for Any One Day os) of mercury co 1.162 0.623 1.204 4.233 0.415 41.500	Maximum for Monthly Average ndensed 0.540 0.249 0.498 1.743 41.500
Pollutant or Pollutant Property mg/kg (lb/million lb Lead Mercury Silver Zinc Gold	Maximum for Any One Day os) of mercury co 1.162 0.623 1.204 4.233 0.415 41.500 62.250	Maximum for Monthly Average ndensed 0.540 0.249 0.498 1.743 41.500 49.800
Pollutant or Pollutant Property mg/kg (lb/million lb Lead Mercury Silver Zinc Gold Oil and grease Total suspended	Maximum for Any One Day os) of mercury co 1.162 0.623 1.204 4.233 0.415 41.500 62.250 Within the r	Maximum for Monthly Average ndensed 0.540 0.249 0.498 1.743 41.500

(h) Condenser Blowdown NSPS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	
mg/kg (lb/million lbs)	of mercury con	densed	
Lead	3.864	1.794	. • • •
Mercury	2.070	0.828	. ,
Silver	4.002	1.656	.'
Zinc	14.080	5.796	
Gold	1.380		,
Oil and grease	138.000	138.000	$\cdot \cdot $
Total suspended solids	207.000	165.600	
pH	Within the ra at	nge of 7.5 to 10.0 all times	0

(i) Mercury Cleaning Bath Water NSPS

Pollutant or Pollutant Property	Maximum for Any One Day		
mg/kg (lb/million lbs)	of mercury	condensed	
Lead	0.392	0.182	
Mercury	0.210	0.084	
Silver	0.406	0.168	
Zinc	1.428	0.588	
Gold	0.140		
Oil and grease	14.000	14.000	+ t.
Total suspended solids	21.000	16.800	
рН		range of 7.5 to 10.0 all times	

EPA is not promulgating PSES for the primary precious metals and mercury subcategory because there are no indirect dischargers. PSNS are promulgated based on the performance achievable by the application of chemical precipitation, sedimentation, and multimedia filtration (lime, settle, and filter) and ion exchange as a polishing step technology, and in-process flow reduction control methods, along with preliminary treatment consisting of oil skimming for selected waste streams. The following pretreatment standards are promulgated for new sources:

(a) Smelter Wet Air Pollution Control PSNS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of gold	and silver smel	lted
Lead Mercury Silver Zinc Gold	0.364 0.195 0.377 1.326 0.130	0.169 0.078 0.156 0.546

(b) Silver Chloride Reduction Spent Solution PSNS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of sil	ver reduced in s	olution
Lead Mercury Silver Zinc Gold	0.112 0.060 0.116 0.408 0.040	0.052 0.024 0.048 0.168

(c) Electrolytic Cells Wet Air Pollution Control PSNS

	tant or t Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy	ounce of gold	refined electr	colytically
Lead Mercury		5.544	2.574
Silver	,	2.970 5.742	1.188 2.376
Zinc		20.200	8.316
Gold		1.980	

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	
mg/troy ounce of si	llver in electrolyte	e produced	
Lead	0.014	0.0065	
Mercury	0.0075	0.0030	
Silver	0.015	0.0060	
Zinc	0.051	0.021	
Gold	0.005		

Calciner Wet Air Pollution Control (e) PSNS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/kg (lb/million lbs	s) of mercury co	ndensed
Lead	6.160	2.860
Mercury	3.300	1.320
Silver	6.380	2.640
Zinc	22.440	9.240
Gold	2.200	

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/kg (lb/million lbs)	of mercury cor	ndensed
Lead Mercury Silver Zinc Gold	4.928 2.640 5.104 17.950 1.760	2.288 1.056 2.112 7.392

(g) Calciner Stack Gas Contact Cooling Water PSNS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/kg (lb/million lbs)	of mercury co	ndensed
Lead Mercury Silver Zinc Gold	1.162 0.623 1.204 4.233 0.415	0.540 0.249 0.498 1.743

(h) Condenser Blowdown

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/kg (lb/million lbs)	of mercury co	ndensed
Lead Mercury Silver Zinc Gold	3.864 2.070 4.002 14.080 1.380	1.794 0.828 1.656 5.796

(i) Mercury Cleaning Bath Water PSNS .

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/kg (lb/million lbs)	of mercury con	densed
Lead	0.392	0.182
Mercury	0.210	0.084
Silver	0.406	0.168
Zinc	1.428	0.588
Gold	0.140	

EPA is not promulgating BCT for the primary precious metals and mercury subcategory at this time.

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SECTION III

SUBCATEGORY PROFILE

This section of the primary precious metals and mercury supplement describes the raw materials and processes used in producing primary precious metals and mercury and presents a profile of the primary precious metals and mercury plants identified in this study. For a discussion of the purpose, authority, and methodology for this study, and a general description of the nonferrous metals manufacturing category, refer to Section III of Vol. I.

DESCRIPTION OF PRIMARY PRECIOUS METALS PRODUCTION

The production of primary precious metals can be divided into three distinct stages - smelting to produce Dore metal, or precipitation of silver from silver ore, separation of gold and silver, and gold and silver purification. The processes used in each stage vary with the type and purity of raw material used. The primary precious metals production process is presented schematically in Figure III-1 (page 2174) and described below.

RAW MATERIALS

Primary precious metals are produced from gold and silver bearing concentrates produced from precious metal ores and as by-products from the beneficiation of base metal ores. A small amount is also produced from placer mining operations. Precious metal ores are mined at various locations in the western United States.

Mining and beneficiation processes for precious metal-bearing ores, including cyanidation, amalgamation, flotation, and gravity concentration are outside of the scope of this subcategory. Both the mining and beneficiation operations are regulated as part of the Ore Mining and Dressing Point Source Category.

Primary precious metals produced as a by-product of primary copper electrolytic refining operations are regulated as part of the primary copper electrolytic refining subcategory. Primary precious metals produced as a by-product of other primary copper operations, such as solvent extraction, are regulated under these limitations.

SMELTING

The gold and silver manufacturing process begins when the precious metals bearing concentrate is sent through a Dore furnace (smelter). In the Dore furnace, the gold, silver, and other precious metals are smelted in the presence of a fluxing agent (commonly soda ash, borax, or silica). This smelting operation produces a slag containing impurities such as copper

and zinc, and a gold base alloy known as Dore, which may also contain silver. The Dore gold may be cast and sold as a product or be further refined.

SILVER PRECIPITATION

The silver manufacturing process may begin when silver is precipitated from a silver-copper ore which has been dissolved into solution. Silver is precipitated using sodium chloride. The silver chloride precipitate is then slurried in dilute acid, and reduced to silver metal in a cementation step. The silver produced may be sold as a product.

GOLD-SILVER SEPARATION

The separation of gold and silver from Dore bars is accomplished through electrolytic refining of the Dore bars or by the Miller process. In the electrolytic method, the Dore metal is cast into anodes and placed into a solution of silver nitrate (AgN3) electrolyte. When a current is applied, fine silver is deposited upon the cathode. This silver is removed, washed, and cast into bars of fine silver for sale. Gold remains as slimes in the canvas anode bags. Gold slimes are washed with acid and rinsed with water before being cast into product ingots. This gold is about 99 percent pure. Silver is recovered in a cementation step from the silver crystals wash water and from the gold slimes acid wash and rinse water. In the cementation process, copper is added to the solution and replaces the silver, causing the silver to precipitate out of solution. The recovered silver is returned to the anode casting stage.

Gold and silver can also be separated from the Dore metal while it is still molten. This purification step is known as the Miller process and consists of bubbling chlorine gas through the molten Dore metal in a parting furnace. This process converts the silver into silver chloride salt and volatilizes base metal impurities. The silver chloride salt rises to the surface and is skimmed off for further processing. The gold produced by the Miller process can be further purified by electrolytic refining or immediately cast as a product. The silver chloride salt which is skimmed off is remelted and cast into slabs. These slabs are reduced to silver metal in an acid solution in a similar process to that described above under silver precipitation. The resulting silver metal is remelted in the presence of borax flux and molten silver is then cast into product ingots.

FURTHER PURIFICATION

After separation, gold and silver can be further refined by various means. One technique to further refine gold is electrolysis. Impure gold is cast into anodes and purified electrolytically by the Wholwill process in a chloride solution. Gold, which is oxidized at the anode, passes into solution and is deposited upon the cathode. The gold cathode is melted and cast into bars with a purity greater than 99.9 percent. As described above, gold slimes can be further purified using an acid wash and water rinse process.

PROCESS WASTEWATER SOURCES

Although a variety of processes are involved in primary precious metals production, the process wastewater sources can be subdivided as follows:

- 1. Smelter wet air pollution control,
- 2. Silver chloride reduction spent solution,
- 3. Electrolytic cells wet air pollution control, and
- 4. Electrolyte preparation wet air pollution control.

DESCRIPTION OF PRIMARY MERCURY PRODUCTION

Primary mercury is produced from mercury ores and gold-bearing ores by roasting or calcining. The primary mercury production process is presented schematically in Figure III-2 (page 2175) and described below.

RAW MATERIALS

The principal source of mercury is cinnabar ore (mercury sulfide). Cinnabar ore is mined primarily in Nevada, California, and Oregon. In addition, a small amount of mercury is recovered as a co-product from gold ore.

ROASTING

After mining and beneficiation, mercury is extracted from mercury-bearing ores by roasting or calcining. In the roasting process, the mercury is vaporized and then recovered in a condenser, while the sulfur is oxidized to SO₂. Some water may condense with the mercury and is discharged as a waste stream. The mercury recovered from the condenser may be washed with water prior to being sold. The mining and beneficiation stage of mercury production is not within the scope of this subcategory.

Sulfur dioxide (SO₂) and other gaseous emissions from the mercury roasting furnace are controlled with a multistage scrubber. Sulfur dioxide emissions are controlled with a wet scrubber. After SO₂ removal, the clean stack gases are cooled with contact cooling water and discharged to the atmosphere. Calciner SO₂ scrubber liquor and stack gas contact cooling water are discharged as waste streams.

PROCESS WASTEWATER SOURCES

Although a variety of processes are involved in primary mercury production, the process wastewater sources can be subdivided as follows:

- 1. Calciner wet air pollution control,
- 2. Calcine quench water,
- 3. Calciner stack gas contact cooling water,
- 4. Condenser blowdown, and
- 5. Mercury cleaning bath water.

OTHER WASTEWATER SOURCES

There are other waste streams associated with the primary precious metals and mercury subcategory. These waste streams may include casting contact cooling water, stormwater runoff, and maintenance and cleanup water. These waste streams are not considered as a part of this rulemaking. EPA believes that the flows and pollutant loadings associated with these waste streams are insignificant relative to the waste streams selected and are best handled by the appropriate permit authority on a case-bycase basis under authority of Section 402 of the Clean Water Act.

AGE, PRODUCTION, AND PROCESS PROFILE

Figure III-3 (page 2176) shows the location of the eight primary precious metals and mercury plants operating in the United States. Four of the eight plants are located in Nevada, with one of the remaining plants each being located in Idaho, Montana, Colorado, and South Dakota.

Table III-1 (page 2169) shows the relative age and discharge status of the primary precious metals and mercury plants. Seven of the eight plants in this subcategory have a zero discharge status, and one plant is a direct discharge facility. The average plant age is less than 12 years. Tables III-2 to III-4 (pages 2170-2172) provide a summary of the current production ranges. It can be seen that production of gold is evenly spread along the ranges with a mean production of 70,000 troy ounces/year. The mean production of silver is 222,500 troy ounces/year.

Table III-5 (page 2173) provides a summary of the number of plants generating wastewater for the waste streams associated with various processes and the number of plants with the process.

Table III-1

INITIAL OPERATING YEAR (RANGE) SUMMARY OF PLANTS IN THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY BY DISCHARGE TYPE

	(Plant Age in Years)					
Type of Plant	1983- 1973 (0-11)	1972- 1968 (12-16)	1967- 1958 (17-26)	1957- 1918 (27-66)	Before 1918 (66%)	Total
Direct	. l	0	0	0	0	1
Indirect	0.	0	0	0	0	0
Zero	4	1	<u>1</u>	<u>0</u>	<u>1</u>	<u>7</u>
TOTAL	5	1	1	0	1	8

Initial Operating Year (Range) (Plant Age in Years)

PRODUCTION RANGES FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

Gold Production Range for 1982				
Type of Plant	0-10,000 (troy oz./yr)	10,001-75,000 (troy oz./yr)	75,001-200,000 (troy oz./yr)	Total Number of Plants
Direct	1	0	0	1
Indirect	0	0	0	0
Zero	1	2	2	5
	· · · ·	· ·		6

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III

PRODUCTION RANGES FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

Silver Production Range for 1982					
Type of Plant	0-10,000 (troy oz./yr)	10,001-50,000 (troy oz./yr)	50,001-500,000 (troy oz./yr)	>500,000 (troy oz./yr)	Total Number of Plants
Direct	0	0	1	0	1
Indirect	0	0	0	0	0
Zero	1	3	0	1 · · · 1	5

2171

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

SECT - III

PRODUCTION RANGES FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

Mercury production ranges are not presented here because the information on which they are based has been claimed confidential.

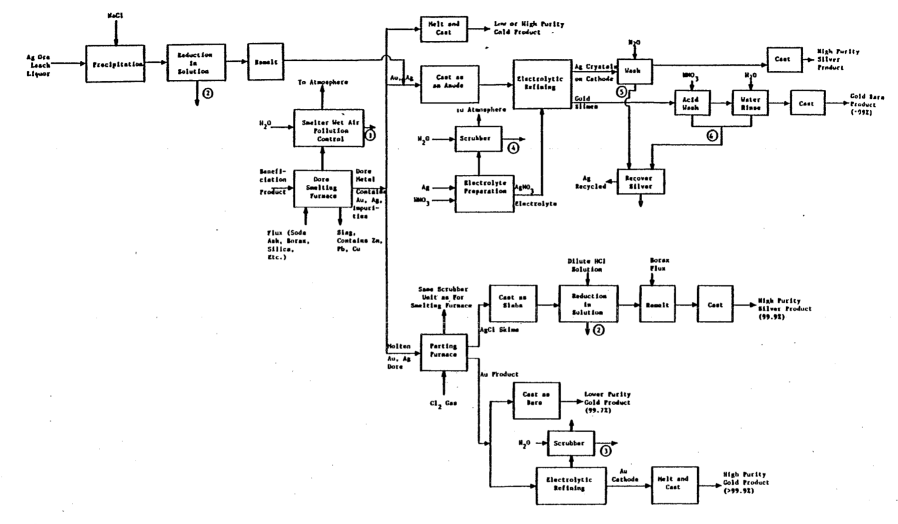
SUMMARY OF PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY PROCESSES AND ASSOCIATED WASTE STREAMS

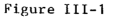
Process or Waste Stream	Number of Plants With Process or Waste Stream	Number of Plants Reporting Generation of Wastewater*
Roasting	8	
 (P.M.) Smelter wet air pollution control (Hg) Calciner wet air pollution control (Hg) Calcine quench (Hg) Calciner stack gas cooling water (Hg) Condenser blowdown 	5 1 1 1 2	3 1 1 1 1
Gold-Silver Separation	3 ¹⁰ ¹⁰	
(P.M.) Electrolytic preparation wet air pollution control	1	· 1
Further Purification	3	
(P.M.) Silver chloride reduction spent solution (P.M.) Electrolytic cells wet air pollution control (Hg) Mercury cleaning bath water	2 1 1	2 1 1

*Through reuse or evaporation practices, a plant may "generate" a wastewater from a particular process but not discharge it. PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SECT 1

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2173





PRIMARY PRECIOUS METALS PRODUCTION PROCESSES

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SECT - III

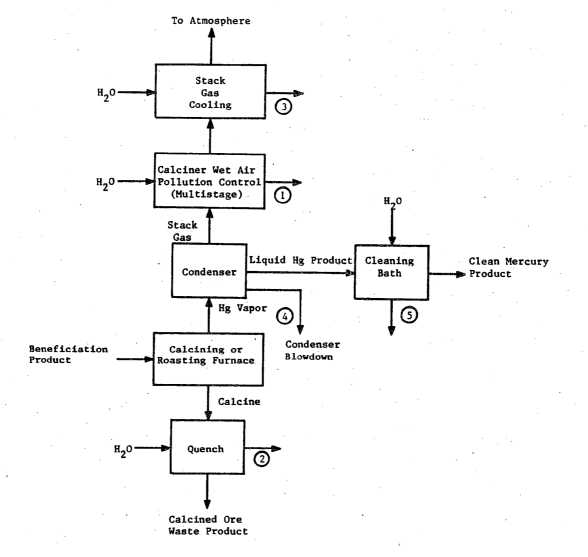


Figure III-2

PRIMARY MERCURY PRODUCTION PROCESS

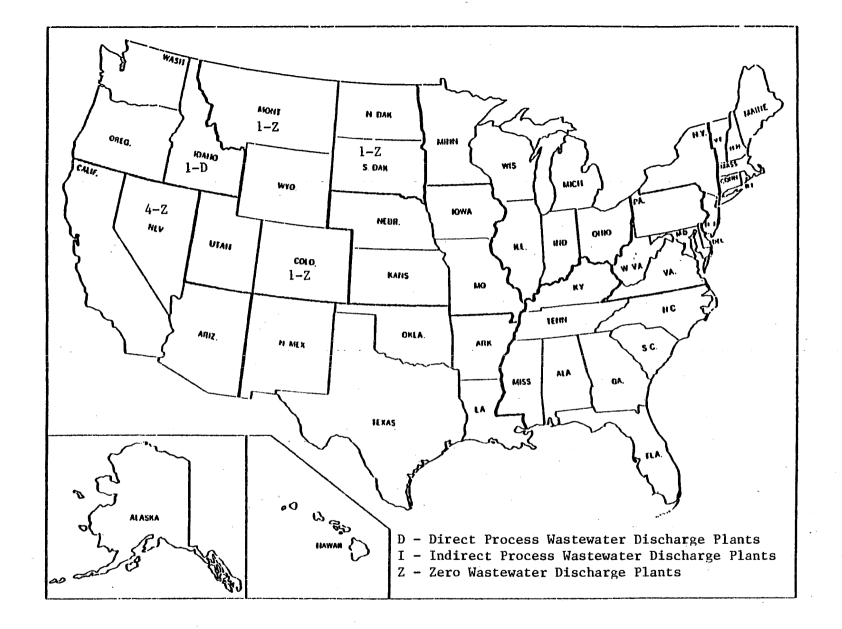


Figure III-3

GEOGRAPHIC LOCATIONS OF THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY PLANTS

2176

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

SECT

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SECTION IV

SECT - IV

SUBCATEGORIZATION

section summarizes the factors considered during This the designation of the primary precious metals and mercury subcategory and its related subdivisions. Production normalizing parameters for each subdivision are also discussed.

FACTORS CONSIDERED IN SUBDIVIDING THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

factors listed for general subcategorization were each The evaluated when considering subdivision of the primary precious metals and mercury subcategory. In the discussion that follows, the factors will be described as they pertain to this particular subcategory.

rationale for considering segmentation of the primary The precious metals and mercury subcategory is based primarily on differences in the production processes and raw materials used. Within this subcategory, a number of different operations are performed, which may or may not have a water use or discharge, and which may require the establishment of separate effluent While primary precious metals and mercury is limitations. considered a single subcategory, a more thorough examination of the production processes has illustrated the need for limitations standards based on a specific set of waste streams. and Limitations will be based on specific flow allowances for the following subdivisions or building blocks.

- 1. Smelter wet air pollution control,
- Silver chloride reduction spent solution, 2.
- Electrolytic cells wet air pollution control, 3.
- Electrolyte preparation wet air pollution control, Calciner wet air pollution control, 4.
- 5.
- 6. Calcine quench water,
- 7. Calciner stack gas contact cooling water,
- 8. Condenser blowdown, and
- Mercury cleaning bath water. 9.

These building blocks follow directly from differences within the three distinct production stages of primary precious metals and mercury.

The smelting of precious metals bearing concentrates to produce Dore metals gives rise to the first subdivision: smelter wet air pollution control wastewater. If any remelt furnaces are used in the process, the resulting off-gases are usually combined with smelter off-gases for air pollution control. Thus, the smelter wet air pollution control subdivision represents the wet air pollution control wastewater for both smelters and remelt

furnaces. The next two subdivisions result from either the Miller process for purifying high silver content Dore metal or the recovery of silver from Ag-Cu ore by silver precipitation. The electrolytic cells scrubber wastewater from gold refining is a principal waste stream, and spent solution from silver reduction is another significant waste stream.

The electrolytic refining of silver from Dore metal gives rise to the next subdivision. The fourth subdivision is created by the wet scrubber used to control air emissions from the electrolyte preparation stage.

The last five subdivisions result from the production of primary mercury. The treatment of calciner off-gases by wet scrubbing gives rise to the first of these subdivisions. Waste streams may also result from the quenching of calciner wastes to reduce their temperature prior to disposal and the cooling of calciner offgases before discharge from the stack. During condensation of the vaporized mercury, the condensation of a water fraction can occur and this condenser blowdown is a possible waste stream. After condensation the liquid mercury may be further purified by use of cleaning baths. This cleaning operation is also a potential source of wastewater.

OTHER FACTORS

The other factors considered in this evaluation were shown to be inappropriate as a bases for further segmentation. Air pollution control methods, treatment costs, and total energy requirements are functions of the selected subcategorization factors--metal product, raw materials, and production processes. Therefore, they are not independent factors and do not affect the subcategorization which has been developed. As discussed in Section IV of the General Development Document, certain other factors, such as plant age, plant size, and the number of employees, were also evaluated and determined to be inappropriate for use as bases for subdivision of nonferrous metals plants.

PRODUCTION NORMALIZING PARAMETERS

As discussed previously, the effluent limitations and standards developed in this document establish mass limitations for the discharge of specific pollutant parameters. To allow these regulations to be applied to plants with various production capacities, the mass of pollutant discharged must be related to a unit of production. This factor is known as the production normalizing parameter (PNP).

In general, for each production process which has a wastewater associated with it, the actual mass of precious metal or mercury product produced will be used as the PNP. Thus, the PNPs for the nine subdivisions are as follows:

Building Block

PNP

- 1. Smelter wet air pollution troy ounce of gold and silver control
- Silver chloride reduction 2. spent solution
- З. Electrolytic cells wet air pollution control
- 4. Electrolyte preparation wet air pollution control
- 5. Calciner wet air pollution control
- 6. Calcine guench water

Condenser blowdown

8

7. Calciner stack gas contact cooling water

troy ounce of silver in

troy ounce of gold refined

troy ounce of silver reduced

electrolyte produced

electrolytically

smelted

in solution

- kkg of mercury condensed
- 9. Mercury cleaning bath water

kkg of mercury condensed

Other PNPs were considered. The use of production capacity instead of actual production was eliminated from consideration because the wastewater generated and the mass of the pollutant produced is more a function of true production than of installed capacity. The use of some common intermediate (i.e., gold and silver cathodes or silver chloride) as a basis for PNPs for all processes was rejected since not all plants follow the same production path to get to the specific end-product. Additionally, some plants divert part of their intermediate products and sell them instead of processing all input raw materials to one final product. If an "end-product" were chosen as the PNP, plants that had these upstream diversions would be allowed to discharge more per mass of product than their competitors who did not.

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SECTION V

WATER USE AND WASTEWATER CHARACTERISTICS

This section describes the characteristics of the wastewaters associated with the primary precious metals and mercury subcategory. Water use and discharge rates are explained and then summarized in tables at the end of this section. Data used to characterize the wastewaters are presented. Finally, the specific source, water use and discharge flows, and wastewater characteristics for each separate wastewater source are discussed.

Two principal data sources were used in the development of effluent limitations and standards for this subcategory: data collection portfolios (dcp) and field sampling results. Data collection portfolios contain information regarding wastewater flows and production levels.

In order to quantify the pollutant discharge from primary precious metals and mercury plants, a field sampling program was conducted. A complete list of the pollutants considered and a summary of the techniques used in sampling and laboratory analyses are included in Section V of Vol. I. Samples were analyzed for 124 of the 126 priority pollutants and other pollutants deemed appropriate. Because the analytical standard TCDD was judged to be too hazardous to be made generally for available, samples were never analyzed for this pollutant. Samples were also not analyzed for asbestos. There is no reason to expect that TCDD or asbestos would be present in nonferrous metals manufacturing wastewater. Two plants were selected for sampling in the primary precious metals and mercury subcategory. In general, the samples were analyzed for three classes of pollutants: priority organic pollutants, priority metal pollutants, and criteria pollutants (which includes both conventional and nonconventional pollutants).

After proposal, EPA gathered additional wastewater sampling data for two of the subdivisions in this subcategory. These data were acquired through a self-sampling program and include data from analyses for the priority metals arsenic, cadmium, chromium, copper, lead, mercury, nickel, silver, thallium, and zinc. These data also include analyses for the nonconventional pollutant qold. These data show pollutant concentrations similar to those by the data which EPA had acquired for indicated these subdivisions prior to proposal. These data also support the assumptions which EPA had made concerning the presence and concentrations of pollutants in those subdivisions where we did not have analytical data for specific pollutants. For this reason, the selection of pollutant parameters for limitation in this subcategory (Section VI) has not been revised based on these new data.

SECT - V PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

As described in Section IV of this supplement, the primary precious metals and mercury subcategory has been divided into nine subdivisions or wastewater sources, so that the promulgated regulation contains mass discharge limitations and standards for nine unit processes discharging process wastewater. Differences wastewater characteristics associated with in the these subdivisions are to be expected. For this reason, wastewater streams corresponding to each subdivision are addressed separately in the discussions that follow. These wastewater sources are:

- Smelter wet air pollution control, 1.
- Silver chloride reduction spent solution, 2.
- 3.
- Electrolytic cells wet air pollution control, Electrolyte preparation wet air pollution control, 4.
- 5. Calciner wet air pollution control,
- Calcine quench water, 6.
- Calciner stack gas contact cooling water, 7.
- Condenser blowdown, and 8.
- 9. Mercury cleaning bath water.

WASTEWATER FLOW RATES

Data supplied by dcp responses were evaluated, and two flow-toproduction ratios, water use and wastewater discharge flow, were calculated for each stream. The two ratios are differentiated by the flow value used in calculation. Water use is defined as the volume of water or other fluid required for a given process per mass of product and is therefore based on the sum of recycle and make-up flows to a given process. Wastewater flow discharged after pretreatment or recycle (if these are present) is used in calculating the production normalized flow--the volume of wastewater discharged from a given process to further treatment, disposal, or discharge per mass of product produced. Differences between the water use and wastewater flows associated with a given stream result from recycle, evaporation, and carry-over on product. The production values used in calculation the correspond to the production normalizing parameter, PNP, assigned to each stream, as outlined in Section IV. As an example, calcine quench water flow is related to the production of refined mercury. As such, the discharge rate is expressed in liters of quench water per metric ton of mercury produced (gallons of quench water per ton of mercury).

The production normalized discharge flows were compiled and statistically analyzed by stream type. These production normalized water use and discharge flows are presented by subdivision in Tables V-1 through V-9 (pages 2189 - 2192) at the end of this section. Where appropriate, an attempt was made to identify factors that could account for variations in water use and discharge rates. These variations are discussed later in this section by subdivision. A similar analysis of factors affecting the wastewater flows is presented in Sections X, XI, and XII where representative BAT, NSPS, and pretreatment flows are selected for use in calculating the effluent limitations.

The water use and discharge rates shown do not include nonprocess wastewater, such as rainfall runoff and noncontact cooling water.

Data used to characterize the various wastewaters associated with primary precious metals and mercury production come from two sources--data collection portfolios and analytical data from field sampling trips.

DATA COLLECTION PORTFOLIOS

In the data collection portfolios, the primary precious metals and mercury plants that generate wastewater were asked to specify the presence or absence of priority pollutants in their wastewater. In most cases, the plants indicated that the priority organic pollutants were believed to be absent. However, two of the plants stated that they either knew or believed priority metals to be present. The responses for asbestos, cyanide, and the priority metals are summarized below:*

<u>Pollutant</u>	Known	Present	<u>Be</u>	lieved	Present
Antimony		0		0	
Arsenic	,	1 •		0	4.
Asbestos		0		. 1	
Beryllium		0		0	
Cadmium		1		0	· ·
Chromium		1	. · · · ·	. 0	×
Copper		1 .		0	÷ .
Cyanide		1		0	×
Lead		1 ·		0	
Mercury		2		1	
Nickel	¢	1		0	
Selenium		1		0	1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -
Silver		1		· 0.	- 4
Thallium	· ·	0		0	1
Zinc		1.	1	0	

*Six plants which produce primary precious metals and mercury have been omitted due to lack of data.

Although asbestos was reported as believed present by one plant, the trip report from this facility stated it was the mineral cummingtonite which was present and not asbestos. While the two minerals have some similarities such as similar chemical formulas, cummingtonite is not listed by EPA as a priority pollutant.

FIELD SAMPLING DATA

In order to quantify the concentrations of pollutants present in wastewater from primary precious metals and mercury plants, wastewater samples were collected at two plants. The analytical results from one of these two plants are not presented here

because they are claimed to be confidential by the plant. A diagram indicating the sampling sites and contributing production processes at the non-confidential plant is shown in Figure V-1 (page 2208).

Raw wastewater data are summarized in Tables V-10 through V-13 (pages 2193 - 2205). Analytical results for the combined stream of smelter scrubber water and Miller electrolysis cell scrubber water as well as spent solution from silver reduction are classified as confidential. Table V-10 presents data for each of the three stages of the calciner scrubber system. Tables V-11, V-12, and V-13 present sampling data for calcine quench water, calciner stack gas cooling water, and mercury cleaning bath water, respectively. Note that the stream numbers listed in the tables correspond to those given in the individual plant sampling site diagram, Figure V-1. Where no data are listed for a specific day of sampling, the wastewater samples for the stream were not collected.

Several points regarding these tables should be noted. The data tables include some samples measured at concentrations considered not quantifiable. The base-neutral extractable, acid extractable, and volatile organics generally are considered not quantifiable at concentrations equal to or less than 0.010 mg/1. Below this concentration, organic analytical results are not quantitatively accurate; however, the analyses are useful to indicate the presence of a particular pollutant. The pesticide fraction is considered not quantifiable at concentrations equal to or less than 0.005 mg/1.

The detection limits shown on the data tables for toxic metals and conventional and nonconventional pollutants are not the same in all cases as the published detection limits for these pollutants by the same analytical methods. The detection limits used were reported with the analytical data and hence are the appropriate limits to apply to the data. Detection limit variation can occur as a result of a number of laboratoryspecific, equipment-specific, and daily operator-specific factors. These factors can include day-to-day differences in machine calibration, variation in stock solutions, and variation in operators.

The statistical analysis of data includes some samples measured concentrations considered not quantifiable. at For data considered as detected but below quantifiable concentrations, a of zero is used for averaging. value Priority organic, nonconventional, and conventional pollutant data reported with a "less than" sign are considered as detected, but not further quantifiable. A value of zero is also used for averaging. If a pollutant is reported as not detected, it is assigned a value of zero in calculating the average. Toxic metal values reported as than a certain value were considered as less below quantification, and consequently were assigned a value of zero in the calculation of the average.

Finally, appropriate source water concentrations are presented with the summaries of the sampling data. The method by which each sample was collected is indicated by number, as follows:

One-time grab

1

- 2 Manual composite during intermittent process operation
- 3 8-hour manual composite
- 4 8-hour automatic composite
- 5 24-hour manual composite
- 6 24-hour automatic composite

WASTEWATER CHARACTERISTICS AND FLOWS BY SUBDIVISION

Since primary precious metals and mercury production involves nine principal sources of wastewater and each has potentially different characteristics and flows, the wastewater characteristics and discharge rates corresponding to each subdivision will be described separately. A brief description of why the associated production processes generate a wastewater and explanations for variations of water use within each subdivision will also be discussed.

Two subdivisions, gold slimes acid wash and water rinse, and silver crystals wash water have been deleted following proposal. These subdivisions have been deleted based on information obtained as a result of a post-proposal request for data. The one facility which was believed to discharge these streams reported that this water is totally reused in other plant processes.

SMELTER WET AIR POLLUTION CONTROL

Six of the eight plants in this subcategory smelt or roast the precious metal-bearing raw material. Only three of those facilities, however, use a wet air pollution control device to control air emissions from the furnace. Two of these devices are scrubbers, while one (at plant 1003) is an electrostatic precipitator (ESP). Two plants practice dry air pollution control, and one plant does not practice any air pollution control. The production normalized water use and discharge rates are presented in Table V-1 (page 2189) in liters per troy ounce of gold and silver smelted.

Analytical data for the combined smelter wet air pollution control and electrolytic cells wet air pollution control waste streams are contained in the confidential record. The data show that this wastewater contains treatable concentrations of toxic metals, suspended solids, and oil and grease.

SILVER CHLORIDE REDUCTION SPENT SOLUTION

Silver metal is produced from silver chloride by a dissolution and cementation process. The silver chloride is dissolved in water and recovered by cementation. The silver is replaced in solution, causing the silver ions to be reduced and precipitated from solution as silver metal. The resulting solution is a

wastewater stream. The silver chloride used as a feed material to this process may be a product of the Miller process or silver chloride precipitated from a silver ore leaching solution. The production normalized water use and discharge flows are presented in Table V-2 (page 2189), in liters per troy ounce of silver reduced in solution.

One plant supplied information in a telephone conversation with EPA concerning a post-proposal plant self-sampling effort which included revised process information. This information shows that the plant recovers silver metal from silver ore using a reduction of silver chloride process. This plant generates a spent solution from this process which it discharges. No flow information was reported for this stream, during the telephone conversation. In the self-sampling effort, information supplied by the facility corroborates the flow selected for BAT for this stream (see Section X), although this was not quantified precisely enough to be used to revise the BAT flow.

Following proposal, sampling data for spent silver chloride reduction solution were acquired at the specific request of EPA through a self-sampling effort. These self sampling data are presented in Table V-14 (page 2208) and show treatable concentrations of antimony, arsenic, cadmium, chromium, copper, lead, nickel, silver, and zinc, thus corroborating the data used at proposal.

ELECTROLYTIC CELLS WET AIR POLLUTION CONTROL

The use of wet scrubbers to control emissions from electrolytic cells is practiced at only one plant in this subcategory. Production normalized water use and discharge rates are presented in Table V-3 (page 2190). Sampling data for the combined smelter wet air pollution control and electrolytic cell wet air pollution control waste stream are contained in the confidential record. The data show this waste stream to contain treatable concentrations of toxic metals, suspended solids, and oil and grease.

ELECTROLYTE PREPARATION WET AIR POLLUTION CONTROL

The silver nitrate electrolyte used in the electrolytic in refining of Dore metal is prepared by dissolving pure silver nitric acid. The facility that uses this process also uses a wet scrubber to control air emissions from the preparation step, thereby generating a waste stream. Production normalized water use and discharge rates are presented in Table V-4 (page 2190) in liters per troy ounce of silver in electrolyte produced. No sampling data were gathered for this waste stream prior to proposal; however, it was expected to have characteristics similar to those of the combined raw wastewaters from smelter wet air pollution control and the electrolytic cell scrubber. This waste stream, therefore, was expected to contain treatable concentrations of suspended solids, toxic metals, and oil and grease.

Following proposal, sampling data for this subdivision were acquired at the specific request of EPA through a self-sampling effort. These data are presented in Table V-14 and show a pH of 1.23 and treatable concentrations of arsenic, cadmium, chromium, copper, lead, mercury, nickel, silver, and zinc, thus corroborating the data used at proposal.

CALCINER WET AIR POLLUTION CONTROL

One of the two plants producing primary mercury uses a water scrubber to control air emissions from the calciner. This plant uses a series of three scrubbers (Venturi, impinger, and SO_2). Sampling data for the wastewater generated by these scrubbers are presented in Table V-10 (page 2202). The scrubber waters have a low pH (2.3 to 2.6) and contain treatable concentrations of priority metals such as lead, mercury, thallium and zinc, and suspended solids. The production normalized water use and discharge rates are shown in Table V-5 (page 2192).

CALCINE QUENCH WATER

One mercury producer uses water to quench the waste calcines from the mercury roaster to allow faster handling and disposal of these materials. Table V-6 (page 2192) presents the production normalized water use and discharge rates for this waste stream. Sampling data are summarized in Table V-11 (page 2205) and show high concentrations of priority metals such as arsenic, mercury and zinc, and suspended solids. This waste stream has a nearly neutral pH of 6.8.

CALCINER STACK GAS CONTACT COOLING WATER

One facility uses contact cooling water to reduce the temperature of the calciner off-gases before releasing them to the atmosphere. Sampling data for this waste stream are summarized in Table V-12 (page 2209). This waste stream has a pH of 2.5 and contains treatable concentrations of mercury and suspended solids. Production normalized water use and discharge rates are given in Table V-7 (page 2192).

CONDENSER BLOWDOWN

When mercury is vaporized in the calciner, some water contained in the Cinnabar or gold ore may also be vaporized. The condensation of mercury for recovery may result in the condensation of some water which is discharged as condenser blowdown. Table V-8 (page 2193) summarizes the production normalized water use and discharge rates for this waste stream.

Although no sampling data were collected for this waste stream, it is expected to be very similar to the discharge from the mercury cleaning bath. The condenser blowdown stream is expected to contain treatable concentrations of mercury and suspended solids.

MERCURY CLEANING BATH WATER

Condensed mercury is processed for the removal of impurities by being passed through a water cleaning bath. This waste stream contains treatable concentrations of mercury and suspended solids and very low concentrations of other toxic metals. The sampling data for this wastewater stream are presented in Table V-13 (page 2205). Production normalized water use and discharge rates are provided in Table V-9 (page 2199).

TABLE V-1

WATER USE AND DISCHARGE RATES FOR SMELTER WET AIR POLLUTION CONTROL

(1/troy ounce of gold and silver smelted)

<u>Plant</u> Code	Percent Recycle	Production Normalized Water <u>Use</u>	Production Normalized Discharge <u>Flow</u>
1131*	76	25.8	6.2
1003	90	5.3	0.53
1137	100	8.41	0
1068	Dry	,	ч
1158	Dry		

*No operations conducted in 1982: water use and discharge rates based on projected 1983 figures.

TABLE V-2

WATER USE AND DISCHARGE RATES FOR SILVER CHLORIDE REDUCTION SPENT SOLUTION

(l/troy ounce of silver reduced in solution)

<u>Plant</u> Code	Percent Recycle	Production Normalized Water Use	Production Normalized Discharge <u>Flow</u>
1003	0	0.4	0.4
1160	0	NR	NR

NR - Data not reported.

TABLE V-3

WATER USE AND DISCHARGE RATES FOR ELECTROLYTIC CELLS WET AIR POLLUTION CONTROL

(1/troy ounce of gold refined electrolytically)

Plant Code	Percent Recycle	Production Normalized Water <u>Use</u>	Production Normalized Discharge <u>Flow</u>
1003	0	198	19

TABLE V-4

WATER USE AND DISCHARGE RATES FOR ELECTROLYTE PREPARATION WET AIR POLLUTION CONTROL

(1/troy ounce of silver in electrolyte produced)

<u>Plant</u> Code	Percent Recycle	Production Normalized Water Use	Production Normalized Discharge <u>Flow</u>
1160	0	0.05	0.05

TABLE V-5

WATER USE AND DISCHARGE RATES FOR CALCINER WET AIR POLLUTION CONTROL

(1/kkg of mercury condensed)

Percent Recycle	Production Normalized Water Use	Production Normalized Discharge <u>Flow</u>
16	4,607	3,870
16	7,536	6,330
<u>16</u>	209,524	176,000
16	221,667	186,200
	<u>Recycle</u> 16 16 <u>16</u>	Percent RecycleNormalized Water164,607167,53616209,524

TABLE V-6

WATER USE AND DISCHARGE RATES FOR CALCINE QUENCH WATER

(1/kkg of mercury condensed)

<u>Plant</u> Code	Percent Recycle	Production Normalized Water <u>Use</u>	Production Normalized Discharge <u>Flow</u>
1124	0	17,600	17,600

TABLE V-7

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WATER USE AND DISCHARGE RATES FOR CALCINER STACK GAS CONTACT COOLING WATER

(1/kkg of mercury condensed)

<u>Plant</u> Code	Percent <u>Recycle</u>	Production Normalized Water <u>Use</u>	Production Normalized Discharge <u>Flow</u>
1124	0	4,150	4,150

TABLE V-8

WATER USE AND DISCHARGE RATES FOR CONDENSER BLOWDOWN

(1/kkg of mercury condensed)

<u>Plant</u> Code	Percent Recycle	Production Normalized Water <u>Use</u>	Production Normalized Discharge <u>Flow</u>
1068	0	13,800	13,800
1124	Dry		

TABLE V-9

WATER USE AND DISCHARGE RATES FOR MERCURY CLEANING BATH WATER

(1/kkg of mercury condensed)

Plant Code	Percent Recycle	Production Normalized Water <u>Use</u>	Production Normalized Discharge <u>Flow</u>
1124	0	1,400	1,400

Table V-10

	RAW WAST	EWATER				
Pollutant	Stream Code	Sample Typet	Con Source	centration Day 1	s (mg/1) Day 2	Day 3
<u>Toxic Pollutants</u>			• ,	·		
114. antimony	40 41 42	1 1 1	<0.003 <0.003 <0.003	<0.003 <0.003 <0.003	• •	
115. arsenic	40 41 42	1 1 1	0.013 0.013 0.013	0.32 0.059 0.013	•	
117. beryllium	40 41 42	1 1 1	<0.01 <0.01 <0.01	<0.01 <0.01 <0.01	:	
118. cadmium	40 41 42	1 • 1 • •	<0.01 <0.01 <0.01	0.04 <0.02 <0.01		- - -
119. chromium (total)	40 41 42	1 1 1	<0.02 <0.02 <0.02	<0.02 <0.02 <0.02	en de la companya de	
120. copper	40 41 42	1	0.31 0.31 0.31	<0.01 <0.01 <0.01		

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SAMPLING DATA CALCINER WET AIR POLLUTION CONTROL RAW WASTEWATER

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PRIMARY PRECIOUS METALS

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SAMPLING DATA CALCINER WET AIR POLLUTION CONTROL RAW WASTEWATER

Pollutant	Stream Code	Sample <u>Typet</u>	Con Source	<u>Centration</u> Day 1	<u>ns (mg/1)</u> Day 2	Day 3
Toxic Pollutants (Continued)						
122. lead	40	1	<0.002	2.2		
	41	1	<0.002	<0.002		
	42	1	<0.002	<0.002		x
123. mercury	40	1	0.016	360		-
	41	1	0.016	130		
	42	1	0.016	0.84		
124. nickel	40	1 · · · ·	<0.05	<0.05		
	41	1	<0.05	<0.05		
	42	1	<0.05	<0.05		
125. selenium	40	1	<0.003	<0.003		
	41	1	<0.003	<0.003		*
	42	1	<0.003	<0.003		
126. silver	40	1	<0.001	<0.001		
	41	1	<0.001	<0.001		
	42	1	<0.001	<0.001		
127. thallium	40	1	<0.002	0.61		-
	41	1	<0.002	0.12		
	42	1	<0.002	<0,002		
128. zinc	40	1	<0.01	0.73		
	41	1	<0.01	<0.01		
··· .	42	1	<0.01	<0.01		

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Pollutant	Stream Code	Sample Typet	Concentration Source Day 1	ns (mg/1) <u>Day 2</u> <u>Day 3</u>
Nonconventional Pollutants				
acidity	40 41 42	1 1 1	<1 490 <1 490 <1 1,430	
alkalinity	40 41 42	1 1 1	190 <1 190 <1 190 <1	
aluminum	40 41 42	1 1 1	<0.05 <0.05 <0.05 <0.05 <0.05 <0.05	
barium	40 41 42	1 1 1	0.05 0.05 0.021 0.05 0.053	
boron	40 41 42	1 1 1	0.041 <0.009 0.041 <0.009 0.041 0.027	
calcium	40 41 42	1 1 1	52 52 52 53 52 51	
chloride	40 41 42	1 1 1	50 270 50 75 50 91	

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SAMPLING DATA CALCINER WET AIR POLLUTION CONTROL RAW WASTEWATER

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SECT

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SAMPLING DATA CALCINER WET AIR POLLUTION CONTROL RAW WASTEWATER

Pollutant	Stream Code	Sample Typet	Conc Source	entrations (mg/l) Day 1 Day 2	Day 3
Nonconventional Pollutants (Continued)				
cobalt	40 41 42	1 1 1	<0.006 <0.006 <0.006	<0.006 <0.006 <0.006	
fluoride	40 41 42	1 1 1	1.1 1.1 1.1	0.77 0.84 1.1	
iron	40 41 42	1 1 1	0.05 0.05 0.05	1.0 0.24 <0.02	
magnesium	40 41 42	1 1 1	8.0 8.0 8.0	7.7 8 0 8.0	
manganese	40 41 42	1 1. 1	<0.01 <0.01 <0.01	<0.01 <0.01 <0.01	
molybdenum	40 41 42	1 1 1	<0.002 <0.002 <0.002	<0.002 <0.002 <0.002	
sodium	40 41 42	1 1 1	53 53 53	83 60 52	

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PRIMARY PRECIOUS METALS

	RAW WASTEWATER	CONTROL
<u>Pollutant</u>	Stream Sample Code Typet	Concentrations (mg/1) Source Day 1 Day 2
Nonconventional Pollutants (Continued)		
sulfate	40 1 41 1 42 1	150 67,000 150 68,000 150 17,000
tin	40 1 41 1 42 1	<0.12 <0.12 <0.12 <0.12 <0.12 <0.12
titanium	40 1 41 1 42 1	<0.005 <0.005 <0.005 <0.005 <0.005 <0.005
total solids (TS)	40 1 41 1 42 1	670 1,300 670 800 670 700
vanadium	40 1 41 1 42 1	<0.003 <0.003 <0.003 <0.003 <0.003 <0.003
yttrium	40 1 41 1 42 1	<0.002 <0.002 <0.002 <0.002 <0.002 <0.002

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SAMPLING DATA CALCINER WET AIR POLLUTION CONTROL

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PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SAMPLING DATA CALCINER WET AIR POLLUTION CONTROL RAW WASTEWATER

Pollutant	Stream Code	Sample Typet	<u>Con</u> Source	centration Day 1	<u>s (mg/1)</u> Day 2	Day 3
Conventional Pollutants						
oil and grease	40 41 42	1 1 1	<1 <1 <1	<1 <1 <1		
total suspended solids (TSS)	40 41 42	1 1 1	<1 <1 <1	80 5 <1		
pH (standard units)	40 41 42	1 1 1	6.9 6.9 6.9	2.3 2.3 2.6		

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tSample Type Code: 1 - One time grab

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PRIMARY PRECIOUS METALS

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Table V-11

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SAMPLING DATA CALCINE QUENCH WATER RAW WASTEWATER

<u>Pollutant</u>	Stream <u>Code</u>	Sample Typet	Con Source	<u>centration</u> Day 1	<u>s (mg/1)</u> Day 2	Day 3
<u>Toxic Pollutants</u>			· .			
114. antimony	45	1	<0.003	<0.007		
115. arsenic	45	1	0.013	17	-	
117. beryllium	45	1	<0.01	<0.01		÷ .
118. cadmium	45	1	<0.01	0.06		
119. chromium (total)	45	1	<0.02	0.09		
120. copper	45	1	0.31	0.30		
122. lead	45	1	<0.002	0.38		
123. mercury	45	1	0.016	1.4		,
124. nickel	45	1	<0.05	<0.05		
125. selenium	45	1.	<0.003	<0.003	· · ·	
126. silver	45	1	<0.001	0.13		
127. thallium	45	1	<0.002	0.19	•	· ·
128. zinc	45	1	<0.001	1.7		

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PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SECT

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PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SAMPLING DATA CALCINE QUENCH WATER RAW WASTEWATER

Pollutant	Stream Code	Sample Typet	Concer Source	ntrations Day 1	(mg/1) Day 2	Day 3
Nonconventional Pollutants			,			
acidity	45	1	< 1 ·	<1		•
alkalinity	45	1	190 4	48		
aluminum	45	1	<0.50	52		
barium	45	1	0.05	1.6		an r a
boron	45	· 1	0.041	1.3		
calcium	45	1	52 1,80	00		
chloride	45	1 .	50 93	30	1	
cobalt	45	1	<0.006	0.044		
fluoride	. 45	. 1	1.1	5.6		
iron	45	1	0.05 1	50	u	-
magnesium	45	. 1	8.0 2	21		
manganese	45	1	<0.01	0.75		
molybdenum	45	1	<0.002	0.66		_

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Table V-11 (Continued)

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SAMPLING DATA CALCINE QUENCH WATER RAW WASTEWATER

Pollutant	Stream Code	Sample Typet	Concentrations Source Day 1	(mg/1) Day 2 Day 3
Nonconventional Pollutants (Continu	ed)			
sodium	45	1	53 2,600	- - -
sulfate	45	1	150 1,900	
tin	45	1	<0.12 <0.12	
titanium	45	1	<0.005 12	
total solids (TS)	45	1	670 13,000	
vanadium	45	1	<0.003 0.55	
yttrium	45	4	<0.002 0.14	
Conventional Pollutants	,			ана страна с Страна страна с
oil and grease	45	. 1	<1 <1	
total suspended solids	45	1	<1 3,700	
pH (standard units)	45	1	6.9 6.8	• •

1 - One-time grab tSample Type Code:

2201

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SECT

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Table V-12

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SAMPLING DATA CALCINER STACK GAS CONTACT COOLING WATER RAW WASTEWATER

Dollar to a t		Stream	Sample	Con			
	Pollutant	Code	<u>Typet</u>	Source	Day 1	(mg/1) Day 2	Day 3
<u>Toxic</u>	e Pollutants						
114.	antimony	43	1	<0.003	<0.003		
115.	arsenic	43	1	0.013	0.017		
117.	beryllium	43	1	<0.01	<0.01		
118.	cadmium	43	1	<0.01	<0.01		. <u>.</u>
119.	chromium (total)	43	[·] 1	<0.02	<0.02		
120.	copper	43	1	0.31	<0.01		- ;
122.	lead	43	. 1	<0.002	<0.002	÷	
123.	mercury	43	1	0.016	2.1		
124.	nickel	43	1 [.]	<0.05	<0.05		
125.	selenium	43	· 1	<0.003	<0.003	,	-
126.	silver	43	1	<0.001	<0.001		
127.	thallium	43	1	<0.002	0.004		
128.	zinc	43	1	<0.01	<0.02	•	

SECT - V

AND MERCURY SUBCATEGORY

PRIMARY PRECIOUS METALS

Table V-12 (Continued)

	Stream	Sample	Conc	Concentrations (mg/1)					
<u>Pollutant</u>	Code	Typet	Source	Day 1	Day 2	Day 3			
Nonconventional Pollutants				•					
acidity	43	1	<1 1,	,800					
alkalinity	43	1	190	<1					
aluminum	43	1	<0.50	<0.50					
barium	43	1	0.050	0.047		- -			
boron	43	1 +	0.041	<0.009					
calcium	43	. 1	52	52	* •				
chloride	43	. 1	50	53					
cobalt	43	1	<0.006	<0.006	·				
fluoride	43	1	1.1	1.1		•			
iron	43	. 1	0.05	0.039		. ·			
magnesium	43	1	8.0	8.0	•				
manganese	43	1	<0.01	<0.01					
molybdenum	43	1	<0.002	<0.002					

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SAMPLING DATA CALCINER STACK GAS CONTACT COOLING WATER RAW WASTEWATER

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

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Table V-12 (Continued)

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SAMPLING DATA CALCINER STACK GAS CONTACT COOLING WATER RAW WASTEWATER

Pollutant	Stream <u>Code</u>	Sample <u>Typet</u>	<u>Concentrations (mg/1)</u> Source <u>Day 1 Day 2 Day 3</u>
Nonconventional Pollutants (Continued)			
sodium	43	. 1	53 53
sulfate	43	1	150 23,600
tin	43	1	<0.12 <0.12
titanium	43	1	<0.005 <0.005
total solids (TS)	43	1	670 880
vanadium	43	1	<0.003 <0.003
yttrium	43	1	<0.002 <0.002
Conventional Pollutants	· ·		
oil and grease	43	, " 1	<1 <1
total suspended solids	43	1	<1 4
pH (standard units)	43	1	6.9 2.5

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SECT I

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Table V-13

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SAMPLING DATA MERCURY CLEANING BATH WATER RAW WASTEWATER

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PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SECT

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Table V-13 (Continued)

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SAMPLING DATA MERCURY CLEANING BATH WATER RAW WASTEWATER

Stream <u>Code</u>	Sample Typet	<u>Concentrations (mg/1)</u> Source Day 1 Day 2 Day 3
44	1	<1 1,700
44	1	190 170
44	1	<0.5 <0.5
44	1	0.050 0.059
44	1	0.041 0.022
44	1	52 52
44	1	50 47
44	1	<0.006 <0.006
44	1	1.1 0.96
44	1	0.05 <0.02
44	1	8.0 7.9
44	1	<0.01 <0.01
44	. 1	<0.002 <0.002
	<u>Code</u> 44 44 44 44 44 44 44 44 44 44 44 44	CodeTypet441441441441441441441441441441441441441441441441441

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PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

Table V-13 (Continued)

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SAMPLING DATA MERCURY CLEANING BATH WATER RAW WASTEWATER

	Stream	Sample		centrations	
<u>Pollutant</u>	Code	<u>Type†</u>	Source	<u>Day 1</u>	<u>Day 2</u> <u>Day 3</u>
Nonconventional Pollutants (Continued)					
sodium	44	1	53	53	
sulfate	44	1	150	74	
tin	44	1	<0.12	<0.12	
titanium	44	1	<0.005	<0.005	
total solids (TS)	44	1	670	690	
vanadium	44	1	<0.003	<0.003	•
yttrium	44	1	<0.002	<0.002	
Conventional Pollutants		•		•	۰ ۲۰ ۲
oil and grease	44	1	<1	<1	
total suspended solids	44	1	<1	4	
pH (standard units)	44	1	6.9	7.5	

tSample Type Code: 1 - One-time grab

2207

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SECT -

TABLE V-14

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY RAW WASTEWATER SELF SAMPLING DATA (A - Silver Chloride Reduction Spent Solution) (B - Electrolyte Preparation Wet Air Pollution Control)

Pollutant	Source A	Source B
Sample Number	88151	88150
Toxic Pollutants		
Antimony Arsenic Beryllium	1.86 0.479 <0.05	28.36 <0.05
Cadmium Chromium Copper	0.43 1.06 62.0	0.36 0.168 534.0
Lead Mercury Nickel	7.02 0.086 2.4	12.6 0.014 0.35
Silver Thallium Zinc	0.011 <0.01 15.0	15.72 <0.01 42.0
Nonconventional Pollu	tants	
Aluminum Cobalt Iron	3.0 0.55 5000.0	1.1 0.55 400.0
Manganese Molybdenum Tin	106.0 1.37 14.0	6.4 <0.5 <5.0
Titanium Vanadium Gold	<0.2 1.6 <0.005	<pre> <0.2 l.0 <0.005 </pre>

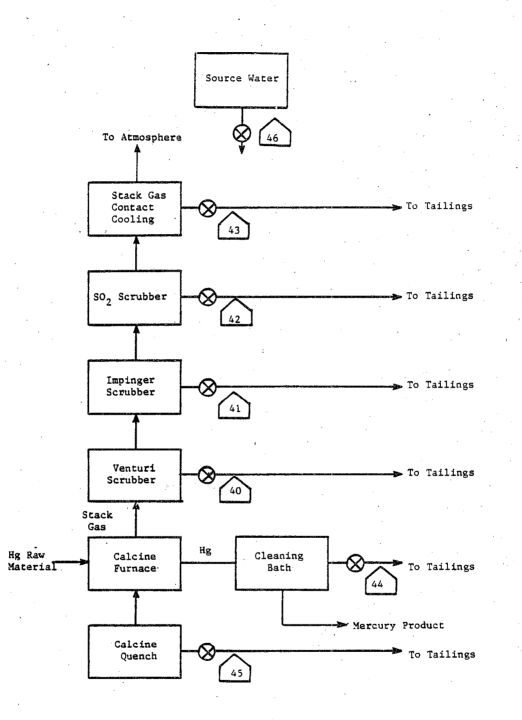


Figure V-1

SAMPLE LOCATIONS AT PRIMARY PRECIOUS METALS AND MERCURY PLANT A .

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SECTION VI

SELECTION OF POLLUTANT PARAMETERS

section examines both the confidential and nonconfidential This chemical analysis data and discusses the selection or exclusion of pollutants for potential limitation. The analytical data from one primary precious metals plant was not presented in Section V The basis for the because it was claimed to be confidential. regulation of toxic and other pollutants, along with a discussion of each pollutant selected for potential limitation is discussed Section VI of Vol. I. That discussion provides information in concerning the nature of the pollutant (i.e., whether it is a naturally occurring substance, processed metal, or a manufactured compound); general physical properties and the form of the toxic effects of the pollutant in humans and other pollutant; behavior of the pollutant in POTW at the animals: and concentrations expected in industrial discharges.

The discussion that follows describes the analysis that was performed to select or exclude toxic pollutants for further consideration for limitations and standards. Also, it describes the analysis that was performed to select or exclude conventional for limitation. Priority pollutants will be pollutants considered for limitation if they are present in concentrations treatable by the technologies considered in this analysis. The treatable concentrations used for the priority metals were the performance achievable chemical values by long-term sedimentation, and filtration. The treatable precipitation, concentrations used for the priority organics were the long-term performance values achievable by carbon adsorption.

This study examined samples from the primary precious metals and mercury subcategory for one nonconventional pollutant (gold) and three conventional pollutant parameters (oil and grease, total suspended solids, and pH).

CONVENTIONAL AND NONCONVENTIONAL POLLUTANT PARAMETERS SELECTED

The nonconventional and conventional pollutants or pollutant parameters selected for limitation in this subcategory are:

gold oil and grease total suspended solids (TSS) pH

Gold was analyzed for and not detected in two samples of raw wastewater from this subcategory. However, gold is expected to be present in the raw wastewater because of its presence in the raw materials and its solubility in the various acids and bases used as raw materials in the refining processes. Gold was presented as being considered for regulation in the Notice of

Data Availability (see 50 FR 10919). For these reasons, gold is selected for limitation in this subcategory.

and grease was detected in two of 10 samples Oil at concentrations above the treatability concentration of 10.0 mg/1. The measured concentrations were 60 and 170 mg/l. These high concentrations occurred in the combined raw wastewater stream from the smelter and electrolytic cells wet air pollution control and in the silver chloride reduction spent solution. Therefore, and grease is selected for limitation in this subcategory. oil was detected at concentrations above the treatability TSS concentration of 2.6 mg/l in eight of the 10 raw waste samples analyzed for this study. These eight TSS concentration values ranged from 4 to 3,700 mg/1. Furthermore, most of the specific methods used to remove toxic metals do so by converting these metals to precipitates, and these toxic metal-containing precipitates should not be discharged. Meeting a limitation on total suspended solids helps ensure that removal of these precipitated toxic metals has been effective. For these reasons, total suspended solids are selected for limitation in this subcategory.

The nine pH values observed during this study ranged from 0.9 to 8.4. Six of the nine values were equal to or less than 2.6, one value was 6.8 and the other two fell within the 7.5 to 10.0 range considered desirable for discharge to receiving waters. Many deleterious effects are caused by extreme pH values or rapid changes in pH. Also, effective removal of toxic metals by precipitation requires careful control of pH. Since pH control within the desirable limits is readily attainable by available treatment, pH is selected for limitation in this subcategory.

TOXIC PRIORITY POLLUTANTS

The frequency of occurrence of the priority pollutants in the raw wastewater samples taken is presented in Table VI-1 (page 2217). Table VI-1 is based on the raw wastewater data presented in Section V (see Tables V-10 through V-13, pages 2793-2205) as well as the primary precious metals analytical data being held confidential. These data provide the basis for the categorization of specific pollutants, as discussed below.

TOXIC POLLUTANTS NEVER DETECTED

The toxic pollutants listed in Table VI-2 (page 2221) were not detected in any raw wastewater samples from this subcategory; therefore, they are not selected for consideration in establishing limitations:

TOXIC POLLUTANTS NEVER FOUND ABOVE THEIR ANALYTICAL QUANTIFICATION CONCENTRATION

The priority pollutants listed below were never found above their analytical quantification concentration in any raw wastewater samples from this subcategory; therefore, they are not selected

for consideration in establishing limitations.

65. phenol

- 66. bis(2-ethylhexyl) phthalate
- 68. di-n-butyl phthalate
- 78. anthracene (a)
- 81. phenanthrene (a)
- 114. antimony

(a) Reported together, as a combined value

TOXIC POLLUTANTS PRESENT BELOW CONCENTRATIONS ACHIEVABLE BY TREATMENT

The pollutants listed below are not selected for consideration in establishing limitations because they were not found in any raw wastewater samples from this subcategory above concentrations considered achievable by existing or available treatment technologies. These pollutants are discussed individually following the list.

117. beryllium
125. selenium

Beryllium was detected at a concentration of 0.15 mg/l in one of the 10 samples analyzed. Available treatment methods can reduce beryllium concentrations only to 0.2 mg/l and this pollutant is, therefore, not considered for limitation.

Selenium was detected in two of 10 samples at concentrations ranging from 0.044 to 0.063 mg/1. These concentrations are below the minimum selenium concentration of 0.2 mg/1 achievable by available treatment methods. Additionally, these concentrations of selenium may be attributable to its presence in the source water at a concentration of 0.10 mg/1. Selenium, therefore, is not considered for limitation.

TOXIC POLLUTANTS DETECTED IN A SMALL NUMBER OF SOURCES

The following pollutants were not selected for limitation because they are detectable in the effluent from only a small number of sources within the subcategory and they are uniquely related to only those sources.

- 4. benzene
- 44. methylene chloride
- 70. diethyl phthalate
- 86. toluene
- 121. cyanide

Although these pollutants were not selected for limitation in establishing nationwide regulations, it may be appropriate, on a case-by-case basis, for the local permitter to specify effluent limitations.

Benzene was detected above its treatable concentration of 0.01 mg/l in one of three samples analyzed at a concentration of 0.016 mg/l. This pollutant is not attributable to specific materials or processes associated with the primary precious metals and mercury subcategory, and is not expected to be present in the wastewater. For this reason, and because very little removal of benzene can be expected with treatment, this pollutant is not considered for limitation.

chloride was detected above its treatabilitv Methvlene concentration of 0.01 mg/l at concentrations ranging from 0.036 to 0.046 mg/l in all three samples analyzed. This pollutant is not attributable to specific materials or processes associated with the primary precious metals and mercury subcategory, but is a common solvent used in analytical laboratories. Because methylene chloride is not expected to be present in the as the high probability of wastewater, as well sample contamination, this pollutant is not considered for limitation.

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Diethyl phthalate was detected above its treatable concentration of 0.01 mg/l in one of three samples analyzed at a concentration of 0.016 mg/l. This pollutant is not attributable to specific materials or processes associated with the primary precious metals and mercury subcategory, and is not expected to be present in the wastewater. For this reason, and because very little removal of diethyl phthalate can be expected with treatment, this pollutant is not considered for limitation.

Toluene was detected above its treatable concentration of 0.01 mg/l in two of three samples analyzed at concentrations of 0.023 and 0.05 mg/l. This pollutant is not attributable to specific materials or processes associated with the primary precious metals and mercury subcategory, and is not expected to be present in the wastewater. For this reason, and because very little removal of toluene can be expected with treatment, this pollutant is not considered for limitation.

Cyanide was measured at concentrations ranging from 0.049 to 0.2 mg/l in three of the four samples for which it was analyzed. These concentrations are above the treatability concentration of 0.047 mg/l, but are suspected to be present because of source water contamination. The source water was found to contain cyanide at a concentration of 8.6 mg/l. Because of its presence in the source water at a high concentration, cyanide is not considered for limitation.

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TOXICPOLLUTANTSSELECTEDFORFURTHERCONSIDERATIONINESTABLISHINGLIMITATIONSANDSTANDARDSIN

The toxic pollutants listed below are selected for further consideration in establishing limitations and standards for this subcategory. The pollutants selected for further consideration for limitation are each discussed following the list.

115. arsenic
118. cadmium
119. chromium
120. copper
122. lead
123. mercury
124. nickel
126. silver
127. thallium
128. zinc

Arsenic was detected in two of 10 samples at concentrations of 0.6 and 17 mg/l. The concentration achievable by treatment methods is 0.34 mg/l. These concentrations were detected in silver chloride reduction spent solution and calcine quench water. Arsenic was detected, but at levels below treatability, in the other eight samples. Therefore, arsenic is selected for further consideration for limitation.

Cadmium was detected above its treatable concentration (0.049 mg/l) in two of 10 raw wastewater samples analyzed. The treatable concentrations were detected in silver chloride reduction spent solution and calcine quench water. Therefore, cadmium is selected for further consideration for limitation.

Chromium was detected above its treatable concentration of 0.07 mg/l in silver chloride reduction spent solution and calcine quench water. The highest concentration was 25 mg/l. All eight other samples indicated that chromium was present, out at a concentration below treatability. Therefore, chromium is selected for further consideration for limitation.

Copper was measured in two samples at concentrations above the treatable concentration of 0.39 mg/l. Copper was also detected in the remaining eight samples, out at concentrations below that achievable by treatment. The highest concentration of copper found was 23,000 mg/l. Therefore, copper is selected for further consideration for limitation.

Lead was detected in six raw waste streams at concentrations above the 0.08 mg/l attainable by identified treatment technology. These concentrations ranged from 0.1 to 600 mg/l. For this reason, lead is selected for further consideration for limitation.

Mercury was detected in six of the 10 samples analyzed at concentrations ranging from 0.84 to 360 mg/1. These concentrations are well above the concentration of 0.036

achievable by current treatment methods In addition, mercury was detected in the remaining four samples, but at values below the treatable concentration. For these reasons, mercury is selected for further consideration for limitation.

Nickel was detected in the silver chloride reduction spent solution at a concentration of 29 mg/1. The treatable concentration for nickel is 0.22 mg/1. Nickel was detected, but below treatable concentrations in all nine of the other samples. Therefore, nickel is selected for further consideration for limitation.

Silver was detected in two samples at concentrations of 0.13 and 6.1 mg/l. These concentrations are above silver's treatable concentration of 0.07 mg/l. Silver is, therefore, selected for further consideration for limitation.

Thallium was detected above its treatable concentration (0.34 mg/l) in two of 10 samples analyzed. The quantifiable concentrations ranged from 0.12 to 2.6 mg/l. Since thallium was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

Zinc was detected above its treatable concentration (0.23 mg/l) in three of 10 samples analyzed. The quantifiable concentrations ranged from 0.10 to 15.0 mg/l. Since zinc was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

Table VI-1

FREQUENCY OF OCCURRENCE OF PRIORITY POLLUTANTS PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY RAW WASTEWATER

Quantification Concentra- Number of Number of Detected Below Below Treat- Above Treat-		Analytical	Treatable		. *		_	2
$ \begin{array}{c c c c c c c c c c c c c c c c c c c $	•	Quantification	Concent ra-		Number of	Detected Below		
2. acrolein 0.010 0.01 2 3 3 3. acrylonitrile 0.010 0.01 2 3 3 4. benzene 0.010 0.01 2 3 3 5. benzidine 0.010 0.01 2 3 3 6. catbon tetrachloride 0.010 0.01 2 3 3 7. chlorobenzene 0.010 0.01 2 3 3 8. i, 2, 4-trichlorobenzene 0.010 0.01 2 3 3 9. hexachlorobenzene 0.010 0.01 2 3 3 11. 1, 1, 1-trichloroethane 0.010 0.01 2 3 3 12. hexachlorobenzene 0.010 0.01 2 3 3 13. 1, 1-dichloroethane 0.010 0.01 2 3 3 13. 1, 1-2ichloroethane 0.010 0.01 2 3 3 14. 1, 1, 2-tric roethane 0.010 0.01 2 3 3 14. 1, 1, 2-trick roethane 0.010 0.01 <	Pollutant				Samples <u>Analyzed ND</u>		able Concen-	able Concen-
2. acrolein 0.010 0.01 2 3 3. acrylonitrile 0.010 0.01 2 3 3 4. benzene 0.010 0.01 2 3 3 5. benzidine 0.010 0.01 2 3 3 7. chlorobenzene 0.010 0.01 2 3 3 8. i, 2, 4-trichlorobenzene 0.010 0.01 2 3 3 9. hexachlorobenzene 0.010 0.01 2 3 3 1 10. i, 2-dichloroethane 0.010 0.01 2 3 3 1 11. 1, 1, 1-trichloroethane 0.010 0.01 2 3 3 1 12. hexachloroethane 0.010 0.01 2 3 3 1 14. i, 1, 2-tric roethane 0.010 0.01 2 3 3 1 14. i, 1, 2, 2-tertachloroethane 0.010 0.01 2 3 3 1 15. i, 1, 2, 2-tertachloroethane 0.010 0.01 2 3 3 1		0.010	0.01	2	3 3		· .	
4. benzeme 0.010 0.01 2 3 3 2 1 5. benzitine 0.010 0.01 2 3 3 2 1 6. carbon tetrachloride 0.010 0.01 2 3 3 2 1 6. carbon tetrachloride 0.010 0.01 2 3 3 3 2 1 6. carbon tetrachloride 0.010 0.01 2 3		0.010		$\overline{2}$	3 3	· .		
5. benzitine 0.000 0.001 2 3 2 1 6. cathon tetrachloride 0.000 0.01 2 3 3 3 7. chlorobenzene 0.010 0.01 2 3 3 3 8. 1,2,4-trtichlorobenzene 0.010 0.01 2 3 3 3 9. bezachlorobenzene 0.010 0.01 2 3 3 3 9. hezachlorobenzene 0.010 0.01 2 3 3 3 10. 1, 2-dichloroethane 0.010 0.01 2 3 3 3 11. 1, 1-trichloroethane 0.010 0.01 2 3 3 3 12. hexachloroethane 0.010 0.01 2 3 3 3 13. 1, 1-dichloroethane 0.010 0.01 2 3 3 3 14. 1, 1, 2, 2-tetractrachoroethane 0.010 0.01 2 3 3 3 3 15. <th></th> <td></td> <td>0.01</td> <td>2</td> <td>3 3</td> <td></td> <td></td> <td>- Fi</td>			0.01	2	3 3			- Fi
1. blanche 0.000 0.01 2 3 3 7. chlorobenzene 0.010 0.01 2 3 3 8. l, 2, 4-trichlorobenzene 0.010 0.01 2 3 3 9. hexachlorobenzene 0.010 0.01 2 3 3 10. l, 2-dichloroethane 0.010 0.01 2 3 3 11. l, 1, 1-trichloroethane 0.010 0.01 2 3 3 12. hexachloroethane 0.010 0.01 2 3 3 13. l, 1-dichloroethane 0.010 0.01 2 3 3 14. l, 1, 2-tric . coethane 0.010 0.01 2 3 3 14. l, 1, 2-tric . coethane 0.010 0.01 2 3 3 15. l, 1, 2, 2-tertanchoroethane 0.010 0.01 2 3 3 3 16. chloroethyl ether 0.010 0.01 2 3 3 3 3 16. chloroophylatellene 0.010 0.01 2 3 3 3 <th></th> <td></td> <td>0.01</td> <td>2</td> <td>3</td> <td>2</td> <td>*</td> <td>, Þ</td>			0.01	2	3	2	*	, Þ
7. chlorobenzene 0.010 0.01 2 3 3 8. 1,2,4-trichlorobenzene 0.010 0.01 2 3 3 9. hexachlorobenzene 0.010 0.01 2 3 3 10. 1,2-dichloroethane 0.010 0.01 2 3 3 11. 1,1,1-trichloroethane 0.010 0.01 2 3 3 12. hexachloroethane 0.010 0.01 2 3 3 13. 1,1-dichloroethane 0.010 0.01 2 3 3 14. 1,1,2-tric roethane 0.010 0.01 2 3 3 15. 1,1,2,2-tetrachloroethane 0.010 0.01 2 3 3 16. chloroethane 0.010 0.01 2 3 3 17. bls(chloromethyl) ether 0.010 0.01 2 3 3 12. 2,4,6-trichloropthol 0.010 0.01 2 3 3 12. 2,4,6-trichloropthol 0.010 0.01 2 3 3 2. 2-chloronaphthaleme 0.010 <			0.01	2	้ จั้ จ	2		
8. 1,2,4-tritichlorobenzene 0,000 0,001 2 3 3 9. hexachlorobenzene 0,010 0,01 2 3 3 10. 1,2-ditchloroethane 0,010 0,01 2 3 3 11. 1,1-trichloroethane 0,010 0,01 2 3 3 12. hexachloroethane 0,010 0,01 2 3 3 13. 1,1-dichloroethane 0,010 0,01 2 3 3 13. 1,1-dichloroethane 0,010 0,01 2 3 3 14. 1,2,2-tetrachloroethane 0,010 0,01 2 3 3 14. 1,1,2-tric croethane 0,010 0,01 2 3 3 15. 1,1,2,2-tetrachloroethane 0,010 0,01 2 3 3 3 16. chloroethane 0,010 0,01 2 3 3 3 3 17. bis(2-chloroethyl) ether 0,010 0,01 2 3 <td< td=""><th></th><td></td><td>0.01</td><td>2</td><td>3 3</td><td></td><td></td><td>U</td></td<>			0.01	2	3 3			U
9. hexachlorobenzene 0.010 0.01 2 3 10. 1,2-dichloroethane 0.010 0.01 2 3 11. 1,1,1-trichloroethane 0.010 0.01 2 3 12. hexachloroethane 0.010 0.01 2 3 3 13. 1,1-dichloroethane 0.010 0.01 2 3 3 14. 1,1,2-tric roothane 0.010 0.01 2 3 3 15. 1,1,2,2-tetrachloroethane 0.010 0.01 2 3 3 3 16. chloroethane 0.010 0.01 2 3 3 3 3 17. bls(chloromethyl) ether 0.010 0.01 2 3 3 3 3 19. 2-chloroethyl vinyl ether 0.010 0.01 2 3 3 3 3 3 20. 2-chloroethyl vinyl ether 0.010 0.01 2 3 3 3 3 3 21. 2,4,6-trichlorophenol 0.010 0.01 2 3 3 3 3 3 3 3 <th></th> <td></td> <td>0.01</td> <td>2</td> <td>3 . 3</td> <td></td> <td></td> <td></td>			0.01	2	3 . 3			
10. 1,2-dichloroethane 0.00 0.01 2 3 11. 1,1,1-trithloroethane 0.010 0.01 2 3 3 12. hexachloroethane 0.010 0.01 2 3 3 13. 1,1-dichloroethane 0.010 0.01 2 3 3 14. 1,1,2-tric coethane 0.010 0.01 2 3 3 15. 1,1,2-tric coethane 0.010 0.01 2 3 3 16. chloroethane 0.010 0.01 2 3 3 3 17. bis(chloromethyl) ether 0.010 0.01 2 3 3 3 18. bis(2-chloroethyl) ether 0.010 0.01 2 3 3 3 3 20. 2-chloroaphthalene 0.010 0.01 2 3 3 3 3 3 21. 2.4.6-trichlorophenol 0.010 0.01 2 3 3 3 3 3 3 3 3 </td <th></th> <td></td> <td>0.01</td> <td>2</td> <td>3 3</td> <td></td> <td></td> <td>Z</td>			0.01	2	3 3			Z
11. 1.1.			0.01	2	3 0			· C
12. hexachloroethane 0.010 0.01 2 3 3 13. 1, 1-dichloroethane 0.010 0.01 2 3 3 14. 1, 1, 2-tric roethane 0.010 0.01 2 3 3 15. 1, 1, 2, 2-tetrachloroethane 0.010 0.01 2 3 3 16. chloroethane 0.010 0.01 2 3 3 16. chloroethane 0.010 0.01 2 3 3 17. bls (chloromethyl) ether 0.010 0.01 2 3 3 18. bls(2-chloroethyl vinyl ether 0.010 0.01 2 3 3 19. 2-chloronapthalarene 0.010 0.01 2 3 3 21. 2,4,6-trichlorophenol 0.010 0.01 2 3 3 22. parachlorometa cresol 0.010 0.01 2 3 3 23. chloroform 0.010 0.01 2 3 3 24. 2-chlorophenol 0.010 0.01 2 3 3 25. 1, 2-dichlorobenzene 0.010	10. 1,2-dichloroethane		0.01	2	3	*		2
13. 1,1-dichloroethane 0.010 0.01 2 3 3 14. 1,1,2-tric roethane 0.010 0.01 2 3 3 15. 1,1,2,2-terizachloroethane 0.010 0.01 2 3 3 16. chloroethane 0.010 0.01 2 3 3 17. bis(chloromethyl) ether 0.010 0.01 2 3 3 18. bis(2-chloroethyl) ether 0.010 0.01 2 3 3 19. 2-chloronaphthalene 0.010 0.01 2 3 3 20. 2-chloronaphthalene 0.010 0.01 2 3 3 21. 2,4,6-trichlorophenol 0.010 0.01 2 3 3 22. parachlorometa cresol 0.010 0.01 2 3 3 23. chloroform 0.010 0.01 2 3 3 24. 2-chlorophenol 0.010 0.01 2 3 3 25. 1, 2-dichlorobenzene 0.010 0.01 2 3 3 26. 1, 3-dichlorobenzene 0.010	11. 1,1,1-trichloroethane			2	3 3			- E
14. 1, 1, 2-tric roethane 0.010 0.01 2 3 3 15. 1, 1, 2, 2-terrachloroethane 0.010 0.01 2 3 3 16. chloroethane 0.010 0.01 2 3 3 17. bis(chloromethyl) ether 0.010 0.01 2 3 3 17. bis(chloromethyl) ether 0.010 0.01 2 3 3 18. bis(2-chloroethyl) unyl ether 0.010 0.01 2 3 3 19. 2-chloroethyl vinyl ether 0.010 0.01 2 3 3 20. 2-chloronaphthalene 0.010 0.01 2 3 3 21. 2, 4, 6-trichlorophenol 0.010 0.01 2 3 3 22. parachlorometa cresol 0.010 0.01 2 3 3 23. chloroform 0.010 0.01 2 3 3 24. 2-chlorophenol 0.010 0.01 2 3 3 25. 1,2-dichlorobenzene 0.010 0.01 2 3 3 26. 1,3-dichlorobenzene	12. nexachloroethane		0.01	2	3 5			۲ ۲
15. 1, 1, 2, 2-tetrachloroethane 0.010 0.01 2 3 3 16. chloroethane 0.010 0.01 2 3 3 17. bis (chloromethyl) ether 0.010 0.01 2 3 3 18. bis (2-chloroethyl) ether 0.010 0.01 2 3 3 19. 2-chloroethyl vinyl ether 0.010 0.01 2 3 3 20. 2-chloronaphthalene 0.010 0.01 2 3 3 21. 2, 4, 6-trichlorophenol 0.010 0.01 2 3 3 22. parachlorometa cresol 0.010 0.01 2 3 3 23. chloroform 0.010 0.01 2 3 3 24. 2-chlorophenol 0.010 0.01 2 3 3 25. 1,2-dichlorobenzene 0.010 0.01 2 3 3 26. 1,3-dichlorobenzene 0.010 0.01 2 3 3 26. 1,3-dichlorobenzidine 0.010 0.01 2 3 3 27. 1,4-dichlorobenzidine 0.010	13. 1,1-dichloroethane		0.01	2	3 3			
16. chloroethane 0.010 0.01 2 3 3 17. bis(chloromethyl) ether 0.010 0.01 2 3 3 18. bis(2-chloroethyl) ether 0.010 0.01 2 3 3 19. 2-chloroethyl vinyl ether 0.010 0.01 2 3 3 20. 2-chloroethyl vinyl ether 0.010 0.01 2 3 3 21. 2, 4, 6-trichlorophenol 0.010 0.01 2 3 3 22. parachlorometa cresol 0.010 0.01 2 3 3 23. chloroform 0.010 0.01 2 3 3 24. 2-chlorophenol 0.010 0.01 2 3 3 25. 1,2-dichlorobe.zene 0.010 0.01 2 3 3 26. 1,3-dichlorobenzene 0.010 0.01 2 3 3 27. 1,4-dichlorobenzene 0.010 0.01 2 3 3 27. 1,4-dichlorobenzidine 0.010 0.01 2 3 3 27. 1,4-dichlorobenzidine 0.010			0.01	2	3 3			··· 5
17. bis(chloromethyl) ether 0.010 0.01 2 3 3 18. bis(2-chloroethyl) ether 0.010 0.01 2 3 3 19. 2-chloroethyl vinyl ether 0.010 0.01 2 3 3 20. 2-chloronethyl vinyl ether 0.010 0.01 2 3 3 21. 2,4,6-trichlorophenol 0.010 0.01 2 3 3 22. parachlorometa cresol 0.010 0.01 2 3 3 23. chloroform 0.010 0.01 2 3 3 24. 2-chlorophenol 0.010 0.01 2 3 3 25. 1,2-dichlorobe.zene 0.010 0.01 2 3 3 26. 1,3-dichlorobenzene 0.010 0.01 2 3 3 26. 1,3-dichlorobenzene 0.010 0.01 2 3 3 27. 1,4-dichlorobenzene 0.010 0.01 2 3 3 27. 1,4-dichlorobenzidine 0.010 0.01 2 3 3 29. 1,1-dichlorophenol 0.010			0.01	2	3 3			K K
18.bis (2-chloroethyl) ether0.0100.012319.2-chloroethyl vinyl ether0.0100.0123320.2-chloronaphthalene0.0100.0123321.2. 4.6 -trichlorophenol0.0100.0123322.parachlorometa cresol0.0100.0123323.chloroform0.0100.0123323.chlorobe:::::::::::::::::::::::::::::::::::			0.01	2	3 3			
19.2-chloroethyl vinyl ether0.0100.0123320.2-chloronaphthalene0.0100.0123321.2,4,6-trichlorophenol0.0100.0123322.parachlorometa cresol0.0100.0123323.chloroform0.0100.0123324.2-chlorophenol0.0100.0123325.1,2-dichlorobenzene0.0100.0123326.1,3-dichlorobenzene0.0100.0123326.1,4-dichlorobenzene0.0100.0123326.1,3-dichlorobenzene0.0100.0123327.1,4-dichlorobenzidine0.0100.0123328.3,3'-dichloroethylene0.0100.0123329.1,1-dichloroethylene0.0100.0123331.2,4-dichloropenol0.0100.0123333.1,3-dichloropropane0.0100.0123333.1,3-dichloropropyletæ0.0100.01233	17. bis(chloromethyl) ether		0.01	2	3 3			<u> </u>
20. 2-chloronaphthalene0.0100.0123321. 2,4,6-trichlorophenol0.0100.0123322. parachlorometa cresol0.0100.0123323. chloroform0.0100.0123324. 2-chlorophenol0.0100.0123325. 1,2-dichlorobe: 0.010 0.0123326. 1,3-dichlorobenzene0.0100.0123327. 1,4-dichlorobenzidine0.0100.0123328. 3,3'-dichlorobenzidine0.0100.0123329. 1,1-dichloroethylene0.0100.0123330. 1,2-trans-dichlorophenol0.0100.0123331. 2,4'-dichlorophenol0.0100.0123333. 1,3-dichlorophylene0.0100.0123333. 1,3-dichlorophylene0.0100.0123333. 1,3-dichlorophylene0.0100.01233	18. Dis(2-chloroethyl) ether		0.01	2	3 3			
21. $2, 4, 6-trichlorophenol$ 0.0100.012322. parachlorometa cresol0.0100.012323. chloroform0.0100.0123324. 2-chlorophenol0.0100.0123325. 1, 2-dichlorobe: Dene0.0100.0123326. 1, 3-dichlorobe: Dene0.0100.0123327. 1, 4-dichlorobe: Dene0.0100.0123328. 3, 3'-dichlorobenzidine0.0100.0123329. 1, 1-dichlorobehylene0.0100.0123330. 1, 2-trans-dichlorophenol0.0100.0123331. 2, 4-dichlorophenol0.0100.0123333. 1, 3-dichlorophenol0.0100.0123333. 1, 3-dichloropropyle:se0.0100.01233	19. 2-chloroethyl vinyl ether		0.01	2	3 3			Č
22. parachlorometa cresol 0.010 0.01 2 3 3 23. chloroform 0.010 0.01 2 3 3 24. 2-chlorophenol 0.010 0.01 2 3 3 24. 2-chlorophenol 0.010 0.01 2 3 3 25. 1, 2-dichlorobe: Dene 0.010 0.01 2 3 3 26. 1, 3-dichlorobe: Dene 0.010 0.01 2 3 3 26. 1, 3-dichlorobenzene 0.010 0.01 2 3 3 27. 1, 4-dichlorobenzene 0.010 0.01 2 3 3 28. 3, 3'-dichlorobenzidine 0.010 0.01 2 3 3 29. 1, 1-dichloroethylene 0.010 0.01 2 3 3 30. 1, 2-trans-dichloroethylene 0.010 0.01 2 3 3 31. 2, 4-dichlorophenol 0.010 0.01 2 3 3 32. 1, 2-dichloropropane 0.010 0.01 2 3 3 33. 1, 3-dichloropropoyleise 0.010	20. 2-chloronaphthalene		0.01	2	3 3			. ♪
23. chloroform 0.010 0.01 2 3 3 24. 2-chlorophenol 0.010 0.01 2 3 3 25. 1,2-dichlorobe: 0.010 0.01 2 3 3 26. 1,3-dichlorobenzene 0.010 0.01 2 3 3 26. 1,3-dichlorobenzene 0.010 0.01 2 3 3 27. 1,4-dichlorobenzene 0.010 0.01 2 3 3 28. 3,3'-dichlorobenzidine 0.010 0.01 2 3 3 29. 1,1-dichloroethylene 0.010 0.01 2 3 3 30. 1,2-trans-dichloroethylene 0.010 0.01 2 3 3 31. 2,4-dichlorophenol 0.010 0.01 2 3 3 32. 1,2-dichloropropane 0.010 0.01 2 3 3 33. 1,3-dichloropropale 0.010 0.01 2 3 3 33. 1,3-dichloropropylexe 0.010 0.01 2 3 3			0.01	2	3 3			
24. 2-chlorophenol 0.010 0.01 2 3 3 25. 1,2-dichlorobe: mene 0.010 0.01 2 3 3 26. 1,3-dichlorobenzene 0.010 0.01 2 3 3 27. 1,4-dichlorobenzene 0.010 0.01 2 3 3 28. 3,3'-dichlorobenzidine 0.010 0.01 2 3 3 29. 1,1-dichloroethylene 0.010 0.01 2 3 3 30. 1,2-trans-dichloroethylene 0.010 0.01 2 3 3 31. 2,4-dichlorophenol 0.010 0.01 2 3 3 32. 1,2-dichlorophenol 0.010 0.01 2 3 3 33. 1,3-dichloropropane 0.010 0.01 2 3 3 33. 1,3-dichloropropylexe 0.010 0.01 2 3 3		0.010	0.01	2	ă ă			- G
25. 1,2-dichlorobe: Bene 0.010 0.01 2 3 3 26. 1,3-dichlorobenzene 0.010 0.01 2 3 3 27. 1,4-dichlorobenzene 0.010 0.01 2 3 3 28. 3,3'-dichlorobenzidine 0.010 0.01 2 3 3 29. 1,1-dichloroethylene 0.010 0.01 2 3 3 30. 1,2-trans-dichloroethylene 0.010 0.01 2 3 3 31. 2,4-dichlorophenol 0.010 0.01 2 3 3 32. 1,2-dichlorophenol 0.010 0.01 2 3 3 33. 1,3-dichloropropane 0.010 0.01 2 3 3 33. 1,3-dichloropropylexe 0.010 0.01 2 3 3	23. chloroform		0.01	2	3 3			Č
26. 1, 3-dichlorobenzene 0.010 0.01 2 3 3 $27. 1, 4-dichlorobenzene0.0100.0123328. 3, 3'-dichlorobenzidine0.0100.0123329. 1, 1-dichloroethylene0.0100.0123330. 1, 2-trans-dichloroethylene0.0100.0123331. 2, 44ichlorophenol0.0100.0123332. 1, 2-dichloropropane0.0100.0123333. 1, 3-dichloropropylexe0.0100.01233$	24. 2-chlorophenol		0.01	$\overline{2}$	ă ă		· ·	<u>.</u>
27. 1,4-dichlorobenzene 0.010 0.01 2 3 3 28. 3,3'-dichlorobenzidine 0.010 0.01 2 3 3 29. 1,1-dichloroethylene 0.010 0.01 2 3 3 30. 1,2-trans-dichloroethylene 0.010 0.01 2 3 3 31. 2,4-dichlorophenol 0.010 0.01 2 3 3 32. 1,2-dichloropropane 0.010 0.01 2 3 3 33. 1,3-dichloropropylexe 0.010 0.01 2 3 3			0.01	2	3 3			· •
28. 3,3'-dichlorobenzidine 0.010 0.01 2 3 3 29. 1,1-dichloroethylene 0.010 0.01 2 3 3 30. 1,2-trans-dichloroethylene 0.010 0.01 2 3 3 31. 2,4-dichlorophenol 0.010 0.01 2 3 3 32. 1,2-dichloropropane 0.010 0.01 2 3 3 33. 1,3-dichloropropylexe 0.010 0.01 2 3 3			0.01	2 .	.3 3	•		
29. 1,1-dichloroethylene 0.010 0.01 2 3 3 30. 1,2-trans-dichloroethylene 0.010 0.01 2 3 3 31. 2,4-dichlorophenol 0.010 0.01 2 3 3 32. 1,2-dichloropropane 0.010 0.01 2 3 3 33. 1,3-dichloropropylexe 0.010 0.01 2 3 3	27. 1,4-dichlorobenzene		0.01	2	3 3			•
30. 1,2-trans-dichloroethylene 0.010 0.01 2 3 3 31. 2,4 dichlorophenol 0.010 0.01 2 3 3 32. 1,2-dichloropropane 0.010 0.01 2 3 3 33. 1,3-dichloropropylese 0.010 0.01 2 3 3		0.010	0.01	2 .	ă ă			70
31. 2,4 dichlorophenol 0.010 0.01 2 3 3 32. 1,2-dichloropropane 0.010 0.01 2 3 3 33. 1,3-dichloropropylese 0.010 0.01 2 3 3	29. I, I-dichloroethylene		0.01	2	3 3		н. Н	
32. 1,2-dichloropropane 0.010 0.01 2 3 33. 1,3-dichloropropylexe 0.010 0.01 2 3 3	30. 1,2-trans-dichloroethylene	0.010	0.01	2	3 3			Ŭ.
33. 1,3-dichloropropylese 0.010 0.01 2 3 3	31. 2,4 dichlorophenol		0.01	2	3 3			Ĥ
	32. 1,2-dichloropropane	0.010	0.01	$\overline{2}$	3 3	•		
34. 2,4-dimethylphenol 0.010 0.01 2 3 3	33. 1,3-dichloropropylesie			2	3 3			
	34. 2,4-dimethylphenol	0.010	0.01	2.	3 3		-	<

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PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

SECT - VI

Table VI-1 (Continued)

FREQUENCY OF OCCURRENCE OF PRIORITY POLLUTANTS PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY RAW WASTEWATER

FREQUENCY OF OCCURRENCE OF PRIORITY POLLUTANTS PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY RAW WASTEWATER									
Pollutant	Analytical Quantification Concentration (mg/l)(a)	Treatable Concentra- tion (mg/1)(b)	Number of Streams Analyzed	Number of Samples Analyzed	<u>ND</u>	Detected Below Quantification Concentration	Detected Below Treat- able Concen- tration	Detected H Above Treat able Concent tration	
35. 2,4-dinitrotoluene	0.010	0.01	2	3	3			METALS	
36. 2,6-dinitrotoluene	0.010	0.01	2	· 3 .	3			E	
37. 1.2-diphenylhydrazine	D.010	0.01	2	3	3			AI	
38. ethylbenzene	0.010	0.01	2 .	3	3	,			
39. fluoranthene	0.010	0.01	2	3	3	· ·			
40. 4-chlorophenyl phenyl ether	0.010	0.01	2	3	3			AND	
41. 4-bromophenyl phenyl ether	0.010	0.01	2	3	3		•	Ä	
42. bis(2-chloroisopropyl) ether	0.010	0.01	2	3	3				
43. bis(2-chloroethoxy) methane	0.010	0.01	2	3	3			MERCURY	
44. methylene chloride	0.010	0.01	2	3				3 🖽	
45. methyl chloride	0.010	0.01	. 2	3	3			õ.	
46. methyl bromide	0.010	0.01	2	3	3			ğ	
47. bromoform	0.010	0.01	2	3	3 3			R	
48. dichlorobromomethane	0.010	0.01	· 2		3			қ	
49. trichlorofluoromethane	0.010	0.01	2	3				ល	
50. dichlorodifluoromethane	0.010	0.01	2	3	3			ġ	
51. chlorodibromomethane	0.010	0.01	2	3 3	3 3			· BO	
52. hexachlorobutadiene	0.010	0.01	2	-	3			A	
 becachlorocyclopentadiene 	0.010	0.01	2	3 3	3	×		Ĥ	
54. isophorone	0.010	0.01	2	3	3			ы Ц	
55. naphthalene	0.010	0.01	2	3	2			ц С	
56. nitrobenzene	0.010	0.01	2	3	3			SUBCATEGORY	
57. 2-nitrophenol	0.010	0.01	2	3	3			Ř	
58. 4-nitrophenol	0.010	0.01	2	3	3				
59. 2,4-dinitrophenol	0.010	0.01	2	3	3				
60. 4,6-dinitro-o-cresol	0.010	0.01	2	3	3	•			
61. N-nitrosodimethylamine	0.010	0.01	2	3	3			ល	
62. N-nitrosodiphenylamine	0.010	0.01	2	3	3			SECT	
63. N-nitrosodi-n-propylamine	0.010	0.01 0.01	2	3	3		•	H H	
64. pentachlorophenol	0.010	0.01		3	2	1			
65. phenol	0.010	0.01	2 2	3	4	3		I	
66. bis(2-ethylhexyl) phthalate	0.010			3	3	, J		· /	
67. butyl benzyl phthalate	0.010	0.01	2		1	2		I V	
68. di-n-butyl phthalate	0.010	0.01	Ζ.	J	•	د ا			

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Table VI-1 (Continued)

FREQUENCY OF OCCURRENCE OF PRIORITY POLLUTANTS PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY RAW WASTEWATER

Pollutant	Analytical Quantification Concentration (mg/l)(a)	Treatable Concentra- tion (mg/1)(b)	Number of Streams Analyzed	Number of Samples Analyzed	ND	Detected Below Quantification Concentration	Detected Below Treat- able Concen- tration	Detected Above Treat- able Concen- tration	RECIOUS
69. di-n-octyl phthalate	0.010	0.01	2	3	3	5	·.		R
70. diethyl phthalate	0.010	0.01	2	3		2		1	METAL
71. dimethyl phthalate	0.010	0.01	2	3	3			•	Ψ
72. benzo(a)anthracene	0.010	0.01	. 2	3	- <u>3</u>	<i>a</i> .			R
73. benzo(a)pyrene	0.010	0.01	2	3	3				้ง
74. 3,4-benzofluoranthene	0.010	0.01	. 2	3	- Š		· · · ·		,
75. benzo(k)fluoranthene	0.010	0.01	2	3	3				AND
76. chrysene	0.010	0.01	2	3	Ĩ	•			A
77. acenaphthylene	0.010	0.01	2	3	3				0
78. anthracene (c)	0.010	0.01	2	3	•	3			М
79. benzo(ghi)perylene	0.010	0.01	· 2	3	3		,		E
80. fluorene	0.010	0.01	2	3	3				õ
81. phenanthrene (c)	0.010	0.01	2	´ 3		3			MERCURY
82. dibenzo(a,h)anthracene	0.010	0.01	2	3	3	, •			77
83. indeno(1,2,3-cd)pyrene	0.010	0.01	2	3	3				R
84. pyrene	0.010	0.01	2	3	3				۲ ۵
85. tetrachloroethylene	0.010	0.01	2	3	3	· ·			ä
86. toluene	0.010	0.01	2	3		· 1		2	μ
87. trichloroethylene	0.010	0.01	2	3	3	·		-	Ω
88. vinyl chloride	0.010	0.01	2	3	3	4	•	~	Ц
114. antimony	0.100	0.47	8	10		10	0	0 -	SUBCATEGORY
115. arsenic	0.010	0.34	8	10		0 .		ž	ଜୁ
117. beryllium	0.010	0.20	8	10		9	1	0	H.
118. cadmium	0.002	0.049	8	10		6	2	2	R
119. chromium	0.005	0.07	8	10		.5	3	2	
120. copper	0.009	0.39	8	10		5	3	2	
121. cyanide (d)	0.02	0.047	2	4		0	1	3	
122. lead	0.020	0.08	8,	10		4	Ó	· 6	S
123. mercury	0.0001	0.036	8 .	10		0	4	6	内
124. nickel	0.005	0.22	8	10		6	3	ĩ	G
125. selenium	0.01	0.20	8	10		8	2	0	-
126. silver	0.02	0.07	8	10	*	5	3	2	1
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				•			•		F , T

Table VI-1 (Continued)

FREQUENCY OF OCCURRENCE OF PRIORITY POLLUTANTS PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY RAW WASTEWATER

Pollutant	Analytical Quantification Concentration (mg/l)(a)	Treatable Concentra- tion (mg/l)(b)	Number of Streams Analyzed	Number of Samples Analyzed	ND	Detected Below Quantification Concentration	Detected Below Treat- able Concen- tration	Detected Above Treat- able Concen- tration
127. thallium	0.100	0. 34	8 [.]	10		6	2	2
128. zinc	0.050	0. 23	ຮ	10		4	3	3
oil and grease	5.0	10.0	8	10		8	υ`	2
total suspended solids (TSS)	1.0	2.6	8	10		2	Ο	ರ

(a) Analytical quantification concentration was reported with the data (see Section V).

(b) Treatable concentrations are based on performance of chemical precipitation, sedimentation, and filtration.

(c) Reported together.

(d) Analytical quantification concentration for EPA Method 335.2, Total Cyanide Methods for Chemical Analysis of Water and Wastes, EPA 600/4-79-020, March 1979.

TABLE VI-2

TOXIC POLLUTANTS NEVER DETECTED

1.	acenaphthene
2.	acrolein
3.	acrylonitrile
5.	benzidine
6.	carbon tetrachloride
7.	chlorobenzene
8.	1,2,4-trichlorobenzene
9.	hexachlorobenzene
10.	1,2-dichloroethane
11.	1,1,1-trichloroethane
12.	hexachloroethane
13.	1,1-dichloroethane
14.	1,1,2-trichloroethane
15.	1,1,2,2-tetrachloroethane
16.	chloroethane
17.	bis(2-chloromethyl) ether (deleted)
18.	bis(2-chloroethyl) ether
19.	2-chloroethyl vinyl ether
20.	2-chloronaphthalene
21.	2,4,6-trichlorophenol
22.	parachlorometa cresol
23.	chloroform
24.	2-chlorophenol
25.	1,2-dichlorobenzene
26.	1,3-dichlorobenzene
27. 28.	1,4-dichlorobenzene
29.	3,3 -dichlorobenzidine
30.	1,1-dichloroethylene
31.	1,2-trans-dichloroethylene 2,4-dichlorophenol
32.	1,2-dichloropropane
33.	1,3-dichloropropylene
34.	2,4-dimethylphenol
35.	2,4-dinitrotoluene
36.	2,6-dinitrotoluene
37.	1,2-diphenylhydrazine
38.	ethylbenzene
39.	fluoranthene
40.	4-chlorophenyl phenyl ether
41.	4-bromophenyl phenyl ether
42.	bis(2-chloroisopropyl)ether
43.	bis(2-chloroethoxy)methane
45.	methyl chloride (chloromethane)
46.	methyl bromide (bromomethane)
47.	bromoform
48.	dichlorobromomethane
49.	trichlorofluoromethane (deleted)

TABLE VI-2 (Continued)

TOXIC POLLUTANTS NEVER DETECTED

- dichlorodifluoromethane (deleted) 50.
- chlorodibromomethane 51.
- hexachlorobutadiene 52.
- hexachlorocyclopentadiene 53.
- 54. isophorone
- naphthalene 55.
- nitrobenzene 56.
- 2-nitrophenol 57.
- 58. 4-nitrophenol
- 59. 2,4-dinitrophenol
- 4,6-dinitro-o-cresol 60.
- N-nitrosodimethylamine 61.
- 62. N-nitrosodiphenylamine
- N-nitrosodi-n-propylamine 63.
- pentachlorophenol 64.
- butyl benzyl phthalate 67.
- 69. di-n-octyl phthalate
- 71. dimethyl phthalate
- 72. benzo(a)anthracene
- 73. benzo(a)pyrene
- 3,4-benzofluoranthene 74.
- benzo(k)fluoranthene 75.
- 76. chrysene
- 77. acenaphthylene
- 79. benzo(ghi)perylene
- fluorene 80.
- dibenzo(a,h)anthracene 82.
- indeno (1,2,3-cd)pyrene 83.
- 84. pyrene
- tetrachloroethylene 85.
- trichloroethylene 87.
- vinyl chloride 88.
- 89. aldrin
- 90. dieldrin
- 91. chlordane
- 92. 4,4'-DDT
- 4,4'-DDE 93.
- 94. 4,4'-DDD
- alpha-endosulfan 95. 96. beta-endosulfan
- 97. endosulfan sulfate
- 98. endrin
- endrin aldehyde 99.
- 100. heptachlor

TABLE VI-2 (Continued)

TOXIC POLLUTANTS NEVER DETECTED

101.	heptachlor	epoxide				
	alpha-BHC	-				
103.	beta-BHC					
104.	gamma-BHC					
105.	delta-BHC					
106.	PCB.1242	(a)	k.			
107.	PCB-1254	(a)				
108.	PCB-1221	(a)				
109.	PCB-1232	(b)				
110.	PCB-1248	(b)				
111.	PCB-1260	(b)	•		*	
112.	PCB-1016	(b)		ι.		
113.	toxaphene					
116.	asbestos					
129.	2,3,7,8-tet	rachloro	libenzo	-p-dioxi	n (TCDD)	
(a),	(b) Reporte	ed togethe	er, as a	a combin	ed value	

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SECTION VII

CONTROL AND TREATMENT TECHNOLOGIES

The preceding sections of this supplement discussed the sources, flows, and characteristics of the wastewaters from primary precious metals and mercury plants. This section summarizes the description of these wastewaters and indicates the treatment technologies which are currently practiced in the primary precious metals and mercury subcategory for each waste stream. section presents the control and treatment Secondly, this technology options which were examined by the Agency for possible the primary precious metals and application to mercurv subcategory.

CURRENT CONTROL AND TREATMENT PRACTICES

This section presents a summary of the control and treatment technologies that are currently being applied to each of the sources generating wastewater in this subcategory. As discussed in Section V, wastewater associated with the primary precious metals and mercury subcategory is characterized by the presence of the toxic metal pollutants, suspended solids, and oil and grease. This analysis is supported by the raw (untreated) wastewater data presented for specific sources. Construction of one wastewater treatment system for combined treatment allows plants to take advantage of economic scale and in some instances to combine streams of different alkalinity to reduce treatment chemical requirements.

All but one of the plants within this subcategory do not discharge wastewater. The one discharging facility discharges to a surface water from a tailings pond. Zero discharge is achieved in most plants through a combination treatment consisting of a tailings pond and recycle or reuse. One of the three plants with a smelter scrubber achieves zero discharge of that waste stream by 100 percent recycle. Partial recycle is used only on two waste streams, the smelter scrubber and the calciner scrubber wastewater. Table VII-1 (page 2227) presents a summary of the number of plants with each wastewater stream and the treatment technologies currently in place.

CONTROL AND TREATMENT OPTIONS

The Agency examined three control and treatment technology options that are applicable to the primary precious metals and mercury subcategory. The options selected for evaluation represent a combination of in-process flow reduction, preliminary treatment technologies applicable to individual waste streams, and end-of-pipe treatment technologies. The effectiveness of these technologies is discussed in Section VII of Vol. I. OPTION A

Option A for the primary precious metals and mercury subcategory requires control and treatment technologies to reduce the discharge of wastewater pollutant mass.

The Option A treatment scheme consists of chemical precipitation and sedimentation technology and ion exchange as a polishing step. Specifically, lime or some other alkaline compound is used to precipitate metal ions as metal hydroxides. The metal hydroxides and suspended solids settle out and the sludge is collected. Vacuum filtration is used to dewater sludge.

Preliminary treatment consisting of oil skimming to remove oil and grease is also included in Option A.

OPTION B

Option B for the primary precious metals and mercury subcategory consists of the Option A (oil skimming, chemical precipitation and sedimentation, ion exchange) treatment scheme plus flow reduction techniques to reduce the discharge of wastewater volume. In-process changes which allow for recycle of electrolytic cells wastewater and calciner scrubber water are the principal control mechanisms for flow reduction.

OPTION C

Option C for the primary precious metals and mercury subcategory consists of all control and treatment requirements of Option B (in-process flow reduction, oil skimming, chemical precipitation and sedimentation, ion exchange) plus multimedia filtration technology added at the end of the Option B treatment scheme. Multimedia filtration is used to remove suspended solids, including precipitates of metals, attainable by gravity sedimentation. beyond the concentration The filter suggested is of the gravity, mixed-media type, although other forms of filters, such as rapid sand filters or pressure filters would perform satisfactorily. The addition of filters also provides consistent removal during periods in which there are rapid increases in flows or loadings of pollutants to the treatment system.

Table VII-1

SUMMARY OF WASTE STREAMS AND TREATMENT PRACTICES IN PRIMARY PRECIOUS METALS AND MERCURY PLANTS

<u>Waste Stream</u>	Number of Plants With Waste Stream	Number of Plants With Tailings Pond Treatment	Number of Plants With <u>Recycle or Reuse</u>
Smelter wet air pollution control	3	2	3
Silver chloride reduction spent solution	2	2	1
Electrolytic cells wet air pollution control	• 1 .	1	
Electrolyte preparation wet air pollu- tion control	1	1	0
Silver crystals wash water	1	1	0
Gold slimes acid wash and water rinse	1	1	0
Calciner wet air pollution control	1	1	1
Calcine quench water	1	1	1
Calciner stack gas contact cooling water	1	1	1
Condenser blowdown	1	1	1
Mercury cleaning bath water	1	1	1

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SECTION VIII

COSTS, ENERGY, AND NONWATER QUALITY ASPECTS

This section presents a summary of compliance costs for the primary precious metals and mercury subcategory and a description the treatment options and subcategory-specific assumptions of Together with the estimated used to develop these estimates. pollutant reduction performance presented in Sections IX, X, XI, and XII of this supplement, these cost estimates provide a basis for evaluating each regulatory option. These cost estimates are also used in determining the probable economic impact of regulation on the subcategory at different pollutant discharge In addition, this section addresses nonwater quality levels. impacts of wastewater treatment and control environmental alternatives, including air pollution, solid wastes, and energy requirements, which are specific to the primary precious metals and mercury subcategory.

TREATMENT OPTIONS FOR EXISTING SOURCES

As discussed in Section VII, three treatment options have been developed for existing primary precious metals and mercury sources. The options are summarized below and schematically presented in Figures X-1 through X-3 (page 2264 - 2266).

OPTION A

Option A consists of preliminary treatment using oil-water separation where required and chemical precipitation and sedimentation and ion exchange end-of-pipe technology.

OPTION B

Option B consists of in-process flow reduction and oil-water separation preliminary treatment where required, and end-of-pipe technology consisting of chemical precipitation and sedimentation and ion exchange. The in-process flow reduction measure consists of the recycle of electrolytic cells scrubber water, and calciner scrubber water through holding tanks.

OPTION C

Option C requires the in-process flow reduction and oil-water separation preliminary treatment measures of Option B, and endof-pipe treatment technology consisting of chemical precipitation, sedimentation, ion exchange and multimedia filtration.

COST METHODOLOGY

A detailed discussion of the methodology used to develop the compliance costs is presented in Section VIII of Vol. I. Plant-

by-plant compliance costs for the nonferrous metals manufacturing category have been revised as necessary following proposal. These revisions calculate incremental costs, above treatment already in place, necessary to comply with the promulgated effluent limitations and standards and are presented in the administrative record supporting this regulation. A comparison of the costs developed for proposal and the revised costs for the final regulation are presented in Table VIII-1 (page 2230) for the direct discharger in this subcategory.

Each of the general assumptions used to develop compliance costs is presented in Section VIII of Vol. I. No subcategory-specific assumptions were used in developing compliance costs for the primary precious metals and mercury subcategory.

NONWATER QUALITY ASPECT

A general discussion of the nonwater quality aspects of the control and treatment options considered for the nonferrous metals category is contained in Section VIII of Vol. I. Nonwater quality impacts specific to the primary precious metals and mercury subcategory, including energy requirements, solid waste and air pollution, are discussed below.

ENERGY REQUIREMENTS

The methodology used for determining the energy requirements for the various options is discussed in Section VIII of the General Development Document. Energy requirements for the three options considered are estimated at 10,900 kwh/yr, 10,900 kwh/yr, and 11,200 kwh/yr for Options A, B, and C, respectively. Option B energy requirements are the same as those for Option A because the one discharging plant has no flow reduction. Option C, which includes filtration, increases energy consumption over Option B by approximately three percent. Option C represents roughly 3.5 percent of a typical plant's electrical energy usage. It is therefore concluded that the energy requirements of the treatment options considered will not have significant impact on total plant energy consumption.

SOLID WASTE

Sludge generated in the primary precious metals and mercury subcategory is due to oily wastes from oil-water separation and the precipitation of metal hydroxides and carbonates using lime.

Sludges associated with the primary precious metals and mercury subcategory will necessarily contain quantities of toxic metal pollutants. These sludges are not subject to regulation as hazardous wastes since wastes generated by primary smelters and refiners are currently exempt from regulation by Act of Congress (Resource Conservation and Recovery Act (RCRA), Section 3001(b)), as interpreted by EPA. If a small (5-10%) excess of lime is added during treatment, the Agency does not believe these sludges would be identified as hazardous under RCRA in any case.

(Compliance costs include this amount of lime.) This judgment is based on the results of Extraction Procedure (EP) toxicity tests performed on similar sludges (toxic metal-bearing sludges) generated by other industries such as the iron and steel industry. A small amount of excess lime was added during treatment, and the sludges subsequently generated passed the toxicity test. See CFR \$261.24. Thus. the Agency believes that the wastewater sludges will similarly not be EP toxic if the recommended technology is applied.

Although it is the Agency's view that solid wastes generated as a result of these guidelines are not expected to be hazardous, generators of these wastes must test the waste to determine if the wastes meet any of the characteristics of hazardous waste (see 40 CFR 262.11).

If these wastes should be identified or are listed as hazardous, they will come within the scope of RCRA's "cradle to grave" hazardous waste management program, requiring regulation, from the point of generation to point of final disposition. EPA's standards would require generators of hazardous gener-ator nonferrous metals manufacturing wastes to meet containerization, labeling, recordkeeping, and reporting requirements; if plants dispose of hazardous wastes off-site, they would have to prepare a manifest which would track the movement of the wastes from the generator's premises to a permitted off-site treatment, storage, or disposal facility. See 40 CFR 262.20 45 FR 33142 (May 19, 1980), as amended at 45 FR 86973 (December 31, 1980). The transporter regulations require transporters of hazardous wastes to comply with the manifest system to assure that the wastes are delivered to a permitted facility. See 40 CFR 263.20 45 FR 33151 (May 19, 1980), as amended at 45 FR 86973 (December 31, 1980). Finally costs for wastewater treatment the cost of hauling and disposing of these wastes. For more details, see Section VIII of the General Development Document.

Sludge generation for BPT of the primary precious metals and mercury subcategory is estimated at 208 metric tons per year. Sludge generation for BAT is not expected to be significantly different.

AIR POLLUTION

There is no reason to believe that any substantial air pollution problems will result from implementation of oil-water separation, chemical precipitation, sedimentation, multimedia filtration and ion exchange. These technologies transfer pollutants to solid waste and are not likely to transfer pollutants to air.

Table VIII-1

COST OF COMPLIANCE FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY DIRECT DISCHARGERS

(March, 1982 Dollars)

Option	Proposa Capital Cost	l Costs Annual Cost	Promulgat Capital Cost	ion Costs Annual Cost
A	27,500	9,000	2,200	26,800
В	27,500	9,000	2,200	26,800
С	30,000	10,000	3,025	27,300

SECTION IX

BEST PRACTICABLE CONTROL TECHNOLOGY CURRENTLY AVAILABLE

This section defines the effluent characteristics attainable through the application of best practicable control technology currently available (BPT). BPT reflects the existing performance by plants of various sizes, ages, and manufacturing processes within the primary precious metals and mercury subcategory, as well as the established performance of the recommended BPT Particular consideration is given to the treatment systems. already in place at plants within the data base.

The factors considered in identifying BPT include the total cost applying the technology in relation to the effluent reduction of benefits from such application, the age of equipment and facilities involved, the manufacturing processes employed, quality environmental impacts (including energy nonwater requirements), and other factors the Administrator considers appropriate. In general, the BPT level represents the average of the existing performances of plants of various ages, sizes, processes, or other common characteristics. Where existing performance is uniformly inadequate, BPT may be transferred from different subcategory or category. Limitations based on a transfer of technology are supported by a rationale concluding that the technology is, indeed, transferable, and a reasonable prediction that it will be capable of achieving the prescribed effluent limits (see Tanner's Council of America v. Train, 540 F.2d 1188 (4th Cir. 11/6). BPT focuses on end-of-pipe treatment rather than process changes or internal controls, except where such practices are common within the subcategory.

TECHNICAL APPROACH TO BPT

The Agency studied the primary precious metals and mercury subcategory to identify the processes used, the wastewaters generated. and the treatment processes installed. Information was collected from the category using data collection portfolios, and specific plants were sampled and the wastewaters analyzed. In making technical assessments of data, reviewing manufacturing processes, and assessing wastewater treatment technology options, indirect and direct dischargers have been considered as a single group.

As explained in Section IV, the primary precious metals and mercury subcategory has been subdivided into nine potential wastewater sources. Since the water use, discharge rates, and characteristics of each of these wastewaters is pollutant potentially unique, effluent limitations will be developed for each of the nine subdivisions or segments.

For each of the subdivisions, a specific approach was followed for the development of BPT mass limitations. The first first requirement to develop these limitations is to account for production and flow variability from plant to plant. Therefore, a unit of production or production normalizing parameter (PNP) was determined for each waste stream which could then be related to the flow from the process to determine a production normalized flow. Selection of the PNP for each process element is discussed in Section IV. Each process within the subcategory was then analyzed to determine (1) which subdivisions were present, (2) the specific flow rates generated for each subdivision, and (3) the specific production normalized flows for each subdivision. This analysis is discussed in detail in Section V. Nonprocess wastewaters such as rainfall runoff and noncontact cooling water are not considered in the analysis.

Production normalized flows for each subdivision were then analyzed to determine the flow to be used as part of the basis for BPT mass limitations. The selected flow (sometimes referred to as a BPT regulatory flow or BPT discharge rate) reflects the water use controls which are common practices within the category. The BPT regulatory flow is based on the average of all applicable data. Plants with normalized flows above the average may have to implement some method of flow reduction to achieve the BPT limitations.

The second requirement to calculate mass limitations is the set of concentrations that are achievable by application of the BPT level of treatment technology. Section VII discusses the various control and treatment technologies which are currently in place for each wastewater source. In most cases, the current control and treatment technologies consist of a combination of tailings ponds and reuse and recycle of process water. Chemical precipitation and sedimentation technology and performance is transferred to this subcategory, because current treatment is inadequate. Oil skimming is applied to streams with treatable concentrations of oil and grease. Ion exchange technology is being added for the removal of gold.

Using these regulatory flows and the achievable concentrations, the next step is to calculate mass loadings for each wastewater source or subdivision. This calculation was made on a stream-bystream basis, primarily because plants in this subcategory may perform one or more of the operations in various combinations.

The mass loadings (milligrams of pollutant per troy ounce or metric ton of production - mg/T.O. or mg/kkg) were calculated by multiplying the BPT regulatory flow (1/T.O. or 1/kkg) by the concentration achievable by the BPT level of treatment technology (mg/l) for each pollutant parameter to be limited under BPT. These mass loadings are published in the Federal Register and in CFR Part 421 as the effluent limitations guidelines.

The mass loadings which are allowed under BPT for each plant will be the sum of the individual mass loadings for the various

wastewater sources which are found at particular plants. Accordingly, all the wastewater generated within a plant may be combined for treatment in a single or common treatment system, but the effluent limitations for these combined wastewaters are based on the various wastewater sources which actually contribute to the combined flow. This method accounts for the variety of combinations of wastewater sources and production processes which may be found at primary precious metals and mercury plants.

The Agency usually establishes wastewater limitations in terms of mass rather than concentration. This approach prevents the use of dilution as a treatment method (except for controlling pH). The production normalized wastewater flow (1/T.0. or 1/kkg) is a link between the production operations and the effluent limitations. The pollutant discharge attributable to each operation can be calculated from the normalized flow and effluent concentration achievable by the treatment technology and summed to derive an appropriate limitation for each plant.

INDUSTRY COST AND POLLUTANT REMOVAL ESTIMATES

In balancing costs in relation to pollutant removal estimates, EPA considers the volume and nature of existing discharges, the volume and nature of discharges expected after application of BPT, the general environmental effects of the pollutants, and the cost and economic impacts of the required pollution control level. The Act does not require or permit consideration of water quality problems attributable to particular point sources or industries, or water quality improvements in particular water quality bodies. Accordingly, water quality considerations were not the basis for selecting the proposed or promulgated BPT. See Weyerhaeuser Company v. Costle, 590 F.2d 1011 (D.C. Cir. 1978).

The methodology for calculating pollutant removal estimates and plant compliance costs is discussed in Section X. The pollutant removal estimates have been revised since proposal based on new flow and production data submitted to EPA through industry comments. Table X-2 (page 2256) shows the estimated pollutant removals for each treatment option for direct dischargers. Compliance costs are presented in Table X-3 (page 2257).

BPT OPTION SELECTION

The technology basis for the promulgated BPT limitations is Option A, chemical precipitation and sedimentation technology to remove metals and solids from combined wastewaters and to control pH, ion exchange as a polishing step to remove gold, and oil skimming to remove oil and grease. This technology is in-place at the discharger in this subcategory. This technology differs from proposed BPT by the addition of ion exchange. The pollutants specifically promulgated for regulation at BPT are arsenic, lead, mercury, silver, zinc, oil and grease, TSS, and pH. Implementation of the promulgated BPT limitations will remove annually an estimated 50,442 kg of priority metals and 3,310 kg of TSS. We project a capital cost of \$2,200 and an annualized cost of \$26,800 (1982 dollars) for achieving promulgated BPT limitations.

More stringent technology options were not selected for promulgated BPT since they require in-process changes or end-ofpipe technologies less widely practiced in the subcategory, and, therefore, are more appropriately considered under BAT.

WASTEWATER DISCHARGE RATES

A BPT discharge rate is calculated for each subdivision based on the average of the flows of the existing plants, as determined from analysis of the dcp. The discharge rate is used with the achievable treatment concentration to determine BPT effluent limitations. Since the discharge rate may be different for each wastewater source, separate production normalized discharge rates for each of the nine wastewater sources are discussed below and summarized in Table IX-1 (page 2246). The discharge rates are normalized on a production basis by relating the amount of wastewater generated to the mass of the intermediate product which is produced by the process associated with the waste stream, in question. These production normalizing parameters are also listed in Table IX-1.

Section V of this supplement further describes the discharge flow rates and presents the water use and discharge flow rates for each plant by subdivision. The proposed and promulgated BPT discharge rates are discussed individually below.

SMELTER WET AIR POLLUTION CONTROL

The BPT wastewater discharge rate proposed for smelter wet air pollution control was 13.2 liters per troy ounce (3.5 gal/troy ounce) to gold and silver smelted, based on zero percent recycle. This rate was allocated only for plants practicing wet air pollution control for the smelter. Three plants reported this waste stream, as shown in Table V-1 (page 2189). The BPT rate was based on the average water use rate for these three plants (25.8, 8.4, and 5.3 liters per troy ounce).

At proposal, EPA was considering a BPT wastewater discharge rate for this waste stream of 1.3 liters per troy ounce, based on 90 percent recycle. Recycle is demonstrated for this waste stream; three plants reporting a smelter scrubber indicated recycle rates of 76 to 100 percent on scrubber liquor. For this reason, EPA considered reducing the discharge allowance for this stream and solicited comments from industry.

The promulgated BPT wastewater discharge rate for smelter wet air pollution control is 1.3 liters per troy ounce (0.343 gal/T.O.) of gold and silver smelted. Since the Agency received no comments on this issue from industry, it decided to incorporate

the 90 percent recycle rate for this waste stream. Consequently, the promulgated BPT flow is based on 90 percent recycle of the average water use reported by the three plants with this waste stream.

SILVER CHLORIDE REDUCTION SPENT SOLUTION

The proposed and promulgated BPT wastewater discharge rate for silver chloride reduction spent solution is 0.4 liters per troy ounce (0.11 gal/troy ounce) of silver reduced in solution. Water use and discharge rates are presented in Table V-2 (page 2189). The proposed BPT discharge rate was based on the flow reported by one plant. Since proposal, EPA received comments from a second plant reporting this waste stream. Flow and production data for this plant were not quantified precisely enough co calculate a production normalized flow. Thus, the promulgated BPT flow rate is based on the values reported by the initial plant and is equal to the proposed flow rate.

ELECTROLYTIC CELLS WET AIR POLLUTION CONTROL

The proposed and promulgated BPT wastewater discharge rate for the electrolytic cells wet air pollution control is 198 liters per troy ounce (52.3 gal/T.O.) of gold refined electrolytically. This normalized flow is based upon the only value reported for this subcategory. The reported water use and discharge rates are presented in Table V-3 (page 2190).

ELECTROLYTE PREPARATION WET AIR POLLUTION CONTROL

The proposed and promulgated BPT wastewater discharge rate for the electrolyte preparation wet air pollution control is 0.05 liters per troy ounce (0.013 gal/troy ounce) of silver in the electrolyte produced. This normalized flow is based upon the only value reported for this subcategory. Water use and discharge rates are provided in Table V-4 (page 2190).

CALCINER WET AIR POLLUTION CONTROL

The proposed and promulgated BPT wastewater discharge rate for the calciner wet air pollution control is 186,200 1/kkg (49,200 gal/kkg) of mercury condensed. This normalized flow is based upon the sum of the flows from three in-series scrubbers at the only facility reporting a calciner scrubber (plant 1124). Table V-5 (page 2291) summarizes the water use and discharge rates for this subdivision. This discharge rate represents 16 percent recycle of scrubber liquor, which is the rate currently achieved by the one plant with this stream.

CALCINE QUENCH WATER

The proposed and promulgated BPT wastewater discharge rate for calcine quench water is 17,600 l/kkg (4,650 gal/kkg) of mercury condensed. This production normalized discharge rate is based

upon the only reported value for this waste stream. Water use and discharge rates are presented in Table V-6 (page 2191).

CALCINER STACK GAS CONTACT COOLING WATER

The proposed and promulgated BPT wastewater discharge rate selected for calciner stack gas contact cooling water is 4,150 l/kkg (1,096 gal/kkg) of mercury condensed. This discharge rate is equivalent to the discharge rate of the only plant reporting this waste stream. Table V-7 (page 2191) presents the reported water use and discharge rates for this waste stream.

CONDENSER BLOWDOWN

The proposed and promulgated BPT wastewater discharge rate for condenser blowdown is 13,800 l/kkg (3,646 gal/kkg) of mercury condensed. Water use and discharge rates for this waste stream are provided in Table V-8 (page 2192). The condenser blowdown normalized discharge rate is based upon the only value reported for this waste stream.

MERCURY CLEANING BATH WATER

The proposed and promulgated BPT wastewater discharge rate for mercury cleaning bath water is $1,400 \ 1/kkg$ (370 gal/kkg) of mercury condensed. This normalized flow is equivalent to the only reported water discharge rate for this waste stream. Table V-9 (page 2192) provides the reported water use and discharge flows for this subdivision.

REGULATED POLLUTANT PARAMETERS

The raw wastewater concentrations from individual operations and the subcategory as a whole were examined to select certain pollutant parameters for limitation. This examination and evaluation is presented in Sections VI and X. Eight pollutants or pollutant parameters are selected for limitation under BPT and are listed below:

122. 123. 126. 128.	lead mercury silver zinc gold	
	oil and grease	
	total suspended solids	(TSS)
	pH	

EFFLUENT LIMITATIONS

The concentrations achievable by application of the promulgated BPT treatment are explained in Section VII of this supplement. The achievable treatment concentrations (both one-day maximum and monthly average values) are multiplied by the BPT normalized discharge flows summarized in Table IX-1 (page 2240) to calculate

the mass of pollutants allowed to be discharged per mass of product. The results of these calculations in milligrams of pollutant per troy ounce or kilogram of product represent the BPT effluent limitations and are presented in Table IX-2 (page 2241) for each individual waste stream.

Table IX-1

BPT WASTEWATER DISCHARGE RATES FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

	BPT Disc		
Waste Stream	or 1/kkg)	(gal/T.O. or gal/kkg)	PNP
Smelter wet air pollution control	1.3	0.343	troy ounce of gold and silver smelted
Silver chloride reduction spent solution	0.4	0.11	troy ounce of silver reduced in solution
Electrolytic cells wet air pollution control	198	52.3	troy ounce of silver refined electrolytically
Electrolyte preparation wet air pollution control	0.05	0.013	troy ounce of silver in elec- trolyte produced
Calciner wet air pollution control	186,200	49,200	kkg of mercury condensed
Calcine quench water	17,600	4,650	kkg of mercury condensed
Calciner stack gas contact cooling water	4,150	1,096	kkg of mercury condensed
Condenser blowdown	13,800	3,646	kkg of mercury condensed
Mercury cleaning bath water	1,400	370	kkg of mercury condensed
	Smelter wet air pollution control Silver chloride reduction spent solution Electrolytic cells wet air pollution control Electrolyte preparation wet air pollution control Calciner wet air pollution control Calcine quench water Calciner stack gas contact cooling water Condenser blowdown	Waste Stream(1/T.0. or 1/kkg)Smelter wet air pollution control1.3Silver chloride reduction spent solution0.4Electrolytic cells wet air pollution control198Electrolyte preparation wet air pollution control0.05Calciner wet air pollution control186,200Calcine quench water cooling water17,600Calciner stack gas contact cooling water4,150Condenser blowdown13,800	Waste Streamor 1/kkg)or gal/kkg)Smelter wet air pollution1.30.343Silver chloride reduction0.40.11Silver chloride reduction0.40.11Electrolytic cells wet air19852.3pollution control0.050.013Electrolyte preparation wet air pollution control0.65Calciner wet air pollution control186,20049,200Calciner quench water17,6004,650Calciner stack gas contact cooling water13,8003,646

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TABLE IX-2

BPT MASS LIMITATIONS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(a) <u>Smelter Wet Air Pollution</u> <u>Control</u> BPT

Pollutant	or	Maximum	for	Maximum	for
pollutant		any one	day	monthly	average

mg/troy ounce of gold and silver smelted

Arsenic	2.717	1.209
Cadmium	0.442	0.195
Chromium	0.572	0.234
Copper	2.470	1.300
*Lead	0.546	0.260
*Mercury	0.325	0.130
Nickel	2.496	1.651
		0.221
*Silver	0.533	
Thallium	2.665	1.183
*Zinc	1.898	0.793
*Gold	0.130	
*Oil and Grease	26.000	15.600
*TSS	53.300	25.350
*pH Within the range		all times

(b) Silver Chloride Reduction Spent Solution BPT

Pollutant	or	Maximum	for	Maximum	for	
pollutant	property	any one	day	monthly	average	

mg/troy ounce of silver reduced in solution

Arsenic	0.836	0.372
Cadmium	0.136	0.060
Chromium	0.176	0.072
Copper	0.760	0.400
*Lead	0.168	0.080
*Mercury	0.100	0.040
Nickel	0.768	0.508
*Silver	0.164	0.068
Thallium	0.820	0.364
*Zinc	0.584	0.244
*Gold	0.040	
*Oil and Grease	8.000	4.800
*TSS	16.400	7.800
*pH Within the range o		times

TABLE IX-2 (Continued)

BPT MASS LIMITATIONS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(c) <u>Electrolytic</u> <u>Cells</u> <u>Wet Air</u> <u>Pollution</u> <u>Control</u> BPT

Pollutant or pollutant prop		imum for one day	Maximum monthly	for average	
mg/troy ounce	of silver re	fined electr	olytical	Ly	· · · ·
Arsenic		413.800		184.100	
Cadmium		67.320		29.700	
Chromium		87.120		35.640	
Copper		376.200	1	198.000	
*Lead		83.160		39.600	
*Mercury		49.500		19.800	ĸ
Nickel		380.200		251.500	
*Silver		81.180		33.660	
Thallium		405.900		180.200	
*Zinc		289.100		120.800	
*Gold		19.800			
*Oil and Grease	e 3.	,960.000	2.	376.000	
*TSS	-	118.000	-	861.000	
	the range of				

(d) Electrolyte Preparation Wet Air Pollution Control BPT

Pollutant or	Maximum for	Maximum for
pollutant property	any one day	monthly average
mg/troy ounce of silve	r in electrolyt	e produced
Arsenic	0.105	0.047
Cadmium	0.017	0.008
Chromium	0.022	0.009
Copper	0.095	0.050
*Lead	0.021	0.010
*Mercury	0.013	0.005
Nickel	0.096	0.064
*Silver	0.021	0.009
Thallium	0.103	0.046
*Zinc	0.073	0.031
*Gold	0.005	
*Oil and Grease	1.000	0.600
*TSS	2.050	0.975
*pH Within the range	e of 7.5 to 10.0	

TABLE IX-2 (Continued)

BPT MASS LIMITATIONS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(e) Calciner Wet Air Pollution Control BPT

Pollutant or	Maximum for	Maximum for	-
pollutant property	any one day	monthly average	

mg/kg (lb/million lbs) of mercury condensed

Arsenic	389.200	173.200
Cadmium	63.310	27.930
Chromium	81.930	33.520
Copper	353.800	186.200
*Lead	78.200	37.240
*Mercury	46.550	18.620
Nickel	357.500	236.500
*Silver	76.340	31.650
Thallium	381.700	169.400
*Zinc	271.900	113.600
*Gold	18.620	
*Oil and Grease	3,724.000	2,234.000
*TSS	7,634.000	3,631.000
*pH Within the	range of 7.5 to 10.0 at	all times
	-	

(f) Calcine Quench Water BPT

<u></u>	· · · · · · · · · · · · · · · · · · ·					
Pollutant	or	Maximum	for	Maximum	for	
			_		-	
pollutant	property	any one	dav	monthly	average	
-		<u>-</u>	<u>-</u>			

mg/kg (lb/million lbs) of mercury condensed

Arsenic	36.780	16.370
Cadmium	5.984	2.640
Chromium	7.744	3.168
Copper	33.440	17.600
*Lead	7.392	3.520
*Mercury	4.400	1.760
Nickel	33.790	22.350
*Silver	7.216	2.992
Thallium	36.080	16.020
*Zinc	25.700	10.740
*Gold	1.760	
*Oil and Grease	352°.000	211.200
*TSS	721.600	343.200
*pH Within the range of	7.5 to 10.0 at all	times

TABLE IX-2 (Continued)

BPT MASS LIMITATIONS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(g) Calciner Stack Gas Contact Cooling Water BPT

Pollutant	or	Məximum	for	Maximum	for
pollutant	property	any one	day	monthly	average

mg/kg (lb/million lbs) of mercury condensed

Arsenic	8.674	3.860
Cadmium	1.411	0.623
Chromium	1.826	0.747
Copper	7.885	4.150
*Lead	1.743	0.830
*Mercury	1.038	0.415
Nickel	7.968	5.271
*Silver	1.702	0.706
Thallium	8.508	3.777
*Zinc	6.059	2.532
*Gold	0.415	
*Oil and Grease	83.000	49.800
*TSS	170.200	80.930
*pH Within the range	of 7.5 to 10.0 at all	times

(h) Condenser Blowdown BPT

1

Pollutant	or	Maximum	for	Maximum	for
pollutant	property	any one	day	monthly	average

mg/kg (lb/million lbs) of mercury condensed

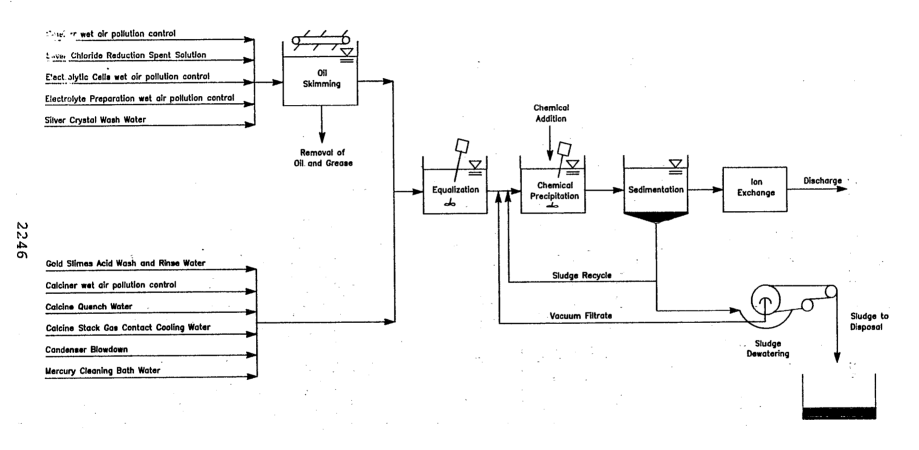
Arsenic	28.840	12.830
Cadmium	4.692	2.070
Chromium	6.072	2.484
Copper	26.220	13.800
*Lead	5.79 ੱ	2.760
*Mercury	3.450	1.380
Nickel	26.500	17.530
*Silver	5.658	2.346
Thallium	28.290	12.560
*Zinc	20.150	8.418
*Golđ	1.380	
*Oil and Grease	276.000	165.600
*TSS	565.800	269.100
*pH Within the range of	of 7.5 to 10.0 at a	ll times

TABLE IX-2 (Continued)

BPT MASS LIMITATIONS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(i) <u>Mercury</u> <u>Cleaning</u> <u>Bath</u> <u>Water</u> BPT

,	the second second	
Pollutant or	Maximum for	Maximum for
pollutant property	any one day	monthly average
· · · · · · · · · · · · · · · · · · ·		
mg/kg (lb/million lbs) of mercury con	ndensed
	• . · · ·	
Arsenic	2.926	1.302
Cadmium	0.476	0.210
Chromium	0.616	0.252
Copper	2.660	1.400
*Lead	0.588	0.280
*Mercury	0.350	0.140
Nickel	2.688	1.778
*Silver	0.574	0.238
Thallium	2.870	1.274
*Zinc	2.044	0.854
*Gold	0.140	0.034
*Oil and Grease	28.000	16.800
*TSS	57.400	
	ge of 7.5 to 10.	
Phi wichin the ran	de or 1.2 ro IO.	V AL AII LIMES



PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

SECT - IX

SECTION X

BEST AVAILABLE TECHNOLOGY ECONOMICALLY ACHIEVABLE

These effluent limitations are based on the best control and treatment technology used by a specific point source within the industrial category or subcategory, or by another category where it is readily transferable. Emphasis is placed on additional treatment techniques applied at the end of the treatment systems currently used, as well as reduction of the amount of water used and discharged, process control, and treatment technology optimization.

The factors considered in assessing best available technology economically achievable (BAT) include the age of equipment and facilities involved, the process used, process changes, nonwater quality environmental impacts (including energy requirements), and the costs of application of such technology (Section 304(b) (2)(B) of the Clean Water Act). BAT represents the best available technology economically achievable at plants of various ages, sizes, processes, or other characteristics. BAT may include feasible process changes or internal controls, even when not in common practice.

The statutory assessment of BAT considers costs, but does not require a balancing of costs against pollutant removals However, in assessing the proposed and promulgated BAT, the Agency has given substantial weight to the economic achievability of the technology.

TECHNICAL APPROACH TO BAT

The Agency reviewed a wide range of technology options and evaluated the available possibilities to ensure that the most effective and beneficial technologies were used as the basis of BAT. To accomplish this, the Agency elected to examine three technology options which could be applied to the primary precious metals and mercury subcategory as alternatives for the basis of BAT effluent limitations.

For the development of BAT effluent limitations, mass loadings were calculated for each wastewater source or subdivision in the subcategory using the same technical approach as described in Section IX for BPT limitations development. The differences in the mass loadings for BPT and BAT are due to increased treatment effectiveness achievable with the more sophisticated BAT treatment technology and reductions in the effluent flows allocated to various waste streams. In summary, the treatment technologies considered for the primary precious metals and mercury subcategory are:

Option A (Figure X-1, page 2264):

- Oil skimming preliminary treatment for streams containing oil and grease at treatable concentrations
- o Chemical precipitation and sedimentation
- o Ion exchange

Option B (Figure X-2 page 2265) is based on

- In-process flow reduction of wet air pollution control water
- Oil skimming preliminary treatment for streams containing oil and grease at treatable concentrations
- o Chemical precipitation and sedimentation
- o Ion exchange

Option C (Figure X-3 page 2266) is based on

- In-process flow reduction of wet air pollution control water
- Oil skimming preliminary treatment for streams containing oil and grease at treatable concentrations
- o Chemical precipitation and sedimentation
- o Multimedia filtration
- o Ion exchange

The three options examined for BAT are further discussed below. The first option considered is the same as the BPT treatment technology which was presented in section IX.

OPTION A

Option A for the primary precious metals and mercury subcategory equivalent to the control and treatment technologies which is The BPT end-of-pipe were analyzed for BPT in Section IX. precipitation and chemical includes treatment scheme sedimentation (lime and settle) technology, with oil skimming treatment of wastewaters containing treatable preliminary concentrations of oil and grease and ion exchange as a polishing The discharge rates for Option A step (see Figure X-1). are equal to the discharge rates allocated to each stream as a BPT discharge flow.

OPTION B

Option B for the primary precious metals and mercury subcategory achieves lower pollutant discharge by building upon the Option A (oil skimming preliminary treatment, chemical precipitation and sedimentation and ion exchange) treatment technology. Flow reduction measures are added to the Option A treatment scheme (see Figure X-2). These flow reduction measures, including inprocess changes, result in the concentration of pollutants in

some wastewater streams. As explained in Section VII of the General Development Document, treatment of a more concentrated effluent allows achievement of a greater net pollutant removal and introduces the possible economic benefits associated with treating a lower volume of wastewater.

Option B flow reduction measures are reflected in the BAT wastewater discharge rates. Flow reduction has been included in determining the promulgated BAT discharge rates for electrolytic cells wet air pollution control and calciner wet air pollution control. Based on available data, the Agency did not feel that further flow reduction over BPT would be feasible for the remaining seven waste streams in the primary precious metals and mercury subcategory. These waste streams are:

- 1. Smelter wet air pollution control,
- 2. Silver chloride reduction spent solution,
- 3. Electrolyte preparation wet air pollution control,
- 4. Calcine quench water,
- 5. Calciner stack gas contact cooling water,
- 6. Condenser blowdown, and
- 7. Mercury cleaning bath water.

Flow reduction measures used in Option B to reduce process wastewater generation or discharge rates include the following:

Recycle of Water Used in Wet Air Pollution Control

There are four wastewater sources associated with wet air pollution control which are regulated under the primary precious metals and mercury subcategory:

- 1. Smelter wet air pollution control,
- 2. Electrolytic cells wet air pollution control,
- 3. Electrolyte preparation wet air pollution control, and
- 4. Calciner wet air pollution control.

(page 2255) presents the number of plants reporting Table X-1 wastewater from the wet air pollution control sources listed above, the number of plants practicing recycle, and the range of recycle values being listed. Recycle of electrolytic cell scrubber water and calciner scrubber water are required for BAT. Recycle of smelter wet air pollution control and electrolyte preparation wet air pollution control is not required for BAT because the BPT discharge flow is close to the minimum possible water discharge from a scrubber. The recycle rate used for the other two sources is based on 90 percent recycle of the average water use reported by all the plants with each waste stream, as will be shown later.

OPTION C

Option C for the primary precious metals and mercury subcategory consists of all control and treatment requirements of Option B (in-process flow reduction, oil skimming preliminary treatment,

chemical precipitation, sedimentation, and ion exchange) plus multimedia filtration technology added at the end of the Option B treatment scheme (see Figure X-3, page 2266). Multimedia filtration is used to remove suspended solids, including precipitates of toxic metals, beyond the concentration attainable by gravity sedimentation. The filter suggested is of the gravity, mixed media type, although other filters, such as rapid sand filters or pressure filters, would perform satisfactorily.

INDUSTRY COST AND POLLUTANT REMOVAL ESTIMATES

As one means of evaluating each technology option, EPA developed estimates of the pollutant removal benefits and the compliance costs associated with each option. The methodologies are described below.

POLLUTANT REMOVAL ESTIMATES

A complete description of the methodology used to calculate the estimated pollutant removal, or benefit, achieved by the application of the various treatment options is presented in Section X of Vol. I. The pollutant removal estimates have been revised from proposal because of additional flow and production information received during the comment period. The methodology for calculating pollutant removals has not changed, and the data used for estimating removals are the same as those used to revise compliance costs.

Sampling data collected during the field sampling program were used to characterize the major waste streams considered for regulation. At each sampled facility, the sampling data was production normalized for each unit operation (i.e., mass of pollutant generated per mass of product manufactured). This value, referred to as the raw waste, was used to estimate the mass of toxic pollutants generated within the primary precious metals and mercury subcategory. The pollutant removal estimates were calculated for each plant by first estimating the total mass of each pollutant in the untreated wastewater. This was calculated by first multiplying the raw waste values by the corresponding production value for that stream and then summing these values for each pollutant for every stream generated by the plant.

Next, the volume of wastewater discharged after the application of each treatment option was estimated for each operation at each plant by comparing the actual discharge to the regulatory flow. The smaller of the two values was selected and summed with the other plant flows. The mass of pollutant discharged was then estimated by multiplying the achievable concentration values attainable with the option (mg/l) by the estimated volume of process wastewater discharged by the subcategory. The mass of pollutant removed is the difference between the estimated mass of pollutant generated within the subcategory and the mass of pollutant discharged after application of the treatment option. The pollutant removal estimates for direct dischargers in the

primary precious metals and mercury subcategory are presented in Table X-2 (page 2256).

COMPLIANCE COSTS

In estimating subcategory-wide compliance costs, the first step was to develop a cost estimation model, relating the total costs associated installation and operation of with wastewater treatment technologies to plant process wastewater discharge. EPA applied the model to each plant. The plant's investment and operating costs are determined by what treatment it has in place and by its individual process wastewater discharge flow. As discussed above, this flow is either the actual or the BAT regulatory flow, whichever is lesser. The final step was to annualize the capital costs, and to sum the annualized capital costs, and the operating and maintenance costs for each plant, yielding the cost of compliance for the subcategory (see Table X-3, page 2257). These costs were used in assessing economic achievability.

BAT OPTION SELECTION - PROPOSAL

EPA selected Option C for the proposed BAT, which included flow reduction, oil skimming preliminary treatment, chemical precipitation and sedimentation, and multimedia filtration.

The pollutants proposed for limitation under BAT were arsenic, lead, mercury, silver, and zinc. Implementation of the proposed BAT limitations was estimated to remove 914.5 kilograms of toxic metals annually. Estimated capital cost for achieving proposed BAT was \$30,000 and annual cost was \$10,000 (1982 dollars).

BAT OPTION SELECTION - PROMULGATION

EPA is promulgating BAT limitations for this subcategory based on flow reduction, oil skimming preliminary treatment, chemical precipitation and sedimentation, ion exchange and multimedia filtration. This preliminary treatment and end-of-pipe technology basis for the promulgated BAT adds ion exchange to the technology used for the proposed BAT limitations. The treatment performance concentrations upon which the mass limitations are based are equal to those used to calculate the proposed mass limitations.

EPA is promulgating multimedia filtration as part of the BAT technology because this technology results in additional removal of toxic metals. Filtration is also presently demonstrated at 25 plants throughout the nonferrous metals manufacturing category. Filtration adds reliability to the treatment system by making it less susceptible to operator error and to sudden changes in raw wastewater flow and pollutant concentrations.

Oil skimming is demonstrated in the nonferrous metals manufacturing category. Although no primary precious metals and mercury plants have oil skimming in place, it is necessary to reduce oil and grease concentrations in the discharge from this subcategory.

EPA has added ion exchange end-of-pipe treatment to the BAT treatment scheme discussed at proposal. Ion exchange is an effective method for removing gold from wastewater generated in the subcategory. EPA has determined that no additional costs will be generated in the secondary precious metals subcategory by adding ion exchange because of the value of the precious metals recovered in the column, and believes this will also be true for primary precious metals subcategory.

Implementation of the control and treatment technologies of Option C will remove annually an estimated 50,443 kilograms of priority metal pollutants from raw wastewater. The estimated capital cost for achieving promulgated BAT is \$3,025 (1982 dollars) and the estimated annual cost is \$27,300 (1982 dollars).

WASTEWATER DISCHARGE RATES

A BAT discharge rate was calculated for each subdivision based upon the flows of the existing plants, as determined from analysis of the data collection portfolios. The discharge rate is used with the achievable treatment concentrations to determine BAT effluent limitations. Since the discharge rate may be production different for each wastewater source, separate normalized discharge rates for each of the nine wastewater sources were determined and are summarized in Table X-4 (page The discharge rates wastewater generated to the mass of 2258). the intermediate product which is produced by the process These production associated with the waste stream in question. normalizing parameters (PNP) are also listed in Table \tilde{X} -4. The discharge rates are normalized on a production basis by relating amount of wastewater generated to the mass of the the intermediate product which is produced by the process associated with the waste stream in question. These production normalizing parameters (PNP) are also listed in Table X-4.

The promulgated BAT discharge rates are the same as the discharge rates proposed for BAT. As discussed previously, the promulgated BAT wastewater discharge rate equals the BPT wastewater discharge rate for seven of the nine waste streams in the primary precious metals and mercury subcategory. Based on the available data, the Agency determined that further flow reduction would not be feasible for these seven wastewater sources. Wastewater streams for which BAT discharge rates differ from BPT are discussed below.

ELECTROLYTIC CELLS WET AIR POLLUTION CONTROL

The promulgated BAT wastewater discharge rate for electrolytic cells wet air pollution control is 19.8 liters per troy ounce of gold refined electrolytically. This rate is based on 90 percent recycle of the water use rate reported by the one plant with this waste stream, as shown in Table V-3 (page 2190). Although

recycle of this stream is not currently demonstrated, the Agency believes that it is achievable.

CALCINER WET AIR POLLUTION CONTROL

The promulgated BAT wastewater discharge rate for calciner wet air pollution control is 22,000 liters per metric ton of mercury condensed. This rate is based on 90 percent recycle of the water use rate reported by the only plant with this waste stream. As shown in Table V-5 (page 2191), the plant reported a flow of 186,000 l/kkg, which represents a 16 percent recycle rate. The BAT rate was determined by the following formula:

 $\frac{(186,000 \ 1/kkg)}{(1.00 \ -} \ \frac{0.90)}{0.16)} = 22,000 \ 1/kkg$

Although 90 percent recycle is not demonstrated for this waste stream, the Agency believes it is achievable.

REGULATED POLLUTANT PARAMETERS

In implementing the terms of the Consent Agreement in NRDC v. Train, Op. Cit., and 33 U.S.C.cl314(b)(2)(A and B) (1976), the Agency placed particular emphasis on the toxic pollutants. The raw wastewater concentrations from individual operations and the subcategory as a whole were examined to select certain pollutant parameters for consideration for limitation. This examination and evaluation, presented in Section VI, concluded that 10 pollutants are present in primary precious metals and mercury wastewaters at concentrations than can be effectively reduced by identified treatment technologies (refer to Section VI).

The high cost associated with analysis for toxic metal pollutants has prompted EPA to develop an alternative method for regulating and monitoring toxic pollutant discharges from the nonferrous metals manufacturing category. Rather than developing specific effluent mass limitations and standards for each of the toxic metals found in treatable concentrations in the raw wastewaters from a given subcategory, the Agency is promulgating effluent mass limitations only for those pollutants generated in the greatest quantities as shown by the pollutant removal analysis. The pollutants selected for specific limitation are listed below:

122. lead
123. mercury
126. silver
128. zinc
gold

By establishing limitations and standards for certain toxic metal pollutants, dischargers will attain the same degree of control over the other toxic metal pollutants as they would have been required to achieve had all the priority metal pollutants been directly limited. This approach is technically justified since the treatment effectiveness concentrations used for chemical precipitation and sedimentation technology are based on optimized treatment for concomitant multiple metals removal. Thus, even though metals have somewhat different theoretical solubilities, they will be removed at very nearly the same rate in a chemical precipitation and sedimentation treatment system operated for multiple metals removal. Filtration as part of the technology basis is likewise justified because this technology removes metals nonpreferentially.

The toxic metal pollutants selected for specific limitation in the primary precious metals and mercury subcategory to control the discharges of other toxic metal pollutants are lead, mercury, silver, and zinc.

Gold is selected for limitation in this subcategory because the methods used to control lead, mercury, silver and zinc are not effective in controlling the discharge of gold.

The following priority pollutants are excluded from limitation on the basis that they are effectively controlled by the limitations developed for lead, mercury, silver, zinc and gold.

115. arsenic
118. cadmium
119. chromium
120. copper
124. nickel
127. thallium

EFFLUENT LIMITATIONS

The concentrations achievable by application of the technology (Option C) are discussed in Section VII of BAT this supplement. These treatment effectiveness concentrations (both one-day maximum and monthly average) are multiplied by the BAT normalized discharge flows summarized in Table X-4 (page 2258) to calculate the mass of pollutants allowed to be discharged per mass of product. The results of these calculations in milligrams of pollutant per troy ounce or kilogram of product represent the promulgated BAT effluent limitations for the primary precious metals and mercury subcategory. BAT effluent limitations based on Option C (oil skimming, chemical precipitation, sedimentation, flow reduction, ion exchange and multimedia in-process filtration) are presented in Table X-5 (page 2259).

TABLE X-1

CURRENT RECYCLE PRACTICES WITHIN THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

Pla	mber of nts With tewater	Number of Plant Practici Recycle	s Range ng of Recycle
			· · · · · ·
Smelter wet air pollution control	3	3	76-100
Electrolytic cells wet air pollution control	1	0	0
Electrolyte preparation wet air pollution control	1	0	0
Calciner wet air pollution control	1	1	16

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s 5 -

TABLE X-2

CURRENT RECYCLE PRACTICES WITHIN THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

	er of s With water	Numb of Pl Pract <u>Rec</u>	ants	Range of Recycle <u>Values (%)</u>
Smelter wet air pollution control	3		3	76 - 100
Electrolytic Cells Wet Air Pollution Control	1	g tostan an 1 an se	0	0
Electrolytic Preparation Wet Air Pollution Control			0	0
Calciner Wet Air Pollution Control	1		1	16

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TABLE X-3

COST OF COMPLIANCE FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY DIRECT DISCHARGERS

(March, 1982 Dollars)

Option	Proposa Capital Cost	l Costs Annual Cost	Promulgat Capital Cost	ion Costs Annual Cost
A	27,500	9,000	2,200	26,800
8 .*	27,500	9.000	2,200	26,800
C	30,000	10,000	3,025	27,300

TABLE X-4

BPT WASTEWATER DISCHARGE RATES FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

			narge Rate	
	Waste Stream	(1/T.O. or 1/kkg)	(gal/T.O. or gal/kkg)	PNP
		<u>/</u>		A LTA
1.	Smelter wet air pollution control	1.3	0.343	troy ounce of gold and silver smelted
2.	Silver chloride reduction spent solution	0.4	0.11	troy ounce of silver reduced in solution
3.	Electrolytic cells wet air pollution control	19.8	5.23	troy ounce of gold refined electrolytically
4.	Electrolyte preparation wet air pollution control	0.05	0.013	troy ounce of silver in elec- trolyte produced
5.	Calciner wet air pollution control	22,000	5,812	kkg of mercury condensed
6.	Calcine quench water	17,600	4,650	kkg of mercury condensed
7.	Calciner stack gas contact cooling water	4,150	1,096	kkg of mercury condensed
8.	Condenser blowdown	13,800	3,646	kkg of mercury condensed
9.	Mercury cleaning bath water	1,400	370	kkg of mercury condensed
7. 8.	Calciner stack gas contact cooling water Condenser blowdown	4,150 13,800	1,096 3,646	kkg of mercury condensed kkg of mercury condensed

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TABLE X-5

BAT MASS LIMITATIONS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(a)	Smelter	Wet	Air	Pollution	Control	BAT
-----	---------	-----	-----	-----------	---------	-----

Pollutant or	Maximum for	Maximum for	
pollutant property	any one day	monthly average	,
mg/troy ounce of gold	and silver smel	ted	
Arsenic	1.807	0.806	
Cadmium	0.260	0.104	
Chromium	0.481	0.195	,
Copper	1.664	0.793	
*Lead	0.364	0.169	
*Mercury	0.195	0.078	•
Nickel	0.715	0.481	
*Silver	0.377	0.156	
Thallium	1.820	0.793	
*Zinc	1.326	0.546	
*Gold	0.130		
	·		;
(b) <u>Silver</u> Chloride	Reduction Spent	Solution BAT	
Pollutant or	Maximum for	Maximum for	
pollutant property	any one day	monthly average	

mg/troy ounce of silver reduced in solution

	•	· · ·
Arsenic	0.556	0.248
Cadmium	0.080	0.032
Chromium	0.148	0.060
Copper	0.512	0.244
*Lead	0.112	0.052
*Mercury	0.060	0.024
Nickel	0.220	0.148
*Silver	0.116	0.048
Thallium	0.560	0.244
*Zinc	0.408	0.168
*Gold	0.040	
•	·	

TABLE X-5 (Continued)

BAT MASS LIMITATIONS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(c) <u>Electrolytic</u> <u>Cells</u> <u>Wet</u> <u>Air</u> <u>Pollution</u> <u>Control</u> BAT

Pollutant or pollutant property	Maximum for any one day	Maximum for monthly average
mg/troy ounce of sil	ver refined elec	trolytically
Arsenic	27.520	12.280
Cadmium	3.960	1.584
Chromium	7.326	2.970
Copper	25.340	12.080
*Lead	5.544	2.574
*Mercury	2.970	1.188
Nickel	10.890	7.326
*Silver	5.742	2.376
Thallium	27.720	12.080
*Zinc	20.200	8.316
*Gold	1.980	· · · · · · · · · · · · · · · · · · ·

Pollutant or Maximum for Maximum for pollutant property any one day monthly average

mg/troy ounce of silver in electrolyte produced

0.070	0.031
0.010	0.004
0.019	0.008
0.064	0.031
0.014	0.007
0.008	0.003
0.028	0.019
0.015	0.006
0.070	0.031
0.051	0.021
0.005	
	0.010 0.019 0.064 0.014 0.008 0.028 0.015 0.070 0.051

TABLE X-5 (Continued)

BAT MASS LIMITATIONS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(e) <u>Calciner Wet Air Pollution Control</u> BAT

Pollutant or	Maximum for	Maximum for
pollutant property	any one day	monthly average
mg/kg (lb/million lk	os) of mercury cor	ndensed
Arsenic	30.580	13.640
Cadmium	4.400	1.760
Chromium	8.140	3.300
Copper	28.160	13.420
*Lead	6.160	2.860
*Mercury	3.300	1.320
Nickel	12.100	8.140
*Silver	6.380	2.640
Thallium	30.800	13.420
*Zinc	22.440	9.240
*Gold	2.200	

(f) Calciner Quench Water BAT

Pollutant or Maximum for Maximum for pollutant property any one day monthly average (1b/million lbs) of mercury condensed	
	je
Augusta 24 460 10 01	
Arsenic 24.460 10.91	.0
Cadmium 3.520 1.40	8
Chromium 6.512 2.64	10
Copper 22.530 10.74	10.
*Lead 4.928 2.28	38
*Mercury 2.640 1.05	66
Nickel 9.680 6.51	2
*Silver 5.104 2.11	.2
Thallium 24.640 10.74	10
*Zinc 17.950 7.39	2
*Gold 1.760	

*Regulated Pollutant

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TABLE X-5 (Continued)

BAT MASS LIMITATIONS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(g) <u>Calciner</u> <u>Stack</u> <u>Gas</u> <u>Contact</u> <u>Cooling</u> <u>Water</u> BAT

Pollutant	or	Maximum	for	Maximum	for
pollutant	property	any one	day	monthly	average
			1		

mg/kg (lb/million lbs) of mercury condensed

Arsenic	5.769	2.573
Cadmium	0.830	0.332
Chromium	1.536	0.623
Copper	5.312	2.532
*Lead	1.162	0.540
*Mercury	0.623	0.249
Nickel	2.283	1.536
*Silver	1.204	0.498
Thallium	5.810	2.532
*Zinc	4.233	1.743
*Gold	0.415	

(h) Condenser Blowdown BAT

Pollutant	or	Maximum	for	Maximum	for
pollutant		any one		monthly	
porrucune	propercy	any one	aaj	monenty	average

mg/kg (lb/million lbs) of mercury condensed

Arsenic Cadmium Chromium Copper *Lead *Mercury Nickel *Silver Thallium *Zinc *Gold	19.180 2.760 5.106 17.660 3.864 2.070 7.590 4.002 19.320 14.080 1.380	8.556 1.104 2.070 8.418 1.794 0.828 5.106 1.656 8.418 5.796
*Gold	1.380	·

*Regulated Pollutant

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TABLE X-5 (Continued)

BAT MASS LIMITATIONS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(i) Mercury Cleaning Bath Water BAT

Pollutant or pollutant property	Maximum for any one day	Maximum for monthly average
mg/kg (lb/million l	bs) of mercury con	ndensed
Arsenic	1.946	0.868
Cadmium	0.280	0.112
Chromium	0.518	0.210
Copper	1.792	0.854
*Lead	0.392	0.182
*Mercury	0.210	0.084
Nickel	0.770	0.518
*Silver	0.406	0.168
Thallium	1.960	0.854
*Zinc	1.428	0.588
*Gold	0.140	````

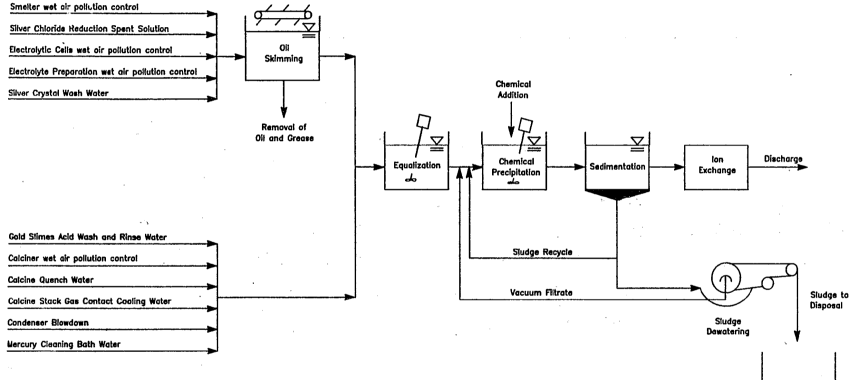


FIGURE X-1

BAT TREATMENT SCHEME FOR OPTION A

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PRIMARY PRECIOUS METALS AND Holding Electrolytic Cells Wet Air Pollution Control Tank Recycle Smelter wat air pollution control Silver Chloride Reduction Spent Solution Oil Skimmina Electrolytic Cells wet air pollution control Backwash Chemical Silver Crystal Wash Water Addition Removal of Y Oil and Grease ∇ V MERCURY SUBCATEGORY ----Multimedia Chemical ion Equalization Sedimentation Filtration Exchange Precipitation T Calciner wet air pollution control Holding Backwash Sludge Tank Sludge Recycle Recycle Vacuum Filtrate Sludge to Gold Slimes Acid Wash and Rinse Water Disposal Sludge Colcine Quench Water Dewatering Calcine Stack Gas Contact Cooling Water Condenser Blowdown Mercury Cleaning Bath Water

> BAT TREATMENT SCHEME FOR OPTION B FIGURE X-2

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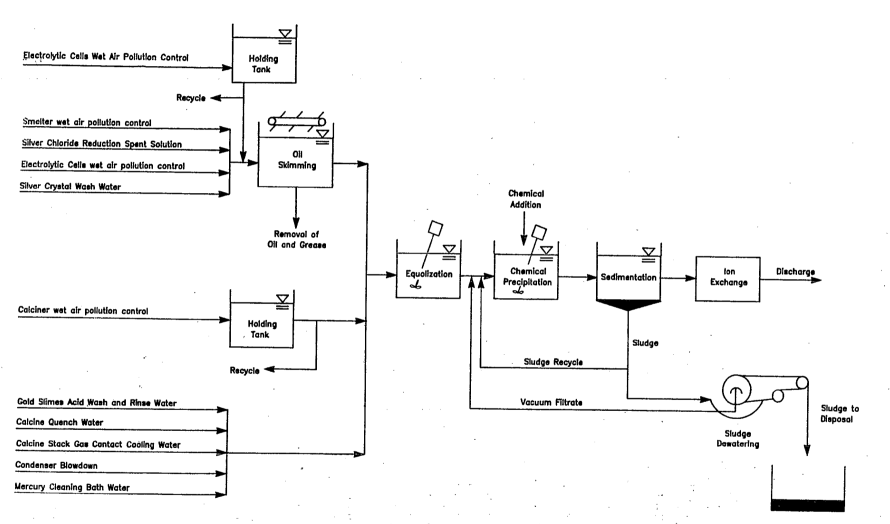


FIGURE X-3 BAT TREATMENT SCHEME FOR OPTION C

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PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

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SECTION XI

NEW SOURCE PERFORMANCE STANDARDS

section describes the technologies for treatment This wastewater from new sources and presents mass discharge standards regulated pollutants for NSPS in the primary precious metals for and mercury subcategory, based on the selected treatment technology. The basis for new source performance standards (NSPS) is the best available demonstrated technology (BDT). New plants have the opportunity to design the best and most efficient processes and wastewater treatment technologies production without facing the added costs and restrictions encountered in retrofitting an existing plant. Therefore, EPA has considered the best demonstrated process changes, in-plant controls, and end-of-pipe treatment technologies which reduce pollution to the maximum extent feasible.

TECHNICAL APPROACH TO NSPS

source performance standards are equivalent to the best New available technology (BAT) selected for currently existing primary precious metals and mercury plants. This result is a consequence of careful review by the Agency of a wide of range technical options for new source treatment systems which is discussed in Section XI of Vol. I. Additionally, there was nothing found to indicate that the wastewater flows and characteristics of new plants would not be similar to those from existing plants, since the processes used by new sources are not to differ from those used at existing sources. expected Consequently, BAT production normalized discharge rates, which are based on the best existing practices of the subcategory, can also be applied to new sources. These rates are presented in Table XI-1 (page 2270).

Treatment technologies considered for the NSPS options are identical to the treatment technologies considered for the BAT options. These options are:

OPTION A

o Preliminary treatment with oil skimming (where required)

o Chemical precipitation and sedimentation

o Ion exchange

OPTION B

o Chemical precipitation and sedimentation

o In-process flow reduction of electrolytic cells and

o Calciner scrubber liquor

o Ion exchange

OPTION C

- o Preliminary treatment with oil skimming (where required)
- o Chemical precipitation and sedimentation
- In-process flow reduction of electrolytic cells and calciner scrubber liquor
- o Multimedia filtration
- o Ion exchange

NSPS OPTION SELECTION - PROPOSAL

EPA proposed that the best available demonstrated technology for the primary precious metals and mercury subcategory be equivalent to Option C (oil skimming preliminary treatment, flow reduction, chemical precipitation and sedimentation, and multimedia filtration).

The wastewater flow rates for NSPS were the same as the proposed BAT flow rates. Flow reduction measures for NSPS beyond the allowances for BAT were not considered feasible as no new demonstrated technologies existed within the subcategory. Therefore, EPA concluded that flow reduction beyond the allowances proposed for BAT were unachievable, and proposed NSPS flow rates should be equal to those for BAT.

NSPS OPTION SELECTION - PROMULGATION

EPA is promulgating best available technology for the primary precious metals and mercury subcategory equivalent to Option C (oil skimming preliminary treatment, flow reduction, chemical precipitation and sedimentation, ion exchange and multimedia filtration).

The wastewater flow rates for NSPS are the same as the BAT flow rates. The Agency does not believe that new plants could achieve any flow reduction beyond the allowances promulgated for BAT. Because NSPS is equal to BAT, the promulgated NSPS will not have a detrimental impact on the entry of new plants into this subcategory.

REGULATED POLLUTANT PARAMETERS

The Agency has no reason to believe that the pollutants that will be found in treatable concentrations in processes within new sources will be any different than with existing sources. Accordingly, pollutants and pollutant parameters selected for limitation under NSPS, in accordance with the rationale of Sections VI and X, are identical to those selected for BAT. The conventional pollutant parameters oil and grease, TSS, and pH are also selected for limitation.

NEW SOURCE PERFORMANCE STANDARDS

The NSPS discharge flows for each wastewater source are the same as the discharge rates for BAT and are shown in Table XI-1 (page 2270). The mass of pollutant allowed to be discharged per mass of product is calculated by multiplying the appropriate treatable concentration (mg/l) by the production normalized wastewater discharge flows (1/T.0. or 1/kkg). The results of these calculations are the production-based new source performance standards. These standards are presented in Tables XI-2 (page 2271 - 2273).

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Table XI-1

NSPS WASTEWATER DISCHARGE RATES FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

	NSPS Discharge Rate			
	Waste Stream	(1/T.O. or 1/kkg)	(gal/T.O. or gal/kkg)	PNP
1.	Smelter wet air pollution control	1.3	0.343	troy ounce of gold and silver smelted
2.	Silver chloride reduction spent solution	0.4	0.11	troy ounce of silver reduced in solution
3.	Electrolytic cells wet air pollution control	19.8	5.23	troy ounce of gold refined electrolytically
4.	Electrolyte preparation wet air pollution control	0.05	0.013	troy ounce of silver in elec- trolyte produced
5.	Calciner wet air pollution control	22,000	5,812	kkg of mercury condensed
6.	Calcine quench water	17,600	4,650	kkg of mercury condensed
7.	Calciner stack gas contact cooling water	4,150	1,096	kkg of mercury condensed
8 .	Condenser blowdown	13,800	3,646	kkg of mercury condensed
9.	Mercury cleaning bath water	1,400	370	kkg of mercury condensed

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TABLE XI-2

NSPS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(a) <u>Smelter</u> <u>Wet</u> <u>Air</u> <u>Pollution</u> <u>Control</u> NSPS

	·				
Pollutant	or	Maximum	for	Maximum	for
pollutant		any one	day	monthly	average

mg/troy ounce of gold and silver smelted

Arsenic	1.807	0.806
Cadmium	0.260	0.104
Chromium	0.481	0.195
Copper	1.664	0.793
*Lead	0.364	0.169
*Mercury	0.195	0.078
Nickel	0.715	0.481
*Silver	0.377	0.156
Thallium	1.820	0.793
*Zinc	1.326	0.546
*Gold	0.130	·
*Oil and Grease	13.000	13.000
*TSS	19.500	15.600
	of 7.5 to 10.0 at a	ll times

(b) Silver Chloride Reduction Spent Solution NSPS

	. ,	·				
Pollutant	or	Maximum	for	Maximum	for	
pollutant		any one	day	monthly	average	·

mg/troy ounce of silver reduced in solution

		0 040
Arsenic	0.556	0.248
Cadmium	0.080	0.032
Chromium	0.148	0.060
Copper	0.512	0.244
*Lead	0.112	0.052
*Mercury	0.060	0.024
Nickel	0.220	0.148
*Silver	0.116	0.048
Thallium	0.560	0.244
*Zinc	0.408	0.168
*Gold	0.040	
*Oil and Grease	4.000	4.000
*TSS	6.000	4.800
*pH Within the rang	e of 7.5 to 10.0 at	all times

TABLE XI-2 (Continued)

NSPS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(c) <u>Electrolytic</u> <u>Cells</u> <u>Wet</u> <u>Air</u> <u>Pollution</u> <u>Control</u> <u>NSPS</u>

Pollutant		Maximum for	Maximum	for
pollutant	property	any one day	monthly	average

mg/troy ounce of silver refined electrolytically

.

Arsenic	27.520	12.280
Cadmium	3.960	1.584
Chromium	7.326	2.970
Copper	25.340	12.080
*Lead	5.544	2.574
*Mercury	2.970	1.188
Nickel	10.890	7.326
*Silver	5.742	2.376
Thallium	27.720	12.080
*Zinc	20.200	8.316
*Gold	1.980	
*Oil and Grease	198.000	198.000
*TSS	297.000	237.600
*pH Within the range	of 7.5 to 10.0 a	at all times

(d) Electrolyte Preparation Wet Air Pollution Control NSPS

Pollutant or	Maximum	for	Maximum for
nollutant property			
pollutant property	any one	uay	monthly average

mg/troy ounce of silver in electrolyte produced

Arsenic	0.070	0.031
Cadmium	0.010	0.004
Chromium	0.019	0.008
Copper	0.064	0.031
*Lead	0.014	0.007
*Mercury	0.008	0.003
Nickel	0.028	0.019
*Silver	0.015	0.006
Thallium	0.070	0.031
*Zinc	0.051	0.021
*Gold	0.005	
*Oil and Grease	0.500	0.500
*TSS	0.750	0.600
*pH Within the range	of 7.5 to 10.0 at all	times

TABLE XI-2 (Continued)

NSPS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(e) Calciner Wet Air Pollution Control NSPS

Pollutant or	Maximum for	Maximum for
pollutant property	any one day	monthly average

mg/kg (lb/million lbs) of mercury condensed

Arsenic	30.580	13.640
Cadmium	4.400	1.760
Chromium	8.140	3.300
Copper	28.160	13.420
*Lead	6.160	2.860
*Mercury	3.300	1.320
Nickel	12.100	8.140
*Silver	6 380	2.640
Thallium	30.800	13.420
*Zinc	22.440	9.240
*Gold	2.200	الحدة معداد •
*Oil and Grease	220.000	220.000
*TSS	330.000	264.000
*pH Within the range	of 7.5 to 10.0 at	all times

(f) Calcine Quench Water NSPS

Pollutant	or	Maximum for	Maximum	for
pollutant	property	any one day	monthly	average

mg/kg (lb/million lbs) of mercury condensed

Arsenic	24.460	10.910
Cadmium	3.520	1.408
Chromium	6.512	2.640
Copper	22.530	10.740
*Lead	4.928	2.288
*Mercury	2.640	1.056
Nickel	9.680	6.512
*Silver	5.104	2.112
Thallium	24.640	10.740
*Zinc	17.950	7.392
*Gold	1.760	
*Oil and Grease	176.000	176.000
*TSS	264.000	211.200
*pH Within the range of	7.5 to 10.0 at al	l times

TABLE XI-2 (Continued)

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NSPS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(g) <u>Calciner Stack</u> <u>Gas</u> <u>Contact</u> <u>Cooling</u> <u>Water</u> NSPS

		and the second			· · · · · · · · · · · · · · · · · · ·
Pollutant or	Max	imum for	Maximum	for	
pollutant pre	onerty any	one day	monthly	average	
porracane pr	opercy any	one day	montentl	average	
			• .	· ·	1

mg/kg (lb/million lbs) of mercury condensed

Arsenic	5.769	2.573
Cadmium	0.830	0.332
Chromium	1.536	0.623
Copper	5.312	2.532
*Lead	1.162	0.540
*Mercury	0.623	0.249
Nickel	2.283	1.536
*Silver	1.204	0.498
Thallium	5.810	2.532
*Zinc	4.233	1.743
*Gold	0.415	
*Oil and Grease	41.500	41.500
*TSS	62.250	49.800
*pH Within the range	of 7.5 to 10.0 at	all times

(h) Condenser Blowdown NSPS

Pollutant	or	Maximum	for	Maximum	for
Pollutant	property	any one	day	monthly	average

mg/kg (lb/million lbs) of mercury condensed

Arsenic	19.180	8.556
Cadmium	2.760	1.104
Chromium	5.106	2.070
Copper	17.660	8.418
*Lead	3.864	1.794
*Mercury	2.070	.828
Nickel	7.590	5.106
*Silver	4.002	1.656
Thallium	19.320	8.418
*Zinc	14.080	5.796
*Gold	1.380	· · · · · · · · · · · · · · · · · · ·
*Oil and Grease	138.000	138.000
*TSS	207.000	165.600
*pH Within the range	of 7.5 to 10.0 at all	times

TABLE XI-2 (Continued)

NSPS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(i) Mercury Cleaning Bath Water NSPS

Pollutant or	Maximum for	Maximum for
pollutant property	any one day	monthly average
mg/kg (1b/million lbs)	of mercury con	densed

Arsenic	1.946	0.868
Cadmium	0.280	0.112
Chromium	0.518	0.210
Copper	1.792	0.854
*Lead	0.392	0.182
	0.210	0.084
	0.770	0.518
*Silver	0.406	0.168
Thallium	1.960	0.854
	1.428	0.588
•	0.140	
*Oil and Grease	14.000	14.000
*TSS	21.000	16.800
*pH Within the range of		L times

*Regulated Pollutant

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PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SECT - XII

SECTION XII

PRETREATMENT STANDARDS

This section describes the control and treatment technologies for pretreatment of process wastewaters from new sources in the primary precious metals and mercury subcategory. Pretreatment standards are designed to prevent the discharge of pollutants which pass through, interfere with, or are otherwise incompatible with the operation of publicly owned treatment works (POTW). The Clean Water Act requires pretreatment for pollutants, such as toxic metals, that limit POTW sludge management alternatives. New indirect discharge facilities, like new direct discharge facilities, have the opportunity to incorporate the best available demonstrated technologies, including process changes, in-plant controls, and end-of-pipe treatment technologies, and to use plant site selection to ensure adequate treatment system installation. Pretreatment standards are to be technology based, analogous to the best available or demonstrated technology for removal of toxic pollutants.

Pretreatment standards for regulated pollutants are presented based on the selected control and treatment technology. EPA is not promulgating pretreatment standards for existing sources in this subcategory because no indirect dischargers exist. Moreover, EPA is promulgating pretreatment standards for new sources because plants may be constructed in the future which may discharge to a POTW.

TECHNICAL APPROACH TO PRETREATMENT

Before proposing and promulgating pretreatment standards, the Agency examines whether the pollutants discharged by the industry pass through the POTW or interfere with the POTW operation or its chosen sludge disposal practices. In determining whether pollutants pass through a well-operated POTW achieving secondary treatment, the Agency compares the percentage of a pollutant removed by POTW with the percentage removed by direct dischargers applying the best available technology economically achievable. A pollutant is deemed to pass through the POTW when the average percentage removed nationwide by well-operated POTW meeting secondary treatment requirements, is less than the percentage removed by direct dischargers complying with BAT effluent limitations guidelines for that pollutant. (See generally, 46 FR at 9415-16 (January 28, 1981)).

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SECT - XII

This definition of pass through satisfies two competing objectives set by Congress: (1) that standards for indirect dischargers be equivalent to standards for direct dischargers while at the same time, (2) that the treatment capability and performance of the POTW be recognized and taken into account in regulating the discharge of pollutants from indirect dischargers.

The Agency compares percentage removal rather than the mass or concentration of pollutants discharged because the latter would not take into account the mass of pollutants discharged to the POTW from non-industrial sources or the dilution of the pollutants in the POTW effluent to lower concentrations due to the addition of large amounts of non-industrial wastewater.

PRETREATMENT STANDARDS FOR NEW SOURCES

Options for pretreatment of wastewaters from new sources are based on increasing the effectiveness of end-of-pipe treatment technologies. All in-plant changes and applicable end-of-pipe treatment processes have been discussed previously in Sections X and XI. The options for PSNS are the same as the BAT and NSPS options discussed in Sections X and XI, respectively.

A description of each option is presented in Sections X and XI, while a more detailed discussion, including pollutants controlled by each treatment process is presented in Section VII of the General Development Document.

Treatment technologies considered for the PSNS options are:

OPTION A

- o Preliminary treatment with oil skimming (where required)
- o Chemical precipitation and sedimentation
- o Ion exchange

OPTION B

- o Preliminary treatment with oil skimming (where required)
- o Chemical precipitation and sedimentation
- In-process flow reduction of electrolytic cells and calciner scrubber liquor
- o Ion exchange

OPTION C

- o Preliminary treatment with oil skimming (where required)
- o Chemical precipitation and sedimentation
- In-process flow reduction of electrolytic cells and calciner scrubber liquor
- o Multimedia filtration
- o Ion exchange

PSNS OPTION SELECTION - PROPOSAL

EPA proposed that the technology basis of the pretreatment standards for new sources in the primary precious metals and mercury subcategory be equivalent to Option C (in-process flow reduction, oil skimming, chemical precipitation, sedimentation, and multimedia filtration).

The wastewater discharge rates for PSNS were equivalent to the proposed BAT discharge rates. No flow reduction measures for PSNS were considered feasible beyond the recycle proposed for BAT.

PSNS OPTION SELECTION - PROMULGATION

EPA has selected Option C (oil skimming, flow reduction, chemical precipitation and sedimentation, ion exchange and multimedia filtration) as the regulatory approach for pretreatment standards for new sources (PSNS). It is necessary to promulgate PSNS to prevent pass-through of lead, mercury, silver, and zinc. These toxic pollutants are removed by a well-operated POTW at an average of 62 percent, while BAT technology removes approximately 93 percent.

The wastewater discharge rates for promulgated PSNS are identical to the promulgated BAT discharge rates for each waste stream. The PSNS discharge rates are shown in Table XII-1 (page 2280). EPA does not believe that new plants could achieve flow reduction beyond the allowances promulgated for BAT.

We believe that the promulgated PSNS are achievable, and that they are not a barrier to entry of new plants into this subcategory.

REGULATED POLLUTANT PARAMETERS

Pollutants selected for limitation, in accordance with the rationale of Sections VI and X, are identical to those selected for limitation for BAT. It is necessary to promulgate PSNS to prevent the pass-through of lead, mercury, silver, and zinc.

PRETREATMENT STANDARDS

Pretreatment standards are based on the treatable concentrations from the selected treatment technology, (Option C), and the discharge rates determined in Sections X and XI for BAT and NSPS, respectively. These discharge rates are presented in Table XII-1 (page 2280). A mass of pollutant per mass of product (mg/troy ounce or mg/kilogram) allocation is given for each subdivision within the subcategory. This pollutant allocation is based on the product of the treatable concentration from the promulgated treatment (mg/l) and the production normalized wastewater discharge rate (1/troy ounce or 1/kkg). The achievable treatment concentrations for BAT are identical to those for PSNS. PSNS are presented in Table XII-2 (page 2281).

Table XII-1

PSNS WASTEWATER DISCHARGE RATES FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

		PSNS Disch (1/T.O.	(gal/T.O.	
	<u>Waste Stream</u>	or l/kkg)	or gal/kkg)	PNP
1.	Smelter wet air pollution control	1.3		troy ounce of gold and silver smelted
2.	Silver chloride reduction spent solution	0.4		troy ounce of gold reduced in solution
3.	Electrolytic cells wet air pollution control	1-9.8	•	troy ounce of gold refined electrolytically
4.	Electrolyte preparation wet air pollution control	0.05		troy ounce of gold in elec- trolyte produced
5.	Calciner wet air pollution control	22,000		kkg of mercury condensed
6.	Calcine quench water	17,600		kkg of mercury condensed
7.	Calciner stack gas contact cooling water	4,150		kkg of mercury condensed
8.	Condenser blowdown	13,800		kkg of mercury condensed
9.	Mercury cleaning bath water	1,400		kkg of mercury condensed

TABLE XII-2

PSNS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(a) <u>Smelter Wet Air Pollution Control</u> PSNS

Pollutant or pollutant property	Maximum for any one day	Maximum for monthly average
mg/troy ounce of gol	d and silver smel	Lted
Arsenic	1.807	0 806

ALSEILC	T.00/	0.806
Cadmium	0.260	0.104
Chromium	0.481	0.195
Copper	1.664	0.793
*Lead	0.364	0.169
*Mercury	0.195	0.078
Nickel	0.715	0.481
*Silver	0.377	0.156
Thallium	1.820	0.793
*Zinc	1.326	0.546
*Gold	0.130	
1	,	•

(b) <u>Silver Chloride</u> <u>Reduction</u> <u>Spent</u> <u>Solution</u> <u>PSNS</u>

Pollutant	or	Maximum	for	Maximum	for
pollutant	property	any one	dav	monthly	average
-					

mg/troy ounce of silver reduced in solution

Arsenic	0.556	0.248
Cadmium	0.080	0.032
Chromium	0.148	0.060
Copper	0.512	0.244
*Lead	0.112	0.052
*Mercury	0.060	0.024
Nickel	0.220	0.148
*Silver	0.116	0.048
Thallium	0.560	0.244
*Zinc	0.408	0.168
*Gold	0.040	
,		

*Regulated Pollutant

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TABLE XII-2 (Continued)

PSNS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(c) Electrolytic Cells Wet Air Pollution Control PSNS

Pollutant or pollutant property	Maximum for any one day	Maximum for monthly average
mg/troy ounce of silv	ver refined elect	crolytically
Arsenic	27.520	12.280
Cadmium	3.960	1.584
Chromium	7.326	2.970
Copper	25.340	12.080
*Lead	5.544	2.574
*Mercury	2.970	1.188
Nickel	10.890	7.326
*Silver	5.742	2.376
Thallium	27.720	12.080
*Zinc	20.200	8.316
*Gold	1.980	:
(d) Electrolyte Pre	paration Wet Air	Pollution Control PSNS
Pollutant or	Maximum for	Maximum for
pollutant property	any one day	monthly average

mg/troy ounce of silver in electrolyte produced

Arsenic	0.070	0.031
Cadmium	0.010	0.004
Chromium	0.019	0.008
Copper	0.064	0.031
*Lead	0.014	0.007
*Mercury	0.008	0.003
Nickel	0.028	0.019
*Silver	0.015	0.006
Thallium	0.070	0.031
*Zinc	0.051	0.021
*Gold	0.005	

TABLE XII-2 (Continued)

PSNS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(e) <u>Calciner Wet Air Pollution Control</u> PSNS

Pollutant or pollutant property	Maximum for y any one day	Maximum for monthly average
mg/kg (lb/million	lbs) of mercury cor	Idensed
Arsenic	30.580	13.640
Cadmium	4.400	1,760
Chromium	8.140	3.300
Copper	28.160	13.420
*Lead	6.160	2.860
*Mercury	3.300	1.320
Nickel	12.100	8.140
*Silver	6.380	2.640
Thallium	30.800	13.420
*Zinc	22.440	9.240
*Gold	2.200	9.240
(f) <u>Calcine</u> <u>Quenc</u>	ch Water PSNS	
Pollutant or	Maximum for	Maximum for

pollutant property	any one day	monthly average
mg/kg (lb/million lbs	s) of mercury con	ndensed
Arsenic	24.460	10.910
Cadmium	3.520	1.408
Chromium	6.512	2.640
Copper	22.530	10.740
*Lead	4.928	2.288
*Mercury	2.640	1.056
Nickel	9.680	6.512
*Silver	5.104	2.112
Thallium	24.640	10.740
*Zinc	17,950	7.392
*Gold	1.760	

TABLE XII-2 (Continued)

PSNS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(g) Calciner Stack Gas Contact Cooling Water PSNS

Pollutant or	Maximum for	Maximum for
pollutant property	any one day	monthly average

mg/kg (lb/million lbs) of mercury condensed

Arsenic	5.769	2.573
Cadmium	0.830	0.332
Chromium	1.536	0.623
Copper	5.312	2.532
*Lead	1.162	0.540
*Mercury	0.623	0.249
Nickel	2.283	1.536
*Silver	1.204	0.498
Thallium	5.810	2.532
*Zinc	4.233	1.743
*Gold	0.415	

(h) Condenser Blowdown PSNS

Pollutant or	Maximum for any one day	Maximum for monthly average
pollutant property	any one day	monenity avoidage
ng/kg (lb/million lbs) of mercury com	ndensed
Arsenic	19,180	8.556
Cadmium	2.760	1.104
Chromium	5.106	2.070
Copper	17.660	8.418
*Lead	3.864	1.794
*Mercury	2.070	0.828
Nickel	7.590	5.106
*Silver	4.002	1.656
Thallium	19.320	8.418
*Zinc	14.080	5,796
*Gold	1.380	

PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SECT - XII

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TABLE XII-2 (Continued)

PSNS FOR THE PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY

(i) Mercury Cleaning Bath Water

×	The second s		<i>b</i>
Pollutant or	Maximum for		for
pollutant property	any one day	monthiy	average
· · · · · · · · · · · · · · · · · · ·	and the second		
mg/kg (lb/million)	lbs) of mercury con	densed	:
	·	N	
Arsenić	1.946	4.	0.868
Cadmium	0.280	*	0.112
Chromium	0.518	,	0.210
Copper	1.792		0.854
*Lead	0.392		0.182
*Mercury	0.210		0.084
Nickel	0.770	·	0.518
*Silver	0.406		0.168
Thallium	1.960		0.854
*Zinc	1.428		0.588
*Gold	0.140	x - 1	

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PRIMARY PRECIOUS METALS AND MERCURY SUBCATEGORY SECT - XIII

SECTION XIII

BEST CONVENTIONAL POLLUTANT CONTROL TECHNOLOGY

EPA is not promulgating best conventional pollutant control technology (BCT) for the primary precious metals and mercury subcategory at this time.

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NONFERROUS METALS MANUFACTURING POINT SOURCE CATEGORY

DEVELOPMENT DOCUMENT SUPPLEMENT

for the

Secondary Precious Metals Subcategory

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Rebecca Hanmer Acting Assistant Administrator for Water

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May 1989

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X-2	BAT Treatment Scheme for Option B	2593
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SECTION I

SUMMARY

document provides the technical basis for promulgating This effluent limitations based on best practicable technology (BPT) best available technology (BAT) for existing direct and pretreatment standards for indirect existing dischargers, dischargers (PSES), pretreatment standards for new indirect dischargers (PSNS), and standards of performance for new source dischargers (NSPS). direct

promulgation of the second phase of the nonferrous metals After manufacturing regulation, petitioners asked the Courts to review the final rule. After reviewing the petitioners complaints, the and petitioners developed a settlement agreement Agency resolving the complaints raised about the secondary precious metals subcategory. In this agreement, the Agency agreed to propose certain changes to the regulation of this subcategory. These amendments were published proposed on April 28, 1989 (54 FR 18412). Details of these amendments are found at that reference and the proposed changes have been incorporated into this document.

The secondary precious metals subcategory is comprised of 49 plants. Of the 49 plants, four discharge directly to rivers, lakes, or streams; 30 discharge to publicly owned treatment works (POTW); and 15 achieve zero discharge of process wastewater.

EPA first studied the secondary precious metals subcategory to determine whether differences in raw materials, final products, manufacturing processes, equipment, age and size of plants, or required the development of separate effluent water usage, and standards for different segments of the limitations subcategory. This involved a detailed analysis of wastewater discharge and treated effluent characteristics, including the sources and volume of water used, the processes used, the sources of pollutants and wastewaters in the plant, and the constituents wastewaters, including toxic pollutants. As a result, 14 of subdivisions or building blocks have been identified for this subcategory that warrant separate effluent limitations. These include:

- 1. Furnace wet air pollution control,
- 2. Raw material granulation,
- 3. Spent plating solutions,
- 4. Spent cyanide stripping solutions,
- 5. Refinery wet air pollution control,
- 6. Gold solvent extraction raffinate and wash water,
- 7. Gold spent electrolyte,
- 8. Gold precipitation and filtration,
- 9. Platinum precipitation and filtration,

10. Palladium precipitation and filtration,

- 11. Other platinum group metals (PGM) precipitation and filtration,
- 12. Spent solution from PGC salt production,
- 13. Equipment and floor wash, and
- 14. Preliminary treatment.

Several distinct control and treatment technologies (both inplant and end-of-pipe) applicable to the secondary precious metals subcategory were identified. The Agency analyzed both historical and newly generated data on the performance of these technologies, including their nonwater quality environmental impacts and air quality, solid waste generation, and energy requirements. EPA also studied various flow reduction techniques reported in the data collection portfolios (dcp) and plant visits.

Engineering costs were prepared for each of the control and treatment options considered for the subcategory. These costs were then used by the Agency to estimate the impact of implementing the various options on the subcategory. For each control and treatment option that the Agency found to be most effective and technically feasible in controlling the discharge of pollutants, the number of potential closures, number of employees affected, and impact on price were estimated. These results are reported in a separate document entitled "The Economic Impact Analysis of Effluent Limitations and Standards for the Nonferrous Metals Manufacturing Industry."

After examining the various treatment technologies, the Agency has identified BPT to represent the average of the best existing technology. Metals removal based on chemical precipitation and sedimentation technology is the basis for the BPT limitations. Steam stripping was selected as the technology basis for ammonia limitations. Cyanide precipitation was selected as the technology basis for cyanide limitations. Ion exchange was selected as the basis for gold, platinum and palladium limitations. To meet the BPT effluent limitations based on this technology, the secondary precious metals subcategory is expected to incur a capital and annual cost. These costs cannot be disclosed because the data on which they are based have been claimed to be confidential.

For BAT, the Agency has built upon the BPT technology basis by adding in-process control technologies which include recycle of process water from air pollution control waste streams and recycle of raw material granulation water. Filtration is added as an effluent polishing step to the end-of-pipe treatment scheme prior to ion exchange. To meet the BAT effluent limitations based on this technology, the secondary precious metals subcategory is expected to incur a capital and annual cost. These costs cannot be disclosed because publication of the costs could reveal the data on which they are based. These data have been claimed confidential.

NSPS is equivalent to BAT. In selecting NSPS, EPA recognizes

that new plants have the opportunity to implement the best and most efficient manufacturing processes and treatment technology. As such, the technology basis of BAT has been determined as the best demonstrated technology.

The technology basis for PSES is equivalent to BAT. To meet the pretreatment standards for existing sources, the secondary precious metals subcategory is estimated to incur a capital cost of \$1,809,400 and an annual cost of \$1,100,500. For PSNS, the Agency selected end-of-pipe treatment and in-process flow reduction control techniques equivalent to NSPS.

Based on comments received after proposal, the Agency believes that it may be necessary for some facilities to use sulfide order to achieve the promulgated effluent polishina in limitations because of high zinc concentrations or complexing problems. Because the Agency believes that these situations will be the exception, rather than the rule, sulfide polishing is not specifically included as part of the model technology on which effluent limitations and performance standards are based. The Agency has, however, evaluated the cost associated with the use of sulfide polishing at secondary precious metals plants. After performing this evaluation, the Agency has concluded that sulfide polishing will result in a very small (less than 5 percent) incremental increase in wastewater treatment costs at a typical secondary precious metals facility.

The best conventional technology (BCT) replaces BAT for the control of conventional pollutants. BCT is not being promulgated because the methodology for BCT has not yet been finalized.

The mass limitations and standards for BPT, BAT, NSPS, PSES, and PSNS are presented in Section II.

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SECTION II

CONCLUSIONS

EPA has divided the secondary precious metals subcategory into 14 subdivisions or building blocks for the purpose of effluent limitations and standards. These building blocks are:

- 1. Furnace wet air pollution control,
- 2. Raw material granulation,
- 3. Spent plating solutions,
- 4. Spent cyanide stripping solutions,
- 5. Refinery wet air pollution control,
- 6. Gold solvent extraction raffinate and washwater,
- 7. Gold spent electrolyte,
- 8. Gold precipitation and filtration,
- 9. Platinum precipitation and filtration,
- 10. Palladium precipitation and filtration,
- 11. Other platinum group metals precipitation and filtration,
- 12. Spent solution from PGC salt production,
- 13. Equipment and floor wash, and
- 14. Preliminary treatment.

BPT is promulgated based on the performance achievable by the application of chemical precipitation and sedimentation (lime and settle) technology, and ion exchange end-of-pipe treatment, along with preliminary treatment consisting of ammonia steam stripping and cyanide precipitation for selected waste streams. The following BPT effluent limitations are promulgated:

(a) Furnace Wet Air Pollution Control BPT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	<u> </u>
mg/troy ounce of pr incinerated or smel		cluding silver,	
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals Total suspended solids pH	136.400 20.820 104.800 9,571.000 21.54 2,944.000 Within the ra	71.800 8.616 43.800 4,207.000 1,400.000 nge of 7.5 to 10.0 11 times	

(b) Raw Material Granulation BPT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	
mg/troy ounce of pr raw material	ecious metal in t	he granulated	
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals Total suspended solids	12.050 1.839 9.256 845.100 1.903 259.900	6.340 0.761 3.867 371.500 123.600	
рн	Within the ra at a	nge of 7.5 to 10.0 11 times	· · ·

(c) Spent Plating Solutions BPT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/liter of spent p raw material	lating solution us	
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals Total suspended solids	1.900 0.290 1.460 133.300 0.300 41.000	1.000 0.120 0.610 58.600 19.500
рн		ange of 7.5 to 10.0 times

(d) <u>Spent</u> Cyanide	Stripping Solutions	DF1
Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of go cyanide stripping	ld produced by	
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals Total suspended solids	7.030 1.073 5.402 493.200 1.110 151.700	3.700 0.444 2.257 216.800 72.150
рН	Within the range at all	e of 7.5 to 10.0 times

(d) Spent Cyanide Stripping Solutions BPT

(e) <u>Refinery Wet</u> <u>Air</u> <u>Pollution</u> <u>Control¹ BPT</u>

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
mg/troy ounce of pre		ncluding silver,
produced in refinery		
Copper	39.900	21.000
Cyanide (total)	6.090	2.520
Zinc	30.660	12.810
Ammonia (as N)	2,799.000	1,231.000
Combined metals	6.300	
Fotal suspended	861.000	409.500
solids		
HC	Within the	range of 7.5 to 10.0
		times

¹This allowance applies to either acid or alkaline wet air pollution control scrubbers. If both acid and alkaline wet air pollution control scrubbers are present in a particular facility the same allowance applies to each. SECONDARY PRECIOUS METALS SUBCATEGORY SECT - II

(f) Gold Solvent Extraction Raffinate and Wash Water BPT

Pollutant or Pollutant Property	Maximum for Any One Day M	Maximum for Aonthly Average	
mg/troy ounce of g	old produced by sol	lvent extraction	
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals Total suspended solids	1.197 0.183 0.920 83.980 0.189 25.830	0.630 0.076 0.38? 36.920 12.290	
pH		nge of 7.5 to 10.0 1 times	

(g) <u>Gold Spent Electrolyte</u> BPT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of gol	d produced by elec	trolysis
Copper	0.017	0.009
Cyanide (total)	0.003	0.001
Zinc	0.013	0.005
Ammonia (as N)	1.160	0.510
Combined metals	0.003	
Total suspended solids	0.357	0.170
pH Within the range of 7. at all times		

(h) Gold Precipitation and Filtration BPT

Pollutant or Pollutant Propert	=y	Maximum fo Any One Da		Maxin Monthly				
mg/troy ounce of	gold p	precipitate	đ			· · · · ·	· .	 - 1
Copper Cyanide`(total) Zinc Ammonia (as N) Combined metals Total suspended solids		8.360 1.276 6.242 586.500 1.320 180.400		0 2 257	.400 .528 .684 .800 .800		•	
pH at		Within all	the	range	of	7.5	to	10.0 times

(i) <u>Platinum Precipitation</u> and <u>Filtration</u> BPT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average		
mg/troy ounce of platinum precipitated				
Copper	9.880	5.200		
Cyanide (total)	1.508	0.624		
Zinc	7.592	3.172		
Ammonia (as N)	693.200	304.700		
Combined metals				
Total suspended solids	213.200	101.400		
pH Within the range of 7.5 to 10 at all times				

(j) <u>Palladium</u> <u>Precipitation</u> and <u>Filtration</u> BPT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	•
mg/troy ounce of pal	ladium precipitat	ted	•
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals Total suspended solids	11.400 1.740 8.760 799.800 1.800 246.0	6.000 0.720 3.660 351.600 117.000	
pH	Within the r at all	ange of 7.5 to 10.0 times	,

(k) Other Platinum Group Metals Precipitation and Filtration BPT

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
mg/troy ounce of other precipitated	platinum group	metals
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	9.880 1.508 7.592 693.200 1.560	5.200 0.624 3.172 304.700
Total suspended	213.200 101.400	
solids	Within the range of 7.5 to 10.0	
pH	at all times	

PRECIOUS METALS SUBCATEGORY SECONDARY SECT - II

(1) Spent Solution fro	om PGC Salt Prod	uction BPT	
Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Averac	
mg/troy ounce of gold	contained in PGC	product	
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals Total suspended solids pH at	1.710 0.261 1.314 120.000 0.270 36.900 Within the all	0.900 0.108 0.549 52.740 17.550 range of 7.5	to 10.0 times
(m) Equipment and Floo	······································		
Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Averag	

mg/troy ounce of precious metals, including silver, produced in refinery

Copper	0 000	0 000
Cyanide (total)	0.000	0.000
Zinc	0.000	0.000
Ammonia (as N)	0.000	0.000
Combined metals	0.000	1
Total suspended solids	0.000	0.000
рН		ge of 7.5 to 10.0 L times

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(n) Preliminary treatment BPT

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
mg/troy ounce of pr	ecious metals pro	duced in refinery
Copper	95.000	50.000
Cyanide (total)	14.500	6.000
Zinc	73.000	30.500
Ammonia (as N)	6665.000	2930.000
Combined metals	15.000	
Total suspended	2050.000	975.000
solids pH		nge of 7.5 to 10.0 11 times

BAT is promulgated based on the performance achievable by the application of chemical precipitation, sedimentation, and multimedia filtration (lime, settle, and filter) technology, ion exchange end-of-pipe treatment, and in-process flow reduction methods, along with preliminary treatment consisting of ammonia steam stripping and cyanide precipitation for selected waste streams. The following BAT effluent limitations are promulgated:

(a) <u>Furnace Wet Air Pollution Control BAT</u>

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	
mg/troy ounce of pr incinerated or smel	ecious metals, ind ted	cluding silver,	a
Copper	5.760	2.745	•
Cyanide (total)	0.900	0.360	
Zinc	4.590	1.890	
Ammonia (as N)	599.900	263.700	

(b) Raw Material Granulation BAT

Pollutant or Pollutant Propert	Maximum for y Any One Day	Maximum for Monthly Average	
mg/troy ounce of raw material	precious metals in in	the granulated	
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	0.819 0.128 0.653 85.310 0.192	0.390 0.051 0.269 37.500	

(c) Spent Plating Solutions BAT

mg/liter of spent plating solution used as a raw material

(d) Spent Cyanide Stripping Solutions BAT

Pollutant or	Maximum for	Maximum for
		Manth la Amorana
Pollutant Property	Any One Day	Monthly Average
· • -		ε.

mg/troy ounce of gold produced by cyanide stripping

Copper	4.736	2.257
Cyanide (total)	0.740	0.296
Zinc	3.774	1.554
Ammonia (as N)	493.200	216.800
Combined metals	1.110	

(e) <u>Refinery Wet</u> <u>Air</u> <u>Pollution</u> <u>Control¹ BAT</u>

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	•
mg/troy ounce of pr produced in refiner		cluding silver,	
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	1 280 0.200 1 020 133.300 0.300	0.610 0.080 0.420 58.600	

¹This allowance applies to either acid or alkaline wet air pollution control scrubbers. If both acid and alkaline wet air pollution control scrubbers are present in a particular facility the same allowance applies to each.

(f) Gold Solvent Extraction Raffinate and Wash Water

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of gold	produced by so	olvent extraction
	r.	· · ·
Copper	0.806	0.384
Cyanide (total)	0.126	0.050
Zinc	0.6?3	0.265
Ammonia (as N)	83.980	36.920
Combined metals	0.189	

(g) Gold Spent Electrolyte BAT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	
mg/troy ounce of gold	l produced by e	lectrolysis	,
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	0.011 0.002 0.009 1.160 0.003	0.005 0.001 0.004 0 510	

SECT - II

(h) Gold Precipitation and Filtration BAT

Pollutant or	Maximum for	Maximum for	
Pollutant Property	Any One Day	Monthly Average	
mg/troy ounce of go	ld precipitated	·····	
Copper	5.632	2.684	
Cyanide (total)	0.880	0.352	
Zinc	4.488	1.848	
Ammonia (as N)	586.500	257.800	
Combined metals	1.320	* 	

(i) Platinum Precipitation and Filtration BAT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Avera	
mg/troy ounce of pla	tinum precipitated		······································
Copper Cyanide (total) Zinc Ammonia (as N) 1.560	6.656 1.040 5.304 693.200	3.172 0.416 2.184 304.700 Co	ombined metals

(j) Palladium Precipitation and Filtration BAT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	
mg/troy ounce of pal	ladium precipita	ted	
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	7.680 1.200 6.120 799.800 1.800	3.660 0.480 2.520 351.600	

(k)	Other Platinum	Group	Metals	Precipitation	<u>and</u>
• •	Filtration BAT		· ·	•	

		1	
Pollutant or	Maximum for	Maximum for	
Pollutant Property	Any One Day	Monthly Average	
Pollucane Propercy	Mily one buy		
	v		
mg/troy ounce of other	nlatinum grou	p metals	
precipitated	pracessan gree	E	
precipitated			
O	6.656	3.172	
Copper		0.416	
Cyanide (total)	1.040		
Zinc	5.304	2.184	
Ammonia (as N)	693.200	304.700	
Combined metals	1.560		
		· · · · · · · · · · · · · · · · · · ·	
(1) Spent Solution fr	om PGC Salt Pr	oduction BAT	
Pollutant or	Maximum for	Maximum for	
Pollutant Property	Any One Day	Monthly Average	
		• • •	
mg/troy ounce of gold	contained in P	GC product	
57 2 0			
Copper	1.152	0.549	
Cyanide (total)	0.180	0.072	
Zinc	0.918	0.378	
Ammonia (as N)	120.000	52.740	
Combined metals	0.270		
Combined metals	0.2/0		
· · · · · · · · · · · · · · · · · · ·			
•			
(m) Equipment and Flo	or Wash BAT		
(m) Equipment and Flo	Masin DAI	с , , , , , , , , , , , , , , , , , , ,	
Pollutant or	Maximum for	Maximum for	
		Monthly Average	
Pollutant Property	Any One Day	Monthly Average	
ma /brow owners of prog	ous metals in	cluding silver.	
mg/troy ounce of preci	ous metais, in	Cruating Street,	
produced in refinery			
	,	0.000	
Copper	0.000	0.000	
Cyanide (total)	0.000	0.000	
Zinc	0.000	0 000	
Ammonia (as N)	0.000	0.000	
Combined metals	0.000	~	
COMPTING MCCATD			

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - II

(n) Preliminary treatment BAT

Pollutant or Pollutant Property	Maximum for Any One Day		imum for ly Average	
mg/troy ounce of t through this opera		metals p	roduced	
Copper Cyanide (total) Zinc Combined metals Ammonia (as N)	64.000 10.000 51.000 15.000 6665.000		30.500 4.000 21.000 2930.000	

NSPS are promulgated based on the performance achievable by the application of chemical precipitation, sedimentation, and multimedia filtration (lime, settle, and filter) technology, ion exchange end-of-pipe treatment, and in-process flow reduction control methods, along with preliminary treatment consisting of ammonia steam stripping and cyanide precipitation for selected waste streams. The following effluent standards are promulgated for new sources:

(a) Furnace Wet Air Pollution Control NSPS

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
	· · · · · · · · · · · · · · · · · · ·	
mg/troy ounce of pro	acious metals inc	luding silver
incinerated or smelt		ruding silver,
indinerated of Bildre		
Copper	5.760	2.745
Cyanide (total)	0.900	0.360
Zinc	4.590	1.890
Ammonia (as N)	599.900	263.700
Combined metals	1.350	
Total suspended	67.500	54.000
solids		
Н		ge of 7.5 to 10.0
	at al	l times

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(b) Raw Material Granulation NSPS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	
mg/troy ounce of pred raw material	cious metals in	the granulated	
Copper	0.819	0.390	
Cyanide (total)	0.128	0.051	
Zinc	0.653	0.269	
Ammonia (as N)	85.310	37.500	
Combined metals	0.192		
Total suspended solids	9.600	7.680	; ,
рН		inge of 7.5 to 10.0 Il times	

(c) Spent Plating Solutions NSPS

Pollutant or	Maximum for	Maximum for Monthly Average
Pollutant Property	Any One Day	Monthly Average
mg/liter of spent	plating solution us	sed as a raw material
Copper	1.280	0.610
Cyanide (total)	0.200	0.080
Zinc	1.020	0.420
Ammonia (as N)	133.300	58.600
Combined metals	0.300	
Total suspended solids	15.000	12.000
рн		nge of 7.5 to 10.0 11 times

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - II

	$(h, f) \in \mathcal{F}$	
(d) <u>Spent</u> <u>Cyanide</u>	Stripping Solution	s NSPS
Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of gol	d produced by cya	nide stripping
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals Total suspended solids	4.736 0.740 3.774 493.200 1.110 55.500	2.257 0.296 1.554 216.800 44.400
pH	Within the range at all i	ge of 7.5 to 10.0 times
(e) <u>Refinery Wet Ai</u>	r Pollution Contro	ol ^l NSPS
Pollutant or Pollutant Property	Maximum for Any One Day Mo	Maximum for onthly Average
mg/troy ounce of pre produced in refinery	cious metals, incl	Luding silver,
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals Total suspended solids	1.280 0.200 1.020 133.300 0.300 15.000	0.610 0.080 0.420 58.600 12.000
	Within the rang at all	ge of 7.5 to 10.0 times

¹This allowance applies to either acid or alkaline wet air pollution control scrubbers. If both acid and alkaline wet air pollution control scrubbers are present in a particular facility the same allowance applies to each.

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT - II

(f) old Solvent Extraction Raffinate and Wash Water NSPS

Pollutant or	Maximum for	Maximum for	
Pollutant Property	Any One Day	Monthly Average	
mg/troy ounce of gold	produced by so	lvent extraction	
Copper	0.806	0.384	
Cyanide (total)	0.126	0.050	
Zinc	0.643	0.265	
Ammonia (as N)	83.980	36.920	
Combined metals	0.189		
Total suspended	9.450	7.560	
solids pH at	Within th all	e range of 7.5	to 10.0 times

 $\{r, f, r\}$

(g) Gold Spent Electrolyte NSPS

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
mg/troy ounce of gold	d produced by el	ectrolysis
Copper	0.011	0.005
Cyanide (total)	0.002	0.001
Zinc	0.009	0.004
Ammonia (as N)	1.160	0.510
Combined metals	0.003	
Total suspended	0.131	0.104
solids		
pH	Within the	range of 7.5 to 10.
ř	at al	l times

(h) Gold Precipitation and Filtration NSPS

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Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
		· -· -· -· -· -· ·· ·· ·· ·· ·· ·· ·· ··
mg/troy ounce of gold	procipitated	
mg/troy ounce or gord	precipicaced	
Copper	5.632	2.684
Cyanide (total)	0.880	0.352
Zinc	4.488	1.848
Ammonia (as N)	586.500	257.800
Combined metals	1.320	
Total suspended	66.000	52.800
solids		
pH	Within the	range of 7.5 to 10.0
· · · · · · · · · · · · · · · · · · ·	at all	times
		. ,

(i) Platinum Precipitation and Filtration NSPS

	•		
Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	
mg/troy ounce of pl	atinum precipitate	ed	
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals Total suspended solids	6.656 1.040 5 304 693.200 1.560 78.000	3.172 0.416 2.184 304.700 62.400	•
pH		ige of 7.5 to 0.0 ill times	

(J) Palladium Precipitation and Filtration NSPS

Maximum for Any One Day	Maximum for Monthly Average	
ladium precipita.	ted	
7.680 1.200 6.120 799.800 1.800 90.000	3.660 0.480 2.520 351.600 72.000	• •
Within the ra at a	nge of 7.5 to 10.0 11 times	<u> </u>
	Any One Day ladium precipita 7.680 1.200 6.120 799.800 1.800 90.000 Within the ra	Any One Day Monthly Average ladium precipitated 7.680 3.660 1.200 0.480 6.120 2.520 799.800 351.600 1.800

(k) Other Platinum Group Metals Precipitation and Filtration NSPS

		-	
Pollutant or Property Any	Maximum One Day Mon	for Maximum hthly Average	for Pollutant
mg/troy ounce of precipitated	other platinum	group metals	
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals Total suspended solids pH	6.656 1.040 5.304 693.200 1.560 78.000 Within	3.172 0.416 2.184 30?.700 62.400 the range of 7.5 to	o 10.0
		at all times	

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - II

(1) Spent Solutions from PGC Salt Production NSPS Maximum for Pollutant or Maximum for Pollutant Property Eny One Day Monthly Average mg/troy ounce of gold contained in PGC product 1.152 0.549 Copper Cyanide (total) 0.180 0.072 0.918 0.378 Zinc Ammonia (as N) 120.000 52.740 Combined metals 0.270 ~ - - -13.500 10.800 Total suspended solids Within the range of 7.5 to 10.0 рĦ at all times (m) Equipment and Floor Wash NSPS Pollutant or Maximum for Maximum for Pollutant Property Any One Day Monthly Average

mg/troy ounce of precious metals, including silver, produced in refinery

1		
Copper	0.000	0.000
Cyanide (total)	0.000	0.000
Zinc	0.000	0.000
Ammonia (as N)	0.000	0.000
Combined metals	0.000	٢
Total suspended solids	0.000	0.000
pH		ge of 7.5 to 10.0 L times

SECT - II

(n) Preliminary treatment NSPS

	Maximum for Any One Day	Maximum for Monthly Average	
mg/troy ounce of tota through this operatio	l precious me n	tals produced	
Copper Cyanide (total) Zinc Combined metals Ammonia (as N) Total suspended solids pH	64.000 10.000 51.000 15.000 6665.000 750.000 Within the at	30.500 4.000 21.000 2930.000 600.000 range of 7.5 to 10 all times).0

PSES are promulgated based on the performance achievable by the application of chemical precipitation, sedimentation, and multimedia filtration (lime, settle, and filter) technology, ion exchange end-of-pipe treatment, and in-process flow reduction control methods, along with preliminary treatment consisting of ammonia steam stripping and cyanide precipitation for selected waste streams. The following pretreatment standards are promulgated for existing sources:

(a) Furnace Wet Air Pollution Control PSES

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of preci incinerated or smelted	ous metals, incl	uding silver,
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	5.760 0.900 4.590 599.900 1.350	2.745 0.360 1.890 263.700

(b) Raw Material Granulation PSES

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
· · · · · · · · · · · · · · · · ·		
	• • • • • • • • • • •	
ng/troy ounce of pre	cious metais in s	the granulated
raw material		
Copper	0.819	0.390
Cyanide (total)	0.128	0.051
	0.653	0.269
Zinc		
Ammonia (as N)	85.310	37.500
Combined metals	0.192	
	· · · · · · · · · · · · · · · · · · ·	
(c) Spent Plating S	olutions PSES	
		·
Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
	·	
ng/libog of group pl	sting colution w	sed as a raw material
mg/liter of spent pr	ating solution a	sed as a law material
Copper	1.280	0.610
Cyanide (total)	0.200	0.080
Zinc	1.020	0.420
Ammonia (as N)	133.300	58.600
	0.300	50:000
Combined metals	0.300	
	· · · · · · · · · · · · · · · · · · ·	
(d) <u>Spent</u> <u>Cyanide</u> <u>S</u>	tripping Solution	ns PSES
Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
· · · · · · · · · · · · · · · · · · ·		
mg/troy ounce of gol	d produced by	· · ·
cyanide stripping		
-	4 776	2 257
Copper	4.736	2.257
Cyanide (total)	0.740	0.296
Zinc	3.774	1.554
Ammonia (as N)	493.200	216.800
Combined metals	1.110	· · · · · · · · · · · · · · · · · · ·

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SECT - II

(e) <u>Refinery Wet</u> <u>Air</u> <u>Pollution</u> <u>Control¹</u> <u>PSES</u>

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of pr produced in refiner	ecious metals, ind Y	cluding silver,
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	1.280 0.200 1.020 133.300 0.300	0.610 0.080 0.420 58.600

¹This allowance applies to either acid or alkaline wet air pollution control scrubbers. If both acid and alkaline wet air pollution control scrubbers are present in a particular facility the same allowance applies to each.

(f) Gold Solvent Extraction Raffinate and Wash Water PSES

Pollutant	or	Maximum	for	Maximum for	,
Pollutant		Any One	Day	Monthly Average	

mg/troy ounce of gold produced by solvent extraction

Copper0.806Cyanide (total)0.126Zinc0.643Ammonia (as N)83.980Combined metals0.189	0.384 0.050 0.265 36.920
--	-----------------------------------

(g) Gold Spent Electrolyte PSES

Pollutant	or	Maximum	for	Maximum for
Pollutant		Any One	Day	Monthly Average

mg/troy ounce of gold produced by electrolysis

Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	0.011 0.002 0.009 1.160 0.003	0.005 0.001 0.004 0.510
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SECONDARY PRECIOUS

(h) Gold Precipitation and Filtration PSES

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of gold	precipitated	
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	5.632 0.880 4.488 586.500 1.320	2.684 0.352 1.848 257.800

(i) Platinum Precipitation and Filtration PSES

Pollutant or Pollutant Property	Maximum for Any One Day	
mg/troy ounce of pla	atinum precipitated	
Copper	6.656	3.172
Cyanide (total)	1.040	0.416
Zinc	5.304	2.184
	693.200	304.700
Ammonia (as N)		

(j) Palladium Precipitation and Filtration PSES

2			
Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	· · · · ·
mg/troy ounce of pal	ladium precipita	ited	
Copper	7.680	3.660	
Cyanide (total)	1.200	0.480	
Zinc	6.120	2.520	
Ammonia (as N)	799.800	351.600	
Combined metals	1.800		
		4. 1	

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(k) Other Platinum Group Metals Precipitation and Filtration PSES

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of oth precipitated	ner platinum grou	up metals
Copper	6.656	3.172
Cyanide (total)	1.040	0.416
Zinc	5.304	2.184
Ammonia (as N)	693.200	304.700
Combined metals	1.560	

ollutant or		or Maximu	m for	Polluta
roperty Any On	Maximum fo e Day Montl	nly Average		
				. <u></u>
g/troy ounce of go	ld contained in	PGC product		
g/croy ounce or go	ra concarnea r.	. roo produoc		
opper	1.152	0.54		
yanide (total)	0.180	0.07		
inc	0.918	0.37		
mmonia (as N)	120.000	52.74	0	
ombined metals	0.270		— .	
m) Equipment and	Floor Wash PSE	S		
() <u>Equipmente</u> dite		· · · · ·		
ollutant or	Maximum for			•
ollutant ProPerty	Any One Day	y Monthly A	verage	•
	· · · · · · · · · · · · · · · · · · ·	······································		
g/troy ounce of pr	ecious metals,	including silv	er,	
roduced in refiner	y		,	
		·		•
opper	0.000	0.00		
yanide (total)	0.000	0.00		
inc	0.000	0.00		
mmonia (as N)	0.000	0.00	0	
ombined metals	0.000		- *	
n) <u>Preliminary</u> tr	eatment PSES			
ollutant or	Maximum for	Maximum for	<u> </u>	
ollutant Property	Any One Day	Monthly Avera		1
			с у 	
g/troy ounce of to		etais produced		
hrough this operat	TOIL			
opper	64.000	30.50	0	
yanide (total)	10.000	4.00		· · ·
inc	51.000	21.00		
ombined metals	15.000			
		2930.00		

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PSNS are promulgated based on the performance achievable by the application of chemical precipitation, sedimentation, and multimedia filtration (lime, settle, and filter) technology, ion exchange end-of-pipe treatment, and in-process flow reduction control methods, along with preliminary treatment consisting of ammonia steam stripping and cyanide precipitation for selected waste streams. The following pretreatment standards are promulgated for new sources:

(a) Furnace Wet Air Pollution Control PSNS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of pre incinerated or smelt		cluding silver,
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	5.760 0.900 4.590 599.900 1.350	2.745 0.360 1.890 263.700

(b) Raw Material Granulation PSNS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	
mg/troy ounce of pre	cious metals in	the granulated raw	material
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	0.819 0.128 0.654 85.310 0.192	0.390 0.051 0.269 37.500	

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - II

(c) Spent Plating Solutions PSNS

Pollutant or	Maximum for	Maximum for	
Pollutant Property	Any One Day	Monthly Average	
g/liter of spent pla	ating solution us	sed as a	
aw material	-	•	
			• *
Copper	1.280	0.610	
Cyanide (total)	0.200	0.080	
linc	1.020	0.420	
Ammonia (as N)	133.300	58.600	•
Combined metals	0.300		
	* · · · ·		
(d) Spent Cyanide S	tripping Solution	n's PSNS	
Pollutant or	Maximum for	Maximum for	
Pollutant Property	Any One Day	Monthly Average	
· · · · · · · · · · · · · · · · · · ·			
	×		•
·			
	d produced by	н. Э	and the second se
	d produced by		
cyanide stripping		2 257	
ng/troy ounce of gold cyanide stripping Copper	4.736	2.257	
cyanide stripping Copper Cyanide (total)	4.736 0.740	0.296	
cyanide stripping Copper Cyanide (total) Zinc	4.736 0.740 3.774	0.296 1.554	
cyanide stripping Copper Cyanide (total) Zinc Ammonia (as N)	4.736 0.740 3.774 493.200	0.296	
cyanide stripping Copper Cyanide (total)	4.736 0.740 3.774	0.296 1.554	
cyanide stripping Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	4.736 0.740 3.774 493.200 1.110	0.296 1.554 216.800	· · · · · · · · · · · · · · · · · · ·
cyanide stripping Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	4.736 0.740 3.774 493.200	0.296 1.554 216.800 col ^l PSNS	
cyanide stripping Copper Cyanide (total) Zinc Ammonia (as N) Combined metals (e) <u>Refinery Wet Ai</u>	4.736 0.740 3.774 493.200 1.110 <u>r Pollution Conc</u> Maximum for	0.296 1.554 216.800 <u>col¹ PSN</u> S <u>Maximum for</u>	· · · · · · · · · · · · · · · · · · ·
cyanide stripping Copper Cyanide (total) Zinc Ammonia (as N) Combined metals (e) <u>Refinery Wet Ai</u> Pollutant or	4.736 0.740 3.774 493.200 1.110 r Pollution Conc	0.296 1.554 216.800 col ^l PSNS	· · · · · · · · · · · · · · · · · · ·
cyanide stripping Copper Cyanide (total) Zinc Ammonia (as N) Combined metals (e) <u>Refinery Wet Ai</u> Pollutant or	4.736 0.740 3.774 493.200 1.110 <u>r Pollution Conc</u> Maximum for	0.296 1.554 216.800 <u>col¹ PSN</u> S <u>Maximum for</u>	· · · · · · · · · · · · · · · · · · ·
Cyanide stripping Copper Cyanide (total) Zinc Ammonia (as N) Combined metals (e) <u>Refinery Wet Ai</u> Pollutant or Pollutant Property	4.736 0.740 3.774 493.200 1.110 <u>r Pollution Conc</u> <u>Maximum for</u> Any One Day	0.296 1.554 216.800 <u>col¹ PSNS</u> <u>Maximum for</u> Monthly Average	
cyanide stripping Copper Cyanide (total) Sinc Ammonia (as N) Combined metals (e) <u>Refinery Wet Ai</u> Collutant or Collutant Property	4.736 0.740 3.774 493.200 1.110 <u>r Pollution Conc</u> <u>Maximum for</u> Any One Day cious metals, in	0.296 1.554 216.800 <u>col¹ PSNS</u> <u>Maximum for</u> Monthly Average	
cyanide stripping Copper Cyanide (total) Sinc Ammonia (as N) Combined metals (e) <u>Refinery Wet Ai</u> Collutant or Collutant Property	4.736 0.740 3.774 493.200 1.110 <u>r Pollution Conc</u> <u>Maximum for</u> Any One Day cious metals, in	0.296 1.554 216.800 <u>col¹ PSNS</u> <u>Maximum for</u> Monthly Average	
cyanide stripping Copper Cyanide (total) Zinc Ammonia (as N) Combined metals (e) <u>Refinery Wet Ai</u> Pollutant or Pollutant Property mg/troy ounce of pre produced in refinery	4.736 0.740 3.774 493.200 1.110 <u>r Pollution Conc</u> <u>Maximum for</u> Any One Day cious metals, in	0.296 1.554 216.800 <u>col¹ PSNS</u> <u>Maximum for</u> Monthly Average cluding silver,	
cyanide stripping Copper Cyanide (total) Cyanide (total) Zinc Ammonia (as N) Combined metals (e) <u>Refinery Wet Ai</u> (e) <u>Refinery Wet Ai</u> Pollutant or Pollutant or Pollutant Property mg/troy ounce of pre produced in refinery Copper	4.736 0.740 3.774 493.200 1.110 <u>r Pollution Conc</u> <u>Maximum for</u> Any One Day cious metals, in 1.280	0.296 1.554 216.800 <u>col¹ PSNS</u> <u>Maximum for</u> Monthly Average cluding silver, 0.610	
cyanide stripping Copper Cyanide (total) Zinc Ammonia (as N) Combined metals (e) <u>Refinery Wet Ai</u> Pollutant or Pollutant property mg/troy ounce of pre produced in refinery Copper Cyanide (total)	4.736 0.740 3.774 493.200 1.110 <u>r Pollution Conc</u> <u>Maximum for</u> Any One Day cious metals, in 1.280 0.200	0.296 1.554 216.800 <u>coll PSNS</u> <u>Maximum for</u> Monthly Average cluding silver, 0.610 0.080	
cyanide stripping Copper Cyanide (total) Zinc Ammonia (as N) Combined metals (e) <u>Refinery Wet Ai</u> Pollutant or Pollutant property mg/troy ounce of pre produced in refinery Copper Cyanide (total) Zinc	4.736 0.740 3.774 493.200 1.110 <u>r Pollution Conc</u> <u>Maximum for</u> Any One Day cious metals, in 1.280 0.200 1.020	0.296 1.554 216.800 <u>coll PSNS</u> <u>Maximum for</u> Monthly Average cluding silver, 0.610 0.080 0.420	
cyanide stripping Copper Cyanide (total) Zinc Ammonia (as N) Combined metals (e) <u>Refinery Wet Ai</u> Pollutant or Pollutant property mg/troy ounce of pre produced in refinery Copper Cyanide (total)	4.736 0.740 3.774 493.200 1.110 <u>r Pollution Conc</u> <u>Maximum for</u> Any One Day cious metals, in 1.280 0.200	0.296 1.554 216.800 <u>coll PSNS</u> <u>Maximum for</u> Monthly Average cluding silver, 0.610 0.080	

¹This allowance applies to either acid or alkaline wet air pollution control scrubbers. If both acid and alkaline wet air pollution control scrubbers are present in a particular facility the same allowance applies to each.

SECT - II

(f) Gold Solvent Extraction Raffinate and Wash Water PSNS

Pollutant or Pollutant Property	Maximum for Any One Day	
mg/troy ounce of gold	produced by	solvent extraction
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	0.806 0.126 0.643 83.980 0.189	0.384 0.050 0.265 36.920

(g) Gold Spent Electrolyte PSNS

Pollutant or	Maximum	for Maxim	um for
Pollutant Proper	ty Any One	Day Monthly	v Average

mg/troy ounce of gold produced by electrolysis

Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	0.011 0.002 0.009 1.160 0.003	0.005 0.001 0.004 0.510
Ammonia (as N)	1.160	

(h) Gold Precipitation and Filtration PSNS

Maximum for	Maximum for
Any One Day	Monthly Average
d precipitated	
5.632	2.684
0.880	0.352
4.488	1.848
586.500	257 . 80 0
1.320	
	Any One Day d precipitated 5.632 0.880 4.488 586.500

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	· ·
mg/troy ounce of plat	inum precipitat	ed	
Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	6.656 1.040 5.304 693.200 1.560	3.172 0.416 2.184 304.700	

(i) Platinum Precipitation and Filtration PSNS

(j) Palladium Precipitation and Filtration PSNS

Pollutant or	Maximum for	Maximum for	
Pollutant Property	Any One Day	Monthly Average	
mg/troy ounce of pall	Ladium precipita	ted	
Copper	7.680	3.660	
Cyanide (total)	1.200	0.480	
Zinc	6.120	2.520	÷.
Ammonia (as N)	799.800	351.600	
Combined metals	1.800		
,		·	
	· · · · · · · · · · · · · · · · · · ·		
	· ,		•
(k) Other Platinum C	Group Metals Pre	cipitation and	•
(k) Other Platinum O Filtration PSNS	Group Metals Pre	cipitation and	۰ ۰
Filtration PSNS	Group Metals Pre Maximum for	cipitation and Maximum for	•
(k) <u>Other Platinum (</u> <u>Filtration</u> PSNS Pollutant or Pollutant Property			• • • •
Filtration PSNS Pollutant or Pollutant Property mg/troy ounce of othe	Maximum for Any One Day	Maximum for Monthly Average	•
Filtration PSNS Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average	•
Filtration PSNS Pollutant or Pollutant Property mg/troy ounce of othe precipitated	Maximum for Any One Day	Maximum for Monthly Average	
Filtration PSNS Pollutant or Pollutant Property mg/troy ounce of othe precipitated Copper	Maximum for Any One Day er platinum grou 6.656	Maximum for Monthly Average p metals	
Filtration PSNS Pollutant or Pollutant Property mg/troy ounce of othe precipitated Copper Cyanide (total)	Maximum for Any One Day er platinum grou	Maximum for Monthly Average p metals 3.172	
Filtration PSNS Pollutant or Pollutant Property mg/troy ounce of othe precipitated	Maximum for Any One Day er platinum grou 6.656 1.040	Maximum for Monthly Average p metals 3.172 0.416	

(1) Spent Solution from PGC Salt Production PSNS

Pollutant or	Maximum for	
Pollutant Property	Any One Day	Monthly Average
mg/troy ounce of go	ld contained in	PGC product
Copper	1.152	0.549
Cyanide (total)	0.180	0.072
Zinc	0.918	0.378
Ammonia (as N)	120.000	52.740
Combined metals	0.270	
	· · · · · · · · · · · · · · · · · · ·	······································
(m) Equipment and	Floor Wash PSNS	
Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
	-	
mg/troy ounce of pr		including silver,
mg/troy ounce of pr produced in refiner		including silver,
produced in refiner	Y	
produced in refiner Copper	У 0 .000	0.000
produced in refiner	9 0.000 0.000	0.000 0.000
produced in refiner Copper Cyanide (total) Zinc Ammonia ∢as N)	У 0 .000	0.000
produced in refiner Copper Cyanide (total) Zinc Ammonia ∢as N)	9 0.000 0.000 0.000	0.000 0.000 0.000
produced in refiner Copper Cyanide (total) Zinc	9 0.000 0.000 0.000 0.000	0.000 0.000 0.000
produced in refiner Copper Cyanide (total) Zinc Ammonia ∢as N)	9 0.000 0.000 0.000 0.000 0.000	0.000 0.000 0.000
produced in refiner Copper Cyanide (total) Zinc Ammonia (as N) Combined metals (n) <u>Preliminary tra</u>	y 0.000 0.000 0.000 0.000 0.000	0.000 0.000 0.000 0.000
produced in refiner Copper Cyanide (total) Zinc Ammonia (as N) Combined metals (n) <u>Preliminary tra</u> Pollutant or	y 0.000 0.000 0.000 0.000 0.000 <u>eatment</u> PSNS Maximum for	0.000 0.000 0.000 0.000 Maximum for
produced in refiner Copper Cyanide (total) Zinc Ammonia (as N) Combined metals	y 0.000 0.000 0.000 0.000 0.000 <u>eatment</u> PSNS Maximum for	0.000 0.000 0.000 0.000
produced in refiner Copper Cyanide (total) Zinc Ammonia (as N) Combined metals (n) <u>Preliminary tra</u> Pollutant or Pollutant Property	y 0.000 0.000 0.000 0.000 0.000 eatment PSNS Maximum for Any One Day	0.000 0.000 0.000 0.000 Maximum for Monthly Average
produced in refiner Copper Cyanide (total) Zinc Ammonia (as N) Combined metals (n) <u>Preliminary tra</u> Pollutant or	y 0.000 0.000 0.000 0.000 0.000 <u>eatment</u> PSNS <u>Maximum for</u> Any One Day cal precious met	0.000 0.000 0.000 0.000 Maximum for Monthly Average
produced in refiner Copper Cyanide (total) Zinc Ammonia (as N) Combined metals (n) <u>Preliminary trans</u> Pollutant or Pollutant Property mg/troy ounce of tot	y 0.000 0.000 0.000 0.000 0.000 <u>eatment</u> PSNS <u>Maximum for</u> Any One Day cal precious met	0.000 0.000 0.000 0.000 Maximum for Monthly Average
produced in refiner Copper Cyanide (total) Zinc Ammonia (as N) Combined metals (n) <u>Preliminary transformer</u> (n) <u>Preliminary transformer</u>	y 0.000 0.000 0.000 0.000 0.000 eatment PSNS Maximum for Any One Day cal precious met	0.000 0.000 0.000 0.000 Maximum for Monthly Average
produced in refiner Copper Cyanide (total) Zinc Ammonia (as N) Combined metals (n) <u>Preliminary transformer</u> (n) <u>Preliminary transformer (n) <u>Prelimina</u></u></u></u></u></u></u></u></u></u></u></u></u></u></u></u></u></u></u></u></u></u></u></u></u></u></u></u>	y 0.000 0.000 0.000 0.000 0.000 <u>eatment</u> PSNS <u>Maximum for</u> Any One Day cal precious met ion 64.000 10.000 51.000	0.000 0.000 0.000 0.000 Maximum for Monthly Average tals produced 30.500
produced in refiner Copper Cyanide (total) Zinc Ammonia (as N) Combined metals (n) <u>Preliminary transformer</u> Pollutant or Pollutant or Pollutant Property mg/troy ounce of tot through this operat: Copper Cyanide (total)	y 0.000 0.000 0.000 0.000 0.000 eatment PSNS Maximum for Any One Day tal precious met ion 64.000 10.000	0.000 0.000 0.000 0.000 Maximum for Monthly Average tals produced 30.500 4.000

EPA is not promulgating best conventional pollutant control technology (BCT) at this time.

SECTION III

SUBCATEGORY PROFILE

This section of the secondary precious metals supplement describes the raw materials and processes used in refining secondary precious metals and presents a profile of the secondary precious metals plants identified in this study. For a discussion of the purpose, authority, and methodology for this study and a general description of the nonferrous metals category, refer to Section III of Vol. I.

DESCRIPTION OF SECONDARY PRECIOUS METALS PRODUCTION

The secondary precious metals industry consists of plants which recover gold and platinum group metals from recycled materials. Platinum group metals, also known as PGM, consist of platinum, iridium, rhodium, osmium, and ruthenium. The palladium, production of secondary precious metals can be divided into two stages: raw material preparation steps and refining steps. Raw crushing, preparation steps include grinding, material incineration, smelting, granulation, cyanide stripping, and precipitation of precious metals from spent plating solutions. Refining steps include dissolution in either strong acid or base, solvent extraction, filtration, recycle, precipitation, salt manufacturing, casting, and electrolytic refining, The secondary precious metals production process is granulation. presented schematically in Figure III-1 (page 2343).

RAW MATERIALS

The principal raw materials used by plants recovering precious metals are jewelry scrap, dental scrap, optical scrap, electrical scrap, impure bullion, spent industrial and automotive catalysts, sweeps, and contaminated or spent electroplating solutions. Sweeps are usually low-grade precious metal-bearing residue generated from various raw materials, including floor sweepings (hence the name); waste treatment sludges and incinerated filter cakes. The various raw material preparation and refining steps a plant uses are dictated by the type and composition of raw materials being processed.

RAW MATERIAL PREPARATION STEPS

Based on the source of raw materials, the raw material preparation steps can be divided into five basic processes for the recovery of precious metals: incineration and smelting (pyrometallurgical steps), raw material granulation, stripping with cyanide solutions, recovering precious metals from spent plating solutions, and other preliminary treatment steps.

Incineration and Smelting

Dental scrap, optical scrap, electrical scrap, and catalysts may be ground and incinerated in a furnace in order to remove the carbonaceous material and volatile fraction. The temperature and rate of burning must be carefully controlled if high efficiency is to be maintained. Air emissions include vapors from the volatilization and decomposition of carbonaceous scrap contaminants, as well as combustion gases and dust. The emissions are usually controlled by afterburners in series with a baghouse or scrubber. Wet air pollution control techniques result in wastewater discharges. Precious metal-bearing residues may then be fed directly to the refinery for recovery of pure metals.

Smelting is generally used to produce a copper-based bullion which can either be sold or further processed to produce a pure metal. The raw material for smelting may be the precious metalbearing residue produced in the incinerator, or it may be groundup raw material. Like the incineration furnace, the smelting furnace may also have emissions which are controlled by a baghouse or scrubber. The furnace or incinerator scrubber results in a wastewater discharge.

Raw Material Granulation

Raw material may be granulated with water in order to make it easier to dissolve in acid in the refinery. Either solid scrap or incinerated residue may be melted in a furnace and granulated with water in a similar manner to shot casting. This operation wastewater discharge, consisting of the spent produces granulation water.

Stripping With Cyanide Solutions

Gold-containing electrical components, strip, or ceramics may be stripped with sodium or potassium cyanide solutions. The raw material may be ground-up prior to stripping in order to increase the exposed surface area. Cyanide attacks the gold which is exposed on the surface of the metal, but does not recover gold which is buried beneath a non-precious metal. Stripping with cyanide has limited application because of the relatively few types of scrap amenable to the process.

After the gold is stripped away from the base metal, it may be precipitated from solution with either sodium hydrosulfite or zinc. An oxidizing agent may be added to destroy the free cyanide. The solids, containing precious metals, are separated from the spent cyanide stripping solution by filtration. Filtration results in a wastewater stream which may be discharged. The product of cyanide stripping is a sludge containing high precious metal values which may be further processed.

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Recovery From Spent Plating Solutions

Precious metals can be recovered from contaminated or spent electroplating solutions, which are cyanide-based, either by sodium hydrosulfite or zinc, or by precipitation with electrolysis. Electroplaters use cyanide solutions for plating precious metals onto base metals. The depleted or contaminated solutions still contain enough precious metal values to make recovery economical. Either gold, palladium, or rhodium can be recovered in this manner. The precipitation process for plating solutions is the same as cyanide stripping. Zinc or sodium hydrosulfite is added and the precious metals are recovered by electrolytic recovery, the spent plating filtration. For solution acts as the electrolyte, and the precious metal is recovered on the cathode. Wastewater may be generated by the discharge of barren solution after either precipitation or electrolytic recovery occurs. The resultant sludge from this process may be routed to the refinery for further processing.

Other Preliminary Treatment Steps

Preliminary treatment steps other than those mentioned above may be used to treat raw material in this subcategory. These processing steps, which have been claimed confidential are used to treat non-combustible, non-metallic-based basis materials. These preliminary treatment steps may produce a wastewater discharge.

REFINING STEPS

Refining steps are taken to recover high-purity precious metals (high-purity generally refers to 99.9 or 99.99 percent pure) from lower purity raw materials, which may or may not have undergone raw material preparation steps. The standard hydrometallurgical process includes dissolution in acid or base, combined with precipitation and filtration. Other hydrometallurgical refining steps include solvent extraction and electrolytic refining. After pure precious metals are produced, they may be further processed into a potassium cyanide-based salt, cast as bars, or granulated.

Hydrometallurgical Processing

Jewelry, dental, optical, electrical, and catalyst scrap, along with sludges generated from spent solutions, containing gold, platinum, palladium, and other platinum group metals (PGM), may be refined using hydrometallurgical processing. The first step usually consists of dissolving the raw material in aqua regia. Aqua regia (one part concentrated nitric acid:three to four parts concentrated hydrochloric acid) is the only known reagent that dissolves gold. Nitric acid alone cannot oxidize gold unless the chloride ion is present to complex the product. The net equation for dissolving gold in aqua regia can be written as below although a variety of nitrogen products may be obtained. $Au(s) + 4NO_3^- + 4Cl^- + 8H_3O^+ ----> AuCl_4 + 4NO_2(g) + 12H_2O$

After dissolving the raw material, the silver chloride solids are filtered away, the nitrates are removed, and the gold is precipitated with sulfur dioxide, ferrous sulfate, or chlorine gas. The filtrate may be sent on for further recovery of platinum group metals, if these are present. The silver chloride solids recovered from the gold dissolution process usually require further purification. Silver recovery and purification is addressed in the secondary silver development document.

Platinum group metals are recovered from raw materials which contain gold and platinum group metals. Platinum is recovered by dissolution in a chlorinated acid solution, yielding soluble platinum chloride. Platinum is precipitated as an insoluble amine called yellow salt which is purified using both alkaline and acid solutions and reduced to platinum metal sponge either thermally or with a strong chemical reducing agent.

Palladium is recovered by dissolution in acid as a soluble chloride salt. Palladium is precipitated from solution as an insoluble amine called red salt. Palladium is purified by redissolving the red salt in an alkaline solution followed by reprecipitation. The red salt is reduced to palladium metal sponge with a strong chemical reducing agent.

Each of the purification processes may be repeated via recycle to increase the purity of the refined metal. After each metal is recovered as either a final product or intermediate, it may be washed with water or an acid or base in order to remove residual acid or base from it and to further purify it. The wash water or solution is generally discharged with the precipitation and filtration water, and is considered as part of the same waste stream.

The various hydrometallurgical processing steps a plant uses to recover precious metals may occur in any order. For example, one plant may recover gold prior to palladium prior to platinum, and another plant may recover platinum first, then gold, and finally palladium. The order of processing does not impact the wastewater generation at a refinery.

Based on the composition of the raw material, and the order of processing, the recovery of each precious metal may result in a wastewater discharge. There is variability in the types of raw materials processed within this subcategory; however, the basic processing steps and wastewaters generated are similar from one plant to another.

Acid fumes generated in the refinery may be controlled with a wet scrubber, resulting in a wastewater stream. In many plants, this scrubber controls the fumes from all the reaction vessels, whether they are acid, alkaline, or cyanide based. The scrubbing

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medium is usually an alkaline solution which neutralizes the acid fumes. Other plants may use separate scrubbers for acid and alkaline fumes.

Solvent Extraction

Solvent extraction may be used to refine impure bullion to high purity gold. Solvent extraction consists of extracting the gold from an acid solution into the organic phase and subsequently recovering it. The aqueous solution which originally contained the dissolved impure bullion may be discharged as a waste stream. After recovery, the gold may be washed with water and the wash water may also be discharged. The aqueous raffinate and wash water may be considered as one waste stream.

Electrolytic Refining

Electrolytic refining is also used as a means of recovering high purity gold from precious metal-containing bullion, jewelry and dental scrap. First, the raw material is melted and cast as an anode. An acidic electrolyte is used, and gold is recovered on the cathode. In the electrolytic method, a current is passed between an anode and a cathode which are suspended in the electrolyte. A portion of the electrolyte is periodically discharged to maintain the purity of the solution.

Further Processing

Once the gold or platinum group metals have been refined to the pure state, they may be further processed. Gold may be reacted with potassium cyanide solution to produce a potassium gold cyanide salt (generally written KAu(CN)₂ or PGC) which is useful in the electroplating industry. There may be a waste stream associated with this process, consisting of excess cyanide solution.

Pure precious metals may either be cast as bars or granulated using a method similar to shot casting. In either case, the metal is melted in a furnace. Molten metal may be poured into molds which may be quenched with water, or it may be poured directly into a container of water, in which case it will be granulated. In either case, a waste stream is generated which may be discharged.

PROCESS WASTEWATER SOURCES

Although a variety of processes are involved in secondary precious metals production, the process wastewater sources can be subdivided into building blocks as follows:

- 1. Furnace wet air pollution control,
- 2. Raw material granulation,
- 3. Spent plating solutions,
- 4. Spent cyanide stripping solutions,
- 5. Refinery wet air pollution control,

- 6. Gold solvent extraction raffinate and wash water,
- 7. Gold spent electrolyte,
- 8. Gold precipitation and filtration,
- 9. Platinum precipitation and filtration,
- 10. Palladium precipitation and filtration,
- 11. Other platinum group metals precipitation and filtration,
- 12. Spent solution from PGC salt production,
- 13. Equipment and floor wash, and
- 14. Preliminary treatment.

The sources of these wastewater streams are identified by their respective numbers in Figure III-1 (page 2343).

OTHER WASTEWATER SOURCES

There are other waste streams associated with the production of secondary precious metals. These waste streams may include casting contact cooling water, final product granulation water, acid storage area wet air pollution control, and pump seal water.

These waste streams are not considered as part of this rulemaking. EPA believes that the flows and pollutant loadings associated with these wastewater streams are insignificant relative to the wastewater streams selected and are best handled by the appropriate permit authority on a case-by-case basis under the authority of Section 403(a) of the Clean Water Act.

Casting contact cooling water is not considered as part of this rulemaking because, although several plants do discharge this stream, sampling data indicate that this wastewater contains little or no pollutants and that the pollutant loadings are insignificant compared with the other waste streams selected. Sampling data for casting contact cooling water are presented in Table V-26 (page 2497).

AGE, PRODUCTION, AND PROCESS PROFILE

Forty-nine secondary precious metals plants were identified in this study. Figure III-2 (page 2344) shows that the plants are concentrated in the Northeast and California, with plants also located in Washington, Arizona, Minnesota, Illinois, Ohio, Virginia, and Florida.

Table III-1 (page 2340) summarizes the relative ages of the secondary precious metals plants by discharge status. Four plants discharge directly, 30 are indirect dischargers, 10 are zero dischargers, and five plants do not generate process wastewater. Most of the Plants began operating within the last 15 years.

Table III-2 (page 2341) shows the production ranges for the 49 secondary precious metals plants. One-third of the plants that reported production data produce less than 10,000 troy ounces of total precious metals per year. All four of the direct

dischargers produce in excess of 50,000 troy ounces per year, as do 10 of the indirect dischargers.

Table III-3 (page 2342) provides a summary of the plants having the various secondary precious metals processes. The number of plants generating wastewater from the processes is also shown.

Table III-1

INITIAL OPERATING YEAR (RANGE) SUMMARY OF PLANTS IN THE SECONDARY PRECIOUS METALS SUBCATEGORY BY DISCHARGE TYPE

Type of Plant Discharge	1983- <u>1973</u> <u>1-10</u>	1972- <u>1968</u> <u>11-15</u>	1967- <u>1958</u> 16-25	1957- <u>1948</u> 26-35	1947- <u>1938</u> 36-45	1937- 1928 46-55	1927- 1918 56-55	1917- 1903 66-80	<u><1903</u> <u>81+</u>	Not Reported	<u>Total</u>
Direct	1	1						1	1		4
Indirect	8	8	4	3	1	ľ	1	2	1	1	30
Zero	4	2	1							3	10
Dry			<u>1</u>	-	<u>1</u>	-	<u>1</u>	-		-	_5
lotal	15	11	6	3	2	1	2	3	2	4	49

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Table III-2

PRODUCTION RANGES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY DURING 1982 (TROY OUNCES OF TOTAL PRECIOUS METALS/YEAR)

Type of Plant	0-1,000	1,001- 10,000	10,001- 50,000	50,001- 100,000	100,001- 500,000	<u>500,000+</u>	Not Reported in_dcp
Direct*							
Indirect	1	9	9	4	6	. 1	
Zero		3	1		4		2
Dry	1	_1	1	· -		<u>2</u>	<u>1</u>
Total	2	13	. 11	4	10	··· 3	3

*Data for these plants are claimed to be confidential.

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Table III-3

SUMMARY OF SECONDARY PRECIOUS METALS SUBCATEGORY PROCESSES AND ASSOCIATED WASTE STREAMS

Process	Number of Plants With Process	Number of Plants Reporting Generation of Wastewater*
Raw Materials Preparation Steps	28	
 Incineration and Smelting (Furnace Air Pollution Control) Raw Material Granulation Stripping With Cyanide Solutions Recovery From Spent Plating Solutions Refining Steps (Hydrometallurgical Processing) 	18 4 6 12 37	7 4 6 12
 Gold Precipitation and Filtration Platinum Precipitation and Filtration Palladium Precipitation and Filtration Other Platinum Group Metals Precipitation and Filtration Solvent Extraction Electrolytic Refining 	28 18 20 3 1 3	28 18 20 3 1 3
PGC Salt Production Equipment and Floor Wash	4 3	4 3

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*Through reuse or evaporation practices, a plant may "generate" wastewater from a particular process but not discharge it.

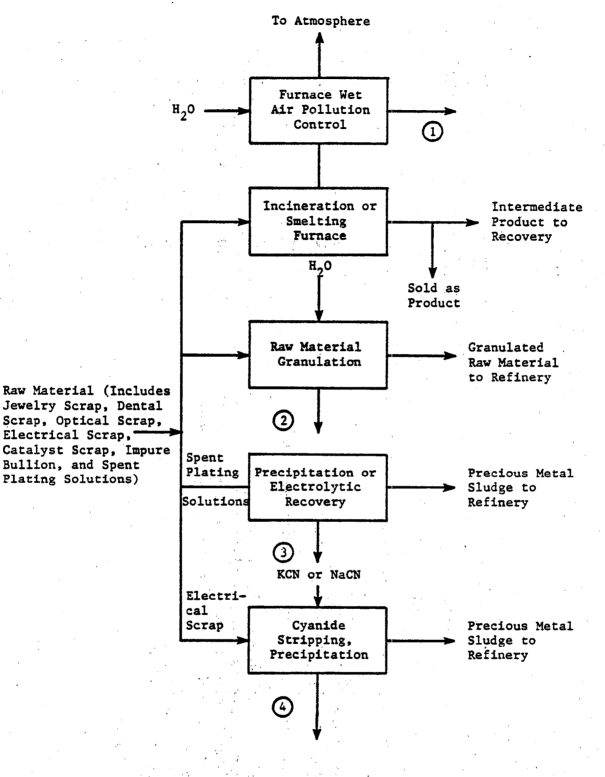
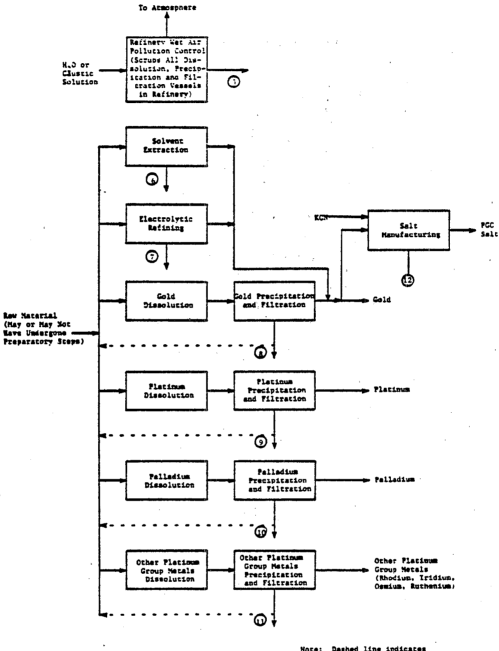


Figure III-1

RAW MATERIAL PREPARATION SECONDARY PRECIOUS METALS PRODUCTION PROCESSES

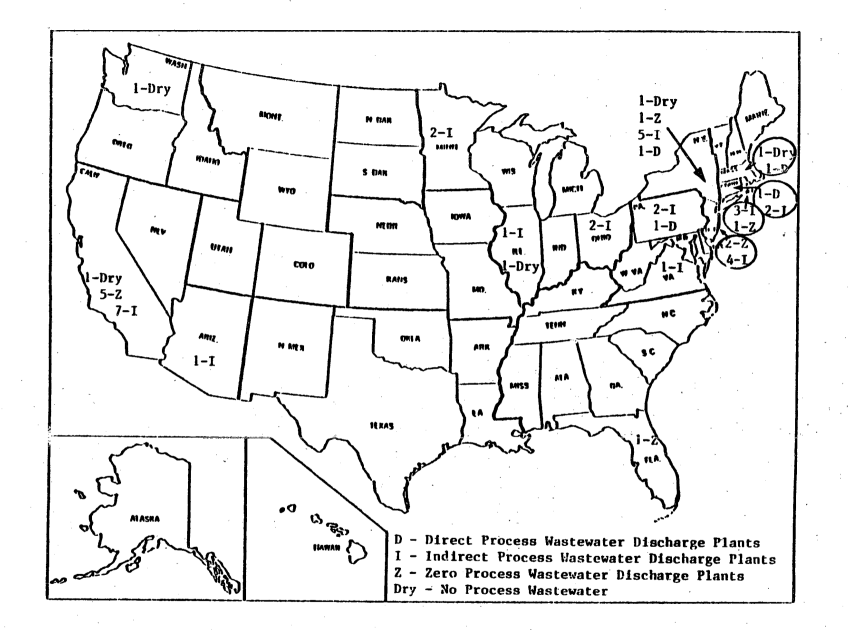
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Nore: Dashed line indicates possible recycle or reuse of filtrate

Figure III-1 (Continued) REFINING STEPS SECONDARY PRECIOUS METALS PRODUCTION PROCESSES



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Figure III-2

GEOGRAPHIC LOCATIONS OF THE SECONDARY PRECIOUS METALS INDUSTRY

SECONDARY PRECIOUS METALS SUBCATEGORY

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SECTION IV

SUBCATEGORIZATION

This section summarizes the factors considered during the designation of the related subdivisions or building blocks of the secondary precious metals subcategory.

FACTORS CONSIDERED IN SUBDIVIDING THE SECONDARY PRECIOUS METALS SUBCATEGORY

The factors listed previously for general subcategorization were each evaluated when considering subdivision of the secondary precious metals subcategory. In the discussion that follows the factors will be described as they pertain to this particular subcategory.

The rationale for considering further subdivision of the secondary precious metals subcategory is based primarily on the production processes used. Within the subcategory a number of different operations are performed which may or may not have a water use or discharge and which may require the establishment of separate effluent limitations and standards. While secondary precious metals is still considered a single Subcategory. a more thorough examination of the production processes, water use anð discharge practices and pollutant generation rates has illustrated the need for limitations and standards based on a set has specific wastewater streams. Limitations and standards will of based on specific flow allowances for the following building be blocks:

- 1. Furnace wet air pollution control.
- 2. Raw material granulation,
- 3. Spent plating solutions.
- 4. Spent cyanide stripping solutions,
- 5. Refinery wet air pollution control,
- 6. Gold solvent extraction raffinate and wash water,
- 7. Gold spent electrolyte,
- 8. Gold precipitation and filtration.
- 9. Platinum precipitation and filtration,
- 10. Palladium precipitation and filtration,
- 11. Other platinum group metals precipitation and filtration,
- 12. Spent solution from PGC salt production,
- 13. Equipment and floor wash, and
- 14. Preliminary treatment.

These building blocks follow directly from differences within the various production stages of secondary precious metals: raw material preparation steps and refining steps. Depending on the type and composition of raw material, a plant may operate one or more raw material preparation or refining steps to recover gold platinum, palladium, or other platinum group metals from scrap. Each of these operations may create a need for a subdivision.

Smelting or incinerating a raw material creates the need for the

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - IV

first subdivision--furnace wet air pollution control. Smelting incineration furnaces produce dust and particulate emissions or which need to be controlled prior to venting to the atmosphere. Other raw material preparation steps which create the need for subdivisions include raw material granulation, recovering gold or other precious metals from spent plating solutions, cyanide stripping of gold from gold-plated scrap, and other confidential preliminary treatment steps. Granulating a raw material involves melting the raw material in a furnace and pouring it into a container of water. This granulates the raw material, and the granulation water may be discharged, thus creating the need for a Spent plating solutions may be treated with a subdivision. precipitating agent such as zinc or sodium thiosulfate in order to precipitate the precious metals. Discharging the depleted solution creates a need for a separate subdivision. Stripping gold away from scrap with a cyanide solution and then precipitating the gold from solution creates a need for the fourth pre-refining subdivision. Other preliminary treatment steps, which are considered confidential are used to treat noncombustible, non-metallic-based basis materials, and create a need for the last subdivision.

Various refining operations create the need for the other nine subdivisions. Recovering gold by a solvent extraction process or electrolytic refining process creates the need for two an subdivisions: gold solvent extraction raffinate and wash water, and gold spent electrolyte. The wet chemistry technique of dissolution and selective precipitation creates the need for four gold precipitation and filtration, platinum subdivisions: precipitation and filtration palladium and precipitation filtration, and other platinum group metals precipitation and filtration. Depending on the composition of the raw material being processed, and the manner in which each metal is recovered, any one or all of the precious metals may result in the discharge of a wastewater stream.

Acid fumes generated during dissolution and precipitation processes are generally controlled with a wet scrubber, creating the need for the seventh refining subdivision: refinery wet air pollution control. Washing the equipment and the floor of the refinery in order to recover any precious metals from spills and leaks creates a need for the equipment and floor wash subdivision. Finally, manufacturing gold into a PGC salt product by reacting it with potassium cyanide solution creates a need for a subdivision: spent solution from PGC salt production.

OTHER FACTORS

The other factors considered in this evaluation were shown to be inappropriate as a bases for further segmentation of the subcategory. Air pollution control methods, treatment costs, nonwater quality aspects, and total energy requirements are functions of the selected subcategorization factors -- raw materials and production processes. As such, they support the method of subcategorization which has been developed. As

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discussed in Section IV of the General Development Document, certain other factors such as plant age, plant size, and the number of employ-ees were also evaluated and determined to be inappropriate for use as bases for subdivision of nonferrous metals plants.

PRODUCTION NORMALIZING PARAMETERS

The effluent limitations and standards developed in this document establish mass limitations for the discharge of specific pollutant parameters. To allow these limitations to be applied to plants with various production capacities, the mass of pollutant discharged must be related to a unit of production. This factor is known as the production normalizing parameter (PNP). In general, the actual precious metals production from the respective manufacturing process is used as the PNP. This is based on the principle that the amount of water generated is proportional to the amount of product made. Therefore, the PNPs for the 14 secondary precious metals subdivisions are as follows:

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PNP Building Block 1. Furnace wet air Troy ounces of precious pollution control metals, including silver, incinerated or smelted بربين المناه المناه 2. Raw material granulation Troy ounces of precious metals in the granulated raw material 3. Spentaplating solutions Liters of spent plating solutions used as a raw material Troy ounces of gold produced 4. Spent cyanide stripping by cyanide stripping solutions Refinery wet air pollution Troy ounces of precious 5. metals, including silver, control produced in refinery Troy ounces of gold produced Gold solvent extraction 6. of solvent extraction raffinate and wash water Troy ounces of gold produced 7. Gold spent electrolyte by electrolysis Troy ounces of gold 8. Gold precipitation and precipitated filtration Troy ounces of platinum Platinum precipitation 9. precipitated and filtration Palladium precipitation Troy ounces of palladium 10. precipitated and filtration 11. Other platinum group Troy ounces of other platinum

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - IV

metals precipitation and filtration

- 12. Spent solution from PGC salt production
- 13. Equipment and floor wash

group metals precipitated

Troy ounces of gold contained in PGC product

- Troy ounces of precious metals, including silver, produced in refinery
- 14. Preliminary treatment

Troy ounces of total precious metals produced through this operation

Other PNPs were considered. The use of production capacity instead of actual production was eliminated from consideration because the mass of pollutant produced is more a function of true production than of installed capacity. The total precious metals produced in the refinery was eliminated from consideration because most of the operations generating wastewater in a refinery do so as a function of one metal being produced, rather than as a function of the total amount of metal produced in a refinery.

The PNP selected for spent plating solutions is liters of spent plating solution used as a raw material. The volumetric PNP was selected rather than the mass of metal processed because a plant cannot control the concentration of precious metals in the raw One plant's material -- the imported spent plating solutions. anothers' in raw material may be many times as concentrated as precious metals, and therefore flow cannot be related to production for this unit operation. Wastewater discharge flow is directly related to volume of spent plating solution used as raw material, and not the quantity of precious metals in the solution.

SECONDARY PRECIOUS METALS SUBCATEGORY

SECTION V

WATER USE AND WASTEWATER CHARACTERISTICS

This section describes the characteristics of the wastewaters associated with the secondary precious metals subcategory. Water use and discharge rates are explained and then summarized in tables at the end of this section. Data used to characterize the wastewaters are presented. Finally, the specific source, water use and discharge flows, and wastewater characteristics for each separate wastewater source are discussed.

The two principal data sources were used in the development of effluent limitations and standards for this subcategory are data collection portfolios (dcp) and field sampling results. Data collection portfolios contain information regarding wastewater flows and production levels.

In order to quantify the pollutant discharge from secondary precious metals plants, a field sampling program was conducted. A complete list of the pollutants considered and a summary of the techniques used in sampling and laboratory analyses are included Section V of the General Development Document. Samples were in analyzed for 124 of the 126 priority pollutants and other pollutants deemed appropriate. Because the analytical standard for TCDD was Judged to be too hazardous to be made generally available, samples were never analyzed for this pollutant. Samples were also never analyzed for asbestos. There is no reason to expect that TCDD or asbestos would be present in nonferrous metals manufacturing wastewater. A total of five plants were selected for sampling in the secondary precious metals subcategory. In general, the samples were analyzed for cyanide and three classes of pollutants: priority organic pollutants, priority metal pollutants, and criteria pollutants (which includes both conventional and nonconventional pollutants). Cyanide was analyzed for because it is present in raw materials for this subcategory.

Additional wastewater characteristics and flow and production data were received through industry comments and an engineering site visit to one facility between proposal and promulgation. This aided EPA in promulgating revised discharge allowances for raw material granulation and spent cyanide stripping solutions waste streams.

Also since proposal, EPA gathered additional wastewater' sampling data for four of the subdivisions in this subcategory through a self-sampling program which was specifically requested by the Agency. The data from this program include data from analyses for the primary metals antimony, arsenic, beryllium, cadmium, chromium, copper, lead, mercury, nickel, selenium, silver, thallium, and zinc. The data also include analyses for cyanide

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - V

and the nonconventional pollutants ammonia, gold, palladium, and platinum. These data support the assumptions which EPA had made concerning the presence and concentrations of pollutants in those subdivisions where we did not have analytical data for specific For this reason, the selection of pollutant pollutants. parameters for limitation in this subcategory (Section VI) has not been revised based on this new data.

As described in Section IV of this supplement, the secondary precious metals subcategory has been further subdivided into 14 building blocks, so that the promulgated regulation contains mass discharge limitations and standards for 14 process wastewater discharging subdivisions. Differences in the wastewater characteristics associated with these subdivisions are to be expected. For this reason, wastewater streams corresponding to each subdivision are addressed separately in the discussions that follow. uses of the monoralise sectors and

The principal wastewater sources in the secondary precious metals subcategory are:

- Furnace wet air pollution control, 1.
- 2. Raw material granulation,
- Spent plating solutions, 3.
- Spent cyanide stripping solutions, 4.
- 5. Refinery wet air pollution control,
- Gold solvent extraction raffinate and wash water, 6.

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- Gold spent electrolyte, 7.
- Gold precipitation and filtration, 8.
- Platinum precipitation and filtration, 9.
- Palladium precipitation and filtration, 10.
- Other platinum group metals precipitation and 11. filtration,
- 12. Spent solution from PGC salt production,
- 13. Equipment and floor wash, and
- 14. Preliminary treatment.

WASTEWATER FLOW RATES

Data supplied by dcp responses were evaluated, and two flow-toproduction ratios were calculated for each stream. The two water use and wastewater discharge flow, are ratios. differentiated by the flow value used in calculation. Water use is defined as the volume of water or other fluid required for a given process per mass of precious metals product and is therefore based on the sum of recycle and make-up flows to a given process. Wastewater flow discharged after preliminary treatment or recycle (if these are present) is used in calculating the production normalized flow -- the volume of in wastewater discharged from a given process to further treatment, disposal, or discharge per mass of precious metals produced. Differences between the water use and wastewater flows associated with a given stream result from recycle, evaporation, and carry-over on the product. The production values used in calculation correspond to the production normalizing parameter,

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - V

PNP, assigned to each stream, as outlined in Section IV. As an example, gold precipitation and filtration wastewater flow is related to gold metal production. As such, the discharge rate is expressed in liters of filtration wastewater discharged per troy ounce of gold produced by precipitation.

The production normalized flows were compiled and statistically analyzed by stream type. These production normalized water use and discharge flows are presented by subdivision in Tables V-1 through V-13 (pages 2360-2372). Where appropriate, an attempt was made to identify factors that could account for variations in water use. This information is summarized in this section. Α similar analysis of factors affecting the wastewater flows is presented in Sections IX, X, XI, and XII where representative BPT, BAT, NSPS, and pretreatment flows are selected for use in calculating the effluent limitations and standards.

WASTEWATER CHARACTERIZATION DATA

Data used to characterize the various wastewaters associated with secondary precious metals production come from two sources data collection portfolios and analytical data from field sampling trips.

DATA COLLECTION PORTFOLIOS

In the data collection portfolios, the secondary precious metals plants which discharge wastewater were asked to specify the presence of toxic pollutants in their effluent. Of the 49 secondary precious metals plants, 12 did not respond to this portion of the questionnaire. No plant responding to this portion of the questionnaire reported that any toxic organic pollutants were known to be or believed to be present in their wastewater.

The responses for the toxic metals and cyanide are summarized below:

Pollutant	Known Present	Believed Present (Based on Raw Materials and Process Chemicals Used)	
Antimony	0	3	
Arsenic	1	5	
Beryllium	2	3	
Cadmium	7	5	
Chromium	9	6	
Copper	20	17	
Cyanide	10	10	
Lead	. 11	8	
Mercury	3	2	
Nickel	16	19	
Selenium	0	3	
Silver	14	18	

Thallium	0
Zinc	20

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FIELD SAMPLING DATA

In order to quantity the concentrations of pollutants present in wastewater from secondary precious metals plants, wastewater samples were collected at five plants. Diagrams indicating the sampling sites and contributing production processes are shown in Figures V-1 through V-5 (pages 2360-2364).

The raw wastewater sampling data for the secondary precious metals subcategory are Presented in Tables V-14 through V-21 (pages 2373-2436). Treated and combined wastewater sampling data are shown in Tables V-22 through V-25 (pages 2440-2484). The stream codes presented in the tables may be used to identify the location of each of the samples on the process flow diagrams in Figures V-1 through V-5. Where no data are listed for a specific day of sampling, the wastewater samples for the stream were not collected.

Several points regarding these tables should be noted. First, the data tables include some samples measured at concentrations considered not quantifiable. The base-neutral extractable, acid traction extractable, and volatile organics are generally considered not quantifiable at concentrations equal to or less than 0.010 mg/l. Below this concentration, organic analytical results are not quantitatively accurate; however, the analyses are useful to indicate the presence of a particular pollutant. The pesticide fraction is considered not quantifiable at concentrations equal to or less than 0.005 mg/l.

Second, the detection limits shown on the data tables for priority metals and conventional and nonconventional pollutants are not the same in all cases as the published detection limits for these pollutants by the same analytical methods. The detection limits used were reported with the analytical data and hence are the appropriate limits to apply to the data. Detection limit variation can occur as a result of a number of laboratoryspecific, equipment-specific, and daily operator-specific factors. These factors can include day-to-day differences in machine calibration, variation in stock solutions, and variation in operators.

Third, the statistical analysis of data includes some samples measured at concentrations considered not quantifiable. For data considered as detected but below quantifiable concentrations, a value of zero is used for averaging. Priority organic, nonconventional, and conventional pollutant data reported with a "less than" sign are considered as detected, but not further quantifiable. A value of zero is also used for averaging. If a pollutant is reported as not detected, it is assigned a value of zero in calculating the average. Finally, priority metal values reported as less than a certain value were considered as not quantifiable, and consequently were assigned a value of zero in the calculation of the average.

Finally, appropriate source water concentrations are presented with the summaries of the sampling data. The method by which each sample was collected is indicated by number, as follows:

- 1. One-time grab
- 2. Manual composite during intermittent process operation
- 3. 38-hour manual composite
- 4. 8-hour automatic composite
- 5. 24-hour manual composite
- 6. 24-hour automatic composite

WASTEWATER CHARACTERISTICS AND FLOWS BY SUBDIVISION

Since secondary precious metals production involves 14 principal sources of wastewater and each has potentially different characteristics and flows, the wastewater characteristics and discharge rates corresponding to each subdivision will be described separately. A brief description of why the associated production processes generate a wastewater and explanations for variations of water use within each subdivision will also be discussed.

FURNACE WET AIR POLLUTION CONTROL

Of the secondary precious metals plants with furnaces, smelters, or incinerators, seven plants use wet scrubbers to control emissions. Five of these discharge wastewater as shown in Table V-1 page 2360). This table shows the water discharge rates in liters per troy ounce of precious metals, including silver, processed through the furnace. Of the seven plants using wet scrubbers, two plants practice 100 percent recycle, two plants practice greater than 90 percent recycle, and three plants do not recycle this water.

The Agency sampled the wastewater from two of the five discharging plants, one of which does not practice recycle. The other practices greater than 90 percent recycle. The Agency also sampled the wastewater at another secondary precious metals plant which did not practice recycle. Furnace wet air pollution control raw wastewater contains priority metals, cyanide, and suspended solids above treatable concentrations. Raw wastewater sampling data are presented in Table V-14 (page 2373).

RAW MATERIAL GRANULATION

Raw material may be melted in a furnace and then poured into a container of standing water in order to granulate it. This process is similar to shot casting. The purpose of this operation is to make it easier to dissolve the raw material in dissolution process. Of the 31 plants acid the which four hydrometallurgically refine precious metals, plants granulate the raw material prior to dissolution. Three plants

discharge this wastewater, as shown in Table V-2 (page 2361). The fourth plant practices 100 percent recycle of granulation water.

Following proposal, sampling data for this subdivision were acquired through a self-sampling effort specifically requested by the Agency. These data show treatable concentrations of cadmium, lead, silver, palladium, and TSS.

SPENT PLATING SOLUTIONS

Spent or contaminated electroplating solutions with a high precious metal content may be recycled to recover the precious metals value. After recovering this value, the depleted solution may be discharged. Twelve plants recover precious metals from spent plating solutions. Discharge rates for these 12 plants are presented in Table V-3 (page 2362), in liters of wastewater per liter of raw material spent plating solution.

The Agency sampled two plants for this waste stream, and the results are presented in Table V-15 (page 2389). This raw wastewater contains priority metals, free and complexed cyanide, and TSS above treatable concentrations.

SPENT CYANIDE STRIPPING SOLUTIONS

Six plants use sodium or potassium cyanide solutions to strip gold away from electronic scrap and other raw materials. After precipitating the gold, the spent cyanide solution may be discharged. Six plants use this technique as shown in Table V-4 (page 2363). Water use and discharge rates are shown in liters per troy ounce of gold produced by cyanide stripping. Gold production is measured as the product from the precipitation operation.

The Agency sampled one plant for this waste stream, and the results are presented in Table V-16 (page 2399). This waste stream contains priority metals, free and complexed cyanide, and TSS above treatable concentrations.

REFINERY WET AIR POLLUTION CONTROL

All of the acid dissolution vessels, alkaline dissolution vessels, cyanide vessels, and precipitation vessels located in the refinery may be vented to a refinery scrubber. A plant can use acid scrubbers, alkali scrubbers, or both types of scrubbers. Of the 29 plants using emissions control, 26 discharge wastewater. The other three plants practice 100 percent recycle. Seventeen of the 26 discharging plants practice recycle of 90 percent or greater. Table V-5 (page 2364) shows water discharge rates in liters per troy ounce of precious metals, including silver, produced in the refinery.

The Agency sampled the wastewater from four discharging plants, three of which practice recycle of at least 90 percent. This raw

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wastewater contains priority metals and suspended solids above treatable concentrations. Raw wastewater sampling data are presented in Table V-17 (page 2402).

GOLD SOLVENT EXTRACTION RAFFINATE AND WASH WATER

Gold can be extracted from an impure raw material using an organic solvent and then recovered from the solvent as pure gold. The raffinate generated by this process can be discharged, and one plant discharges this waste stream as shown in Table V-6 (page 2366). After the pure gold is recovered, it is washed with water and this wash water is also discharged.

Following proposal, sampling data for this subdivision were acquired through a self-sampling effort specifically requested by the Agency. These data show treatable concentrations of antimony, cadmium, chromium, copper, lead, nickel, silver, thallium, zinc, ammonia, TSS, gold, palladium, and platinum.

GOLD SPENT ELECTROLYTE

Three plants use electrolytic refining as a purification step in secondary gold processing and discharge the spent electrolyte wastewater associated with this process. Water use and discharge rates are shown in Table V-7 (page 2366). No samples were taken of this waste stream however, the Agency believes it should be similar to gold precipitation and filtration wastewater because of contact with similar raw materials. This wastewater is expected to contain priority metals and TSS above treatable concentrations.

GOLD PRECIPITATION AND FILTRATION

Gold may be recovered by dissolving the raw material in strong acid such as aqua regia, filtering away the silver chloride, and precipitating the gold with a strong reducing agent such as chlorine, ferrous sulfate or sulfur dioxide gas. Gold sponge is recovered by filtering away the wastewater and washing the sponge with water one or more times to remove residual acid. This combined filtrate and wash water waste stream may be discharged via a cementation tank where either zinc or iron is added to recover additional precious metals, and then to treatment. The 28 plants with this waste stream are shown in Table V-8 (page 2367).

The Agency sampled this waste stream at four plants, one prior to cementation, and all four as combined wastewater after cementation. Only the plant sampled prior to cementation is presented in Table V-18 to characterize this raw wastewater because of the metallic replacement reactions and commingling of wastewater taking place in the cementation tank. As shown in Table V-22 (page 2440), the post-cementation data support the general characterization of gold precipitation and filtration wastewater data. Both show high priority metal concentrations, along with ammonia and TSS above treatable concentrations. If a

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - V

plant cements this wastewater with zinc, the effluent from cementation will contain high zinc concentrations.

PLATINUM PRECIPITATION AND FILTRATION

Platinum may be recovered by dissolving the raw material in acid, filtering away the impurities, and precipitating the platinum as an amine. The insoluble amine (yellow salt) is then separated from the solution by filtration. The filtrate may be combined with wash water, and sent to cementation or treatment. Eighteen plants recover platinum in this manner as shown in Table V-9 (page 2369).

Following proposal, sampling data for this subdivision were acquired through a self-sampling effort specifically requested by the Agency. These data show treatable concentrations of arsenic, cadmium, chromium, copper, lead, nickel, selenium, zinc, and ammonia.

PALLADIUM PRECIPITATION AND FILTRATION

Palladium may be recovered by dissolving the raw material in strong acid or base, filtering away impurities, precipitating the palladium as an amine, and filtering away the solution. The insoluble amine (red salt) may be reduced with a strong reducing agent to the pure metal sponge. The filtrate may be combined with wash water, and sent to cementation or treatment. Twenty plants recover palladium in this manner as shown in Table V-10 (page 2370).

The Agency sampled one of the discharging plants for three palladium batch discharges, as shown in Table V-19 (page 2426). The raw wastewater shows priority metals, ammonia, and TSS above treatable concentrations.

OTHER PLATINUM GROUP METALS PRECIPITATION AND FILTRATION

Three plants use a wet chemistry process similar to the type used to recover either platinum or palladium, to recover other platinum group metals including rhodium and iridium. All three plants discharge wastewater as shown in Table V-11 (page 2371).

Following proposal, sampling data for this subdivision were acquired through a self-sampling effort specifically requested by the Agency. These data show treatable concentrations of antimony, arsenic, cadmium, chromium, copper, lead, nickel, selenium, zinc, and ammonia.

SPENT SOLUTION FROM PGC SALT PRODUCTION

Four plants manufacture potassium gold cyanide (PGC) salt from pure gold and potassium cyanide solution. Excess cyanide solution may be discharged from this process. Water use and discharge rates are shown in Table V-12 (page 2371).

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The Agency sampled one plant for this waste stream, and the results are presented in Table V-20 (page 2436). Raw wastewater contains toxic metals, and free and complexed cyanide above treatable concentrations.

EQUIPMENT AND FLOOR WASH

Three plants reported an equipment and floor wash waste stream. This waste stream is discharged via cementation, to treatment. Table V-13 (page 2372) shows water use and discharge rates in liters per troy ounce of Precious metals, including silver, produced in the refinery. The Agency sampled this waste stream at one plant, and the data are presented in Table V-21 (page 2436). This wastewater contains priority metals, ammonia, and TSS above treatable concentrations.

PRELIMINARY TREATMENT

Based on information provided to the Agency after promulgation of the regulation for this subcategory, EPA agreed to add a new building block for the preliminary treatment process. This building block was omitted from the promulgated rule because EPA believed that the processing of basis materials was accounted for by the furnace wet air pollution building block (FWAP). However, subsequent to promulgation, EPA found that the FWAP building block does not reflect the raw material processing steps required for non-combustible, non-metallic-based basis materials. As discussed in Section III, details of the preliminary treatment steps are claimed confidential, therefore water use and discharge rates are not presented in this document. SECONDARY PRECIOUS METALS SUBCATEGORY SECT - V

TABLE V-1

WATER USE AND DISCHARGE RATES FOR FURNACE WET AIR POLLUTION CONTROL

(1/troy ounce of precious metals, including silver, incinerated or smelted)

Plant <u>Code</u>	Percent Recycle	Production Normalized Water <u>Use</u>	Production Normalized Discharge Flow
1081	98.2	7.26	0.131
1038	0	116	116
1138		27.6	27.6
1105	<u>></u> 90	NR	4.5
1112	0	137	137
1094	100	NR	. 0
1084	100 ·	NR	0
1095	Dry		
1153	Dry		
1163	Dry	٢	
1020	Dry		
1019	Dry		
1082	Dry		
1134	Dry		
1071	Dry		
1088	Dry		
1051	Dry		
1045	Dry		

NR - Data not reported.

SECONDARY PRECIOUS METALS SUBCATEGORY

TABLE V-2

WATER USE AND DISCHARGE RATES FOR RAW MATERIAL GRANULATION

(liters/troy ounce of precious metals in the granulated raw material)

Plant Code	Percent Recycle	Production Normalized Water Use	Production Normalized Discharge Flow
1008	0	8.67	8.67
1094	0	4.0	4.0
1112	0	0.30	0.30
1082	100	Unknown	0

TABLE V-3

WATER USE AND DISCHARGE RATES FOR SPENT PLATING SOLUTIONS

(1/liter of raw material spent plating solution)

Plant Code	Percent Recycle	Production Normalized Water Use	Production Normalized Discharge Flow
1002	0	1.0	1.0
1163	0	1.0	1 0
1094	0	1.0	1.0
1092	0	1.0	1.0
1023	0	1.0	1.0
1128	0	1.0	1.0
1083	0	1.0	1.0
1167	NR	NR	NR
1071	0	1.0	1.0
1034	0	1.0	1.0
1067	. 0	1.0	1.0
1065	0	1.0	1.0

NR - Data not reported.

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - V

TABLE V-4

WATER USE AND DISCHARGE RATES FOR SPENT CYANIDE STRIPPING SOLUTIONS

(1/troy ounce of gold produced by cyanide stripping)

Plant Code	Percent Recycle	Production Normalized Water Use	Production Normalized Discharge Flow
1100	0	78.3	78.3
1034	0	7.63	7.63
1163	0	6.03	6.03
1067	0	2.92	2.92
1083	0	1.14	1.14
1026	. 0	0.631	0.631

TABLE V-5

WATER USE AND DISCHARGE RATES FOR REFINERY WET AIR POLLUTION CONTROL

(l/troy ounce of precious metals, including silver, produced in refinery)

Plant Code	Percent Recycle	Production Normalized Water Use	Production Normalized Discharge Flow
1100	0	107	107
1117	~ 0	42	42
1029	0	32.8	32.8
1020	<u>></u> 90	NR	14.2
1051	0	13.2	13.2
1147	75	39.4	9.85
1065	0	6.8	6.8
1067	90	46.4	4.64
1112	0	3.4	3.4
1091	NR	NR	3.32
1071	0	2.4	2.4
1105	<u>></u> 90	NR	2.3
1080	<u>></u> 90	NR	1.75
1115	<u>></u> 90	NR	1.665
1069	<u>></u> 90	NR	1.41
1008	<u>></u> 90	NR	1.1
1164	90	7.0	0.7
1083	<u>></u> 90	NR	0.67
1104	<u>></u> 90	NR	0.234
1138	<u>></u> 90	NR	0.21

SECONDARY PRECIOUS METALS SUBCATEGORY

TABLE V-5 (Continued)

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WATER USE AND DISCHARGE RATES FOR REFINERY WET AIR POLLUTION CONTROL

Plant <u>Code</u>	Percent Recycle	Production Normalized Water Use	Production Normalized Discharge Flow
1094	<u>></u> 90	NR	0.19
1165	<u>></u> 90	NIR	0.172
1082	99	7.2	0.072
1026	<u>></u> 90	NR	0.06
1072	<u>></u> 90	NR	0.036
1167	95	0.6	0.03
1053	100	NR	0
1128	100	NR	0
1034	100	NR	0

NR - Data not reported.

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - V

TABLE V-6

WATER USE AND DISCHARGE RATES FOR GOLD SOLVENT EXTRACTION RAFFINATE AND WASH WATER

(1/troy ounce of gold produced by solvent extraction)

Plant Code	Percent Recycle	Production Normalized Water Use	Production Normalized Discharge Flow	
1094	0	0.63	0.63	

TABLE V-7

WATER USE AND DISCHARGE RATES FOR GOLD SPENT ELECTROLYTE

(1/troy ounce of gold produced by electrolysis)

Plant <u>ode</u>	Percent Recycle	Production Normalized Water Use	Production Normalized Discharge Flow
1071	0	0.294	0.294
108?	0	0.0087	0.0087
1088	NR	NR	NR

NR - Data not reported.

TABLE V-8

WATER USE AND DISCHARGE RATES FOR GOLD PRECIPITATION AND FILTRATION WASTEWATER

(1/troy ounce of gold precipitated)

Plant ode	Percent Recycle	Production Normalized <u>Water</u> <u>Use</u>	Production Normalized Discharge Flow
1034	0	560.5	560.5
1100	0	404	404
1091	• • 0	69.1	69.1 -
1053	0	24.3	24.3
1165	0	7.98	7.98
1083	0	4.1	4.1
1067	0	3.34	3.34
1063	0	2.65	2.65
1082	0	2.5	2.5
1147	0	1.86	1.86
1110	0	0.815	0.815
1008	0	0.63	0.63
1138	0	0.341	0.341
1 0 65	0	0.312	0.312
1117	0	0.27	0.27
1153	0	0.144	0.144
1026	0	0.05	0.05
1020	NR	NR	0
1069	NR	NR	0

TABLE V-8 (Continued)

WATER USE AND DISCHARGE RATES FOR GOLD PRECIPITATION AND FILTRATION WASTEWATER

(1/troy ounce of gold precipitated)

Plant <u>Code</u>	Percent <u>Recycle</u>	Production Normalized <u>Water</u> <u>Use</u>	Norma	action alized rge Flow
1018	NR	NR		NR
1104	NR	NR		NR
1128	NR	NR		NR
1 164	NR	NR		NR
1029	NR	NR		NR
1167	NR	NR		NR
1072	NR	NR		NR
1115	NR	NR	- - 	NR
1071	NR	NR	1 - 4	NR

14

NR - Data not reported.

TABLE V-9

WATER USE AND DISCHARGE RATES FOR PLATINUM PRECIPITATION AND FILTRATION

(1/troy ounce of platinum precipitated)

Plant <u>Code</u>	Percent Recycle	Production Normalized <u>Water Use</u>	Production Normalized Discharge Flow
1020	0	354	354
1082	0	30.2	30.2
1069	0	10.4	10.4
1105	• 0	4.5	4.5
1147	0	0.58	0.58
10?1	NR	NR	NR
1018	NR	NR	NR
1063	NR	NR	NR
1072	NR	NR	NR
1115	NR	NR	NR
1117	NR	NR	NR
1104	NR	NR	NR
1156	NR	NR	NR
1138	NR	NR	NR
1080	NR	NR	NR
1088	NR	NR	NR
1153	NR	NR	NR
1134	NR	NR	NR

NR - Data not reported.

TABLE V-10

WATER USE AND DISCHARGE RATES FOR PALLADIUM PRECIPITATION AND FILTRATION

(1/troy ounce of palladium precipitated)

Plant <u>Code</u>	Percent Recycle	Production Normalized <u>Water</u> Use	Production Normalized <u>Discharge</u> Flow
1069	0	15.8	15.8
1147	0	4.58	4.58
1105	Ο	4.4	4.4
1112	0	3.9	3.9
1082	0	3.4	3.4
1138	0	1.53	1.53
1020	NR	NR	0
1153	NR	NR	NR
1018	NR	NR	NR
1128	NR	NR	NR
1029	NR	NR	NR
1072	NR	NR	NR
1115	NR	NR	NR
1117	NR	NR	NR
1104	NR	NR	NR
1156	NR	NR	NR
1080	NR	NR	NR
1071	NR	NR	NR
1088	NR	NR	NR
1051	NR	NR	NR
1051	NR	NR	NR

NR - Data not reported.

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - V

TABLE V-11

WATER USE AND DISCHARGE RATES FOR OTHER PLATINUM GROUP METALS PRECIPITATION AND FILTRATION

(1/troy ounce of other platinum group metals precipitated)

Plant <u>Code</u>	Percent Recycle	Production Normalized Water Use	Production Normalized Discharge Flow
1115	NR	NIR	NR
1051	NR	NIR	NR
1156	NR	NR	NR

NR - Data not reported.

TABLE V-12

WATER USE AND DISCHARGE RATES FOR SPENT SOLUTION FROM PGC SALT PRODUCTION

(1/troy ounce of gold contained in PGC product)

Plant Code	Percent Recycle	Production Normalized Water <u>Use</u>	Production Normalized Discharge Flow
1100	0	260	260
1112	0	1.5	1.5
1034	0	0.90	0.90
1128	NR	NR	NR

NR - Data not reported

TABLE V-13

WATER USE AND DISCHARGE RATES FOR EQUIPMENT AND FLOOR WASH

		Production	Production
Plant Code	Percent RecYcle	Normalized Water Use	Normalized Discharge Flow
1020	0	14.2	14.2
1105	0	1.0	1.0
1138	0	0.97	0.97

Table V-14

SECONDARY PRECIOUS METALS SAMPLING DATA FURNACE WET AIR POLLUTION CONTROL RAW WASTEWATER

			Stream Sample		Conc	Concentrations (mg/1)			
	Pollutant		Code	Typet	Source	Day 1	Day 2	<u>Dav 3</u>	
Toxic	Pollutants		• •				•	•	
1.	acenaphthene		189 5	1 2	ND ND	ND	ND	ND N D	
2.	acrolein	•	189 5	1	ND ND	ND	ND	N D ND	
3.	acrylonitrile		189 5	1	ND ND	ND	ND	ND ND	
4.	benzene		189 5	1 1	ND ND	ND	ND	N D ND	
5.	benzidine		189 5	1 2	ND ND	ND	ND	ND ND	
6.	carbon tetrachloride		189 5	1 1 1	ND ND	ND	ND	N D ND	
7.	chlorobenzene	-	189 5	1 1	ND ND	ND	ND	ND <0.010	
8.	1,2,4-trichlorobenzene		189	1 2	ND ND	ND	ND	N D ND	
9.	hexachlorobenzene	. .	189	1 2	ND ND	ND	ND	ND N D	

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT - V

SECONDARY PRECIOUS METALS SAMPLING DATA FURNACE WET AIR POLLUTION CONTROL RAW WASTEWATER

	Pollutant	Stream Code	Sample Typet	<u>Conc</u> Source	entration Day 1	ns (mg/1) Day 2	Day 3
		<u> </u>	<u></u>	bource		<u>Duy z</u>	<u>Day J</u>
Toxic	Pollutants (Continued)	•					
10.	1,2-dichloroethane	189	1	ND			ND
	-	5	1	ND	ND	ND	ND
11.	1,1,1-trichloroethane	189	1	0.01			0.015
		5	1	ND	ND	ND	ND
12.	hexachloroethane	189	. 1	ND			ND
• = •		5	2	ND	ND	ND	ND
13.	1,1-dichloroethane	189	1	ND			N D
		5	1	ND	ND	ND	ND
14.	1,1,2-trichloroethane	189	1	ND			ND
	, , ,	5	1.	ND	ND	ND	ND
15.	1,1,2,2-tetrachloroethane	189	1	ND			ND ·
		5	1	ND	ND	ND	ND
16.	chloroethane	189	1	ND			ND
		5	1	ND	ND	ND	ŇD
17.	bis(chloromethyl)ether	189	1	ND			ND
		5	1 .	ND	ND	ND	ND
18.	bis(2-chloroethyl)ether	189	1	ND			ND
		5	2	ND	ND	ND ·	ND

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SECONDARY PRECIOUS METALS SAMPLING DATA FURNACE WET AIR POLLUTION CONTROL RAW WASTEWATER

	Stream	Sample	Conc	entratio	n s (mg/1)	
Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3
<u>Foxic Pollutants</u> (Continued)					2 C	
19. 2-chloroethyl vinyl ether	189 5	1 1	ND ND	ND	ND	ND ND
20. 2-chloronaphthalene	189 5	1 2	ND ND	ND	ND	N D ND
2 . 2,4,6-trichlorophenol	- 189 5	1 2	ND ND	ND	ND	ND ND
22. p-chloro-m-cresol	189 5	1 2	ND ND	ND	ND	N D ND
23. chloroform	189 5	1 1	ND 0.050	0.010	<0.010	ND 0.020
24. 2-chlorophenol	189 5	1 2	ND ND	ND	ND	N D ND
25. 1,2-dichlorobenzene	189 5	1. 2	ND ND	ND	ND	ND ND
26. 1,3-dichlorobenzene	189 5	1 2	ND ND	ND	ND	N D ND
27. 1,4-dichlorobenzene	189 5	1 2	ND ND	ND	ND /	ND ND

SECONDARY PRECIOUS METALS SAMPLING DATA FURNACE WET AIR POLLUTION CONTROL RAW WASTEWATER

		Stream	Sample	Cong	entration	e (mg/l)		E S S S S S S
	<u>Pollutant</u>	<u>Code</u>	Typet	Source	Day 1	Day 2	Day 3	CONI
<u>Toxic</u>	Pollutants (Continued)							SECONDARY
28.	3,3'-dichlorobenzidine	189 5	1 2	ND ND	ND	ND	ND ND	
29.	1,1-dichloroethylene	189 5	1	ND ND	ND	ND	N D ND	PRECIOUS
30.	1,2- <u>trans</u> -dichloroethylene	18 <u>9</u> 5	1 1	ND ND	ND	ND	ND ND	METALS
31.	2,4-dichlorophenol	189 5	1 2	ND ND	ND	ND	N D ND	
32.	1,2-dichloropropane	189 5	1 · 1	ND ND	ND	ND	ND ND	SUBCATEGORY
33.	1,3-dichloropropene	189 5	1	ND ND	ND	ND	N D ND	ORY
34.	2,4-dimethylphenol	189 5	1 2	ND ND	ND	ND	ND ND	SECT
35.	2,4-dinitrotoluene	189 5	1 2	ND ND	ND	ND	N D ND	Г - V
36.	2,6-dinitrotoluene	189 5	1 2	ND ND	ND	ND	ND ND	٤

SECONDARY PRECIOUS METALS SAMPLING DATA FURNACE WET AIR POLLUTION CONTROL RAW WASTEWATER

Pollut	ant	Stream Code	Sample Typet	Conc Source	<u>entration</u> Day 1	ns (mg/1) Day 2	Day 3
oxic Pollutants	(Continued)		•				
37. 1,2-dipheny	lhydrazine	189 5	1 2	ND ND	ND	ND	ND ND
38. ethylbenzen	e	189 5	1	ND ND	ND	ND	N D ND
9. fluoranthen	e	189 5	1 2	ND ND	ND	ND	ND ND
0. 4-chlorophe	nyl phenyl ether	189 5	1 2	ND ND	ND	ND	N D ND
1. 4-bromophen	yl phenyl ether	189 5	1 2	ND ND	ND	ND	ND ND
. bis(2-chlor	oisopropyl)ether	189 5	1 2	ND ND	ND	ND	N D ND
bis(2-choro	ethoxy)methane	189 5	1 2	ND ND	ND	ND	ND N D
4. methylene c	hloride	189 5	1]-	ND <0.01	<0.01	<0.01	ND <0.01
45. methyl chlo	ride (chloromethane)	189 5	1	ND ND	ND	ND	ND ND

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SECONDARY PRECIOUS METALS SUBCATEGORY

SECONDARY PRECIOUS METALS SAMPLING DATA FURNACE WET AIR POLLUTION CONTROL RAW WASTEWATER

Pollutant	Stream Code	Sample Typet	Conc Source	<u>Day 1</u>	<u>ns (mg/1)</u> Day 2	Day 3
Toxic Pollutants (Continued)						
6. rethyl bromide (bromomethane)	189 5	1 1	ND ND	ND	ND	ND ND
47. bromoform (tribromomethane)	189 5	1 1	ND ND	ND -	ND	N D ND
48. iichlorobromomethane	189 5	1	ND <0.01	<0.01	<0.01	ND ND
49. trichlorofluoromethane	189 5	1 . 1	ND ND	ND	ND	N D ND
50. dichlorodifluoromethane	189 5	1	ND ND	ND	ND	ND ND
51. chlorodibromomethane	189 5	1	ND ND	ND	ND	N D ND
52. hexachlorobutadiene	189 5	1 2	ND ND	ND	ND	ND ND
53. hexachlorocyclopentadiene	189 5	1 2	ND ND	ND	ND	N D ND
54. isophorone	189 5	1 2	ND ND	ND	ND	ND ND

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT

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SECONDARY PRECIOUS METALS SAMPLING DATA FURNACE WET AIR POLLUTION CONTROL RAW WASTEWATER

	·			• .			
		Stream	· · · · · · · · · · · · · · · · · · ·	Concentrations (mg/l)			
	Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3
oxic	Pollutants (Continued)						
55.	naphthalene	189 5	1 2	ND ND	ND	ND	ND ND
6.	nitrobenzene	189	1	ND		нD	ND
		5	2	ND	ND	ND	ND
57.	2-nitrophenol	189 5	1 2	ND ND	ND	ND	ND ND
8.	4-nitrophenol	189 5	1 2	ND ND	ND	ND	N D ND
9.	2,4-dinitrophenol	189 5	1 2	ND ND	ND	ND	ND ND
0.	4,6-dinitro-o-cresol	189 5	1 2	ND ND	ND	ND	N D ND
1.	N-nitrosodimethylamine	189 5	1 2	ND ND	ND	ND	ND ND
2.	N-nitrosodiphenylamine	189 5	1 2	ND ND	ND	ND	0.01 ND
3.	N-nitrosodi-n-propylamine	189 5	1 2	ND ND	ND	ND	ND ND

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SECONDARY PRECIOUS METALS SUBCATEGORY

SECONDARY PRECIOUS METALS SAMPLING DATA FURNACE WET AIR POLLUTION CONTROL RAW WASTEWATER

		Stream	Sample	Concentrations (mg/1)				BCC
	Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3	ECONDARY
Toxic	<u>Pollutants</u> (Continued)					•		
64.	pentachlorophenol	189 5	1 2	ND ND	ND	ND	ND ND	PRECIOUS
65.	phenol	189 5	1 2	ND ND	<0.01	<0.01	0.013 ND	
66.	bis(2-ethylhexyl) phthalate	189 5	1 2	0.026 0.02	<0.01	<0.01	0.034 <0.01	METALS
67.	butyl benzyl phthalate	189	1 2	ND ND	ND	ND	N D ND	SUBC
68.	di-n-butyl phthalate	189 5	1 2	ND <0.01	<0.01	<0.01	0.002 <0.01	SUBCATEGORY
69.	di-n-octyl phthalate	189 5	1 2	ND ND	ND	ND	0.003 ND	RY
70.	diethyl phthalate	189 5	1 2	ND <0.01	<0.01	<0.01	ND <0.01	SECT
71.	dimethyl phthalate	189 5	1 2	ND ND	ND	ND	0.006 ND	י ל
72.	benzo(a)anthracene	189	1 2	ND ND	ND	ND	ND ND	

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SECONDARY PRECIOUS METALS SAMPLING DATA FURNACE WET AIR POLLUTION CONTROL RAW WASTEWATER

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	Stream	Sample	Conc	Concentrations (mg/l)		
Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3
Toxic Pollutants (Continued)	ي د م ۲۰ في ۲۰	•	مع آباً م مع من قرار مع م م	21 की 75 8 96 ₩ 152 - 1		
73. benzo(a)pyrene	189 5	1	ND ND	ND	ND	ND ND
74. benzo(b)fluoranthene	189 5	1 2	ND ND	ND	ND	N D ND
5. benzo(k)fluoranthane	189 5	1 . 2 .	ND. ND.	ND	ND	ND ND
76. chrysene	189 5	1 2	ND NDT	ND	ND	N D ND
7. acenaphthylene	189 5	1 2	ND ND.	ND	ND	ND ND
78. anthracene (a)	189 5	1 2	ND ND	ND	ND	N D ND
79. benzo(ghi)perylene	189 5	1	ND ND	ND	ND	ND ND
80. fluorene	189		ND ND	ND	ND	N D ND
81. phenanthrene (a)	189	1	ND		• • • •	ND

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SECONDARY PRECIOUS METALS SUBCATEGORY

SECT

SECONDARY PRECIOUS METALS SAMPLING DATA FURNACE WET AIR POLLUTION CONTROL RAW WASTEWATER

	<u>Pollutant</u>	Stream Code	Sample Typet	Conce Source	Day 1	s (mg/1) Day 2	Day 3
" xic	Pollutants (Continued)						
. 2.	dibenzo(a,h)anthracene	189 5	1 2	ND ND	ND	ND	ND ND
83.	indeno (1,2,3-c,d)pyrene	189 5	1 2	ND ND	ND	ND	N D ND
84.	pyrene	189 5	1 2	ND ND	ND	ND	ND . ND
85.	tetrachloroethylene	189 5	1 1	ND ND	ND	ND	N D ND
86.	toluene	189 5	1 1	ND ND	ND	ND	ND ND
87.	trichloroethylene	189 5	1 · · · · · · · · · · · · · · · · · · ·	ND ND	ND	ND	N D ND
88.	vinyl chloride (chloroethylene)	189 5	1 1	ND ND	ND	ND	ND ND
114.	antimony	200 189 5	2 1 2	<0.01 <0.003 <0.003	0.004	<0.01 <0.003	0.19 <0.093

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SECONDARY PRECIOUS METALS SUBCATEGORY

SECT -

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SECONDARY PRECIOUS METALS SAMPLING DATA FURNACE WET AIR POLLUTION CONTROL RAW WASTEWATER

	Stream	Sample	Conc	entration	s (mg/l)		
Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3	
Toxic Pollutants (Continued)				• ,			
	-						
115. arsenic	200	2 1	<0.01		<0.01		
	189	1 N 1 N 1 N	<0.003			0.025	
	5	2	<0.005	0.004	<0.005	0.005	
117. beryllium	200	2	<0.01		<0.01		
	189	1	<0.01		•••••	<0.01	
	5	2	č0.0002	0.0006	<0.0002	<0.0002	
118. cadmium	200	2	<0.05		<0.05		
	189	··· 1	<0.01	-	10.05	0.61	
	. 5	2	0.0002	0.001	0.020	0.003	
		-					
119. chromium (total)	200	2	<0.05		<0.05		
	189	Î	<0.01			1.1	
	5	2	0.003	0.003	0.001	0.003	
120. copper	200	2	<0.05		0.05		
	189	1	<0.01			11.0	
	5	2	0.017	0.026	0.140	0.040	
121. cyanide (total)	200	1	0.05	*	0.095		
(LII Cyanide (Cocai)	189	1	<0.02		0.075	<0.02	
	5	1	0.052	0.008	0.090	0.150	
20 1	200	0	10 10		(0 10 ¹		
122. lead	200	2 1	<0.10		<0.10	· • • •	
	189	1 2	<0.10	0 0 0 7	0 00	3.0	
	5	Z	0.030	0.037	0.80	0.092	

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SECONDARY PRECIOUS METALS SUBCATEGORY

SECT -

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SECONDARY PRECIOUS METALS SAMPLING DATA FURNACE WET AIR POLLUTION CONTROL RAW WASTEWATER

Pollutant	Stream Code	Sample Typet	Concentration Source Day 1	ons (mg/1) Day 2	Day 3
Toxic Pollutants (Continued)			•		
123. mercury	200 189 5	2 1 2	0.0002 0.002 0.0002 0.0003	0.015 3 <0.0001	<0.0002 <0.0001
124. nickel	200 189 5	2 1 2	<0.2 0.075 0.020 0.017	<0.2 0.016	30.0 0.014
125. selenium	200 189 5	2 1 2	<0.1A <0.003 <0.002 120.0	<0.1A <0.002	0.007 <0.002
6. silver	200 189 5	2 1 2	<0.01 <0.0005 <0.0002 0.001	0.05 0.003	0.13 0.004
27. thallium	200 189 5	2 1 2	<0.01 <0.002 <0.001 <0.001	<0.01 0.004	<0.002 <0.30
28. zinc	200 189 5	2 1 2	0.10 2.5 <0.010 0.110		,100 0.160

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT -<

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SECONDARY PRECIOUS METALS SAMPLING DATA FURNACE WET AIR POLLUTION CONTROL RAW WASTEWATER

	Stream Code	Sample Typet	Concentrations (mg/l)			
<u>Pollutant</u>			Source	Day 1	Day 2	Day 3
Nonconventional Pollutants			•			· · · · ·
acidity	200 189	2 1	<1 <1	• • •	<1	<1
alkalinity	200 189 5	2 1 2	98 127 16	31	75 0	91 33
aluminum	200 189	2 1	0.20 <0.050		0.3	8.3
ammonia nitrogen	200 189	2 1	0.04 <0.01	. ·	0.24	<0.01
barium	200 189	2 1	<0.050 0.070		<0.05	0.37
boren	200 189	1 1	<0.10 <0.009	·	<0.10	2.1
calcium	200 189 5	2 1 2	37.7 11 13	9.9	38.4 14	710 9.7
chemical oxygen demand (COD)	200 189	2 1	<5		100	490

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT -

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SECONDARY PRECIOUS METALS SAMPLING DATA FURNACE WET AIR POLLUTION CONTROL RAW WASTEWATER

<u>Pollutant</u>	Stream Code	Sample Typet	Conc Source	entrations Day 1	(mg/1) Day 2	Day 3 SECONDARY
Nonconventional Pollutants (Continued)					DAR
chloride	200 189	2 1	14 52		63 1	
cobalt	200 189	2 1	<0.050 <0.006		<0.05	,000 PRECIOUS
fluoride	200 189	2 1	0.28 1.1		0.21	1.8 METALS
gold	5	1	0.025	0.086	0.48	0.42
iron	200 189 5	2 1 2	<0.050 0.31 0.29	0.32	0.2 0.52	SUBCATEGORY
magnesium	200 189 5	1 1 1	8.50 2.4 3.1	3.2	8.6 3.4	GORY 15 3.2
manganese	200 189	2 1	<0.050 <0.01		<0.05	1.2 CF
molybdenum	200 189	2 1	<0.050 <0.002		<0.05	، 0.061 <
phenolics	5	1	0.15	0.25	0.061	0.005

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SECONDARY PRECIOUS METALS SAMPLING DATA FURNACE WET AIR POLLUTION CONTROL RAW WASTEWATER

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	Straom	Sample	Concentrations (mg/l)				Ë
Pollutant	Stream <u>Code</u>	Sample Typet	Source	Day 1	(mg/1) Day 2	Day 3	ECONDARY
Nonconventional Pollutants (Continue	ed)		r			•	ARY
phosphate	200 189	2 1	14	•	<0.9	<4	PRECIOUS
sodium	200 189	2 1	9 54	- - -	13.3	, 100	
sulfate	200 189	2 1	57 13		27	920	METALS
tin	200 189	2 1	<0.050 <0.12	· ·	<0.05	<0.12	· .
titanium	200 189	2 1	<0.005 <0.005		<0.05	<0.005	SUBCATEGORY
total organic carbon (TOC)	200 189	2 1	4.3 43		4	94	ORY
total solids (TS)	200 189	2 1	- 380 410		6,000 35	,000	SECT
vanadium	200 189	2	<0.050 <0.003		<0.05	<0.003	1
	<i>,</i>						

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SECONDARY PRECIOUS METALS SAMPLING DATA FURNACE WET AIR POLLUTION CONTROL RAW WASTEWATER

	Stream	tream Sample <u>Conce</u>		Concentrations (mg/1)			
Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3	
Nonconventional Pollutants (Continued)						
yttrium	200 189	2 1	<0.50 <0.002		<0.05	0.004	
Conventional Pollutants		•					
oil and grease	200 189	1	<1 <1		2	<1	
	5	1	1.6	<1	<1	< 1.	
total suspended solids (TSS)	200 189	2 1	60 8	0		,100 13	
	5 *	Z	0	~ U	8	13	
pH (standard units)	200 189	2	7.5		7.3	5.96	
	5	2	6.8	6.6	3.4	7.1	

iSample Type Code: 1 - One-time grab 2 - Manual composite during intermittent process operation

(a) Reported together.

A - Detection limit raised due to interference.

SECONDARY PRECIOUS METALS SUBCATEGORY SECT

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Table V-15

SECONDARY PRECIOUS METALS SAMPLING DATA SPENT PLATING SOLUTIONS RAW WASTEWATER

	Stream Sample		Concentrations (mg/1)				
Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3	
Te c Pollutants		· · ·			."		
acenaphthene	6	1	ND	ND	ND	ND I	
2 acrolein	6	1	ND	ND	ND	ND	
3. acrylonitrile	6	. 1	ND	ND	ND	ND	
4. benzene	6	1	ND	ND.	<0.01	ND	
5. benzidine	6	1 1	ND	ND	ND	ND	
6. carbon tetrachloride	6	1	ND	ND	ND	ND	
7. chlorobenzene	6	1	ND	ND	ND	ND	
8. 1,2,4-trichlorobenzene	6	. 1	ND	ND	ND	ND	
9. hexachlorobenzene	6	1	ND	ND	ND	ND	
10. 1,2-dichloroethane	6	1.	ND	ND	ND	ND	
11. 1,1,1-trichloroethane	6	1	ND	ND	ND	ND	
12. hexachloroethane	6	1	ND	ND	ND	ND	
13. 1,1-dichloroethane	6	1	ND	ND	ND	ND	
14. 1,1,2-trichloroethane	6	1	ND	ND	ND	ND	

SECONDARY PRECIOUS METALS SAMPLING DATA SPENT PLATING SOLUTIONS RAW WASTEWATER

	Pollutant	Stream Code	Sample Typet	Conc Source	entrations Day 1	B (mg/1) Day 2	Day 3	SECONDARY
<u>Toxic</u>	Pollutants (Continued)							DAR
15.	1,1,2,2-tetrachloroethane	6	1	ND	ND	ND	ND	
16.	chloroethane	6	1	ND	ND	ND	ND	REC
:7.	bis(chloromethyl)ether	6	1	ND	ND	ND	ND	PRECIOUS
18.	bis(2-chloroethyl)ether	6	1	ND	ND	ND	ND	
19.	2-chloroethyl vinyl ether	6	1	ND	ND	ND	ND	METALS
20	2-chloronaphthalene	6	1	ND	ND	ND	ND	
21.	2,4,6-trichlorophenol	6	1	ND	ND	ND	ND	SUBCATEGORY
22.	p-chloro-m-cresol	6	1 .	ND	ND	ND	ND	TEGC
23.	chloroform	6	1	ND	ND	ŇD	ND	ORY
24.	2-chlorophenol	6	. 1	ND	<0.01	ND	ND	
25.	1,2-dichlorobenzene	6	1	ND	ND	ND	ND	SECT
26.	1,3-dichlorobenzene	6	1	ND	ND	ND	ND	ł
27.	1,4-dichlorobenzene	6	1	ND	ND	ND	ND	4
28.	3,3'-dichlorobenzidine	6	1	ND	ND	ND	ND	

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SECONDARY PRECIOUS METALS SAMPLING DATA SPENT PLATING SOLUTIONS RAW WASTEWATER

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SECONDARY PRECIOUS METALS SUBCATEGORY NECH

SECONDARY PRECIOUS METALS SAMPLING DATA SPENT PLATING SOLUTIONS RAW WASTEWATER

•	Pollutant	Stream <u>Code</u>	Sample Typet	<u>Conc</u> Source	centration Day 1	<u>ns (mg/l)</u> Day 2	Day 3	SECONDARY
Toxi	Pollutants (Continued)		•					ZUNC
43.	bss(2-choroethoxy)methane	6	1	ND	ND	ND	ND	NRY
44.	methylene chloride	6	1	<0.01	<0.01	<0.01	<0.01	PRE
45.	methyl chloride (chloromethane)	6	1	ND	ND	ND	ND	PRECIOUS
46.	methyl bromide (bromomethane)	6	1	ND	ND	ND	ND	
47.	bromoform (tribromomethane)	6	1	ND	ND	ND	ND .	METALS
48.	dichlorobromomethane	6	1	NĎ	ND	ND	ND	
49.	trichlorofluoromethane	6	1	ND	ND	ND	ND	SUBCATEGORY
50.	dichlorodifluoromethane	6	1	ND	ND	ND	ND	ATEO
51.	chlorodibromomethane	6	1	ND	ND	ND	ND	JORY
5 2.	hexachlorobutadiene	6	1	ND	ND	ND	ND	
53.	hexachlorocyclopentadiene	6	. 1	ND	ND	ND	ND	SECT
54.	isophorone	6	1	ND	ND	ND	ND	Ц Г
55.	naphthalene	6	1	ND	ND	ND	ND	4
56.	nitrobenzene	6	1	ND	ND	ND	ND	

SECONDARY PRECIOUS METALS SAMPLING DATA SPENT PLATING SOLUTIONS RAW WASTEWATER

	Stream	Sample	the second s	entratio			E N
<u>Pollutant</u>	Code	Typet	Source	<u>Day 1</u>	Day 2	Day 3	CO
oxic Pollutants (Continued)			•	• • •		•	ECONDARY
57. 2-nitrophenol	6	1	ND	ND	0.01	<0.01	
58. 4-nitrophenol	6	1	ND	ND	ND	ND	PREC
59. 2,4-dinitrophenol	6	1	ND	ND	ND	ND	PRECIOUS
60. 4,6-dinitro-o-cresol	6	1 -	ND	ND	ND	NID	
61. N-nitrosodimethylamine	6	1.	ND	ND	ND	ND	METALS
62. N-nitrosodiphenylamine	6	`1	ND	ND	ND	· NĐ	
63. N-nitrosodi-n-propylamine	6	1	ND	ND	ND	ND	JBCA
64. pentachlorophenol	6	1	ND	ND	ND	ND	SUBCATEGORY
65. phenol	6	. 1	ND	0.17	0.45	0.65	ORY
66. bis(2-ethylhexyl) phthalate	6	1	0.02	0.06	0.10	0.02	
67. butyl benzyl phthalate	6	1	ND	ND	ND	ND	SECT
68. di-n-butyl phthalate	6	1	<0.01	<0.01	<0.01	ND	۲, ۱,
69. di-n-octyl phthalate	6	1	ND	ND	ND	ND	4
70. diethyl phthalate	6	1	<0.01	<0.01	<0.01	<0.01	

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SECONDARY PRECIOUS METALS SAMPLING DATA SPENT PLATING SOLUTIONS RAW WASTEWATER

Pollutant		Stream Code	Sample Typet	<u>Conc</u> Source	entration Day 1	<u>s (mg/l)</u> Day 2	Day 3	SEC
Toxic Pollutants (Co	ntinued)		<u>=7F=:</u>					SECONDARY
71. dimethyl phtha		6	1	ND	NĎ	ND	ND	
72. benzo(ą)anthra		6	1	ND	ND	ND	ND	PRECIOUS
73. benzo(a)pyrene		6	1	ND	ND	ND	ND	CIO
74. benzo(b)fluora		6	1	ND	ND	ND	ND	
75. benzo(k)fluora		6	1	ND	ND	ND	ND	METALS
		6'	1	ND	ND	ND	ND	
76. chrysene			. I 1			ND	ND	SUBCATEGORY
77. acenaphthylene		6		ND	ND			JATE
78. anthracene	(a)	6	1	ND .	ND	ND	ND	GOE
79. benzo(ghi)pery	lene	6	1	ND	ND	ND	ND	A2
80. fluorene		6	1	ND	ND	ND	ND	τn
81. phenanthrene	(a)	6	1	ND	ND	ND	ND	SECT
82. dibenzo(a,h)an	thracene	6	1	ND	ND	ND	ND	I
83. indeno (1,2,3-	c,d)pyrene	6	1	ND	ND	ND	ND	4
84. pyrene		6	1	ND	ND	ND	ND	

SECONDARY PRECIOUS METALS SAMPLING DATA SPENT PLATING SOLUTIONS RAW WASTEWATER

		Stream	Sample	e Concentrations (mg/l)					
<u>Po1</u>	lutant	Code	Typet	Source	Day 1	Day 2	Day 3		
"oxic Pollutan	<u>ts</u> (Continued)				· · ·				
85. tetrachl	oroethylene	6	- 1	ND	ND	ND	ND		
36. toluene		6	1	ND	<0.01	<0.01	<0.01		
87. trichlor	oethylene	6	1	ND	ND	ND	ND		
38. vinyl ch	loride (chloroethylene)	6	1	ND	ND	ND	ND		
4. antimony		6 701	1	<0.003 <0.003	3.5 <0.003	1.0	5.2		
15. arsenic		6 701	1 	<0.005 <0.002	0.25 <0.003	2.2	1.0		
117. berylliu	m	6 701	1	<0.0002 0.002	0.09 0.005	0.46	0.17		
118. cadmium		6 701	1	0.0002 0.014	0.74	1.6	0.48		
119. chromium	(total)	6 701	1 1	0.003 0.015	20.0 0.14	22.0	14.0		
chromium	(hexavalent)	701	1		<0.02	•	- 2		

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SECONDARY PRECIOUS METALS SUBCATEGORY

SECONDARY PRECIOUS METALS SAMPLING DATA SPENT PLATING SOLUTIONS RAW WASTEWATER

	Stream	Sample						
Pollutant	Code	<u>Typet</u>	Source	<u>Day 1</u>	Day 2	Day 3	CO	
Oxic Pollutants (Continued)							SECONDARY	
20. copper	6 701	1 1	0.017 2.3	12.0	340.0	130.0		
21. cyanide (total)	6 701	1 - 1	0.052 0.41	170.0 100	11.0	25.0	PRECIOUS	
cyanide (free)	701	1	0.26	29				
122. lead	6 701	1 1	0.03 <0.08	3.4 <0.084	9.7	2.1	METALS	
123. mercury	6 701	1 1	0.0002 0.0007	<0.0001 0.0004	<0.0001	<0.0002	SUBCA	
124. nickel	6 701	1 1	0.02 0.25	56.0 1.1	700.0	1.8	SUBCATEGORY	
125. selenium	6 701	1 1	<0.002 <0.002	<0.002 0.18	<0.002	<0.002		
126. silver	6 701	1 1	<0.0002 <0.0005	0.26 <0.0005	0.27	0.28	SECT	
127. thallium	6 701	1	<0.001 <0.002	1.2 <0.002	0.91	0.90	- <	

SECONDARY PRECIOUS METALS SUBCATECORV

SECONDARY PRECIOUS METALS SAMPLING DATA SPENT PLATING SOLUTIONS RAW WASTEWATER

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Pollutant	Stream Sam Code Typ		crations (mg/1) M Day 1 Day 2 Day 3
Toxic Pollutants (Continued)	<u>Code</u> <u>Typ</u>	<u>et Source I</u>	Contactions (mg/1) Cay 1 Day 2 Day 3 CONDARY
128. zinc	6 1 701 1	<0.01 9,600 0.055 <0	
Nonconventional Pollutants			
alkalinity	6 1	16 66,000	J 39,000 12,000
aluminum	701 1	0.86 2,100	METALS
barium	701 1	0.056 0	1.012
boron	701 1	<0.009 <0	.009
calcium	6 1 701 1		0.009 BBCA 2.2 7.6 4.4 田田GORY 2.1 GORY
cobalt	701 1	0.044 2	. 1
old	6 1 701 1	0.025 5 0.15 15	
iron	6 1 701 1		•.7 6.0 0.53 H
magr. sium	6 1 701 1		 .8 1.6 0.47 .99
mangaaese	701 1	0.013 0	.043

SECONDARY PRECIOUS METALS SAMPLING DATA SPENT PLATING SOLUTIONS RAW WASTEWATER

<u>Pollutant</u>	Stream <u>Code</u>	Sample Typet	<u>Concentrations (mg/1)</u> Source Day 1 Day 2 Day 3
Nonconventional Pollutants (Continued))		· · · · · · · · · · · · · · · · · · ·
molybdenum	701	1	0.029 <0.002
phenolics	6	1	0.15 0.02 0.07 0.37
sodium	701	1	13 1,500
c in	701	1	<0.12 <0.12
tanium	701	1	0.12 0.13
anadium	701	1	0.073 0.094
ystrium	701	1	<0.002 <0.002
nventional Pollutants	-		
1 and grease	6	1	1.6 <1 <1
tal suspended solids	6	1	600 1,200 520
pH (standard units)	6 701	1	6.8 12.0 12.3 12.4 7 10

TSample Type Code: 1 - One-time grab

(a) Reported together.

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Table V-16

SECONDARY PRECIOUS METALS SAMPLING DATA SPENT CYANIDE STRIPPING SOLUTION RAW WASTEWATER

		Stream Sample			centrations	(mg/1)	
	Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3
<u>Toxic</u>	Pollutants						•
114.	antimony	702	1	<0.003	<0.003		
115.	arsenic	702	. 1 -	<0.002	0.11		
117.	beryllium	702	1	0.002	2.4		
118.	cadmium	702	1	0.014	7.6		•
î19.	chromium (total)	702	1.	0.015	0.12		
.0.	copper	702	1	2.3	5,000		: :
i21.	cyanide (total) cyanide (free)	702 702	1	0.41 0.26	9,897 40		
122.	lead	702	1	<0.08	<0.08		•
123.	mercury	702	e 1	0.0007	0.0004		
124.	nickel	702	1	0.25	890	· · · · · · · · · · · · · · · · · · ·	
125.	selenium	702	1	<0.002	0.18	•	
126.	silver	702	· 1	<0.0005	<0.0005		
127.	thallium	702	1	<0.002	<0.002	, ·	
128.	zinc	702	1	0.055	56		

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SECONDARY PRECIOUS METALS SAMPLING DATA SPENT CYANIDE STRIPPING SOLUTION RAW WASTEWATER

<u>Pollutant</u>	Stream Code	Sample Typet	Con Source	centrations (mg/1) Day 1 Day 2	Day 3
Nonconventional Pollutants					
aluminum	702	1	0.86	280	
barium	702	1	0.056	0.072	
boron	702	1	<0.009	<0.009	
calcium	702	1	4.2	0.42	
cobalt	702	1	0.044	73	
gold	702	. 1	0.15	8.9	
iron	702	. 1	0.94	23	
magnesium	702	1	1.3	0.19	
manganese	702	1	0.013	0.045	
molybdenum	702	1	0.029	12	
sodium	702	1	13	2,500	
tin	702	1	<0.12	<0.12	
"itanium	702	1	0.12	0.59	
anadium	702	1	0.073	0.51	
yttrium	702	1	<0.002	<0.002	

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT

SECONDARY PRECIOUS METALS SAMPLING DATA SPENT CYANIDE STRIPPING SOLUTION RAW WASTEWATER

	Stream	Sample	Sample <u>Concentrations (mg/l)</u>			
Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3
Conventional Pollutants						
pH (standard units)	702	1	7	10	•	* .
				· · · ·	·	
		- 		2 54 -		
•		•		-		
			ی رو - ب			
	· • • •	n				
			· · · · · ·			
to an in man Code, 1 - One-time gra	h			н. ¹ н	2	4

SECONDARY PRECIOUS METALS SUBCATEGORY

SECT -

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Sample Type Code: 1

2401

- One-time gra

Table V-17

SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

D-11	Stream			<u>Concentrations (mg/l)</u>				
<u>Pollutant</u>	Code	<u>Typet</u>	Source	Day 1	<u>Day 2</u>	Day 3		
Toxic Pollutants		<i>4</i> ,						
1. acenaphthene	187	1	ND			ND		
-	4	/ 2 1	ND	ND	ND	ND		
	821	1		ŅD		,		
. acrolein	187	1	ND			ND		
	. 4	1	ND	ND	ND	ND		
	821	• 1	'.	ND	*			
acrylonitrile	187	· 1	ND			ND		
	4	. 1	ND	ND	ND	ND		
	821	1		ND	1. 1. se a			
4. benzene	187	1	ND			ND		
	4	1	ND	<0.01	ND	ND		
	821	1		ND				
5. benzidine	187	1	ND			ND		
·	4	2	ND	ND	ND	ND		
	821	1		ND	112	ЦЪ		
6. carbon tetrachloride	187	1	ND	-	· · · ·	0.210		
	4	1	ND	ND	ND	ND		
	821	1.		ND		11-		
7. chlorobenzene	187	1	ND			ND		
	4	1	ND	ND	ND	ND		
	821	1		ND				

SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

ι.	<u>Pollutant</u>	Stream Code	Sample <u>Typet</u>	Conc Source	entration Day 1	<u>s (mg/l)</u> Day 2	Day 3
<u>Toxic</u>	Pollutants (Continued)		· · · · ·		•		
8.	1,2,4-trichlorobenzene	187 4 821	1 2 1	ND ND	ND ND	ND	ND ND
9.	hexachlorobenzene	187 4 821	1 2 1	ND ND	ND ND	ND	ND ND
10.	1,2-dichloroethane	187 4 821	1 1 1	ND ND	ND ND	ND	ND <0.01
1.,	1,1,1-trichloroethane	187 4 821	1 1 1 1	0.01 ND	ND ND	ND	ND ND
12.	hexachloroethane	187 4 821	1 2 1	ND ND	ND ND	ND	ND ND
13.	1,1-dichloroethane	187 4 821	1 1 1	ND ND	ND ND	ND	ND ND
14.	1,1,2-trichloroethane	187 4 821	1 1 1 1 1	ND ND	ND ND	ND	ND ND

2403

SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

Dollutant	Stream	Sample	<u>Concentrations (mg/l)</u>			
<u>Pollutant</u>	Code	<u>Typet</u>	Source	Day 1	<u>Day 2</u>	<u>Day 3</u>
<u>Toxic Pollutants</u> (Continued)						
15. 1,1,2,2-tetrachloroethane	187	1	ND			ND
	4	1	ND	ND	ND	ND
	821	1		ND		
16. chloroethane	187	1	ND	•		ND
	4	1	ND ·	ND	ND	ND
	821	1		ND		
17. bis(chloromethyl)ether	187	1	ND			ND
	4	1	ND	ND	ND	ND
	821	1		ND		
18. bis(2-chloroethyl)ether	187	1	ND			ND
	4	2	ND	ND	ND	ND
	821	1		ND		ND
19. 2-chloroethyl vinyl ether	187	1	ND	·		ND
	4	2	ND	ND	ND	ND ,
	821	1		ND		ПD
20. 2-chloronaphthalene	187	1	ND			ND t
•	4	2	ND	ND	ND	ND +
	821	-1		ND	ЦD	ND
21. 2,4,6-trichlorophenol	187	1	ND			ND
•	4	2	ND	ND	ND	ND
	821	1		<0.01	112	11 D

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT -<

SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

	Pollutant	Stream Code	Sample Typet_	<u>Conc</u> Source	entration Day 1	ns (mg/1) Day 2	Day 3
	<u></u>					<u></u>	
Taxic	Pollutants (Continued)						
22.	r-chloro-m-cresol	187	1	ND	¢.		ND
ta L o		4	2	ND	ND	ND	ND
		821	1:		ND		
0.0	11	187	1	ND			ND.
23.	chloroform	4	i	0.05	0.02	0.02	0.02
		821	1		ND		
					n an	, *	1
24.	2-chlorophenol	187	- 1	ND			ND
		· 4	2	ND	ND .	ND	ND
		821	1.		ND		·:
25.	1,2-dichlorobenzene	187	1	ND		· · ·	ND
23.	1,2-uICHIOLODCH2CHC	4	2	ND	ND	ND	ND
		821	1 .		ND		
96	1.2. Heblanchennene	187	1	ND		· •	ND
26.	1,3-dichlorobenzene	4	2	ND	ND	ND	ND
- ¹ 14 - 1		821	1		ND		
						•	ND
2.	1,4-dichlorobenzene	187	11 C	ND	ND	ND	ND
		4	2	ND	ND .	ND	ND
	· · ·	821	. 8	· · ·	ND		· · ·
28.	3,3'-dichlorobenzidine	187	1	ND		•	ND
		2 · . · · · · · · · · · · · · · · · · ·	2	ND	ND	ND	ND
		821	1		ND	-	

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT -

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SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

Pollutant	Stream Code	Sample Typet	Conc Source	entrations Day 1	(mg/1) Day 2	Day 3	SECONDARY
xic Pollutants (Continued)							NDAJ
<pre>. 1,1-dichloroethylene</pre>	187 4 821	1 1 1	ND ND	ND ND	ND	ND ND	
.0. 1,2- <u>trans</u> -dichloroethylene	187 4 821	1 1 1	ND ND	ND ND	ND	ND ND	PRECIOUS M
31. 2,4-dichlorophenol	187 4 821	1 2 1	ND ND	ND ND	ND	ND N D	METALS 3
32. 1,2-dichloropropane	187 4	1	ND ND	ND	ND	ND ND	SUBCATEGORY
33. 1,3-dichloropropene	821 187 4	1 1 1	ND ND	ND ND	ND	ND N D	BORY
34. 2,4-dimethylphenol	821 187 4	1 1 2	ND ND	ND ND	ND	ND ND	SECT
35. 2,4-dinitrotoluene	821 187 4	1	ND ND	ND ND	ND	ND ND	l V
	821	2 1	1112	ND	11 <i>1</i>	ЦD	·

SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

Pollutant	Stream <u>Code</u>	Sample Typet	Conc Source	entration Day 1	ns (mg/1) Day 2	Day 3	SECONDARY
Toxic Pollutants (Continued)				. *			NDA
56. 2,6-dinitrotoluene	187 4 821	1 2 1	ND ND	ND ND	ND	ND ND	
37. 1,2-diphenylhydrazine	187 4 821	1 2 1	ND ND	ND ND	ND	ND ND	PRECIOUS N
38. ethylbenzene	187 4 821	1	ND ND	ND ND	ND	ND ND	METALS
39. fluoranthene	187	1 2	ND ND	ND	ND	ND ND	SUBCATEGORY
40. 4-chlorophenyl phenyl ether	821 187 4	1 1 2	ND ND	ND ND	ND	ND ND	EGORY
41. 4-bromophenyl phenyl ether	821 187 4	1 1 2	ND ND	ND ND	ND	ND ND	SECT
42. bis(2-chloroisopropyl)ether	821 187 4 821	1 1 2 1	ND ND	ND ND ND	ND	ND ND	- V

2407

SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

	Stream	Sample	Concentrations (mg/l)			
<u>Pollutant</u>	Code	Typet	Source	Day 1	<u>Day 2</u>	Day 3
Toxic Pollutants (Continued)				·* 0		
43. bis(2-chloroethoxy)methane	187	1,	ND			ND
	4 821	2	ND	ND ND	ND	ND
4. methylene chloride	187	1	ND			ND
4. methylene chioride	4	i	<0.01	<0.01	<0.01	<0.01
· · · · ·	821	1	1000 - 1000 1000 - 1000 1000 - 1000	ND		2
5. methyl chloride (chloromethane)	187	1	ND			ND
·	4 821	1	ND	ND ND	ND	ND
6. methyl bromide (bromomethane)	187	. 1	ND			ND
	4	1	ND	ND	ND	ND
· ·	821	1.		ND		ά.
7. tromoform (tribromomethane)	187	1	ND			ND
	4 821	1	ND	<0.01 ND	ND	ND
48. di hlorobromomethane	187	ì	ND			ND
	4	1	ND .	ND	ND	ND
	821	1 1		ND		,
49. trichlorofluoromethane	187	1	ND			ND
	4 821	1	ND	ND ND	ND	ND
	VL:	•				

SECONDARY PRECIOUS METALS SUBCATEGORY

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SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

		Stream	Sample Concentration			s (mg/l)		
-	Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3	
Toxic	Pollutants (Continued)	-						
.:0.	dichlorodifluoromethane	187 4 821	1 1 1	ND ND	ND ND	ND	ND ND	
51.	chlorodibromomethane	187 4 821	1 1 1	ND <0.01	<0.01 ND	ND	ND ND	
2.	hexachlorobutadiene	187 4 821	1 2 1	ND ND	ND ND	ND	ND ND	
53.	hexachlorocyclopentadiene	187 4 821	1 2 1	ND ND	ND ND	ND	ND ND	
54.	isophorone	187 4 821	1 2 1	ND ND	<0.01 ND	ND	ND ND	
55.	naphthalene	187 4 821	1 2 1	ND ND	N D ND	ND	ND ND	
56.	nitrobenzene	187 4 821	1 2 1	ND ND	NÐ ND	ND	ND ND	

2409

SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

		Stream	Sample		entration	ns (mg/1)	
	<u>Pollutant</u>	Code	<u>Typet</u>	Source	Day 1	Day 2	Day 3
Toxic	Pollutants (Continued)						
57.	2-nitrophenol	187 4 821	1 2 1	ND ND	<0.01 ND	<0.01	ND <0.01
58.	4-nitrophenol	187 4 821	1 2 1	ND ND	ND ND	ND	ND ND
59.	2,4-dinitrophenol	187 4 821	1 2 1	ND ND	ND ND	ND	ND N D
60.	4,6-dinitro-o-cresol	187 4 821	1 2 1	ND ND	ND ND	ND	ND ND
·ő1.	N-nitrosodimethylamine	187 4 821	1 2 1	ND ND	ND ND	ND	ND ND
62.	N-nitrosodiphenylamine	187 4 821	1 2 1	ND ND	ND ND	ND	ND ND
¢3.	N-nitrosodi-n-propylamine	187 4 821	1 2 1	ND ND	ND ND	ND	ND ND

SECONDARY PRECIOUS METALS SUBCATEGORY SECT ł

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2410

SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

	G .	·	··· ·	,		
Pollutant	Stream <u>Code</u>	Sample Typet	<u>Con</u> Source	centration		Day 2
		Type	Durce	Day 1	Day 2	Day 3
Toxic Pollutants (Continued)	· · · · ·		ý	s. N	•	
64. pentachlorophenol	187	1	ND			ND
off peneaentorophenor	4	2	ND ND	ND	ND	ND N D
	821	1		ND.		ND
65. phenol	187	•	ND			
ost prenot	4	2 I	ND ND	<0.01	<0.01	ND ND
	821	1	ŇĎ	ND		ND
66. bis(2-ethylhexyl) phthalate	187		0 000	· · ·		
oo: bis(2-ethylnexyl) phthalate	4	2	0.026	<0.01	<0.01	0.033
、	821	1	0.02	0.073		<0.01
	-					• .
67. butyl benzyl phthalate	187	1	ND ND	ND		ND
	4 821	1	ND	ND ND	ND	ND
		•		ND	.*	
©8. di-n-butyl phthalate	187	1	ND	1		0.002
	4 821	2	<0.01	<0.01	<0.01	ND
	021			<0.01	-	
di-n-octyl phthalate	187	1	ND		•	ND
	4	2 -	ND	ND	ND	ND
	821	.		ND		
3. diethyl phthalate	187	1	ND			ND
$(1,1,2,\dots,n_{n-1}) = (1,1,2,\dots,n_{n-1}) = (1,1$	4	2	<0.01	<0.01	<0.01	<0.01
	821	1	1 - D	<0.01		-

2411

SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

	Stream	Sample		entration		
<u>Pollutant</u>	Code	<u>Typet</u>	Source	Day 1	Day 2	Day 3
Toxic Pollutants (Continued)		-				
71. dimethyl phthalate	187	1	ND			ND
	4	2	ND	ND	ND	ND
-	821	1	_	ND		
72. benzo(a)anthracene	187	1	ND		· •	ND
/=• -•	4	2	ND	ND	ND	ND
	, 821	1	<u>م</u>	ND		
73. benzo(a)pyrene	187	1 .	ND			ND
751 Denzo(a)pyrene	4	2	ND	ND	ND	ND
	821	1		ND		
74. benzo(b)fluoranthene	187	1	ND		• 	ND
	4	2	ND	ND	ND	ND
	821	1		ND		
5. benzo(k)fluoranthane	187	1	ND		•	ND
	4	2	ND	ND	ND	ND
· · · · · · · · · · · · · · · · · · ·	821	1		ND		
76. chrysene	187	1	ND			ND
/or enrysence	4	2	ND	ND	ND	ND
	821	1		ND		
77. acenaphthylene	187	1	ND			ND
(,, modiapition) and	4	2	ND	ND	ND	ND
	821	1		ND		

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT

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SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

			0 1	Gome	entration	o (ma/l)	·
ж.н. С	Pollutant	Stream Code	Sample Typet	Source	Day 1	$\frac{\text{Day } 2}{\text{Day } 2}$	Day 3
Toxic	Pollutants (Continued)		- - 	رس ب ۶۰۰	1000 1000 1000 1000		•
78,	anthracene (a)	187	1 2	ND ND	ND	ND	ND ND
,		821	1		ND	• •	
79.	benzo(ghi)perylene	187	1	ND ND	ND	ND	ND ND
		4 821	1		ND		
80.	fluorene	187	1	ND	ND	NN	ND ND
-		4 821	2	ND	ND ND	ND	-
81.	phenanthrene (a)	187	1	ND	29 F		ND
• _ `		821	2	ND	ND ND	ND	ND
82.	dibenzo(a,h)anthracene	187	1	ND	97 (c) 		ND
میں ۲۰ به		4 821	2 1	ND	ND ND	ND	ND
83.	indeno (1,2,3-c,d)pyrene	187		ND	erten en e		ND
		821	2 1	ND	ND ND	ND	ND
84.	pyrene	187		ND			ND
		4 821	2	ND	ND ND	ND	ND

SECONDARY PRECIOUS METALS SUBCATEGORY SECT -

2413

SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

		Stream	Sample	Conc	entration	s (mg/1)	
	Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3
Toxic	Pollutants (Continued)						
85.	tetrachloroethylene	187 4	1 1	ND ND	ND	ND	ND ND
×		821	- 1		ND		
86.	toluene	187 4 821	1 1 1	ND ND	<0.01 <0.01	<0.01	ND <0.01
87.	trichloroethylene	187 4 821	1 1 1	ND ND	ND ND	ND	ND ND
88.	vinyl chloride (chloroethylene)	187 4 821	1 2 1	ND ND	ND ND	ND	ND ND
114.	antimony	201 187 4 821	2 1 2 1	<0.01 <0.003 <0.003	1.8 <0.003 1.7	3.2 <0.003	0.32 <0.003
115.	arsenic	201 187 4 821	2 1 2 1	<0.01 <0.003 <0.005	2.4 0.027 0.061	0.6 <0.005	0.12 <0.005

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SECONDARY PRECIOUS METALS SUBCATEGORY

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SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

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Pollutant	Stream Code	Sample Type†	Conc	entration Day 1	ns (mg/1) Day 2	Day 3	SECONDARY
							DA
Toxic Pollutants (Continued)				-			RY
117. beryllium	201	2	<0.01	<0.01	<0.01		PI
, , , , , , , , , , , , , , , , , , ,	187	1	<0.01			<0.01	Æ
	4	2	<0.0002	0.001	<0.0002	<0.0002	C
	821	1		<0.001			PRECIOUS
	021	•	· ·				SD
118. cadmium	201	2	<0.05	<0.05	<0.05		M
	187	1	<0.01	-		<0.01	臣
	4	2	0.0002	0.001	0.001	0.001	METALS
	821	1		4.4			2 2
						*	τΩ
11°, chromium (total)	201	2	<0.05	0.75	0.70		SUBCATEGORY
	187	1	<0.01			<0.01	ũ.
	4	2	0.003	0.012	0.003	0.004	AI
	821	1		0.012			Ë
		-	-			-	ö
120. copper	201	2	<0.05	2.3	2.7	•	RY
inor copper	187	1	<0.01			0.15	•
	4	2	0.017	0.016	0.038	0.017	
	•						N
121. cyanide (total)	201	1	0.05	0.37	0.15		SECT
1211 Oyuntue (cocur)	187	1	<0.02			<0.02	Ĥ
· · ·	4	1	0.052	0.930	0.840	0.98	1
	821	1		0.29	· · · · · ·		<
	.	. •	•				7

SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

5 11	Stream Code	Sample	Conc. Source	entration Day 1	<u>s (mg/1)</u> Day 2	Day 3
Pollutant	<u> </u>	<u>Typet</u>	Douroe			
Toxic Poilutants (Continued)		•				
122. lead	201	2	<0.10	0.1	0.2	
122. Ieau	187	1	<0.10			<0.10
en e	4	2	0.030	0.020	0.050	0.250
	821	1		2.7		
10.2 — — — — — — — — — — — — — — — — — — —	201	2	0.0002	<0.0002	<0.0002	
123. mercury	187	1	0.002	4		<0.0002
	4	2	0.0002	<0.0001	<0.0001	<0.0001
· ·	821	1		<0.001		
124. nickel	201	2	<0.20	1.8	2.0	
124. HICKEL	187	1	0.075			0.43
- · · ·	4	2	0.020	0.009	0.008	0.018
	821	1		4.6		
125. selenium	201	2	<0.10A	B	<0.1A	
12J, Selenium	187	1	<0.003			0.019
	4	2	<0.002	7.0	<0.002	<0.002
	821	.1		7.4		
126. silver	201	2	<0.01	1.9	1.6	
126. silver	187	Ī	<0.0005			0.18
·	4	2	<0.0002	0.003	0.002	0.001
-	821	1	۳. ۳. ۲.	0.53		

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT -<

SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

•	Stream	Sample	Con	centratio	<u>ns (mg/1)</u>	
Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3
Toxic Pollutants (Continued)						
127. thallium	201 187 4 821	2 1 2 1	<0.01 <0.002 <0.001	B <0.001 0.82	B <0.004	<0.002 <0.004
128. zinc	201 187 4 821	2 1 2 1	0.10 2.5 <0.010	1.8 0.59 2.3	2.5 0.26	6.9 0.39
Nonconventional Pollutants						
acidity	201 187	2 1	<1 <1	960	<1	<1
alkalinity	201 187 4	2 1 2	98 127 16	<1 280	47 93 360	,000 330
aluminum	201 187	2 1	0.2 <0.05	<10	<10	<0.05
ammonia nitrogen	201 187 821	2 1 1	0.04 <0.01	4	6.5	<0.01

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SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

Pollutant	Stream _Code_	Sample Typet	Concentrations (mg/1) Source Day 1 Day 2 Day 3
Nonconventional Pollutants (Continued)		
barium	201 187	2 1	
boron	201 187	2 1	<0.05
calcium	201 187 4	2 1 2	37.7 40 40 11 49 13 13 15 13
chemical oxygen demand (COD)	201 187	2 1	
chloride	201 187	2 1	<pre><5 1,800 16,000 1,100 14 69,000 80,000 52 1,200</pre>
cobalt	201 187	2 1	<0.05 <5 <5 <0.006 <0.006
fluoride	201 187	2 1	0.28 1.9 1.6 1.1 1.3
gold	4	2	0.025 0.48 0.42 0.42
irön	201 187 4 821	2 1 2 1	<0.05

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SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

	-	Stream	Sample	Con	centratio	ons (mg/1)	
Pollutant	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	Day 3					
aconventional Pollutants	(Continued)		•				-
magnesium	•	187	2 1 2	2.4		· .	4.3 3.1
"Maganese			2 1		0.1	0.1	0.06
ybdenum		201 187	2 1		<5	<5	<0.002
phenolics	• •		1 1	0.15		0.067	<0.001
phosphata			2 1	14	5	<0.9	<4
sodium		201 187	2 1	9.0 53 54	,000 6	5,000 72	,000
sulfate		201 187	2 1	57 7 13	,000 1	5,000	260
tin			2 1		59.4	74.9	<0.12

SECONDARY PRECIOUS METALS SUBCATEGORY SECT -2

SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

Pollutant	Stream Code	Sample Typet	Conce Source	Day 1	ns (mg/1) Day 2	Day 3
Conconventional Pollutants (Continued)						
titanium	201 187	2 1	<0.005 <0.005	<5	<5	<0.005
total organic carbon (TOC)	201 187	2 1	4.3 43	20	29	80
total solids (TS)	201 187	2 1	380 140, 410	000 180	, 000 27	7,000
vanadium	201 187	2 1	<0.05 <0.003	<5	<5	<0.003
yttrium	201 187	2 1	<0.05 <0.002	<5	<5	<0.002
Conventional Pollutants						
oil and grease	201 187 4 821	1 1 1	<1 <1 1.6	3.5 <1 37	14 <1	<1 <1

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SECONDARY PRECIOUS METALS SAMPLING DATA REFINERY WET AIR POLLUTION CONTROL RAW WASTEWATER

Pollutant	Stream	Sample	Concentrations $(mg/1)$				
Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3	
Conventional Pollutants (Continued)	-				• • • •		
cotal suspended solids (TSS)	201 187 4	2 1 2	60 8 0	5,500 90	390 14	690 21	
pH (standard units)	201 187 4	- 2 1 2	7.5 7.36 6.8	1.6	8.5	12.59 10.9	

tSample Type Code: 1 - One-time grab
2 - Manual composite during intermittent process operation

Reported together.

A - Detection limit raised due to interference.

B - Chemical interference.

SECONDARY PRECIOUS METALS SUBCATEGORY SECT

Table V-18

SECONDARY PRECIOUS METALS SAMPLING DATA GOLD PRECIPITATION AND FILTRATION RAW WASTEWATER

	Pollutant	Stream Code	Sample Typet		trations (mg/1) Day 1 Day 2 Day 3
<u>Toxic</u>	Pollutants				
114.	antimony	233	1	<0.010	<0.50A
115.	arsenic	233	1	<0.010	<0.20
1.7.	beryllium	233	1	<0.010	0.150
113.	cadmium	233	1	<0.050	0.10
9.	chromium (total)	233	1	<0.050	3.40
·	copper	233	1	<0.050	100.0
· · · ·	cyanide (total)	233	1	0.05	<0.02
. 2.	lead	233	. 1	<0.10	6.5
	mercury	233	1	0.0002	<0.0002
	nickel	233	1	<0.20	46.0
	selenium	233	1	<0.10A	В
125.		233	1	<0.010	26.0
126.	silver	233	1	<0.010	<0.5A
127.	thallium .	233	1	0.10	340.0
128.	zinc	200	I		

2422

SECONDARY PRECIOUS METALS SUBCATEGORY

SECT - V

SECONDARY PRECIOUS METALS SAMPLING DATA GOLD PRECIPITATION AND FILTRATION RAW WASTEWATER

	Stream	Sample	Conc	entration	s (mg/1)	
Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3
Nonconventional Pollutants						
acidity	233	1	<1		<1	
alkalinity	233	1	98		850	
aluminum	233	1	0.20		109	
ammonia nitrogen	233	1	0.04		570	
barium	233	1	<0.050		<0.5	
boron	233	1	<0.10	•	<1.0	
calcium	233	1	37.7		44.0	
chemical oxygen demand (COD)	233	1	<5	3	7,000	• •
chloride	233	1	14	1	6,000	
cobalt	233	1	<0.050	•	1.0	• •
fluoride	233	1	0.28		0.65	
iron	233	1	<0.050		10.0	
magnesium	233	1 · · · ·	8.50	• •	10.0	·
manganese	233	1	<0.050		0.30	X

SECONDARY PRECIOUS METALS SUBCATEGORY SECT -

SECONDARY PRECIOUS METALS SAMPLING DATA GOLD PRECIPITATION AND FILTRATION RAW WASTEWATER

Pollutant	Stream Code	Sample Typet	Conce Source	entrations (mg/1) Day 1 Day 2 Day 3
Nonconventional Pollutants (Continued)				
molybdenum	233	1	<0.050	2.0
phosphate	233	1	14	130
sodium	233	1	9	2, 390
sulfate	233		57	30,000
tin	233	. 1	<0.050	<0.5
titanium	233	1	<0.05 0	<0.5
total organic carbon (TOC)	233	× 1	4.3	140
total solids (TS)	233	1	380	240,000
vanadium	23 3	1	<0.050	<0.5
yttrium	233	1	<0.50	<0.5
Conventional Pollutants				
oil and grease	233	1 .	<1	3

SECONDARY PRECIOUS METALS SUBCATEGORY SECT I

SECONDARY PRECIOUS METALS SAMPLING DATA GOLD PRECIPITATION AND FILTRATION RAW WASTEWATER

	Stream	Sample	Conc	entrations (mg/1)	·
Pollutant	Code	Typet	Source	Day 1 Day 2	Day 3
Conventional Pollutants (Continued)				• •	
total suspended solids (TSS)	233	1	60	1,670	
pH (standard units)	233	1	7.5	9.3	

tSample Type Code: 1 - One-time grab

Detection limit raised due to interference.

Chemical interference.

2425

SECT

Table V-19

SECONDARY PRECIOUS METALS SAMPLING DATA PALLADIUM PRECIPITATION AND FILTRATION RAW WASTEWATER

	Stream	Sample	Con	centration	ns (mg/l)	
Pollutant	<u>Code</u>	<u>Typet</u>	Source	Day 1	Day 2	Day 3
Toxic Pollutants						
114. antimony	230	1	<0.01	0.50		
	230	1	<0.01		<0.50A	
•	230	1	<0.01		<0.20A	
11 ^g ansenic	230	1	<0.01	<0.2A		i.
	230	1	<0.01		<0.01	
	2,30	1	<0.01		0.07	
117 beryllium	230	1	<0.01	<0.01		
	230	1	<0.01		0.02	
	230	1	<0.01		0.02	
118. cadmium	230	1	<0.05	0.05		•
	230	1	<0.05		0.25	
	230	1	<0. 05		0.4	
119. chromium (total)	230	1	<0.05	1.2		
	230	1	<0.05		1.8	
	230	1	<0.05		0.65	
120. copper	230	1	<0.05	300.0		
	230	1	<0.05		72.0	ì
	230	1	<0.05		72.0	
121. cyanide (total)	230	1	0.05	<0.02		
	230	1	0.05		<0.02	
	230	1	0.05	,	<0.02	

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SECONDARY PRECIOUS METALS SUBCATEGORY

SECT -

SECONDARY PRECIOUS METALS SAMPLING DATA PALLADIUM PRECIPITATION AND FILTRATION RAW WASTEWATER

	Pollutant	: · · ·		Stream Code	Sample Typet	<u>Conc</u> Source	entration Day 1	ns (mg/1) Day 2	Day 3
Toxic	c Pollutants (Continued)	-	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	بر میں بر کر کر ک ی کی بندر			ŬŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢŢ	، بارورویک دورور د ر	
		· · ·		220	1	20.10	1.8		
122.	lead			230 230	• I 1 *	<0.10 <0.10	1.0	6.8	
				230	1	<0.10		6.2	
123.	mercury	~	-	230	1	0.0002	<0.0002	· · ·	
1234	mereary			230	1	0.0002	(010004	<0.0002	
	·	•		230	1	0.0002	• •	<0.0002	
124.	nickel			230	1	<0.2	30.0		
				230	1	<0.2	-	8.0	
				230	1	<0.2	* 7	8.8	
125.	selenium			230	. 1	<0.1A	B	· .	
		• • •	`	230	1	<0.1A	-	<0.1A	· .
	•			230	1	<0.1A	· · · ·	<0.1A	·
126.	silver			230	1	<0.01	9.4		
	•			230	1	<0.01		10.0	
	а, сторо с -	, , , , , , , , , , , , , , , , , , ,		230	1	<0.01		1.8	
127.	thallium			230	1	<0.01	<0.1A		
				230	1	<0.01		<0.01	*
		· · ·	۰. ب	230	1	<0.01	• * •	<0.04A	· ·
128.	zinc	X		230	⊆ 1 *	0.10	170		
		. •	`	230	1	0.10		180	
			*	230	1	0.10		270	

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SECONDARY PRECIOUS METALS SUBCATEGORY

SECT -

SECONDARY PRECIOUS METALS SAMPLING DATA PALLADIUM PRECIPITATION AND FILTRATION RAW WASTEWATER

	Stream	Sample	Conc	entrations		
Pollutant	Code	Typet	Source	Day 1	Day 2 Day	3
Nonconventional Pollutants						
acidity	230	1	<۱	2,500		
•	230	1	<1 <1	•	9,000	
	230	1	<1		9,000 9,500	
alkalinity	230	1	98	<1		
-	230	1	98		<1	
• • • • • •	230	1	98		<1 <1	
aluminum	230	1	0.2	<1		
	230	1	0.2		23.3	
	230	1	0.2		9.8	
ammonia nitrogen	230	1	0.04	5,060		
	230	1	0.04	•	1.4	
	230	1	0.04		2,700	
barium	230	1	<0.05	<0.5		
•	230	1	<0.05		<0.05	
	230	1	<0.05		<0.05	
boron	230	1	<0.1	<1		
· ·	230	1	<0.1		1.4	
	230	1	<0.1		0.5	
calcium	230	1	37.7	10		
	230	1	37.7		33 18.3	
	230	1 L L	37.7		18.3	

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT

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SECONDARY PRECIOUS METALS SAMPLING DATA PALLADIUM PRECIPITATION AND FILTRATION RAW WASTEWATER

<u>Pollutant</u> <u>Monconventional Pollutants</u> (Continued chemical oxygen demand (COD)	Stream	Sample	Concentrations (mg/1)			
Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3
Monconventional Pollutants (Continued)) (1) (2) (2)	•			1 M	
chemical oxygen demand (COD)	230 230 230	1 1 1	<5 <5 <5	34,000 2 4	4,000 0,000	-
chloride	230 230 230	1 1 1	14 14 14		2,000	
cobalt	230 230 230	1 1 1	<0.05 <0.05 <0.05	<0.5	0.3 0.4	:
fluoride	230 230 230	1 1 1	0.28 0.28 0.28	0.28	0.8 0.37	•
iron	230 230 230	1 1 1	<0.05 <0.05 <0.05	4.8	21 31	-
magnesium	230 230 230	1 1 1	8.5 8.5 8.5	2	9 4	

SECT

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SECONDARY PRECIOUS METALS SAMPLING DATA PALLADIUM PRECIPITATION AND FILTRATION RAW WASTEWATER

	Stream	Sample	Concentrations (mg/1)			
Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3
Negeonventional Pollutants (Continued))					
salaganese	230	1	<0.05	0.1		
-	230	1	<0.05		0.3	
	230	1	<0.05		0.3	
molybdenum	230	1	<0.050	<0.5		
	230	1	<0.050		0.7	
	230	1	<0.050		0.6	
phosphate	230	1	14	<4		
FF.	230	1	14	•••	<1	
	230	1	14		<1	
sodium	230	1	9.00	54		
	230	1	9.00		25.7	
	230	1	9.00		160	
sulfate	230	1	57	5,500		
	230	1	57		00,000	
	230	1	57		2,700	
tin	230	1	<0.050	3.4		
	230	1	<0.050		18.4	
•	230	1	<0.050		5.6	

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SECONDARY PRECIOUS METALS SAMPLING DATA PALLADIUM PRECIPITATION AND FILTRATION RAW WASTEWATER

· · ·		Stream	Sample	Conc		(mg/1)	
Pollutant		Code	Typet	Source	Day 1	Day 2	Day 3
Nonconventional Pollutants	(Continued)						
titanium		230	1	<0.050	<0.5	0 0 F	
		230	• • 1 • • •	<0.050		0.25	
		230	. 1	<0.050		0.4	
total organic carbon (TOC)		230	1 .	4.3	2,700		·
total organic carbon (100)	:	230	1	4.3		34	
		230	· 1	4.3		2,600	、
total solids (TS)		230	1	380	170,000		
		230	1	380		0,000	
		230	1	380	5	5,000	
vanadium		230	1	<0.050	<0.5		·
Vallautum		230	1	<0.050		<0.05	
· · · · · · · · · · · · · · · · · · ·		230	1	<0.050		<0.05	
whether from		230	1	<0.50	<0.5		
yttrium		230	1	<0.50		<0.05	,
	· 	230	i	<0.50		<0.05	

SECONDARY PRECIOUS METALS SUBCATEGORY

SECT -

SECONDARY PRECIOUS METALS SAMPLING DATA PALLADIUM PRECIPITATION AND FILTRATION RAW WASTEWATER

Pollutant	Stream _Code	Sample Typet	<u>Conc</u> Source	entration Day 1	s (mg/1) Day 2	Day 3
Conventional Pollutants			•			
oil and grease	230 230 230	1 1 1	<1 <1 <1	2	<1 5	
total suspended solids (TSS)	230 230 230	1 1 1	60 60 60	200	630 210	,
pH (standard units)	230 230 230	1 1 1	7.5 7.5 7.5	1.6	0.1	

tSample Type Code: 1 - One-time grab

A - Detection limit raised due to interference.

B - Chemical interference.

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Table V-20

SECONDARY PRECIOUS METALS SAMPLING DATA SPENT SOLUTION FROM PGC SALT PRODUCTION RAW WASTEWATER

	Stream	Sample	Concentrations				
	<u>Pollutant</u>	Code	<u>Typet</u>	Source	Day 1	Day 2	Day 3
<u> Zexic</u>	Pollutants				· · · · ·		
	antimony	703	1	<0.003	<0.003		¥
\$	arsenic	703	1	<0.002	<0.003		
7.	beryllium	703	1	0.002	0.006		•
118.	cadmium	703	1	0.014	0.037	•	
179.	ch.comium (total)	703	1	0.015	0.32		
120.	opper	703	1.	2.3	5.9		· ·
121.	cyanide (total) cyanide (free)	703 703	1	0.41 0.26	5,000 42		· · · ·
122.	lead	703	1	<0.08	0.33		► .
123.	mercury	703	1	0.0007	<0.0002		
124.	nickel	703	1	0.25	0.61		•
125.	selenium	703	8	<0.002	<0.003		
126.	silver	703	1	<0.0005	<0.0005		• •
127.	thallium	703	1	<0.002	<0.002		
128.	zinc	703	1	0.055	0.98		

SECONDARY PRECIOUS METALS SUBCATEGORY SECT 1

SECONDARY PRECIOUS METALS SAMPLING DATA SPENT SOLUTION FROM PGC SALT PRODUCTION RAW WASTEWATER

Pollutant	Stream Code	Sample Typet	Conc Source	entrations (mg/l) Day 1 Day 2	Day 3
aconventional Pollutants					
a,um inum	703	1	0.86	4.0	
esrium	703	1	0.056	<0.001	
boron	703	1	<0.009	0.67	
calcium	703	1	4.2	2.6	:
cobalt	703	1	0.044	0.11	· .
gold	703	1	0.15	1,4	,
iron	703	1	0.94	27	
magnesium	703	1	1., 3	0.77	
manganese	703	1	0.013	0.041	ļ
molybdenum	703	. 1	0.029	0.1	
sodium	703	1	13	520	
tin	703	1	<0.12	<0.12	
titanium	703	. 1	0.12	0.84	
vanadium	703	1	0.073	0.18	
yttrium	703	1	<0.002	0.06	

SECONDARY PRECIOUS METALS SAMPLING DATA SPENT SOLUTION FROM PGC SALT PRODUCTION RAW WASTEWATER

Pollutant	Stream Code	Sample Typet	<u>Conc</u> Source	entrations (mg/l) Day 1 Day 2	Day 3 C
Gonventional Pollutants					NDAR
pH (standard units)	703	1	7	10	JA A
	•	Р 	• • •		ECIO
					US MET
					TALS

SUBCATEGORY

SECT

tSample Type Code: 1 - One-time grab

Table V-21

SECONDARY PRECIOUS METALS SAMPLING DATA EQUIPMENT AND FLOOR WASH RAW WASTEWATER

	ñ	Stream	Sample		entration		
	Pollutant	Code	<u>Typet</u>	Source	Day 1	Day 2	Day 3
Tric	Pollutants						
114.	antimony	228	1	<0.01	0.08	0.06	
1/5.	arsenic	228	1	<0.01	<0.05A	<0.05A	
[]7.	beryllium	228	1	<0.01	<0.01	<0.01	
118.	cadmium	228	1 .	<0.05	0.6	0.1	
119.	chromium (total)	228	1	<0.05	1.1	0.35	
120.	copper	228	1	<0.05	280.0	21.0	
121.	cyanide (total)	228	1	<0.05	0.13	0.11	
122.	lead	228	1	<0.10	8.0	1.3	
123.	mercury	228	1 .	0.0002	<0.0002	<0.0002	
124.	nickel	228	. 1	<0.20	12.0	1.8	
125.	ɛelenium	228	1	<0.1A	<0.1A	<0.1A	
126.	silver	228	1	<0.01	0.26	0.09	
127.	thallium	228	1	<0.01	<0.05A	<0.01	
128.	zinc	228	1	0.10	440.0	9.2	

SECONDARY PRECIOUS METALS SUBCATEGORY SECT i

SECONDARY PRECIOUS METALS SAMPLING DATA EQUIPMENT AND FLOOR WASH RAW WASTEWATER

	Stream	Sample	Conc	entrations	(mg/1)	-
<u>Pollutant</u>	Code	Typet	Source	Day 1	Day 2	Day 3
Nonconventional Pollutants	*					
acidity	228	1	۰ ۰	1,080	<1	
askalinity	228 [±]	1	98	<1	360	- ,
aminum	228	1	0.20	5.5	4.9	
comonia nitrogen	228	- 1	0.04	120	75	
barium	228	1	<0.050	<0.05	<0.05	
boron	228	1	<0.10	0.2	0.2	\$
calcium	228	1	37.7	43.7	34.2	
chemical oxygen demand (COD)	228	1	<5	37,000	6,200	·
chloride	228	1	14	<1	490	L
cobalt	228	1	<0.050	0.45	0.05	
fluoride	228	1	0.28	0.47	0.4	
iron	228	1	<0.050	39.8	12.0	
magnesium	228	1	8.50	9.3	7.9	
manganese	228	1	<0.050	0.3	0.15	

SECONDARY PRECIOUS METALS SUBCATEGORY SECT

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SECONDARY PRECIOUS METALS SAMPLING DATA EQUIPMENT AND FLOOR WASH RAW WASTEWATER

Pollutant	Stream Code	Sample Typet	Conce Source		(mg/l) Day <u>2</u> Day	3
Nonconventional Pollutants (Continued)	·					-
molybdenum	228	1	<0.050	0.05	0.45	
phosphate	228	1	14	<4	<1	
sodium	228	1	9	44	478	
sulfate	228	1	57	2,900	3,000	
tin	228	1	<0.050	7.55	3.3	
citanium	228	1	<0.050	<0.05	<0.5	
total organic carbon (TOC)	228	1	4.3	15	15	
total solids (TS)	228	1	380	120	2,600	
anadium	228	1	<0.050	<0.05	, <0.05	I
*ttrium	228	1	<0.50	<0.05	<0.05	.
Conventional Pollutants						l
_il and grease	228	1	<1	2	6	

SECONDARY PRECIOUS METALS SUBCATEGORY SECT

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SECONDARY PRECIOUS METALS SAMPLING DATA EQUIPMENT AND FLOOR WASH RAW WASTEWATER

	Stream	Sample	Conc			
Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3
Conventional Pollutants (Continued)				, ,		
totak suspended solids (TSS)	228	1	60	20	760	
pH (standard units)	228	1	7.5	1.8	10.9	

SECONDARY PRECIOUS METALS SUBCATEGORY

SECT

Sample Type Code: 1 - One-time grab

- Detection limit raised due to interference.

3 - Chemical interference.

Table V-22

SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

	Stream	Sample		entration			Ŋ
Pollutant	<u>Code</u>	<u>Typet</u>	Source	Day 1	Day 2	Day 3	EC C
<u>T</u> (ic Pollutants					~		SECONDARY
acenaphthene	185 3	1 3	ND ND	ND .	N D ND	ND	
· · ·	822	1	ND	ND			RE
2 acrolein	185 3	1 1	ND ND	ND	N D ND	ND	PRECIOUS
· ·	822	1	ND	ND			
3. acrylonitrile	185 3	1	ND ND	ND	N D ND	ND	METALS
	822	1	ND	ND			
4. benzene	185 3	. 1 1	ND ND	ND	N D ND	ND	SUBCATEGORY
	822	1	ND	ND			IEO
5. benzidine	185 3	1 3	ND ND	ND	N D ND	ND	JORY
· · · ·	822	1	ND.	ND.	¥		
6. carbon tetrachloride	185	1 1	N D ND	ND	N D ND	ND	SECT
	822	1	ND	ND			1
7. chlorobenzene	185	1 3	ND ND	ND	N D ND	ND	۲
	822	1	ND .	ND			

SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

	Stream	Sample	Conc	entration	<u>s (mg/l)</u>		70
Pollutant	Code	Typet	Source	Day 1	Day 2	<u>Day 3</u>	ECC
Toxic Pollutants (Continued)					r ar	• •	SECONDARY
8. 1,2,4-trichlorobenzene	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND	
9. hexachlorobenzene	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND	PRECIOUS N
10. 1,2-dichloroethane	185 3 822	1	ND ND ND	ND ND	N D ND	ND	METALS S
11. 1,1,1-trichloroethane	185 3 822	1 1 1	0.01 ND ND	ND ND	N D ND	ND	SUBCATEGORY
12. hexachloroethane	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND	GORY
13. 1,1-dichloroethane	185 3 822	1 1 1	ND ND ND	ND ND	N D ND	ND	SECT -
14. 1,1,2-trichloroethane	185 3 822	1 1 1	ND ND ND	ND ND	N D ND	ND	V

SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

Pollutant	Stream <u>Code</u>	Sample Typet	Conc Source	entrations Day 1	a (mg/1) Day 2	Day 3	SECONDARY
<u>Toxic Pollutants</u> (Continued)							IDAR
15. 1,1,2,2-tetrachloroethane	185 3 822	1 1 1	ND ND ND	ND ND	N D ND	NĎ	
16. chloroethane	185 3	1 1	ND ND ND	ND ND	N D ND	ND	PRECIOUS N
17, bis(chloromethyl)ether	822 185 3	1 1	ND ND	ND	N D ND	ND	METALS
18. bis(2-chloroethyl)ether	822 185 3	1 1 3	ND ND ND	ND	N D ND	ND	SUBCATEGORY
	822 185	1	ND	ND	N D		EGORY
19. 2-chloroethyl vinyl ether	822	1	ND ND	ND ND	ND	ND	•
20. 2-chloronaphthalene	185 3 822	1 3 1	ND ND ND	<0.01 ND	N D ND	ND	SECT -
21. 2,4,6-trichlorophenol	185 3 822	1 3	ND ND ND	0.020 ND	ND 0.030	0.020	۷.
	044	I	110				

SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

	Stream	Sample	Concentrations (mg/1)				
Pollutant	Code	Typet	Source	Day 1	<u>Day 2</u>	<u>Day 3</u>	
Toxic Pollutants (Continued)	• • • •				-		
22. p-chloro-m-cresol	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND	
23. chloroform	185 3 822	1 <u>.</u> 1 [°] 1	ND 0.050 ND	<0.010 ND	ND <0.01	<0.01	
24. 2-chlorophenol	185 3 822	1 3 1	ND ND ND	<0.01 ND	ND <0.01	<0.01	
25. 1,2-dichlorobenzene	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND	
2. 1,3-dichlorobenzene	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND	
27. 1,4-dichlorobenzene	185 3 822	1 3 1	ND ND ND	ND ND	ND ND	ND	
28. 3,3'-dichlorobenzidine	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND	

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT

SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

Pollutant	Stream Code	Sample Typet	Conc Source	entration Day 1	<u>s (mg/1)</u> Day 2	Day 3
<u>xic Pollutants</u> (Continued)						
29. 1,1-dichloroethylene	185 3 822	1 1 1	ND ND ND	ND ND	N D ND	ND
30. , ² - <u>trans</u> -dichloroethylene	185 3 822	- 1 1 1	ND ND ND	ND ND	N D ND	ND
31. 2,4-dichlorophenol	185 3 822	1 3 1	ND ND ND	0.030 ND	ND 0.030	0.020
32. 1,2-dichloropropane	185 3 822	1 1 1	NÐ ND ND	ND ND	N D ND	ND
33. 1,3-dichloropropene	185 3 822	1 1 1	ND ND ND	ND ND	N D ND	ND
34. 2,4-dimethylphenol	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND
35 2,4-dinitrotoluene	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND

SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

	Stream	Sample		entration	s (mg/l)	
Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3
oxic Pollutants (Continued)	•	•	-11. 2		•	
36. 2,6-dinitrotoluene	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND
37. 1,2-diphenylhydrazine	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND
38. ethylbenzene	185 3 822	1 1 1	ND ND ND	ND ND	N D ND	ND
39. fluoranthene	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND
40. 4-chlorophenyl phenyl ether	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND
41. 4-bromophenyl phenyl ether	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND
2. bis(2-chloroisopropyl)ether	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND

SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

Pollutant	Stream Code	Sample Typet	<u>Conc</u> Source	entration Day 1	<u>s (mg/l)</u> Day 2	Day 3
<u>Texic Pollutants</u> (Continued)						
43. bis(2-chloroethoxy)methane	185	1	ND		N D	
43. bis(2-chloroethoxy)methane	3	3	ND	ND	ND	ND
	822	1	ND	ND		
	185	1	ND		N D	
44. methylene chloride	3	1	<0.01	ND	0.010	<0.01
	822	1	ND	0.040		
45. methyl chloride (chloromethane)	185	⁻ 1	ND	۴	N D	
45. methyl chloride (chloromethane)	3	1	ND	ND	ND	ND
	822	1	ND	ND		
() 1 1 to a day (because then a)	185	1	ND		ND	
46. methyl bromide (bromomethane)	3	i	ND	ND	ND	ND
· · · ·	822	1	ND	ND		
(7) (not know on the no)	185	1	ND		N D	
47. bromoform (tribromomethane)	3	1	ND	ND	ND	ND
	822	i	ND	ND		
	185	1	ND		N D	
48. dichlorobromomethane	3	1	<0.01	<0.01	<0.01	<0.01
	822	1	ND	ND		
49. trichlorofluoroemthane	185	1	ND	· ,	N D	
49. trichlorofluoroemthane	3	1	ND	ND	ND	ND
<u>.</u>	822	1	ND	ND		

SECONDARY PRECIOUS METALS SUBCATEGORY

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SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

		Stream	Sample	Concentrations (mg/l)			
	Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3
ic	Pollutants (Continued)		• • • •			Ŧ	
÷	dichlorodifluoromethane	185	1	ND		ND	
• 2 •	dientorodifiuoromeenane	3	1	ND	ND	ND	ND
	•	822	1	ND	ND	-	
-	chlorodibromomethane	185	1	ND		ND	
•		3	j 1	ND	ND	ND .	ND
		822	1	ND	ND	• -	
	hexachlorobutadiene	185	1	ND	, .	N D	
		3	3	ND	ND	ND	ND
		822	1	ND	ND	•	
53.	hexachlorocyclopentadiene	185	1	ND		ND	
		3	3	ND	ND	ND	ND
		822	1	ND	ND		
54.	isophorone	185	1	ND		ND	
		3	3	ND	ND	ND	ND
		822	1	ND	ND		•
55.	naphthalene	185	1	ND		ND	w
	•	3	3	ND	<0.01	ND	ND
		822	1	ND	ND		
56.	nitrobenzene	185	1	ND		ND	
		3	3	ND	ND	ND	ND
		822	1	ND	ND		

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SECONDARY PRECIOUS METALS SUBCATEGORY

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SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

Pollutant	Stream Code	Sample Typet	Conc Source	entration Day 1	<u>s (mg/1)</u> Day 2	Day 3
Taxic Pollutants (Continued)						
57. 2-nitrophenol	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND
58. 4-nitrophenol	185 3 822	1 3 1	ND ND ND	0.040 ND	N D ND	ND
59. 2,4-dinitrophenol	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND
60. 4,6-dinitro-o-cresol	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND
51. N-nitrosodimethylamine	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND
52. N-nitrosodiphenylamine	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND
63. N-nitrosodi-n-propylamine	185 3 822	1 3 1	ND ND ND	<0.01 ND	N D ND	ND

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT -

SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

	Stream	Sample	Concentrations (mg/1)				
Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3	
Toxic Pollutants (Continued)			-				
64. pentachlorophenol	185 3 822	1 3 1	ND ND ND	<0.01 ND	ND <0.01	<0.01	
65. phenol	185 3 822	1 3 1	ND ND ND	0.020 ND	0.041 0.010	<0.01	
66. bis(2-ethylhexyl) phthalate	185 3 822	1 3 1	0.026 0.020 ND	0.040 <0.010	0.007 0.030	0.040	
7. butyl benzyl phthalate	185 3 822	13	ND ND ND	ND ND	N D ND	ND	
.8. di-n-butyl phthalate	185 3 822	1 3 1	ND <0.01 ND	<0.01 <0.010	0.002 <0.01	ND	
9. di-n-octyl phthalate	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND	
0. diethyl phthalate	185 3 822	1 3 1	ND <0.01 ND	<0.01 ND	ND <0.01	<0.01	

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SECONDARY PRECIOUS METALS SUBCATEGORY

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SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

Pollutant	Stream Code	Sample Typet	<u>Conc</u> Source	<u>entration</u> Day 1	s (mg/1) Day 2	Day 3	SEC
Toxic Pollutants (Continued)							SECONDARY
71. dimethyl phthalate	185 3 822	1 3 1	ND ND ND	ND ND	N D ND	ND	
72. benzo(a)anthracene	185 3 822	1 3 1	ND ND ND	ŅD ND	N D ND	ND	PRECIOUS
⁷ 3. benzo(a)pyrene	185 3 822	1 3	ND ND ND	ND ND	N D ND	ND	METALS
74. benzo(b)fluoranthene	185 3	1 3	ND ND	ND	N D ND	ND	SUBCATEGORY
75. benzo(k)fluoranthane	822 185 3	1 3	ND ND ND	ND ND	N D ND	ND	EGORY
76. chrysene	822 185 3	1 1 3	ND ND ND	ND ND	N D ND	ND	SECT
77. acenaphthylene	822 185 3 822	1 3 1	ND ND ND ND	ND ND ND	N D ND	ND	ו ל

SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

		Stream	Sample	Conc	entration	s (mg/1)		л С
P	ollutant	Code	Typet	Source	Day 1	Day 2	Day 3	SECONDARY
m. 'a Dallut	ants (Continued)	•				2		
16. C POLLUL	alles (continued)							JAF
78 anthra	cene (a)	185	1	ND		N D	•	R
		3	3	ND	ND	ND	ND ·	ተ
	· · · · ·	822	1	ND	ND		<i>b</i>	PRECIOUS
79. benzo(ghi)perylene	185	1	ND		N D		CIC
/9. Delizo(guryperyrene	3	3	ND	ND	ND	ND	Ŭ
		822	1	ND	ND			
500 61		185	- 1	ND		ND		METALS
80. fluore	ine	/ . 3-	3	ND	ND	ND	ND	Ę
4		822	1	ND	ND			
04	(a)	185	1	ND		ND		SUB
81. phenar	threne (a)	2	-3	ND	ND	ND	ND	C A
		822	1	ND	ND			SUBCATEGORY
						ND		က် ပြ
82. dibenz	o(a,h)anthracene	185	1	ND	ND	ND	ND	RY
		3	3	ND		ND -	ND	
		822		ND	ND			
83. indend	(1,2,3-c,d)pyrene	185	1 1	ND		ND	-	SECT
	(112)3 010/2/2/2000	3	3	ND	ND	ND	ND	្អ
		822	· · 1 · · ·	ND	ND	**	•	د. ۱
84. pyrene	3	185	- 1	ND	•	N D		<
04. Pyrene	-	3	3	ND	ND	ND	ND	
		822	- 1	ND	ND			

SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

Pollutant	Stream <u>Code</u>	Sample Typet	Conc Source	entration Day 1	ns (mg/1) Day 2	Day 3	SECONDARY
Texic Pollutants (Continued)							ND₽
5. tetrachloroethylene	185 3 822	1 1 1	ND ND ND	ND ND	N D ND	ND	
86. toluene	185 3 822	1 1 1	ND ND ND	<ଲ.01 0.046	ND <0.01	<0.01	PRECIOUS M
87. trichloroethylene	185 3 822	1 1 1	ND ND ND	<0.01 ND	ND <0.01	<0.01	METALS
88. vinyl chloride (chloroethylene)	185 3 822	1 1 1	ND ND ND	ND ND	N D ND	ND	SUBCATEGORY
114. antimony	203 185 3 822	2 1 3	<0.01 <0.003 <0.003	0.13	1.70 0.26 0.55	0.340 1.2	GORY
15. arsenic	203 185 3 822	2 1 3 1	<0.010 <0.003 <0.005	0.095 0.19	<0.20A 0.068 0.190	<0.20A 0.140	SECT - V

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SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

<u>Pollutant</u>	Stream Code	Sample Typet	Conc. Source	entration Day 1	ns (mg/1) Day 2	Day 3
Foxic Pollutants (Continued)				به ۲۰۰۰	· · · · ·	
117. beryllium	203 185	2	<0.010 <0.01		0.36 <0.01	0.05
	3 822	3	<0.0002	1.4 0.012	1.32	1.9
1 . cadmium	203	2	<0.050 <0.01	•	28.0 3.4	16.0
	185 3 822	1 3 1	0.0002	6 0.21	5.8	8.8
19. chromium (total)	203	2	<0.050		25.0	2.8
	185 3	1 3	<0.01 0.0003	31.0 <0.001	11.1 15.0	19.0
20. copper	822 203	1 2	<0.050		,800	150.0
20. copper	185	1 3	<0.01 0.017	220	55.0 210	320
	822	. 1		14		
121. cyanide (total)	203 185	1 1 1	0.05 <0.02	(0.001	<0.02 <0.02	<0.02
	3 822	1	0.052	<0.001 0.67	<0.0001	<0.001

SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

<u>Pollutant</u>	Stream _Code	Sample Typet	Concentrations (mg/1) Source Day 1 Day 2 Day 3	SECONDARY
Toxic Pollutants (Continued)				DAF
122. lead	203 185 3 822	2 1 3 1	<0.10 490.0 90.0 <0.10 10 0.020 100 80 110 5.0	RY PRECIOUS
123. mercury	203 185 3 822	2 1 3 1	0.0002 0.002 0.002 0.002 0.003 (0.0002 (0.	OUS METALS
124. nickel	203 185 3 822	2 1 3 1	<0.10A	S SUBCATEGORY
125. selenium	203 185 3 822	2 1 3 1	<0.10A B B <0.003	GORY
126. silver	203 185 3 822	2 1 3 1	$\begin{array}{cccccccccccccccccccccccccccccccccccc$	SECT - V

SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

<u>Pollutant</u>	Stream Code	Sample <u>Typet</u>	<u>Concentra</u> Source Day	and the second secon	
Toxic Pollutants (Continued)					
127. thallium	203 185 3	2 1 3	<0.010 <0.002 <0.001 0.3		
	822	1	0.5	1	
128. zinc	203 185 3 822	2 1 3 1	0.100 2.5 <0.010 3,400 1.0	4,500 2,800	2,000 3,400
Nonconventional Pollutants					
acidity	203 185	2 1	<1	8,000 85	930
alkalinity	203 185 3	2 1 3	98 127 16	<1 <1	<1
alum. num	203 185	2 1	0.20 <0.050	1,070.0 44	50.0
ammonia nitrogen	203 185 822	2 1 1	0.04 <0.01 340	3,300 980	8.5

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SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

Pollutant	Stream <u>Code</u>	Sample Typet	Concentra Source Day	
Nonconventional Pollutants (Continued)			
barium	203	2	<0.050	<5.0 <5.0
	185	1	0.070	1.9
boron	203 -	2	<0.10	10.0 <10.0
	185	1	<0.009	35
calcium	203	2	37.7	80.0 50.0
	185	1	11	150
	3	3	15 16	19 20
chemical oxygen demand (COD)	203 185	2 1	<5	20,000 >50,000
chloride	203	2	14	78,000 11,000
	185	1	52	56,000
cobalt	203	2	<0.050	15.0 <5
	185	1	<0.006	14.5
fluoride	203	2	0.28	5.0 1.1
	185	1	1.1	3.1
gc d	3	3	0.025 2.9	4.4 4.2
iros:	203 185 3 822	2 1 3 1	<0.050 0.31 0.29 2,100 5,400	790.0 160.0 825 1,900 2,600

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SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

	Stream Sample Concentration					۰۰. د. انتظار بر این در این در این در
Pollutant	<u>Code</u>	Typet	Source	Day 1	Day 2	Day 3
aconventional Pollutants (Continued))		. * ••			· .
gnesium	203 185	2 1	8.50 2.4		100.0 28	20.0
	3	3	3.1	33	34	56
agane 3e	203 185	2 1	<0.050 <0.01		6.60 6.2	1.40
molyi eram	203 185	2 1	<0.050 <0.002	· · ·	10.0 0.89	<5.0
phenol.cs	3 822	1	0.015	0.78 0.092	0.44	0.51
phosphate	203 185	2 1	14			210
sodium	203 185	2 1	9.00 54		,090.0 ,000	940.0
sulfate	203 185	2 1	57 13	82	2,000 41 195	,000
tin	203 185	2 1	<0.050 <0.12		54.9 <0.12	9.9

SECONDARY PRECIOUS METALS SUBCATEGORY SECT

SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

Pollutant	Stream	Sample	Conc.	entrations (mg/1)
	Code	Typet	Source	Day 1 Day 2 Day 3
onconventional Pollutants (Continued))			
¢itanium	203	2	<0.050	125.0 <5.0
	185	1	<0.005	0.16
total organic carbon (TOC)	203	2	4.3	450 590
	185	1	43	166
total solids (TS)	203	2	380	17,000 120,000
	185	1	410	77,000
vanadium	203	2	<0.050	<5.0 <5.0
	185	1	<0.003	0.090
yttrium	203	2	<0.50	<5.0 <5.0
	185	1	<0.002	0.012
Conventional Pollutants				
oil and grease	203 185 3 822	1 1 1	<1 <1 1.6	4 3 <1 33 35 14 30

SECONDARY PRECIOUS METALS SUBCATEGORY SECT

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SECONDARY PRECIOUS METALS SAMPLING DATA CEMENTATION TANK EFFLUENT

. *	Stream	Sample	Concentrations (mg/1)			
<u>Pollutant</u>	Code	Typet	Source	Day 1	Day 2	Day 3
Conventional Pollutants (Continued)						
total suspended solids (TSS)	203 185	2	60 8		570 895	70
	3	. 3		48	100	61
pH (standard units)	203 185	2	7.5		0.40 3.75	1.00
	3	3	6.8	0.3	0.5	1.2

Frample Type Code: 1 - One-time grab 2 - Manual composite during intermittent process operation 3 - 8-hour manual composite

(a) Reported together.

A - Detection limit raised due to interference.

B - Chemical interference.

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Table V-23

SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT A

Pollutant	Stream <u>Code</u>	Sample Typet	Conc Source	entration Day 1	ns (mg/1) Day 2	Day 3	SECONDARY
		and the second second	6	ann an the second s			ÖN
<u>s xic Polletants</u>							DAI
4. antimony	209	5 5	<0.01	<0.40A	0.40	0.120	
	212	5	<0.01	<0.20A	<0.50A	<0.50A	PRECIOUS
	215	5 5	<0.01	<0.50A	0.18	<0.01	Ĕ
	218	5	<0.01	<0.20A	<0.01	<0.20A	Ĥ
	221	5	<0.01	<0.50A	<0.20A	<0.20A	g
	224	6	<0.01	<0.01	<0.20A	0.390	
5. arsenic	209	5	<0.01	0.04	0.06	0.03	METALS
	212	5	<0.01	<0.05A	<0.05A	<0.05A	ΓA
	215	5 5	<0.01	<0.05A	<0.10A	<0.05A	Ľ
,	218	5 5	<0.01	<0.10A	<0.10A	<0.10A	
•	221	5	<0.01	<0.20A	<0.20A	<0.20A	US US
	224	6	<0.01	<0.40A	<0.05A	<0.05A	BCł
7. beryllium	209	5	<0.01	0.01	0.10	0.05	SUBCATEGORY
	212	5 5	<0.01	0.01	0.05	0.07	GO
	215	5	<0.01	0.01	0.05	0.07	1X
	218	5 5 5 6	<0.01	0.01	0.05	0.08	• •
	221	5	<0.01	0.02	0.06	0.07	
	224	6	<0.01	0.01	0.02	0.08	S
118. cadmium	209	5	<0.05	1.70	7.80	5.90	SECT
	212	5 5 5	<0.05	2.20	3.90	5.70	1
	215	5	<0.05	2.40	3.90	5.30	
	218	5	<0.05	2.30	4.0	5.60	<
	221	5	<0.05	2.30	4.1	5.7	
	224	6	<0.05	3.6	2.3	5.7	

SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT A

		Stream	Sample	Concentrations (mg/l)			
Pollutant		Code	Typet	Source	Day 1	Day 2	Day 3
<u>Foxic Pollutants</u>					•		1. n
19. chromium (total)	`	209	5	<0.05	2.0	6.30	3.50
	•	212	5	<0.05	1.50	3.20	3.20
		215	5	<0.05	1.40	2.80	3.0
		218	5	<0.05	1.40	2.80	2.80
		221	5	<0.05	1.6	2.8	2.9
		224	6	<0.05	1.8	1.4	3.5
20. copper		209	5	<0.05	200.0	520.0	230.0
	•	212	5	<0.05	370.0	300.0	320.0
**		215	5	<0.05	390.0	320.0	310.0
	· · · ·	218	5	<0.05	400.0	330.0	330.0
		221	5	<0.05	430	330	340
· · · · ·		224	6	<0.05	210	100	190
21. cyanide (total)		209	5	0.05	<0.02	<0.02	<0.02
· · · · · · · · · · · · · · · · · · ·		212	5	0.05	<0.02	<0.02	<0.02
		215	5	0.05	<0.02	<0.02	<0.02
		218	5	0.05	<0.02	<0.02	<0.02
- 	4	221	5	0.05	<0.02	<0.02	<0.02
	• • •	224	6	0.05	<0.02	<0.02	<0.02
22. lead		209	5	<0.10	24.0	110.0	72.0
	·	212	່ 5	<0.10	25.0	39.0	44.0
	an a	215	5	<0.10	26.0	33.0	36.0
· · ·	1990 - A.	218	2. 5 2.	<0.10	24.0	33.0	34.0
		221	5	<0.10	25.0	32.0	35.0
		224	6	<0.10	23.0	19.0	33.0

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SECONDARY PRECIOUS METALS SUBCATEGORY

SECT -

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT A

Pollutant	Stream Code	Sample Typet_	Conc Source	entratio Day 1	ns (mg/1) Day 2	Day 3
<u>I O I I d'ant</u>		<u></u>	boulou	<u></u>	<u></u>	<u></u>
<u>Toxic Pollutants</u>						-
123. mercury	209	5 5	0.0002	0.001	0.002	0.008
	212	5	0.0002	0.005	0.005	0.009
	215	5 5 5	0.0002	0.003	0.009	0.004
	218	5	0.0002	0.009	0.04	0.008
	221	5	0.0002	0.008	0.018	0.021
	224	6	0.0002	0.007	0.051	0.009
124. nickel	209	5	<0.20	24.0	130.0	57.0
	212	5	<0.20	34.0	58.0	68.0
	215	5 5 5 5	< 0. 20	34.0	62.0	66.0
	218	5	<0.20	34.0	64.0	68.0
	221	5	<0.20	36.0	64.0	68.0
	224	-6	<0.20	48.0	36.0	90.0
125. selenium	209	5	<0.10A	<0.10A	<0.10A	<0.10A
	212	5 5	<0.10A	<0.10A	В	В
	215	5	X0.10A	<0.10A	В	В
	218	5 5 5 6	<0.10A	<0.10A	<0.10A	<0.10A
	221	5	<0.10A	<0.10A	В	В
	224	6	<0.10A	<0.10A	<0.10A	В
26. silver	209	5	<0.01	1.30	4.30	2.70
	212	5	<0.01	1.80	2.20	2.90
	215	5 5	<0.01	1.30	1.90	2.0
	218	5	<0.01	1.50	2.40	1.40
	221	5 5	<0.01	1.40	2.10	1.30
,	224	6	<0.01	0.94	0.74	1.0

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SECONDARY PRECIOUS METALS SUBCATEGORY

SECT -

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT A

		Stream	Sample	ncentratio	ons (mg/l)		
Pollutant		Code	Typet	Source	Day 1	Day 2	Day 3
Toxic Pollutants				. .			а т Ал
127. thallium		209 212	5	<0.01 <0.01	B <0.50A	<0.20A <0.20A	B <0.20A
	· .	215	5 5 5 5 5	<0.01	<0.20A	B	<0.50A
		218 221	5	<0.01 <0.01	<0.02A <0.50A	<0.50A <0.02A	В <0.5А
		224	6	<0.01	<0.5A	B	B
128. zinc		209	5	0.10		3,400	5,200
		212	5 5 5 5 5 6	0.10			5,200 5,000
		215 218	·) 5	0.10	2,600 4	+,400 (+,500 (5,100
		221	5	0.10	2,800	i,500 (5,300
	. 1	224	6	0.10	4,100 2	2,600	5,100
N conventional Pollutants			×			···*	
a dity	· · · ·	209	5			3,400	980
•		212	5	<1	<1	170	180
		215 218	5 5 5 5 5 6	<1 <1	<pre>< <1 </pre> <pre></pre>	29 28	18 29
	-	221	5		<1	17	29
		224	6`	<1	12	53	48
a .alinity		209	5	98	<1	<1	`_<1
		212	5 5	98	23 23	<1 <1	<1 <1
	- - -	215 218	5	98 98	25	<1 <1	<1
• • • • • • • • • • •		221	5	98	41	<1	<1
		224	6	98	<1	<1	<1 .

SECONDARY PRECIOUS METALS SUBCATEGORY

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT A

n 11	Stream	Sample	Conc	1)		
Pollutant	Code	<u>Typet</u>	Source	Day 1	Day	2 Day 3
Nor. conventional Pollutants (Continued))					
al. inum	209	5	0.20	1.5.0	230.0	112.0
	212	5	0.20	15.0	96.0	121.0
	215	5 5	0.20	15.0	103.0	121.0
	218	5	0.20	17.0	107.0	126.0
	221	5	0.20	16.0	105.0	128.0
	224	6	0.20	28.0	34.0	138.0
ammonia nitrogen	209	. 5	0.04	470	110	1,100
	212	5	0.04	670	670	1,100
	215	5	0.04	580	700	1,100
	218	5 5 5 6	0.04	590	760	910
	221	5	0.04 1,	080	760	1,150
	224	6	0.04 1,	700	640	770
barium	209	5	<0.05	1.0	<0.5	<0.5
	212	5	<0.05	<0.5	<0.5	<0.5
•	215	5 5 5 5 5 6	<0.05	<0.5	<0.5	<0.5
	218	5	<0.05	<0.5	<0.5	<0.5
	221	. 5	<0.05	<0.5	<0.5	<0.5
	224	6	<0.05	<0.5	<0.5	<0.5
						· ř
boron	209	5	<0.10	<1.0	3.0	2.0
	212	5	<0.10	<1.0	1.0	2.0
	215	5 5 5 5 5	<0.10	<1.0	2.0	2.0
	218	5	<0.10	<1.0	2.0	2.0
	221	-	<0.10	<1.0	2.0	2.0
	224	6	<0.10	<1.0	<1.0	2.0

SECONDARY PRECIOUS METALS SUBCATEGORY

SECT - V

SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT A

Nonconventional Pollutants (Continued) calcium 209 5 212 5 215 5 218 5 221 5 218 5 224 6 chemical oxygen demand (COD) 209 5 218 5 218 5 218 5 212 5 212 5 212 5 213 5 214 6 chloride 209 5 215 5 214 6 215 5 216 5 217 5 218 5 215 5 215 5 215 5 215 5 218 5	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
215 5 218 5 221 5 224 6 chemical oxygen demand (COD) 209 5 212 5 215 5 218 5 221 5 218 5 224 6 chloride 209 5 212 5 213 5 214 5 215 5 216 5 217 5 218 5 218 5 218 5 218 5 217 5 218 5 217 5 212 5 215 5	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
215 5 218 5 221 5 224 6 chemical oxygen demand (COD) 209 5 212 5 215 5 218 5 221 5 218 5 224 6 chloride 209 5 212 5 213 5 214 5 215 5 216 5 217 5 218 5 218 5 218 5 218 5 217 5 218 5 217 5 212 5 215 5	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
218 5 221 5 224 6 chemical oxygen demand (COD) 209 5 212 5 215 5 218 5 218 5 218 5 218 5 218 5 224 6 chloride 209 5 212 5 215 5 215 5 215 5 215 5	$\begin{array}{rrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrrr$
221 5 224 6 chemical oxygen demand (COD) 209 5 212 5 215 5 218 5 221 5 218 5 224 6 chloride 209 5 212 5 213 5 224 6 212 5 212 5 212 5 212 5 215 5	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
224 6 chemical oxygen demand (COD) 209 5 212 5 215 5 218 5 221 5 218 5 221 5 218 5 221 5 218 5 221 5 218 5 221 5 212 5 212 5 212 5 215 5	37.7 36.0 37.0 40.0 <5
chemical oxygen demand (COD) 209 5 212 5 215 5 218 5 218 5 221 5 221 6 224 6 chloride 209 5 212 5 225 5 215 5	<5 4,000 >50,000 3,100 <5 6,100 2,400 2,400
212 5 215 5 218 5 221 5 221 5 221 5 224 6 chloride 209 5 212 5 212 5 215 5	<5 6,100 2,400 2,400
212 5 215 5 218 5 221 5 221 5 224 6 chloride 209 5 212 5 215 5	
215 5 218 5 221 5 224 6 chloride 209 5 212 5 215 5	<5 3,900 3,700 3,400
218 5 221 5 224 6 chloride 209 5 212 5 215 5	
224 6 chloride 209 5 212 5 215 5	<5 5,000 4,900 3,600
224 6 chloride 209 5 212 5 215 5	<5 3,200 5,500 5,200
212 5 215 5	<5 3,600 6,200 5,100
212 5 215 5	
212 5 215 5	14 12,000 13,000 17,000
	14 28,000 21,000 18,000
010 5	14 15,000 14,000 18,000
	14 32,000 14,000 14,000
221 5	14 15,000 15,000 18,000
224 6	14 18,000 14,000 19,000
cobalt 209 5	<0.05 1.50 3.50 2.0
212 5	<0.05 1.50 2.0 2.0
215 5	<0.05 1.50 2.0 2.0
218 5	<0.05 1.50 2.0 2.0
221 5	
221 5	<0.05 1.50 2.0 2.0

SECONDARY PRECIOUS METALS SUBCATEGORY

SECT -

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT A

	Stream	Sample	<u>Concentrations (mg/l)</u>				
Pollutant	<u>Code</u>	<u>Typet</u>	Source	Day 1	Day 2	Day 3	
Nonconventional Pollutants (Continued	i)		. · · · ·			:	
fluoride	209	5	0.28	2.4	0.13	0.73	
	212	5	0.28	2.3	0.85	0.88	
	215	5	0.28	0.48	0.88	0.87	
	218	5	0.28	0.61	-0.86	0.89	
	221	5	0.28	0.32	.0.87	0.92	
	224	.6	0.28	0.68	∽ 0.63	1.01	
iron	209	5	<0.05	20.0	200.0	110.0	
· · · · · · · · · · · · · · · · · · ·	212	5	<0.05	29.0	79.0	120.0	
	215	5	<0.05	28.0	84.0	110.0	
	218	5	<0.05	29.0	86.0	110.0	
	221	5	<0.05	.30.0	83.0	120.0	
· · ·	224	6	<0.05	250.0	190.0	340.0	
magnesium	209	5 5	8.5	9.0	28.0	19.0	
	212		8.5	9.0	16.0	19.0	
	215	5	8.5	9.0	17.0	19.0	
	218	5	8.5	9.0	17.0	19.0	
	221	5	8.5	8.0	17.0	19.0	
	224	6	8.5	8.0	11.0	21.0	
manganese	209	5	<0.05	0.15	0.85	0.90	
- 6	212	5	<0.05	0.30	0.75	1.20	
	215	× 5	<0.05	0.30	0.85	1.10	
	218	5	<0.05	0.25	0.85	1.20	
•	221	5	<0.05	0.25	0.85	1.10	
	224	6	<0.05	1.30	1.20	2.60	
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SECONDARY PRECIOUS METALS SUBCATEGORY SECT ſ

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT A

· · · · · ·	м.	Stream	Sample	C	oncentra	tions (mg/	1)
Pollutant		Code	Typet_	Sourc	e Day		
Nonconventional Pollutants	(Continued)		•	•			
molybdenum		209	5	<0.05		3.0	1.50
		212	. 5	<0.05	<0.5	0 1.0	<0.050
		215	5	<0.05		0.50	<0.50
•		218	5 5	<0.05	<0.5	0 <0.50	<0.50
		221	5	<0.05	<0.5) <0.50	<0.50
		224	6	<0.05	<0.5	0 <0.50	<0.50
phosphate	•	209	5	14	13,000	600	310
		212	5		37,000	340	360
		215	5	14	3,000	360	360
		218	5	- 14	1,300	380	410
	•	221	5	14	680	380	600
	· .	224	6	14	500	130	600
soctum	•	209	5 -	9.0	1,250	2,650	799.0
		212	5	9.0	9,540	8,150	6,910
· · · · · · · · · · · · · · · · · · ·	-	215	5	9.0	9,480	8,300	6,860
		218	5		10,500	8,920	7,290
· · ·		221	5		10,200	8,640	7,410
		224	6		11,600	8,780	9,390
sulface		209	5	57	6,000	15,000	11,000
	x	212	5	57	9,000	11,000	15,000
		215	5	57	9,000	15,000	11,000
		218	5		10,000	10,000	13,000
	· .	221	5		12,000	12,000	15,000
· · · · ·		224	6	57	9,000	11,000	18,000

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT A

	Stream	Sample	Con	centrations (mg/l)				
Pollutant	_Code	<u>Typet</u>	Source	Day 1	Day 2	Day 3		
acon entional Pollutants (Contin	nued)							
tín	209	5	<0.05	56.9	22.4	7.90		
	212	5	<0.05	5.70	31.4	9.10		
	215	5 5 5 5 5	<0.05	<0.5	6.4	<0.5		
	218	5	<0.05	<0.5	<0.5	<0.5		
	221	5	<0.05	<0.5	<5.0	<0.5		
	224	6	<0.05	<5.0	<5.0	<5.0		
titanium	209	5	<0.05	<0.50	24.5	11.50		
	212	5	<0.05	<0.50	<0.50	1.50		
	215	5	<0.05	<0.50	<0.50	<0.50		
·	218		<0.05	<0.50	<0.50	<0.50		
	221	5	<0.05	<0.50	<0.50	<0.50		
	224	6	<0.05	<0.50	<0.50	<0.50		
total organic carbon (TOC)	209	5	4.3	100	300	330		
	212	5 5	4.3	150	240	280		
	215	5	4.3	140	240	300		
	218	5	4.3	130	240	300		
	221	5	4.3	150	200	270		
	224	6	4.3	150	230	240		

SECT - V

SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT A

	Stream	Sample	Concentrations (mg/l)			
Pollutant	Code	Typet	<u>Source</u> D	ay 1 Day		
		, P	The second se			
Nonconventional Pollutants (Continued)	÷ • •	· · · ·			- - -	
total solids (TS)	209	5	380	40,000	30,000	
	212	5 °	380 °	34,000	41,000	
	215	5	380 37,000		,42,000	
	218	5.	380 38,000	41,000	43,000	
	221	5 °	380 42,000		44,000	
	224	. 6	380 53,000	35,000	46,000	
vanadium	209	5	<0.05 <0	.50 <0.5	0 <0.50	
Vallaulum	212	5		.50 <0.5		
	215	- 5	• • • • • •	.50 <0.5	-	
	218	5		.50 <0.5		
	221	5		.50 <0.5	0 <0.50	
	224	6		.50 <0.5		
	209	5	<0.5 <0	.50 <0.5	0 <0.50	
yttrium	209	5		.50 <0.5		
		5		.50 <0.5		
	215	2		.50 <0.5		
	218	5		.50 <0.5		
	221 224	5		.50 <0.5		

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT A

Pollutant	Stream -	Sample	Cor	Concentrations (mg/l)				
Pollutant	Code	<u>Typet</u>	Source	Day	<u>1 Day 2</u>			
<u>Criventional Pollutants</u>								
🚈 i and grease	209	1	<1	3	6	5		
	212	1	<1	3 2	6	<1		
	215	1	<1	1.5	Š	či		
	218	1	<1	2	5 2	λί		
	221	1 .	<1	<1	<1	3		
	224	1	< 1	2	4	6		
actal suspended solids (TSS)	209	5	60	160	140	140		
	212	5	60	140	290	140		
	215	5	60	88	160	180		
• • • • • • • • • • • • • • • • • • •	218	5 5	60	160	5,700	120		
	221	5	60	360	130	110		
	224	6	60	580	120	190		
pH (st neard units)	209	5	7.5	1.0	1.1	1.2		
	212	5 5	7.5	4.7	4.1	3.9		
	215	5	7.5	4.8	4.2	4.3		
	218	5	7.5	4.81	4.21	4.1		
	221	5	7.5	4.91	4.4	4.1		
	224	6	7.5	4.37	4.1	3.92		
<pre>tSample Type Code: 1 - One-time grab</pre>		Ň						

mpl	е Туре	Code:	1	-	One-time	grab	
					0/ 1.		

5 - 24-hour manual composite6 - 24-hour automatic composite

A - Detection limit raised due to interference.

B - Chemical interference.

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Table V-24

SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT B

	Stream	Sample	الشوارية استحصي ويستعدا المشاوية المتناز المتكر المتحاكم والمرجوع والبراج	entrations Day 1	3 (mg/1) Day 2	Day 3
Pollutant	Code	Typet	Source	Day		
ic Pollutants						·
, acenaphthene	184	1	ND	ND		
acrolein	184	1	ND	N D		
acrylonitrile	184	1·	ND	ND		
, benzene	184	1	ND	ND		•
5. benzidine	184	1	ND	N D	• • •	
6. carbon tetrachloride	184	1	ND	N D		, ,
7. chlorobenzene	184	1	ND	ND		•
8. 1,2,4-trichlorobenzene	184	1	ND	N D		
9. hexachlorobenzene	184	1	ND	ND		•
10. 1,2-dichloroethane	184	1	ND .	ND	- -	
11. 1,1,1-trichloroethane	184	1	0.01	0.01		. *
12. hexachloroethane	184	1	ND	ND		•
13. 1,1-dichloroethane	184	1	ND	ND		
14. 1,1,2-trichloroethane	184	1	ND	ND		, , , ,

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SECONDARY PRECIOUS METALS SUBCATEGORY

SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT B

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	Pollutant	Stream Code	Sample Typet	Conce Source	entrations Day 1	<u>mg/1)</u> Day 2	Day 3
<u>Toxic</u>	Pollutants (Continued)						
15.	1,1,2,2-tetrachloroethane	184	1	ND	ND		
16.	chloroethane	184	1	ND	ND		
17.	bis(chloromethyl)ether	184	1	ND	ND		
18.	bis(2-chloroethyl)ether	184	1	ND	N D		
19.	2-chloroethyl vinyl ether	184	1	ND	N D		
20.	2-chloronaphthalene	184	1	ND	ND		
21.	2,4,6-trichlorophenol	184	1	ND	ND		
22.	p-chloro-m-cresol	184	1	ŇD	N D		
23.	chloroform	184	1	ND	N D		
24.	2-chlorophenol	184	1	ND	N D		
23.	1,2-dichlorobenzene	184	1	ND	ND		
26.	1,3-dichlorobenzene	184	1	ND	ND		
27.	1,4-dichlorobenzene	184	1	ND	N D		
28.	3,3'-dichlorobenzidine	184	1	ND	N D		
28.	3,3'-dichlorobenzidine	184	I	ND	ND		

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT B

					,		
	Stream	Sample	Conc	s (mg/1)	ig/1)		
Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3	
Toxic Pollutants (Continued)			5. .	•			
29. 1,1-dichloroethylene	184	1	ND	N D			
30. 1,2- <u>trans</u> -dichloroethylene	184	1	ND	N D			
31. 2,4-dichlorophenol	184	• 1	ND	N D			
2. 1,2-dichloropropane	184	- 1	ND	ND		-	
3. 1,3-dichloropropene	184	1	ND	N D			
34. 2,4-dimethylphenol	184	1	ND	ND			
5. 2,4-dinitrotoluene	184	1	ND	ND		-	
. 2,6-dinitrotoluene	184	1	ND	ND			
'. 1,2-diphenylhydrazine	184	1	ND	N D			
3. ethylbenzene	184	- 1	ND	ND		•	
. fluoranthene	184	1	ND	ND	- -		
. 4-chlorophenyl phenyl ether	184	1	ND	ND			
41. 4-bromophenyl phenyl ether	184	1	ND	ND		· · .	
42. bis(2-chloroisopropyl)ether	184	1	ND	ND		-	
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SECONDARY PRECIOUS METALS SUBCATEGORY SECT -

SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT B

	Pollutant	Stream <u>Code</u>	Sample Typet	Conce Source	entrations Day 1	(mg/1) Day 2	Day 3
<u>Toxic</u>	Pollutants (Continued)						
43.	bis(2-choroethoxy)methane	184	1	ND	ND		
44.	methylene chloride	184	1	ND	ND		
45.	methyl chloride (chloromethane)	184	1	ND	N D		,
5.	methyl bromide (bromomethane)	184	1	ND	N D		
•	bromoform (tribromomethane)	184	1-	ND	N D		
3.	dichlorobromomethane	184	1	ND	N D		
4. 9.	trichlorofluoromethane	184	1	ND	N D		
30.	dichlorodifluoromethane	[^] 184	. 1 ·	ND	N D		
51.	chlorodibromomethane	184	1	ND	ND		
52.	hexachlorobutadiene	184	1	ND	N D		
53.	hexachlorocyclopentadiene	184	- 1	ND	N D		
54.	isophorone	184	1	ND	N D		-
55.	naphthalene	184	[`] 1	ND	ND	- '	•
56.	nitrobenzene	184	1	ND	ND)	

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT -

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT B

		Stream	Sample	وجرواء وترجيعه ويرجيه والكارة فالتحط الرجية ويستبدوا بدوارة	entrations		
-	<u>Pollutant</u>	Code	Typet	Source	Day 1	Day 2	Day 3
Toxic	Pollutants (Continued)	•	- 	, -			
57.	2-nitrophenol	184	1	ND	ND		
58.	4-nitrophenol	184	1	ND	ND		
59.	2,4-dinitrophenol	184	1	ND	ND		
60.	4,6-dinitro-o-cresol	184	1 *	ND	ND	r	
61.	N-nitrosodimethylamine	184	1.	ND	ND		•
62.	N-nitrosodiphenylamine	184	1	ND	ND		÷.
63.	N-nitrosodi-n-propylamine	184	1	ND	ND	·	
64.	pentachlorophenol	184	1	ND	N D		-
65.	phenol	184	1	ND	0.028		
66.	bis(2-ethylhexyl) phthalate	184	. 1	0.026	0.030		
67.	butyl benzyl phthalate	184	1	ND	N D		
68.	di-n-butyl phthalate	184	1	ND	0.002	· · · · ·	
÷9.	di-n-octyl phthalate	184		ND	0.001		
70.	diethyl phthalate	184	1 1 1	ND	ND	•	Х

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - V

SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT B

<u>Pollutant</u>	Stream <u>Code</u>	Sample <u>Type</u> t	Conc Source	entrations Day 1	(mg/1) Day 2	Day 3
Toxic Pollutants (Continued)						
71. dimethyl phthalate	184	1	ND	0.02		
72. benzo(a)anthracene	184	1	ND	ND		
73. benzo(a)pyrene	184	1	ND	ND		
74. benzo(b)fluoranthene	184	1	ND	N D		
75. benzo(k)fluoranthane	184	1	ND	ND		
76. chrysene	184	1	ND	N D		
77. acenaphthylene	184	1	ND	ND		
78. anthracene (a)	184	1	ND	ND		
79. benzo(ghi)perylene	184	1	ND	ND		
80. fluorene	184	. 1	ND	ND		
D. phenanthrene (a)	184	1	ND	ND		
•	184	1	ND	ND		
	184	1	ND	ND		
<pre>indeno (1,2,3-c,d)pyrene</pre>	184	1	ND	ND		
84. pyrene	104	8	125	- F4		

SECONDARY PRECIOUS METALS SUBCATEGORY

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT B

		Stream	Sample	Conc	entration	s (mg/l)	
• • •	Pollutant	<u>Code</u>	Typet	Source	Day 1	Day 2	Day 3
Toxic	Pollutants (Continued)					• •	
85.	tetrachloroethylene	184	1	ND	N D		
86.	toluene	184	1	ŇD	ND		
87	trichloroethylene	184	1	ND	ND		
88.	vinyl chloride (chloroethylene)	184	1	ND	ND	-	
114.	antimony	184 011 012	1 1 1	<0.003 <0.003 <0.003	2.6 0.20 0.35	0.21 0.097	1.8 0.29
115.	arsenic	184 011 012	1 1 1	<0.003 <0.003 <0.003	0.84 0.12 0.061	0.073 0.023	0.43 0.043
117.	beryllium	184 011 012	1 1 1	<0.01 <0.01 <0.01	<0.01 <0.01 <0.01	<0.01 <0.01	<0.01 <0.01
118.	cadmium	184 011 012	1 1 1	<0.01 <0.01 <0.01	<0.01 1.0 <0.01	0.53 <0.01	0.81 <0.01
119.	chromium (total)	184 011 012	1 1 1 1	<0.01 <0.01 <0.01	0.42 0.22 0.11	0.04 <0.01	0.21 <0.01

SECONDARY PRECIOUS METALS SUBCATEGORY SECT

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT B

Pollutant	Stream Code	Sample Typet	<u>Conc</u> Source	entration Day 1	ns (mg/1) Day 2	Day 3	SECONDARY
Toxic Pollutants (Continued)							NDA
120. copper	184	1	<0.01	0.40	• •	0.0	
	011 012	1	<0.01 <0.01	2.0 1.1	1.8 0.12	23 <0.01	PRECIOUS
121. cyanide (total)	184	1	<0.02	<0.02			CIO
	011 012	1 1	<0.02 <0.02	<0.02 <0.02	<0.02 <0.02	<0.02 <0.02	
122. lead	184	1	<0.01	3.2			METALS
	011 012	1	<0.01 <0.01	0.59 0.28	1.5 <0.10	2.0 <0.10	
123. mercury	184	1	0.002	0.38			SUBO
1238 mercury	011 012	· 1	0.002	0.14 0.002	0.005 <0.0002	0.175 <0.0002	SUBCATEGORY
104 stabal	184	• 1	0.075	101	(010002	(00000-	IGOR
124. nickel	011	1	0.075	90	30 0.61	60 1.7	Ř
· · · · · · · · · · · · · · · · · · ·	012	1	0.075	0.60	0.01	1/	ល
125. selenium	184 011	1	<0.003 <0.003	0.18 0.010	0.013	0.082	SECT
	012	1	<0.003	0.014	0.010	0.012	י ל
126. silver	184 011	1	<0.0005 <0.0005	0.041 0.081	0.069	0.056	
	012	1	<0.0005	0.035	0.049	0.077	

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT B

	Stream	Sample	Concentrations (mg/1)				
Pollutant	Code	Typet	Source	Day 1	Day	2 Day 3	
Toxic Pollutants (Continued)		. (* .	-		
127. thallium	184	1	<0.002			2 <0.002	
	011 012	1	<0.002 <0.002				
128. zinc	184	1		1,900	0.00	1 200	
	011 012	1	2.5 2.5	2,800 56	6.9	1,300 41	
Nonconventional Pollutants			•			н 1973 — Малан 1974 — Малан Алан	
acidity	184	1	<1	<1 <1		. <1	
	011 012	1	<1 <1	<1	<1 <1	< <u>1</u>	
alkalinity	184	1		8,000	4 000		
	011 012	1		8,000 1,200	1,090 1,000	8,400 1,460	
aluminum	184	1	<0.05	7.7			
	011 012	1	<0.05 <0.05	49 2.8	24 1.8	11 <0.05	
ammonia nitrogen	184	1	<0.01	2,300			
	011 012	1	<0.01 <0.01	2,300 670	220 1,070	<0.01 <0.01	

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT B

	Stream	Sample	Conc	entratio	ons (mg/l	
Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3
Noncoventional Pollutants (Continued)						
barium	184	1	0.07	0.66		
	011	1	0.07	1.2	1.1	0.71
	012	1	0.07	0.24	0.85	0.05
boron	184	1	<0.009	3.8		
	011	1	<0.009	2.9	2.7	3.4
	012	1	<0.009	3.4	2.2	3.3
calcium	184	1	11	430		
	011	1		600 (5,000	1,100
• • •	012	1		730	5,200	19
chemical oxygen demand (COD)	011	1	٤,			990
	012	1	· · ·		•	640
chloride	184	1.	52 40,	000	7 10	÷,
	011	1	52 45,		3,000	1,000
	012	1	52 21,	000	200	1,200
cotalt	184	1	<0.006	4.6		
	011	1	<0.006	5.7	2.7	2.9
· · · · · ·	012	1 1	<0.006	0.70	0.17	0.18
fluoride	184	1	1.1	11.0		
	011	1	1.1	0.81	1.9	1.5
	012	1	1.1	7.9	2.0	7.9

SECONDARY PRECIOUS METALS SUBCATEGORY ş ۰, SECT I

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT B

				<u>Concentrations (mg/l)</u>				
Poll	utant	Code	Typet	Source	Day 1	Day 2	Day 3	
coventional	Pollutants (Continued)			-	•		· .	
n		10/	•	0.01		5 5 N. 	an a	
18		184 011		0.31	310	000	000	
	· · ·	012	1	0.31	460	200	200	
•	· · · · · · · · · · · · · · · · · · ·	012	•	0.31	- 1.1	0.15	0.21	
ae gnesium	· · · · · · · · · · · · · · · · · · ·	184	1	2.4	15.0			
		011	1	2.4	21.0	12.0	11.0	
		012	1	2.4	1.0	1.3	2.2	
manganese		184	1	` <0.01	3.0			
		011	1	<0.01	5.0	2.4	2.2	
•		012	1	<0.01	<0.01	<0.01	<0.01	
molybdenum		184	1	10 000		-		
		011	- 4	<0.002 <0.002	0.23	0 075	0 10	
		012	1	<0.002 <0.002	0.15 0.039	0.075	0.13	
		012		10.002	0.039	0.019	0.066	
phosphate		011	1.	1			<4	
		012	1	r.			K4	
		- -					N-V	
sodium		184	1	54 26	,000			
		011	1 w 1			,000 22	.000	
	1 -	012	1	54 14	,000 11		,000	
sulfate		10/	1	1.0 0				
		184	1	13 3	,400			
		011 012	· · · · · · · · · · · · · · · · · · ·	13 13 2	<0.6		,500	
		V12	. •	13 2	,100	830 1	,500	

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT B

Pollutant	Stream Code	Sample Typet	Concentrations (mg/1) Source Day 1 Day 2 Day 3
onconventional Pollutants (Continued)		DAR R
tin	184	1	<0.12 <0.12
	011 012	1	(0.12 (0.12 1.3 (0.12 (0.12 (0.12 6.9 (0.12
titanium	184	1	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
	011 012	1 1	
total organic carbon (TOC)	184	- 1	<0.005 0.056 0.11 <0.005 m H → H → H → H → H → H → H → H → H → H →
cocur orBanno oncom (===)	011 012	1 1	
total solids (TS)	184	1	410 87,000
	011 012	1 1	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
vanadium	184	1	
Vallautum	011 012	1	<pre><0.003 0.17 0.078 <0.003 <0.003 0.063 0.09 <0.003</pre>
· · · · · · · · · · · · · · · · · · ·	184	1	く0.002 0.015 く0.002 0.014 0.064 0.006 日
yttrium	011 012	1	<0.002 0.014 0.064 0.006 ∺ <0.002 0.014 0.046 <0.002 .
	012	U	

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT B

	Stream	Sample	Concentrations (mg/1)				
Pollutant	Code	<u>Typet</u>	Source	Day 1	Day 2	<u>Day 3</u>	
Conventional Pollutants	-	•		· ·		•	
oil and grease	184 011	1	<1 <1 <1	<1 6 <1	۲۱ ۲۱	<1 <1	
total suspended solids (TSS)	012 184 011 012	1	8 8 8	5,700 2,300 2,500	3,900 5,700	4,500 53	
pl (standard units)	184 011 012	1	7.36 7.36 7.36	8.61	10.00	8.76 8.65	

tSample Type Code: 1 - One-time grab

(a) Reported together.

Table V-25

SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT C

Pollutant	Stream <u>Code</u>	Sample Typet	<u>Conc</u> Source	entration Day 1	s (mg/1) Day 2	Day 3
Toxic Pollutants						
acenaphthene	7 2	3 6	ND ND	ND ND	ND ND	N D ND
2. acrolein	7 2	1 1	ND ND	nd Nd	ND ND	ND ND
3. acrylonitrile	7	1 1	ND ND	ND ND	ND ND	N D ND
. benzene	7 2	1	ND ND	ND ND	ND ND	ND ND
j. benzidine	7	3 6	ND ND	ND ND	ND ND	N D ND
ó. carbon tetrachloride	7 2	1	ND ND	ND ND	ND ND	ND ND
7. chlorobenzene	7 2	1	ND ND	ND ND	ND ND	N D ND
8. 1,2,4-trichlorobenzene	7 2	3 6	ND ND	ND ND	ND ND	ND N D
9. hexachlorobenzene	7 2	3	ND ND	ND ND	ND ND	N D ND

SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT C

		Stream	Sample	Concentrations (mg/l)			
	Pollutant	Code	Typet	Source	Day 1	Day 2	Day 3
ſoxic	Pollutants (Continued)			1. jan 1.	•	a**	
							, . ,
10.	1,2-dichloroethane	7	1	ND	ND	<0.01	ND
		2	1	ND	ND	ND	ND
ł.	1,1,1-trichloroethane	7	1	ND	ND	ND	ND
		2	1	ND	ND	ND	ND
12.	hexachloroethane		2		ND	ND	ND
12.	nexacitoroetnane	2	- J - L	ND ND	ND ND	ND ND	
		· ∠	Ŭ	ND	ND	ND	ND
13.	1,1-dichloroethane	· 7.	1	ND	ND	ND	ND
		2	1	ND	ND	ND	ND
1 2	1 1 0 traichleneathana	7	1	ND	NÐ	ND	ND
14.	1,1,2-trichloroethane	. 2.	· I 1	ND	ND	ND	ND
		• 6	J	ND	ND	ΠD	ND
15.	1,1,2,2-tetrachloroethane	7	· 1. 🔆	ND	ND	ND ¹	ND
		2	1	ND	ND	ND	ND
16.	chloroethane	7	1	ND	ND	ND	ND
	chiolocchanc	2	1	ND	ND	ND	ND
· .		· •	•		ne		
17.3	bis(chloromethyl)ether	7.	1	ND	ND	ND	ND
e .		2	1	ND	ND	ND	ND
18.	bis(2-chloroethyl)ether	7	3	ND	ND	ND	N D
	ore/a curorocular/ecuer	2	с	ND	ND	ND	ND

SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT C

Pollutant	Stream <u>Code</u>	Sample Type†	Conce Source	<u>Day 1</u>	<u>s (mg/l)</u> Day 2	Day 3
Toxic Pollutants (Continued)						
19. 2-chloroethyl vinyl ether	7	1	ND	ND	ND	N D
	2	1	ND	ND	ND	ND
20. 2-chloronaphthalene	7	3	ND	ND	ND	ND
	2	6	ND	ND	ND	ND
21. 2,4,6-trichlorophenol	7	3	ND	ND	<0.01	N D
	2	6	ND	ND	ND	ND
22. p-chloro-m-cresol	7	3	ND	ND	ND	ND
	2	6	ND	ND	ND	ND
23. chloroform	7 2	1 1	0.05 0.05	0.06 0.13	0.02	0.02
24. 2-chlorophenol	7	3	ND	ND	ND	ND
	2	6	ND	ND	ND	ND
25. 1,2-dichlorobenzene	7	3	ND	ND	ND	N D
	2	6	ND	ND	ND	ND
25. 1,3-dichlorobenzene	7	3	ND	ND	ND	ND
	2	6	ND	ND	ND	ND
27. 1,4-dichlorobenzene	7	3	ND	ND	ND	N D
	2	6	ND	ND	ND	ND

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT C

•		Stream	Conc	Concentrations (mg/l)			
	<u>Pollutant</u>	Code	Typet	Source	Day 1	Day 2	Day 3
Toxic	Pollutants (Continued)						
28.	3,3'-dichlorobenzidine	7 2	3 6	ND ND	ND ND	ND ND	N D ND
29.	1,1-dichloroethylene	7 2	1	ND ND	ND ND	ND ND	ND ND
30.	1,2- <u>trans</u> -dichloroethylene	7 2	1 1	ND ND	ND ND	ND ND	N D ND
31.	2,4-dichlorophenol	. 7 2	3 6	ND ND	ND ND	<0.01 ND	ND ND
32.	1,2-dichloropropane	7 2	1	ND ND	ND ND	ND ND	N D ND
•	1,3-dichloropropene	7 2	1	ND ND	ND ND	ND ND	ND ND
24.	2,4-dimethylphenol	72	3 6	ND ND	ND ND	ND ND	N D ND
35.	2,4-dinitrotoluene	7 2	3 6	ND ND	ND ND	ND ND	ND ND
36.	2,6-dinitrotoluene	7 2	3 6	ND ND	ND ND	ND ND	N D N D

SECONDARY PRECIOUS METALS SUBCATEGORY SECT -

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT C

	Pollutant	Stream Code	Sample Typet	Conc Source	centration Day 1	ns (mg/l) Day 2	Day 3
Tox [†] e_Pollutants (Continued)							
37.	1,2-diphenylhydrazine	7 2	3 6	ND ND	ND ND	ND ND	N D ND
38.	ethylbenzene	7 2	1	ND ND	ND ND	ND ND	ND ND
39.	fluoranthene	7 2	3 6	ND ND	ND ND	ND ND	N D ND
40.	4-chlorophenyl phenyl ether	7 2	3 6	ND ND	ND ND	ND ND	ND ND
41.	4-bromophenyl phenyl ether	7 2	3 6	ND ND	ND ND	ND ND	N D ND
42.	bis(2-chloroisopropyl)ether	7 2	3 · 6	ND ND	ND ND	ND ND	ND ND
43.	bis(2-choroethoxy)methane	7 2	3 6	ND ND	ND ND	ND ND	N D ND
44.	methylene chloride	7 2	1 1	<0.01 <0.01	<0.01 <0.01	<0.01 <0.01	<0.01 <0.01
45.	methyl chloride (chloromethane)	7 2	1	ND ND	ND ND	ND ND	N D ND

SECONDARY PRECIOUS METALS SUBCATEGORY SECT ł 4

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT C

	Stream	Sample	ومراد والمسادي والمراجع فالمتحدث والمتحدث والمتحدث	centration		
Pollutant	Code	<u>Typet</u>	Source	Day 1	Day 2	Day 3
Toxic Pollutants (Continued)				• •		
46. methyl bromide (bromomethane)	7 2	- 1	ND ND	ND ND	ND ND	N D ND
47. bromoform (tribromomethane)	7 2-	1 1	ND ND	ND <0.01	ND ND	ND ND
48. dichlorobromomethane	7 2	1 - 1 - 1 1	<0.01 <0.01	<0.01 <0.01	<0.01 <0.01	<0.01 <0.01
49. trichlorofluoromethane	7 2	· 1 1	ND ND	<0.01 ND	ND ND	ND ND
50. dichlorodifluoromethane	7 2	1	ND ND	ND ND	ND ND	N D ND
51. chlorodibromomethane	7 2	1	<0.01 <0.01	<0.01 0.02	ND ND	ND ND
52. hexachlorobutadiene	7 2	3	ND ND	ND ND	ND ND	N D ND
53. hexachlorocyclopentadiene	7 2	3	ND ND	ND ND	ND ND	ND ND
34. isophorone	7 2	3	ND ND	ND ND	ND <0.01	N D ND

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT

SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT C

		Stream	Sample	Conc			
	Pollutant	Code	<u>Typet</u>	Source	Day 1	Day 2	Day 3
Toxic	Pollutants (Continued)						
55.	naphthalene	7 2	3 6	ND ND	ND ND	ND ND	N D ND
56.	nitrobenzene	7. 2	3 6	ND ND	ND ND	ND ND	ND ND
57.	2-nitrophenol	7 2	3 6	ND ND	<0.01 <0.01	<0.01 <0.01	N D ND
58	4-nitrophenol	7 2	3 6	ND ND	<0.01 ND	0.019 <0.01	ND ND
59,	2,4-dinitrophenol	7 2	3 6	ND ND	ND ND	ND 0.110	N D ND
60.	4,6-dinitro-o-cresol	7 2	3 6	ND ND	ND ND	ND ND	ND ND
61.	N-nitrosodimethylamine	7 2	3 6	ND ND	ND ND	ND ND	N D N D
62.	N-nitrosodiphenylamine	7	3 6	ND ND	ND ND	ND ND	ND ND
63.	N-nitrosodi-n-propylamine	7 2	3 6	ND ND	ND ND	ND ND	N D ND

SECONDARY PRECIOUS METALS SUBCATEGORY SECT

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT C

	<u>Pollutant</u>	Stream <u>Code</u>	Sample Typet	<u>Conc</u> Source	<u>Day 1</u>	ns (mg/1) Day 2	Day 3
<u>Toxic</u>	Foliutants (Continued)		. ·		•		
64.	pentachlorophenol	7 2	3 6	ND ND	ND ND	<0.01 ND	N D ND
65.	phenol	7 2	3 6	ND ND	<0.01 0.03	0.018 0.01	<0.01 0.01
66.	bis(2-ethylhexyl) phthalate	7 2	3 6	0.02	0.01 <0.01	<0.01 <0.01	0.01 <0.01
67.	butyl benzyl phthalate	7 2	36	ND ND	ND ND	ND ND	ND ND
68	di-n-butyl phthalate	7 2	3 [.] 6	<0.01 <0.01	<0.01 <0.01	<0.01 <0.01	<0.01 <0.01
69.	di-n-octyl phthalate	7 2	3 6	ND ND	ND ND	ND ND	ND ND
70.	diethyl phthalate	7 2	3 6	<0.01 <0.01	<0.01 <0.01	<0.01 <0.01	<0.01 <0.01
71.	dimethyl phthalate	72	3 6	ND ND	ND ND	ND ND	ND ND
72.	benzo(a)anthracene	7 2	3 6	ND ND	ND ND	ND ND	N D ND

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SECONDARY PRECIOUS METALS SUBCATEGORY

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SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT C

<u>Pollutant</u>	Stream Code	Sample Typet	<u>Conc</u> Source	entration Day 1	ns (mg/1) Day 2	Day 3
<u>iciic Pollutants</u> (Continued)						
73. benzo(a)pyrene	7	3	ND	ND	ND	N D
	2	6	ND	ND	ND	ND
74. benzo(b)fluoranthene	7	3	ND	ND	ND	ND
	2	6	ND	ND	ND	ND
75. benzo(k)fluoranthane	7	3	ND	ND	ND	N D
	2	6	ND	ND	ND	ND
76. chrysene	7 2	3 6	ND ND	ND ND	ND ND	ND ND
77. acenaphthylene	7	3	ND	ND	ND	N D
	2	6	ND	ND	ND	ND
78. anthracene (a)	7	3	ND	ND	ND	ND
	2	6	ND	ND	ND	ND
79. benzo(ghi)perylene,	7	3	ND	ND	ND	N D
	2	6	ND	ND	ND	ND
80. fluorene	7	3	ND	ND	ND	ND
	2	6	ND	ND	ND	ND
81. phenanthrene (a)	7	3	ND	ND	ND	N D
	2	6	ND	ND	ND	ND

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT -4

SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT C

-,		Stream	Sample	Cónc	entration	s (mg/l)	
1.1 .	Pollutant	Code	Typet_	Source	Day 1	Day 2	Day 3
Toxic	Pollutants (Continued)					-	- - -
82.	dibenzo(a,h)anthracene	7 2	3 6	ND ND	ND ND	ND ND	N D ND
83.	indeno (1,2,3-c,d)pyrene	7 2	3	ND ND	ND ND	ND ND	ND ND
84.	pyrene	7 2	3 6	ND ND	ND ND	ND ND	N D ND
85.	tetrachloroethylene	72	1 1 -	ND ND	ND ND	ND ND	ND ND
86.	toluene	7 2	1	ND ND	ND <0.01	ND <0.01	ND <0.01
87.	trichloroethylene	7 2	1	ND ND	<0.01 ND	ND ND	<0.01 ND
88.	vinyl chloride (chloroethylene)	7 2	1 1	ND ND	ND ND	ND ND	<0.01 ND
114.	antimony	7 2	3 6	<0.003 <0.003	0.15 0.016	0.03 0.021	0.022 0.014
115.	arsenic	7 2	3 6	<0.005 <0.005	0.013 0.016	0.004 0.013	0.005

SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT C

		Stream	Concentrations (mg/l)					
	Pollutant	Code	<u>Type</u> t	Source	Day 1	Day 2	Day 3	
Texic	Pollutants (Continued)							
° † 7 .	beryllium	7 2	3 6	<0.0002 <0.0002	0.006	0.013 0.007	0.003	
27 3 .	cadmium	7 2	3 6	0.0002 0.0002	0.11	0.10	0.04 0.06	
•	chromium (total)	7 2	36	0.003	0.76 0.35	0.45 0.71	0.16 0.17	
: Zina	copper	72	3 6	0.017 0.017	2.10 1.80	0.57 2.90	0.73	
121.	yanide (total)	7 2	1	0.053	9.0 8.0	70.0 16.0	190.0 140.0	
122.	lead	7 2	3 6	0.03	0.63 0.19	0.51 0.41	0.75	
123.	mercury	7 2	3 6	0.0002	0.0002	<0.0001 <0.0001	<0.0001 <0.0001	Į
124.	nickel	7 2	3 6	0.02	3.70 0.39	7.20 4.20	0.720 2.80	[(}
125.	selenium	7 2	3 6	<0.002 <0.002	740.0 <0.002	<0.002 <0.002	<0.002 <0.002	•

SECONDARY PRECIOUS METALS SUBCATEGORY

SECT - V

SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT C

Pollutant	Stream <u>Code</u>	Sample Typet	Conc Source	<u>entratio</u> Day 1	ns (mg/1) Day 2	Day 3	SEC
ic Pollutants (Continued)						•	SECONDARY
. silver	7 2	3 6	<0.0002 <0.0002	0.05 0.049	0.06 0.05	0.03 0.10	
. thallium	7 2	3 6	<0.001 <0.001	<0.004 <0.003	<0.004 <0.004	0.012 0.30	PRECIOUS
28. zinc	7 2	3 6	<0.01 <0.01	180.0 91.0	170.0 160.0	140.0 150.0	
Nonconventional Pollutants		• 			. ¹		METALS
alkalinity	7 2	3 6	16 1 16	,900 23	0 300	870 18	
calcium	7 2	3 6	13 13	8.2 10.0	9.6 9.0	7.3 9.5	SUBCATEGORY
iron	7 2	3 6	0.29	2.3 3.4	48 14	4.5 3.6)RY
magnesium	7 2	3 6	3.1 3.1	3.3 3.5	3.4 3.7	2.6 2.9	SECT
phenolics	7 2	1 1	0.15	0.110 0.088	0.017 0.063	0.008 0.054	- - -

SECONDARY PRECIOUS METALS SAMPLING DATA TREATMENT PLANT SAMPLES - PLANT C

	Stream	Sample	<u>Concentrations (mg/1)</u>				
Pollutant	Code	<u>Typet</u>	Source	Day 1	Day 2	Day 3	
Conventional Polluants							
oil and grease	7	1	1.6	2.4	<1.7	<1.0	
	2	1	1.6	<1	<1	<1	
total suspended solids (TSS)	7	3	0	140	6	24	
	2	· 6	0	0	59	3	
pH (standard units)	7	3	6.8	1.9	2.0	8.9	
	2	6	6.8	10.0	10.5	6.7	

tSample Type Code: 1 - One-time grab
3 - 8-hour manual composite
6 - 24-hour automatic composite

(a) Reported together.

SECONDARY PRECIOUS METALS SUBCATEGORY SECT ł

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Table V-26

SECONDARY PRECIOUS METALS SAMPLING DATA CASTING CONTACT COOLING WATER

	Stream	Sample	Concentrations (mg/1)				
<u>Pollutant</u>	Code	Typet	Source	<u>Day 1</u>	Day 2	Day 3	
Toxic Pollutants				- -			
114. antimony	204	1	<0.01	<0.01			
115. arsenic	204	1	<0.01	<0.01			
117. beryllium	204	1	<0.01	<0.01			
118. cadmium	204	1	<0.05	<0.05			
119. chromium (total)	204	1	<0.05	<0.05	· ·		
120. copper	204	1	<0.05	0.05	· ·		
121. cyanide (total)	204	1	0.05	0.36			
2. lead	204	· 1	<0.10	<0.10	· · · · ·		
3. mercury	204	. 1	0.0002	0.0004	-		
124. nickel	204	1	<0.2	0.20	· · ·		
15. selenium	204	1	<0.1A	<0.1A	•		
	204	1	<0.01	<0.01		÷	
127. thallium	204	1	<0.01	<0.01			
128. zinc	204		0.10	0.15		-	

SECONDARY PRECIOUS METALS SUBCATEGORY SECT

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SECONDARY PRECIOUS METALS SAMPLING DATA CASTING CONTACT COOLING WATER

D 11	Stream	Sample Typet	<u>Concentrations (mg/l)</u>			
Pollutant	Code		Source	Day 1	Day 2	Day 3
Nonconventional Pollutants						
acidity	204	1	<1	<1		
alkalinity	204	1	98	120		
aluminum	204	1	0.2	0.2		
ammonia nitrogen	204	1	0.04	0.06	se ist	
ba ium	204	1	<0.05	<0.05	- -	
bozon	204	1	<0.1	1.8		
calcium	204	1	37.7	37.3		
chemical oxygen demand (COD)	204	1	<5	33		
chloride	204	1	14	250	•	
cobalt	204	1	<0.05	<0.05		
fluoride	204	1	0.28	0.31		
iron	204	1	<0.05	1.5		
magnesium	204	1	8.5	8.3		-

SECONDARY PRECIOUS METALS SUBCATEGORY SECT I

SECONDARY PRECIOUS METALS SAMPLING DATA CASTING CONTACT COOLING WATER

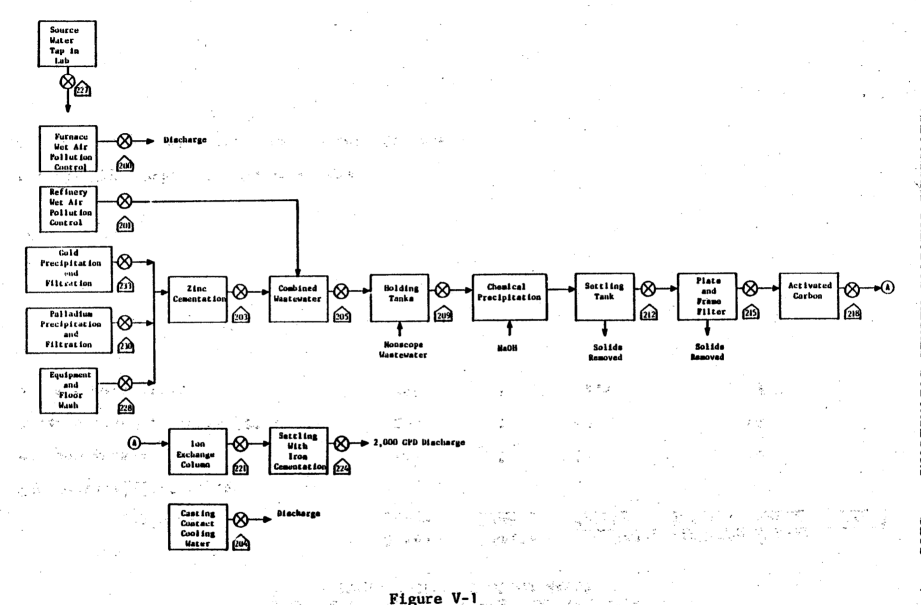
Dellutent	Stream Code	Sample Twpet	Concentrations (mg/1) Source Day 1 Day 2 Day 3				
<u>Pollutant</u>	······································	Typet	JUILE	Day 1	Day 2	<u>Day J</u>	
Nonconventional Pollutants (Continued)							
manganese	204	1			<0.05		
molybdenum	204	1	2 · · · ·		<0.05	•	
phosphate	204	1	· · · ·		36		
sodium	204	[1		• • • • • •	22		
sulfate	204	1		- - -	79		
tin	204	1			<0.05	• •	
titanium	204	. 1			<0.05		
total organic carbon (TOC)	204	1	• 		24	-	
total solids (TS)	204	1			410		
vanadium	204	1		• • •	<0.05		
yttrium	204	1			<0.05		

SECONDARY PRECIOUS METALS SAMPLING DATA CASTING CONTACT COOLING WATER

	Stream	Sample	Conc			
<u>Pollutant</u>	<u>Code</u>	<u>Typet</u>	Source	Day 1	Day 2	Day 3
Conventional Pollutants						
oil and grease	204	1	17		20	
total suspended solids (TSS)	204	1	60		44	
pH (standard units)	204	1	7.5		7.3	

tSample Type Code: 1 - One-time grab

A - Detection limit raised due to interference.



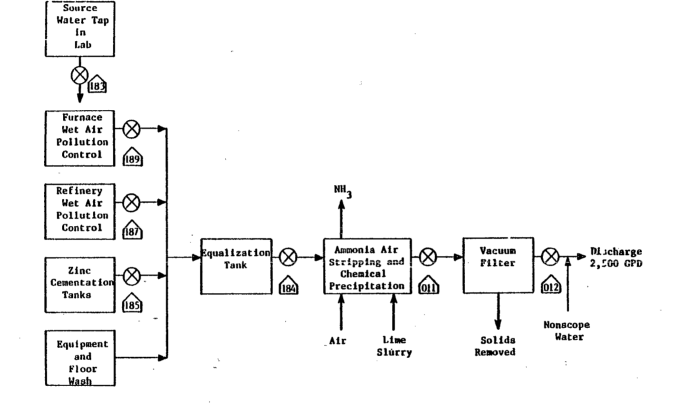
2501

CIOUS METALS PLANT A SAMPLING SITES SECONDARY AT 10.200

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT

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SAMPLING SITES AT SECONDARY PRECIOUS METALS PLANT B

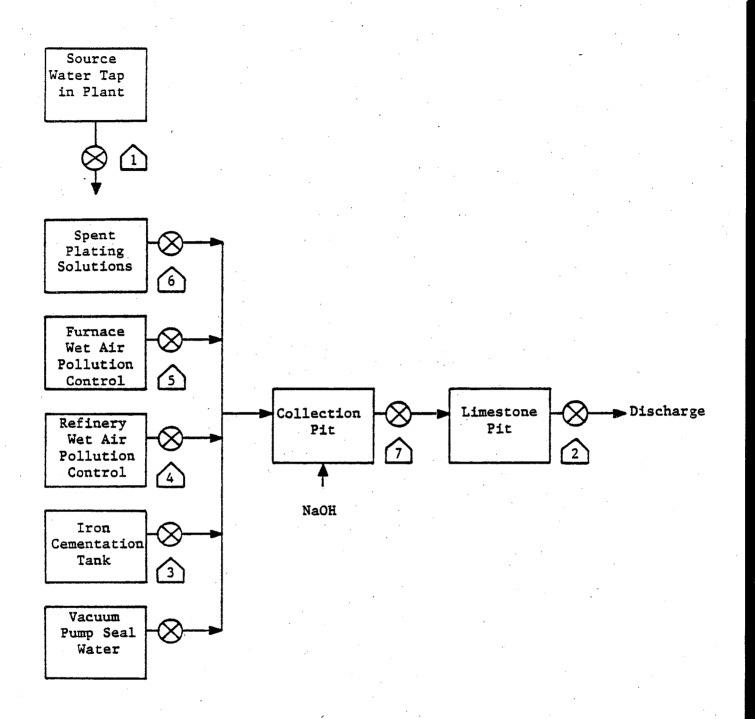
SECONDARY PRECIOUS METALS SUBCATEGORY

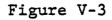
SECT

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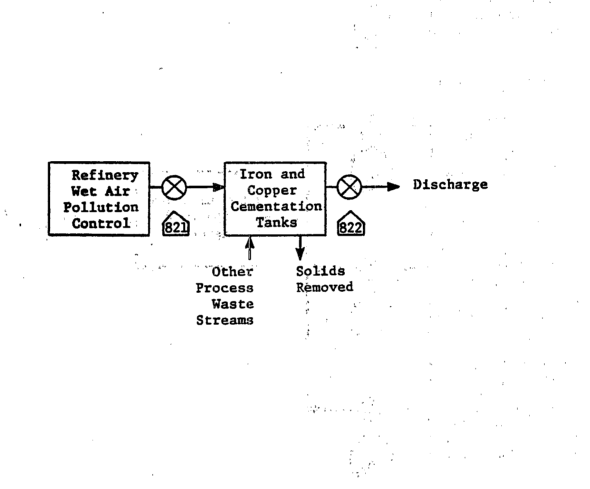
SECONDARY PRECIOUS METALS SUBCATEGORY

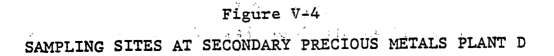
SECT - V

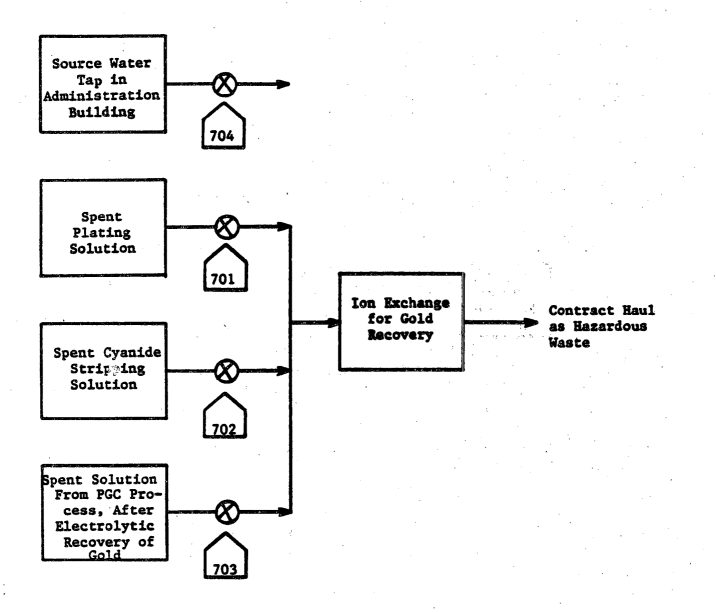


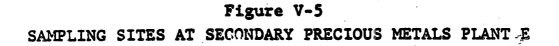


SAMPLING SITES AT SECONDARY PRECIOUS METALS PLANT C









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SECONDARY PRECIOUS METALS SUBCATEGORY SECT VI

SECTION VI

SELECTION OF POLLUTANT PARAMETERS

This section examines the chemical analysis data presented in Section V and discusses the selection or exclusion of pollutants for potential limitation. The basis for the regulation of toxic and other pollutants along with a discussion of each pollutant selected for potential limitation is discussed in Section VI of Vol. I. That discussion provides information concerning where the pollutant originates (i.e., whether it is a naturally occurring substance, processed metal, or a manufactured compound); general physical properties and the form of the pollutant toxic effects of the pollutant in humans and other animals; and behavior of the pollutant in POTW at the concentrations expected in industrial discharges.

The discussion that follows describes the analysis that was performed to select or exclude toxic pollutants for further consideration for limitations and standards. Pollutants will be considered for limitation if they are present in concentrations treatable by the technologies considered in this analysis. Also described is the analysis performed to select or exclude conventional and nonconventional pollutants for limitation. The treatment effectiveness concentrations used for the toxic metals long-term performance values achievable by chemical were the sedimentation, and filtration (lime, settle precipitation, on, sedimentation, and filtration (lime, settle and The achievable concentrations used for the toxic and filter). were the long-term values achievable organics by carbon adsorption (see Section VII of Vol. I -- Combined Metals Data Base).

CONVENTIONAL AND NONCONVENTIONAL POLLUTANT PARAMETERS

This study examined samples from the secondary precious metals subcategory for three conventional pollutant parameters (oil and grease, total suspended solids, and pH) and two nonconventional pollutant parameters (ammonia, and combined metals (gold, platinum, and palladium)).

CONVENTIONAL AND NONCONVENTIONAL POLLUTANT PARAMETERS SELECTED

The conventional and nonconventional pollutants and pollutant parameters selected for limitation in this subcategory are:

ammonia combined metals (the sum of gold, platinum, and palladium) total suspended solids (TSS) pH

Ammonia was found in 10 of 12 samples analyzed in concentrations ranging from 0.24 to 5,060 mg/l. Five of the values recorded are well above the treatable concentration of 32.2 mg/l, attainable

by the available treatment technology. In addition, ammonia is expected to be present based on its use in the raw materials. For these reasons, ammonia is selected for limitation in this subcategory.

Combined metals consists of the sum of gold, platinum, and palladium. This nonconventional pollutant parameter results from the information made available after promulgation of this rule. Petitioners were concerned with the variable nature of the production of gold, platinum, and palladium and the accuracy of analytical procedures for detecting these pollutants. Therefore, gold, platinum, and palladium were grouped into the one nonconventional pollutant parameter "combined metals".

Gold was analyzed for in 12 raw wastewater samples and was detected above its treatable concentration (0.01 mg/l) in all 12 samples. The concentrations ranged from 0.086 to 40 mg/l. Six of the 12 samples were found to contain more than 1 mg/l of gold. addition, gold is expected to be present in wastewaters from In subcategory due to its presence in the raw materials and this Platinum and palladium were not analyzed for in any products. raw wastewater samples from this subcategory. However, they are expected to be present in the raw wastewater in concentrations exceeding that achievable by treatment (0.01 mg/l). The reason for expecting treatable concentrations of platinum and palladium is they are both present in the raw materials and in the products produced by plants in this subcategory. For these reasons, combined metals are selected for limitations in this subcategory.

Oil and grease was analyzed for in 20 samples and was detected below quantifiable levels 11 times. In only two cases was oil and grease detected above its treatable concentration of 10 mg/l. The two treatable values are 14 mg/1 and 37 mg/1 and they are both for samples of refinery wet air pollution control. However, five other samples of this waste stream show oil and grease well below treatability. Because of the small number of in sources and grease was detected above its treatable which oil concentration, oil and grease is not selected for limitation in this subcategory.

Total suspended solids (TSS) concentrations ranging from 0 to 5,600 mg/l were observed in the 20 samples analyzed for this study. Nineteen of 20 samples exhibited concentrations above the concentration attainable by the identified treatment technology (2.6 mg/l). Furthermore, most of the specific methods for removing priority metals do so by precipitation, and the metal precipitates should not be discharged. Meeting a limitation on TSS also aids in removal of precipitated priority metals. For these reasons, total suspended solids are selected for limitation in this subcategory.

The pH values observed in 14 of 20 samples were outside the 7.5 to 10.0 range considered desirable for discharge to receiving waters. Six pH values ranged from 0.1 to 3.4. Six samples ranged from 5.9 to 9.3. The remaining eight samples ranged from

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - VI

10.9 to 12.6. Effective removal of priority metals by chemical precipitation requires careful control of pH. Therefore, pH is selected for limitation in this subcategory.

TOXIC PRIORITY POLLUTANTS

The frequency of occurrence of the toxic pollutants in the raw wastewater samples taken is presented in Table VI-1 (page 2514). These data provide the basis for the categorization of specific pollutants, as discussed below. Table Vi-1 is based on the raw wastewater data from streams 200, 189, 5, 201, 187, 4, 821, 233, 230, 228, 6, 701, 702, and 703 (see Section V). Treatment plant samples were not considered in the frequency count.

TOXIC POLLUTANTS NEVER DETECTED

The priority pollutants listed in Table VI-2 (page 2518) were not detected in any wastewater samples from this subcategory; therefore, they are not selected for consideration in establishing limitations:

TOXIC POLLUTANTS NEVER FOUND ABOVE THEIR ANALYTICAL QUANTIFICATION LIMIT

The priority pollutants listed below were never found above their analytical quantification concentration in any wastewater samples from this subcategory; therefore, they are not selected for consideration in establishing limitations.

4. benzene

- 7. chlorobenzene
- 10. 1,2-dichloroethane
- 21. 2,4,6-trichlorophenol
- 24. 2-chlorophenol
- 34. 2,4-dimethylphenol
- 44. methylene chloride (dichloromethane)
- 47. bromoform (tribromomethane)
- 48. dichlorobromomethane
- 51. chlorodibromomethane
- 54. isophorone
- 62. N-nitrosodiphenylamine
- 68. di-n-butyl phthalate
- 69. di-n-octyl phthalate
- 70. diethyl phthalate
- 71. dimethyl phthalate
- 86. toluene

PRIORITY POLLUTANTS PRESENT BELOW CONCENTRATIONS ACHIEVABLE BY TREATMENT

The pollutants listed below are not selected for consideration in establishing limitations because they were not found in any wastewater samples from this subcategory above concentrations considered achievable by existing or available treatment technologies. These pollutants are discussed individually SECONDARY PRECIOUS METALS SUBCATEGORY SECT - VI

following the list.

57. 2-nitrophenol 123. mercury

2-Nitrophenol was found in only one sample at its quantification limit. The reported concentration was 0.01 mg/l, which is also the treatable concentration. Since the pollutant was not detected above the concentration attainable by identified treatment technology, 2-nitrophenol is not considered for limitation.

Mercury was detected below its quantification limit in 20 out of 24 samples analyzed. The four values reported above the quantification limit ranged from 0.0003 mg/l to 0.015 mg/l, which are all below the concentration attainable by identified treatment technology, which is 0.036 mg/l. Therefore, mercury is not considered for limitation.

PRIORITY POLLUTANTS DETECTED IN A SMALL NUMBER OF SOURCES

The following pollutants were not selected for limitation on the basis that they are detectable in the effluent from only a small number of sources within the subcategory and they are uniquely related to only those sources.

- 6. carbon tetrachloride
- 11. 1,1,1-trichloroethane
- 23. chloroform
- 65. phenol
- 66. bis(2-ethylhexyl) phthalate
- 117. beryllium

Although these pollutants were not selected for consideration in establishing nationwide limitations, it may be appropriate, on a case-by-case basis, for the local permittee to specify effluent limitations.

Carbon tetrachloride was detected in only one of 12 samples analyzed, at a concentration of 0.21 mg/1. The treatability concentration is 0.01 mg/1 for this pollutant. Since it was not detected in 11 other samples, the measurement may be regarded as specific to the site and not characteristic of the subcategory as a whole. Also, carbon tetrachloride cannot be attributed to specific materials and processes used in the secondary precious metals subcategory. Therefore, carbon tetrachloride is not considered for limitation.

1,1,1-Trichloroethane was detected in only one of 12 samples analyzed, at a concentration of 0.015 mg/1. The treatability concentration is 0.01 mg/1 for this pollutant. Since it was not detected in 11 other samples, the measurement may be regarded as specific to the site and not characteristic of the subcategory as a whole. Also, 1,1,1-trichloroethane cannot be attributed to specific materials and processes used in the secondary precious metals subcategory. Therefore, 1,1.1-trichloroethane is not

considered for limitation.

Chloroform was detected in four of 12 samples above its treatable concentration of 0.01 mg/1. The four concentrations are all 0.02 mg/1. All four samples have a lower concentration of chloroform than the source water at the plant (0.05 mg/1). Chloroform cannot be attributed to specific materials or processes used in the subcategory, and very little removal of this pollutant can be expected with treatment. Therefore, chloroform is not considered for limitation.

Phenol was detected in only four of 12 samples above its treatable concentration of 0.01 mg/1. The four concentrations are 0.013 mg/1, 0.17 mg/1, 0.45 mg/1, and 0.65 mg/1. The three samples with concentrations above 0.10 mg/1 were all taken at one plant which was shut down indefinitely subsequent to being Since phenol was not detected above its treatable sampled. concentration in eight other samples, the measurements may be regarded as specific to the site and not characteristic of the subcategory as a whole. Phenol cannot be attributed to specific materials and processes used in the secondary precious metals subcategory. Also, because of the relatively low concentrations of phenol in the raw waste compared with its treatable concentration, very little removal of phenol can be expected with treatment. Therefore, phenol is not considered for limitation.

Bis(2-ethylhexyl) phthalate was found above its treatable concentration of 0.01 mg/l in six of 12 samples. The concentrations ranged from 0.02 mg/l to 0.1 mg/l. This pollutant is not associated with specific processes used in the secondary precious metals subcategory, but is commonly used as а plasticizer in laboratory and field sampling equipment. Since the presence of this pollutant may be attributed to sample contamination, bis(2-ethylhexyl) phthalate is not considered for limitation.

Beryllium was found in only one out of 24 samples analyzed above its treatable concentration of 0.20 mg/l. The sample had a concentration of 0.46 mg/l. Since it was not found above its treat. able concentration in 23 other samples, the measurement may be regarded as site-specific and not characteristic of the subcategory as a whole. Although beryllium may be part of a raw material, such as jewelry scrap, used in the secondary precious metals industry, all the wastewater samples analyzed from plants which process these raw materials showed beryllium present below treatable concentrations. Therefore, beryllium is not considered for limitation.

PRIORITY POLLUTANTS SELECTED FOR FURTHER CONSIDERATION IN ESTABLISHING LIMITATIONS AND STANDARDS

The priority pollutants listed below are selected for further consideration in establishing limitations and standards for this subcategory. The priority pollutants selected for further consideration for limitation are each discussed following the SECONDARY PRECIOUS METALS SUBCATEGORY

list.

114.	antimony
115.	arsenic
118.	cadmium
119.	chromium
120.	copper
121.	cyanide
122.	lead
124.	nickel
125.	selenium
126.	silver
127.	thallium
128.	zinc

Antimony was detected above its treatable concentration (0.47 mg/l) in seven of 24 samples. The quantifiable concentrations ranged from 0.19.mg/l to 5.2 mg/l. Since antimony was present in concentrations exceeding the concentrations achievable by identified treatment technology, it is selected for consideration for limitation.

SECT

Arsenic was detected above its treatable concentration (0.34 mg/l) in four of 24 samples. The quantifiable concentrations ranged from 0.025 mg/l to 2.4 mg/l. Since arsenic was present in concentrations exceeding the concentrations achievable by identified treatment technology, it is selected for consideration for limitation.

Cadmium was detected above its treatable concentration (0.049 mg/l) in 12 of 24 samples. The quantifiable concentrations ranged from 0.0029 mg/l to 7.6 mg/l. Since cadmium was present in concentrations exceeding the concentrations achievable by identified treatment technology, it is selected for consideration for limitation.

Chromium was detected above its treatable concentration (0.07 mg/l) in 15 of 24 samples. The quantifiable concentrations ranged from 0.012 mg/l to 22 mg/l. Since chromium was present in concentrations exceeding the concentrations achievable by identified treatment technology, it is selected for consideration for limitation.

Copper was detected above its treatable concentration (0.39 mg/l) in 15 of 23 samples. The quantifiable concentrations ranged from 0.016 mg/l to 5,000 mg/l. Since copper was present in concentrations exceeding the concentrations achievable by identified treatment technology, it is selected for consideration for limitation.

Cyanide (total) was detected above its treatable concentration (0.047 mg/l) in 17 of 24 samples. The quantifiable concentrations ranged from 0.09 mg/l to 9,897 mg/l. Since cyanide is used as a raw material, and was present in concentrations exceeding the concentrations achievable by

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - VI

identified treatment technology, it is selected for consideration for limitation.

Lead was detected above its treatable concentration (0.08 mg/l) in 17 of 24 samples. The quantifiable concentrations ranged from 0.02 mg/l to 0.7 mg/l. Since lead was present in concentrations exceeding the concentrations achievable by identified treatment technology. it is selected for consideration for limitation.

Nickel was detected above its treatable concentration (0.22 mg/l) in 17 of 24 samples. The quantifiable concentrations ranged from 0.008 mg/l to 890 mg/l. Since nickel was present in concentrations exceeding the concentrations achievable by identified treatment technology, it is selected for consideration for limitation.

Selenium was detected above its treatable concentration (0.20 mg/l) in three of 24 samples. The quantifiable concentrations ranged from 0.019 mg/l to 120 mg/l. Since selenium was present in concentrations exceeding the concentrations achievable by identified treatment technology, it is selected for consideration for limitation.

Silver was detected above its treatable concentration (0.07 mg/l) in 14 of 24 samples. The quantifiable concentrations ranged from 0.05 mg/l to 26 mg/l. Since silver was present in concentrations exceeding the concentrations achievable by identified treatment technology, it is selected for consideration for limitation.

Thallium was detected above its treatable concentration (0.34 mg/l) in four of 22 samples. The quantifiable concentrations ranged from 0.82 mg/l to 1.2 mg/l. Since thallium was present in concentrations exceeding the concentrations achievable by identified treatment technology, it is selected for consideration for limitation.

Zinc was detected above its treatable concentration (0.23 mg/l) in 18 of 23 samples. The quantifiable concentrations ranged from 0.11 mg/l to 10,000 mg/l. Since zinc is used in the cementation process, and was present in concentrations exceeding the concentrations achievable by identified treatment technology, it is selected for consideration for limitation.

Table VI-1

FREQUENCY OF OCCURRENCE OF PRIORITY POLLUTANTS SECONDARY PRECIOUS METALS SUBCATEGORY RAW WASTEWATER

<u>Pollutant</u>	Analytical Quantification Concentration (mg/l)(a)	Treatable Concentra- tion (¤g/1)(b)	Number of Streams <u>Analyzed</u>	Number of Samples Analyzed	ND	Detected Below Quantification Concentration	Detected Below Treat- able Concen- tration	Detected Above Treat- able Concen- tration	SECONDARY
1. acenaphthene	0.010	0.01	6	12	12				R
2. acrolein	0.010	0.01	6	12	12				7
3. acrylonitrile	0.010	0.01	6	12	12	•			Ы
4. benzene	0.010	0.01	6	12	10	2			PRECIOUS
5. benzidine	0.010	0.01	6	12	12			•	Ö
6. carbon tetrachloride	0.010	0.01	6.	12				I	Ĥ
7. chlorobenzene	0.010	0.01	6	12	11	I			g
8. 1,2,4-trichlorobenzene	0.010	0.01	6	12	12				J.C.
9. hexachlorobenzene	0.010	0.01	6	12	12				V1 .
10. 1,2-dichloroethane	0.010	0.01	6	12	11	I		1	M
11. 1,1,1-trichloroethane	0.010	0.01	0	12 12	11 12			·	METAL
12. hexachloroethane	0.010	0.01	0	12	12	-			Ā
13. 1,1-dichloroethane	0.010	0.01	0	12	12				H
4. 1,1,2-trichloroethane	0.010	0.01	0	12	12				ũ
15. 1,1,2,2-tetrachloroethane	0.010	0.01	0	12	12				ທ່
6. chloroethane	0.010	0.01	0.	12	12				q
7, bis(chloramethyl) ether	0.010	0.01 <i>~</i> 0.01	0 4	12	12				B
8. bis(2-chloroethyl) ether	0.010		U 4	12	12				D D
19. 2-chloroethyl vinyl ether	0.010	0.01	0 4	12	12				ĥ
20. 2-chloronaphthalene	0.010 0.010	0.01 0.01	0	12	11	1			E
21. 2,4,6-trichlorophenol 22. parachlorometa cresol	0.010	0.01	6	12	12	•			မ္မ
23. chloroform	0.010	0.01	6	12	6	1	1	4	SUBCATEGORY
24. 2-chlorophenol	0.010	0.01	6	12	пĭ	i	•	•	R
25. 1,2-dichlorobenzene	0.010	0.01	6	12	12				
26. 1.3-dichlorobenzene	0.010	0.01	6	12	12				
17. 1.4-dichlorobenzene	0.010	0.01	6	12	12	,			
8. 3.3'-dichlorobenzidine	0.010	0.01	6	12	12				С Н
29. 1,1-dichloroethylene	0.010	0.01	6	12	12				Ö
30. 1,2-trans-dichloroethylene	0.010	0.01	6	12	12		•		CT
31. 2.4-cichlorophenol	0.010	0.01	6	12	12				
32. 1,2-dichloropropane	0.010	0.01	6	12	12				•
33. 1, 3-dichloropropylene	0.010	0.01	6	12	12				۲V
34. 2 4-dimethylphenol	0.010	0.01	6	12	11	1			н
35. 2 dinitrotoluene	0.010	0.01	6	12	12				

FREQUENCY OF OCCURRENCE OF PRIORITY POLLUTANTS SECONDARY PRECIOUS METALS SUBCATEGORY RAW WASTEWATER

<u>Pollutant</u>	Analytical Quantification Concentration (mg/l)(a)	Treatable Concentra- tion (mg/1)(b)	Number of Streams Analyzed	Number of Samples Analyzed	ND	Detected Below Quantification Concentration	Detected Below Treat- able Concen- tration	Detected Above Treat- able Concen- tration	SECONDARY
36. 2,6-dinitrotoluene	0.010	0.01	6	12	12		•		AR
37. 1,2-diphenylhydrazine	0.010	0.01	0	12	12				К
38. ethylbenzene	0.010	0.01	0	12	12				н
39. fluoranthene	0.010	0.01	6	12	.12 12			۳.	PRECIOUS
40. 4-chlorophenyl phenyl ether	0.010	0.01		12 12	12		• •		Ĕ
41. 4-bromophenyl phenyl ether	0.010	0.01	0			•			H
42. bis(2-chloroisopropyl) ether	0.010	0.01	0 (12	12				ö
43. bis(2-chloroethoxy) methane	0.010	0.01	°,	12	12	9	•		g
	0.010 0.010	0.01	D K	12 12	12	9	ν.		0
	0.010	0.01	6	12	12				Z
46. methyl bromide 47. bromoform	0.010	0.01	6	12	11	1			METALS
3. dichlorobromomethane	0.010	0.01	- 0	12	10				Ц,
49. trichlorofluoromethane	0.010	0.01	6	12	12	2			H
50. dichlorodifluoromethane	0.010	0.01	6	12	12				លី
1. chlorodibromomethane	0.010	0.01	6	12	ii	1	•		70
2. hexachlorobutadiene	0.010	0.01	6	12	12				č
3. hexachlorocyclopentadiene	0.010	0.01	6	12	12				SUBCATEGORY
54. isophorone	0.010	0.01	6	12	ii	1			ß
5. naphthalche	0.010	0.01	6	12	12	•			A L
. nitrobenzene	0.010	0.01	6	12	12		:		Ē
7. 2-nitrophenol	0.010	0.01	6	12	7	4	1		ភ្ន
8. 4-nitrophenol	0.010	0.01	6	12	12	•	• •		С Н
.9. 2.4-dinitrophenol	0.010	0.01	ň	12	12				Ř
50. 4,6-dinitro-o-cresol	0.010	0.01	6	12	12			1. Sec. 1. Sec	
61. N-nitrosodimethylamine	0.010	0.01	ő	12	12	-		÷	
62. N-nitrosodiphenylamine	0.010	0.01	6	12	11	1			
63. N-nitrosodi-n-propylamine	0.010	0.01	6	12	12	*			S
64. pentachlorophenol	0.010	0.01	6	12	12				E
65. phenol	0.010	0.01	6	12	4	4	н. 1	4	CT
66. bis(2-ethylhexyl) phthalate	0.010	0.01	6	12	Ó	··· 6 · · · ·		6	L,
67. butyl benzyl phthalate	0.010	0.01	6 ·	12	12		•	-	I
68. di-n-butyl phthalate	0.010	0.01	6	12	. 2	10	•	- '	~
69. di-n-octyl phthalate	0.010	0.01	6	12	11	1 .			Ĥ
70. diethyl phthalate	0.010	0.01	6	12	2	10			••
				· · · ·					

FREQUENCY OF OCCURRENCE OF PRIORITY POLLUTANTS SECONDARY PRECIOUS METALS SUBCATEGORY RAW WASTEWATER

<u>Pollutant</u>	Analytical Quantification Concentration (mg/l)(a)	Treatable Concentra- tion (mg/1)(b)	Number of Streams Analyzed	Number of Samples <u>Analyzed</u>	ND	Detected Below Quantification Concentration	Detected Below Treat- able Concen- tration	Detected Above Treat- able Concen- tration	
71. dimethyl phthalate	0.010	0.01	6	12	11	1			- 7 F
72. benzo(a)anthracene	0.010	0.01	6	12	12				L
73. benzo(a)pyrene	0.010	0.01	6	12	12				् म्
74. 3,4-benzofluoranthene	0.010	0.01	6	12	12				- 6
75. benzo(k)fluoranthene	0.010	0.01	6	12	12				Ţ
76. chrysene	0.010	0.01	6	12	12				F
77. acenaphthylene	0.010	0.01	6	12	12				2
78. anthracene (c)	0.010	0.01	6	12	12				Ū
79. benzo(gh1)perylene	0.010	0.01	6	12	12				
80. fluorene	0.010	0.01	6	12	12				Ē
81. phenanthrene (c)	0.010	0.01	6	12	12				- F
82. dibenzo(a,h)anthracene	0.010	0.01	6	12	12		•		Þ
83. Indeno(1,2,3-cd)pyrene	0.010	0.01	6	12	12				Ę
84. pyrene	0.010	0.01	6	12	12				
85. tetrachloroethylene	0.010	0.01	6	12	12				Ū
86. toluene	0.010	0.01	6	12	5	7			S
87. trichloroethylene	0.010	0.01	6	12	12				2
88. vinyl chloride	0.010	0.01	6	12	12				ģ
114. antimony	0.100	0.47	14	24	0	15	2	7	Ē
115. arsenic	0.010	0.34	14	24	0	13	. 7	4	2
116. asbestos	10 MFL	10 MFL	0			•			Ľ
117. beryllium	0.010	0.20	14	24	0	17	6	1	Þ
118. cadmium	0.002	0.049	14	24	0	8	4	12	H
119. chromium	0.005	0.07	14	24	0	7	2	15	
120. copper	0.009	0.39	13	23	0	0	8	15	
121. cyanide (d)	0.02	0.047	. 14	24	0	7	Ũ	17	_
122. lead	0.020	0.08	14	24	0	4	3	17	U E
123. mercury	0.0001	0.036	14	24	0	20	4	0	Č
124. nickel	0.005	0.22	14	24	0	1	6	17	È
125. selenium	0.01	0.20	13	21	0	15	3	3	,
126. silver	0.02	0.07	14	24	0	9	1	14	i
127. thallium	0.100	0.34	13	22	0	18	0	4	<
128. zine	0.050	0.23	13	23	0	1	4	18	H
129, 2.3.7.8-tetrachlorodibenzo-			0						

129. 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDM)

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT

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FREQUENCY OF OCCURRENCE OF PRIORITY POLLUTANTS SECONDARY PRECIOUS METALS SUBCATEGORY RAW WASTEWATER

- (a) Analytical quantification concentration was reported with the data (see Section V).
- (b) Treatable concentrations are based on performance of lime precipitation, sedimentation, and filtration.
- (c) Reported together.
- (d) Analytical quantification concentration for EPA Method 335.2, Total Cyanide Methods for Chemical Analysis of Water and Wastes, EPA 600/4-79-020, March 1979.

SECONDARY PRECIOUS METALS SUBCATEGORY

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- VI

TABLE VI-2

TOXIC POLLUTANTS NEVER DETECTED

1.	acenaphthene
2.	acrolein
3.	acrylonitrile
5.	benzidene
8.	1,2,4-trichlorobenzene
9.	hexachlorobenzene
12.	hexachloroethane
13.	l,l-dichloroethane
14.	1,1,2-trichloroethane
15.	1,1,2,2-tetrachloroethane
16.	chloroethane
17.	bis(2-chloromethyl) ether (Deleted)
18.	bis (2-chloroethyl) ether
19.	2-chloroethyl vinyl ether
20.	2-chloronaphthalene
22.	parachlorometa cresol
25.	1,2-dichlorobenzene
26.	1,3-dichlorobenzene
27.	1,4-dichlorobenzene
28.	3,3'-dichlorobenzidine
29.	1,1-dichloroethylene
31.	2,4-dichlorophenol
32.	1,2-dichloropropane
33.	1,2-dichloropropylene (1,3-dichloropropene)
35.	2,4-dinitrotoluene
36.	2,6-dinitrotoluene
37.	1,2-diphenylhydrazine
38.	ethylbenzene
39.	fluoranthene
40.	4-chlorophenyl phenyl ether
41.	4-bromophenyl phenyl ether
42.	bis(2-chloroisopropyl) ether
43.	bis(2.choroethoxy) methane
45.	methyl chloride (chloromethane)
46.	methyl bromide (bromomethane)
49.	trichlorofluoromethane (Deleted)
50.	dichlorodifluoromethane (Deleted)
52.	hexachlorobutadiene
53.	hexachlorocyclopentadiene
55.	naphthalene
56.	nitrobenzene
58.	4-nitrophenol
59.	2,4-dinitrophenol
60.	4,6-dinitro-o-cresol
61.	N-nitrosodimethylamine
63.	N-nitrosodi-n-propylamine
64.	pentachlorophenol

67. butyl benzyl phthalate

TABLE VI-2 (Continued)

TOXIC POLLUTANTS NEVER DETECTED

72. 73. 74. 75. 76. 77. 78. 79. 80. 81. 82. 83.	<pre>benzo (a)anthracene (1,2-benzanthracene) benzo (a)pyrene (3,4-benzopyrene) 3,4-benzofluoranthene benzo(k)fluoranthene (11,12-benzofluoranthene) chrysene acenaphthylene anthracene benzo(ghi)perylene (1,11-benzoperylene) fluorene phenanthrene dibenzo (a,h)anthracene (1,2,5,6-dibenzanthracene) indeno (1,2,3-cd)pyrene (w,e,-o-phenylenepyrene)</pre>
84.	pyrene tetrachloroethylene
85. 87.	trichloroethylene
88.	vinyl chloride (chloroethylene)
89.	aldrin*
90.	dieldrin*
91.	chlordane*
92.	4,4'-DDT*
93.	4,4'-DDE(p,p'DDX)*
94.	4,4'-DDD(p,p TDE)*
95.	a-endosulfan-Alpha*
96.	b-endosulfan-Beta*
97.	endosulfan sulfate*
98.	endrin* endrin aldehyde*
99. 100.	heptachlor*
101.	heptachlor epoxide*
102.	a-BHC-Alpha*
103.	b-BHC-Beta*
104.	r-BHC (lindane)-Gamma*
105.	g-BHC-Delta*
106.	PCB-1242 (Arochlor 1242)*
107.	PCB-1254 (Arochlor 1254)*
108.	PCB-1221 (Arochlor 1221)*
109.	PCB-1232 (Arochlor 1232)* PCB-1248 (Arochlor 1248)*
110.	PCB-1260 (Arochlor 1240) PCB-1260 (Arochlor 1260)*
111.	PCB-1016 (Arochlor 1016)*
113.	toxaphene*
115.	asbestos
129.	2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD)

+We did not analyze for these pollutants in samples of raw wastewater from this subcategory. These pollutants are not believed to be present based on the Agency's best engineering judgment which includes consideration of raw materials and process operations. THIS PAGE INTENTIONALLY LEFT BLANK

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SECTION VII

CONTROL AND TREATMENT TECHNOLOGIES

The preceding sections of this supplement discussed the sources, flows, and characteristics of the wastewaters from secondary precious metals plants. This section summarizes the description of these wastewaters and indicates the level of treatment which is currently practiced by plants in the secondary precious metals subcategory for each waste stream.

CURRENT CONTROL AND TREATMENT PRACTICES

This section presents a summary of the control and treatment technologies that are currently being applied to each of the sources generating wastewater in this subcategory. As discussed in Section V, wastewater associated with the secondary precious metals subcategory is characterized by the presence of the toxic metal pollutants. free and complexed cyanide, ammonia, combined metals (gold, platinum, and palladium), and suspended solids. The raw (untreated) wastewater data for specific sources as well as combined waste streams are presented in Section V. Generally, these pollutants are present in each of the waste streams at concentrations above treatability, and these waste streams are commonly combined for treatment. Construction of one wastewater treatment system for combined treatment allows plants to take advantage of economies of scale and, in some instances, to combine streams of differing alkalinity to reduce treatment chemical requirements. Twenty-four plants in this subcategory currently have combined wastewater treatment systems, 20 have chemical precipitation and sedimentation, and one of these has chemical precipitation, sedimentation and pressure filtration. One plant currently strips ammonia with air, and eight plants currently treat for cyanide. Seven of the eight use alkaline oxidation, and one plant precipitates cyanide with ferrous sulfate. Three plants currently practice ion exchange for removal of gold, platinum, and palladium. Three options have selected for consideration for BPT, BAT, NSPS, and been pretreatment in this subcategory, based on combined treatment of these compatible waste streams.

FURNACE WET AIR POLLUTION CONTROL

Air emission sources in secondary precious metals furnace operations include incinerator and smelting furnaces. Eighteen secondary precious metals producers control air emissions using various methods. These are:

Dry baghouse - 11 plants, and
 Wet scrubber - seven plants.

Priority organics, metals, cyanide, combined metals (gold, platinum, and palladium), and suspended solids are present at

treatable concentrations in the wastewater produced by furnace wet air pollution control. Two plants producing this wastewater practice complete recycle. Two practice partial recycle (>90 percent). Three practice no recycle. Treatment methods used are:

- 1. No treatment two plants, and
- 2. Chemical precipitation and sedimentation three plants.

One plant producing this wastewater practices ion exchange endof-pipe treatment.

RAW MATERIAL GRANULATION

Three of four plants reporting this waste stream discharge it. The three discharging plants do not practice recycle or treatment of this waste stream. The non-discharging plant practices total recycle.

SPENT PLATING SOLUTIONS

Spent or contaminated cyanide solutions from electroplating shops may have the precious metal values recovered by a precipitation or electrolytic process. The waste stream is characterized by treatable concentrations of priority organics and metals, free and complexed cyanide, combined metals (gold, platinum, and palladium), and TSS. Treatment methods for this wastewater consist of:

- Total cyanide precipitation using ferrous sulfate one plant,
- 2. Free cyanide destruction using alkaline oxidation six plants,
- 3. Chemical precipitation and sedimentation one plant, and
- 4. Contractor disposal four plants.

Four plants that have cyanide pretreatment also have chemical precipitation and sedimentation end-of-pipe treatment. The plant which uses ferrous sulfate also uses alkaline oxidation for cyanide treatment.

One plant producing this wastewater practices ion exchange endof-pipe treatment.

SPENT CYANIDE STRIPPING SOLUTIONS

Six plants use potassium or sodium cyanide solution to strip gold away from scrap. Four plants employ contractor disposal methods to achieve zero discharge of spent stripping solution. This wastewater contains priority metals, free and complexed cyanide, combined metals (gold, platinum, and palladium), and TSS above treatable concentrations. One of the two discharging plants destroys the free cyanide with chlorine gas (alkaline oxidation). The other plant destroys the free and complexed SECONDARY PRECIOUS METALS SUBCATEGORY

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cyanide with ferrous sulfate, and then practices chemical precipitation and sedimentation.

One plant producing this wastewater practices ion exchange endof-pipe treatment.

REFINERY WET AIR POLLUTION CONTROL

Scrubbers are used at 29 plants to control fumes from precipitation and filtration processes. Facilities can use acid scrubbers, alkali scrubbers, or both types of scrubbers. This wastewater contains treatable concentrations of priority metals, combined metals (gold, platinum, and palladium) and suspended solids. Twenty-one plants discharge this waste-water, six of which practice no recycle, and 15 of which practice recycle of 75 percent or more. Eight plants do not discharge this wastewater. Three of these plants practice 100 percent recycle, and five of them have this wastewater disposed of by a contractor.

At the 21 discharging plants, scrubber water is commonly combined with other process wastewater and treated in a central treatment facility. Treatment methods used are:

- 1. Chemical precipitation and sedimentation seven plants;
- 2. Chemical precipitation, sedimentation, and filtration one plant; and
- 3. No treatment 13 plants.

Two plants producing this wastewater practice ion exchange endof-pipe treatment.

GOLD SOLVENT EXTRACTION RAFFINATE AND WASH WATER

One plant recovers gold by a solvent extraction process, and generates a raffinate waste stream and a wash water waste stream. Priority metals and TSS are expected to be found at treatable levels in the raffinate and wash water. This waste stream is not recycled. Treatment before discharge consists of neutralization with caustic, but no solids are removed.

GOLD SPENT ELECTROLYTE

Wastewater discharges from electrolytic refining consist of spent electrolyte solution. Of the three plants practicing electrolytic refining, one discharges wastewater. This wastewater is expected to contain treatable concentrations of priority metals, ammonia, and TSS. This waste stream is for recycled. The one discharging plant practices chemical precipitation and sedimenta-tion of the spent electrolyte prior to discharge. The other two plants are zero discharge by means of contractor disposal.

GOLD PRECIPITATION AND FILTRATION

Nineteen of 28 plants which produce gold by dissolving goldcontaining raw material in acid and then selectively precipitating it from solution discharge this waste stream. This wastewater contains priority metals, combined metals (gold, platinum, and palladium), ammonia and TSS above treatable concentrations. No plants reported recycling this waste stream. Treatment methods for this wastewater consist of:

- 1. Chemical precipitation and sedimentation 10 plants;
- Chemical precipitation, sedimentation, and filtration one plant;
- 3. Contractor disposal seven plants;
- 4. One hundred percent reuse two plants; and
- 5. No treatment eight plants.

Two plants producing this wastewater practice ion exchange endof-pipe treatment.

PLATINUM PRECIPITATION AND FILTRATION

Fourteen of 18 plants which produce platinum by a dissolution and selective precipitation process discharge this waste stream. This wastewater is expected to contain priority metals, combined metals (gold, platinum, and palladium), ammonia, and TSS above treatable concentrations. No plants reported recycling this waste stream. Treatment methods for this wastewater consist of:

- 1. Chemical precipitation and sedimentation 10 plants (one with ammonia air stripping);
- Chemical precipitation, sedimentation, and filtration one plant;
- 3. No treatment three plants; and
- 4. Contractor disposal four plants.

Two plants producing this wastewater practice ion exchange endof-pipe treatment.

PALLADIUM PRECIPITATION AND FILTRATION

Fifteen of 20 plants which produce palladium by a dissolution and selective precipitation process discharge this waste stream. This wastewater should contain priority metals, combined metals (gold, platinum, and palladium), ammonia, and TSS above treatable concentrations. No plants reported recycling this waste stream. Treatment methods for this wastewater consist of:

- 1. Chemical precipitation and sedimentation nine plants (one with ammonia air stripping);
- 2. Chemical precipitation, sedimentation, and filtration one plant;
- 3. No treatment five plants;
- 4. One hundred percent reuse one plant; and
- 5. Contractor disposal four plants.

Two plants producing this wastewater practice ion exchange endof-pipe treatment.

OTHER PLATINUM GROUP METALS PRECIPITATION AND FILTRATION

Two of three plants using a wet chemistry technique to produce platinum group metals such as rhodium and iridium discharge this waste stream. This waste stream is expected to contain priority metals, ammonia and TSS. Treatment methods for this wastewater consist of:

- 1. Chemical precipitation and sedimentation one plant,
- 2. No treatment one plant, and
- 3. Contractor disposal one plant.

SPENT SOLUTION FROM PGC SALT PRODUCTION

Three of the four plants which produce PGC salt from pure gold and potassium cyanide discharge excess cyanide solution. Two of the three dischargers chlorinate the wastewater to destroy free cyanide, one has no treatment in-place, and one practices chemical precipitation and sedimentation. The non-discharging plant achieves this status by contractor disposal. The untreated wastewater contains priority metals, free and complexed cyanide, combined metals (gold, platinum, and palladium) and TSS above treatable concentrations.

One plant producing this wastewater practices ion exchange endof-pipe treatment.

EQUIPMENT AND FLOOR WASH

Three plants reported an equipment and floor wash waste stream and two of these plants discharge it. This wastewater contains priority metals, ammonia, and TSS above treatable concentrations.

No plants reported recycling this waste stream. Both discharging plants practice chemical precipitation and sedimentation. One of the two plants air strips ammonia. The nondischarging plant uses contractor disposal to achieve this status.

PRELIMINARY TREATMENT

As discussed in Section V, EPA agreed to add a new building block for the preliminary treatment process. This building block was not included in the promulgated rule because the Agency believed that the furnace wet air pollution control (FWAP) building block accounted for flows generated by the preparatory processing of basis materials required before these materials can be introduced into the main hydrometallurgical refining system. Wastewaters from this building block should include priority organics, metals, cyanide, combined metals (gold, platinum, and palladium), and suspended solids, all at treatable concentrations.

CONTROL AND TREATMENT OPTIONS CONSIDERED

Based on an examination of the wastewater sampling data, three control and treatment technologies that effectively control the pollutants found in secondary precious metals wastewaters were selected for evaluation. The effectiveness of these technologies is detailed in Section VII of Vol. I, and the technology options are discussed below.

OPTION A

Option A for the secondary precious metals subcategory requires treatment technologies to reduce pollutant mass. The Option A treatment scheme consists of ammonia steam stripping preliminary treatment applied to the combined stream of gold precipitation and filtration, platinum precipitation and filtration, palladium precipitation and filtration, other platinum group metal precipitation and filtration, and equipment and floor wash water; and cyanide precipitation preliminary treatment applied to the combined stream of spent plating solution, spent cyanide stripping solution, and spent solutions from PGC salt production. cyanide Preliminary treatment is followed by chemical precipitation and sedimentation (lime and settle) treatment applied to the combined stream of steam stripper effluent, cyanide precipitation effluent, and the combined stream of all other wastewater. precipitation Chemical precipitation is used to remove metals by the addition of lime or caustic followed by gravity sedimentation. Suspended solids are also removed by the process. End-of-pipe treatment consisting of ion exchange is included for removal of gold, platinum and palladium.

OPTION B

Option B for the secondary precious metals subcategory consists of ammonia steam stripping, cyanide precipitation, chemical precipitation, sedimentation and ion exchange technology considered in Option A plus control technologies to reduce the discharge of wastewater volume. Recycle of furnace and refinery scrubber water as well as raw material granulation water are the principal control mechanisms for flow reduction.

OPTION C

Option C for the secondary precious metals subcategory consists of the ammonia steam stripping, cyanide precipitation, in-process flow reduction, chemical precipitation, sedimentation and ion exchange technology considered in Option B plus multimedia filtration technology added in between the sedimentation and ion exchange steps of the Option B treatment scheme. Multimedia filtration is used to remove suspended solids, including precipitates of metals, beyond the concentration attainable by sedimentation. The filter suggested is of the gravity gravity, mixed media type, although other forms of filters such rapid sand filters or pressure filters would as perform

satisfactorily. The addition of filters also provides consistent removal during periods in which there are rapid increases in flows or loadings of pollutants to the treatment system.

The Agency believes that it may be necessary for some facilities to use sulfide polishing in order to achieve the promulgated effluent limitations because of high zinc concentrations or complexing problems. Because the Agency believes that these situations will be the exception, rather than the rule, sulfide polishing is not specifically included as part of the model technology on which effluent limitations and performance standards are based. The Agency has, however, evaluated the cost associated with the use of sulfide polishing at secondary precious metals plants. After performing this evaluation, the Agency has concluded that sulfide polishing will result in a very small (less than 5 percent) incremental increase in wastewater treatment costs at a typical secondary precious metals facility

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SECONDARY PRECIOUS METALS SUBCATEGORY

SECTION VIII

COSTS, ENERGY, AND NONWATER QUALITY ASPECTS

This section presents a summary of compliance costs for the secondary precious metals subcategory and a description of the treatment options and subcategory-specific assumptions used to develop these estimates. Together with the estimated pollutant reduction performance presented in Sections IX. X, XI, and XII of this supplement, these cost estimates provide a basis for evaluating each regulatory option. These cost estimates are also used in determining the probable economic impact of regulation on the subcategory at different pollutant discharge levels. In addition, this section addresses nonwater quality environmental impacts of wastewater treatment and control alternatives, including air pollution, solid wastes, and energy requirements, which are specific to the secondary precious metals subcategory.

TREATMENT OPTIONS FOR EXISTING SOURCES

As discussed in Section VII, three treatment options have been developed for existing secondary precious metals sources. The treatment schemes for each option are summarized below and schematically presented in Figures X-1 through X-3 (page 2592).

OPTION A

Option A consists of ammonia steam stripping and cyanide precipitation preliminary treatment (where required), and chemical precipitation, sedimentation and ion exchange end-ofpipe technology.

OPTION B

Option B consists of in-process flow reduction measures, ammonia steam stripping and cyanide precipitation preliminary treatment (where required), and chemical precipitation, sedimentation and ion exchange end-of-pipe technology. The in-process flow reduction measures consists of the recycle of furnace scrubber water and refinery scrubber water as well as raw material granulation water through holding tanks.

OPTION C

Option C requires the in-process flow reduction measures of Option B, ammonia steam stripping and cyanide precipitation preliminary treatment, and end-of-pipe treatment technology consisting of chemical precipitation, sedimentation, multimedia filtration, and ion exchange.

The Agency believes that it may be necessary for some facilities to use sulfide polishing in order to achieve the promulgated effluent limitations because of high zinc concentrations or complexing problems. Because the Agency believes that these situations will be the exception, rather than the rule, sulfide polishing is not specif-ically included as part of the model technology on which effluent limitations and performance standards are based. The Agency has, however, evaluated the cost associated with the use of sulfide polishing at secondary precious metals plants. After performing this evaluation, the Agency has concluded that sulfide polishing will result in a very small (less than 5 percent) incremental increase in wastewater treatment costs at a typical secondary precious metals facility.

COST METHODOLOGY

A detailed discussion of the methodology used to develop the compliance costs is presented in Section VIII of the General Development Document. Plant-by-plant compliance costs for the nonferrous metals manufacturing category calculate incremental costs above treatment already in place, necessary to comply with promulgated effluent limitations and standards and are presented . in the administrative record supporting this regulation. The costs developed for the final regulation are presented in Tables VIII-1 and VIII-2 (page 2343) for direct and indirect dischargers in this subcategory, respectively.

- (1) For overlap plants (i.e., secondary precious metals secondary silver or secondary precious metals - secondary tungsten plants), costs and removal estimates are apportioned on a flow-weighted basis. The total flow used for flow-weighting costs includes recycled floor wash water, whereas the total flow used for flow-weighting removals does not include floor wash water.
- (2) A flow allowance for floor washing is assumed for each plant of the basis of 1.0 liter per troy ounce of precious metals, including silver, produced in the refinery. The flow allowance is based on the rates reported by the three plants supplying information about this stream. Table V-13 (page 2372) shows water use rates of 14.2, 1.0, and 0.97 liters per troy ounce. The highest rate was omitted because it is more than 10 times the next highest rate. The flow allowance was based an the average of the two lower rates.
- (3) Floor wash water is obtained by recycling wastewater treated by chemical precipitation and sedimentation for all options. The recycle ratio is equal to the flow of floor wash water divided by the total flow to treatment.
- (4) If a plant has a precipitation and filtration operation for platinum, palladium, other platinum group metals (PGM), or silver (from photographic raw materials), we assume floor wash water requires ammonia stripping to meet the proposed ammonia limitations.

- sludge produced from lime precipitation is expected to (5)All cyanide nonhazardous. All sludge produced from be precipitation is hazardous, under RCRA regulations.
- and filtration wastewater (aold, precipitation (6) All platinum, palladium, or other PGM) are assumed to undergo cementation prior to entering waste treatment. Zinc cementation is assumed unless iron cementation is specifically noted as in-place. Costs for installing operating a cementation system are not included in and cost estimates because cementation is not considered a the wastewater treatment operation. Only the zinc or iron raw waste values are changed by operating a cementation The revised raw waste values impact a plant's process. waste treatment cost.
- 50 Ammonia stripping costs for plants having less than (7) liters per hour of water requiring stripping are based on air stripping via agitation-aeration in the batch chemical precipitation tank instead of steam stripping. These costs include a blower, sparger and hood.
- Costs for ion exchange end-of-pipe treatment are completely (8) offset by the gold, platinum, and palladium values recovered by this system.

NONWATER QUALITY ASPECTS

A general discussion of the nonwater quality aspects of the control and treatment options considered for the nonferrous metals category is contained in Section VIII of Vol. I. Nonwater quality impacts specific to the secondary precious metals subcategory, including energy requirements, solid waste and air pollution are discussed below.

ENERGY REQUIREMENTS

The methodology used for determining the energy requirements for the various options is discussed in Section VIII of the General Development Document. Energy requirements for the three options considered are estimated at 5.30 x 106 kwh/yr, 5.31 x 106 kwh/yr, and 5.48 x 106 kwh/yr for Options A, B, and C, respectively. Option B energy requirements are similar to those for Option A. Because less water is being treated, energy costs for lime and settle treatment are less; however, recycle equipment such as holding tanks and pumps require additional energy, offsetting the Option C, which includes filtration, is savings. energy estimated to increase energy consumption over Option B by approximately 1 percent. Option C represents roughly 8 percent of a typical plant's electrical energy usage. It is therefore concluded that the energy requirements of the treatment options considered will not have a significant impact on total plant energy consumption.

SOLID WASTE

Sludge generated in the secondary precious metals subcategory is due to the precipitation of metal hydroxides and cyanide using lime and other chemicals. Sludges associated with the secondary precious metals subcategory will necessarily contain quantities of priority metal pollutants. Wastes generated by secondary metal industries can be regulated as hazardous. However, the Agency examined the solid wastes that would be generated at secondary nonferrous metals manufacturing plants by the suggested treatment technologies and believes they are not hazardous wastes under the Agency's regulations implementing Section 3001 of the Resource Conservation and Recovery Act. The one exception to this is solid wastes generated by cyanide precipitation. These sludges are expected to be hazardous and this judgment was included in this study. None of the non-cyanide wastes are listed specifically as hazardous. Nor are they likely to exhibit a characteristic of hazardous waste. This judgment is made based on the recommended technology of lime precipitation, sedimentation, and filtration. By the addition of a small excess (5-10%) of lime during treatment, similar sludges, specifically priority metal bearing sludges, generated by other industries such as the iron and steel industry passed the Extraction Procedure (EP) toxicity test. (See 40 CFR \$261.24.) Thus, the Agency believes that the wastewater sludge will similarly not be EP toxic if the recommended technology is applied.

Although it is the Agency's view that solid wastes generated as a result of these guidelines are not expected to be hazardous, generators of these wastes must test the waste to determine if the wastes meet any of the characteristics of hazardous waste (see 40 CFR \$262.11).

If these wastes identified should be or are listed as hazardous, they will come within the scope of RCRA s "cradle to grave" hazardous waste management program, requiring regulation from the generation to point of final disposition. point of EPA's standards would require generators of generator hazardous nonferrous metals manufacturing wastes to meet containerization, labeling, recordkeeping, and reporting requirements; if plants dispose of hazardous wastes off-site, they would have to prepare a manifest which would track the movement of the wastes from the generator's premises to a permitted off-site treatment, storage, disposal facility. See 40 CFR \$262.20 [45 FR 33142 (May or 19, 1980), as amended at 45 FR 86973 (December 31, 1980)]. The transporter regulations require transporters of hazardous waste to comply with the manifest system to assure that the wastes are delivered to a permitted facility. See 40 CFR \$263.20 [45 FR 33151 (May 19, 1980), as amended at 45 FR 86973 (December 31, 1980)]. Finally, RCRA regulations establish standards for hazardous waste treatment, storage, and disposal facilities allowed to receive such wastes. See 40 CFR Part 464 [46 FR 2802 (January 12, 1981), 47 FR 32274 (July 26, 1982)].

Even if these wastes are not identified as hazardous, they still

must be disposed of in compliance with the Subtitle D open dumping standards, implementing \$4004 of RCRA. See 44 FR 53438 (September 13, 1979). The Agency has calculated as part of the costs for wastewater treatment the cost of hauling and disposing of these wastes.

The Agency estimates that the promulgated BPT regulation for secondary precious metals manufacturing facilities will generate 523 metric tons of solid wastes (wet basis) in 1982 as a result of wastewater treatment. Promulgated BAT will not significantly increase sludge generation, however promulgated PSES will add 1,585 metric tons of solid waste per year which includes 344 tons per year of cyanide bearing sludges from treatment of cyanide bearing wastewaters by cyanide precipitation.

AIR POLLUTION

There is no reason to believe that any substantial air pollution problems will result from implementation of ammonia steam stripping, cyanide precipitation, chemical precipitation, sedimentation, multimedia filtration and ion exchange. These technologies transfer pollutants to solid waste and are not likely to transfer pollutants to air.

TABLE VIII-1

COST OF COMPLIANCE FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY DIRECT DISCHARGERS

The costs for this subcategory are not presented here because the data on which they are based are claimed to be confidential.

TABLE VIII-2

COST OF COMPLIANCE FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY INDIRECT DISCHARGERS

(March, 1982 Dollars)

Option	Total Required Capital Cost	Total Annual Cost
A	1,774,400	1,078,000
В	1,707,000	1,034,000
С	1,809,400	1,100,500

SECONDARY PRECIOUS METALS SUBCATEGORY

SECTION IX

BEST PRACTICABLE CONTROL TECHNOLOGY CURRENTLY AVAILABLE

This section defines the effluent characteristics attainable through the application of best practicable control technology currently available (BPT), Section 301(b)(a)(A). BPT reflects the existing performance by plants of various sizes, ages, and manufacturing processes within the secondary precious metals subcategory, as well as the established performance of the model BPT systems. Particular consideration is given to the treatment already in place at plants within the data base.

The factors considered in identifying BPT include the total cost of applying the technology in relation to the effluent reduction benefits from such application, the age of equipment and involved, the manufacturing processes facilities employed, quality environmental impacts (including energy nonwater requirements), and other factors the Administrator considers. appropriate. In general the BPT level represents the average to the existing performances of plants of various ages, sizes, processes, or other common characteristics. Where existing performance is uniformly inadequate, BPT may be transferred from different subcategory or category. Limitations based on а transfer of technology are supported by a rationale concluding that the technology is indeed, transferable, and a reasonable prediction that it will be capable of achieving the prescribed effluent limits (see Tanner's Council of America v. Train. 540 F.2d 1188 (4th Cir. 1176)). BPT focuses on end-of-pipe treatment rather than process changes or internal controls, except where such practices are common within the subcategory.

TECHNICAL APPROACH TO BPT

The Agency studied the nonferrous metals category to identify the processes used the wastewaters generated, and the treatment processes installed. Information was collected from the category using data collection portfolios, and specific plants were sampled and the wastewaters analyzed. In making technical assessments of data, reviewing manufacturing processes, and assessing wastewater treatment technology options, both indirect and direct dischargers have been considered as a single group. An examination of plants and processes did not indicate any process differences based on the type of discharge, whether it be direct or indirect.

As explained in Section IV, the secondary precious metals subcategory has been subdivided into 14 potential wastewater sources. Since the water use, discharge rates, and pollutant characteristics of each of these wastewaters is potentially unique, effluent limitations have been developed for each of the 14 building blocks.

each of the subdivisions, a specific approach was followed For the development of BPT mass limitations. The first for requirement to calculate these limitations is to account for production and flow variability from plant to plant. Therefore, a unit of production or production normalizing parameter (PNP) was determined for each waste stream which could then be related to the flow from the process to determine a production normalized flow. Selection of the PNP for each process element is discussed Each plant within the subcategory was then Section IV. in analyzed to determine which subdivisions were present, the specific flow rates generated for each subdivision, the and specific production normalized flows for each subdivision. This analysis is discussed in detail in Section V. Nonprocess wastewaters such as rainfall runoff and noncontact cooling water are not considered in the analysis.

Production normalized flows were then analyzed to determine the flow to be used as part of the basis for BPT mass limitations. The selected flow (sometimes referred to as the BPT regulatory flow or BPT discharge rate) reflects the water use controls which are common practices within the category. The BPT regulatory flow is based on the average to all applicable data. Plants with normalized flows above the average may have to implement some method of flow reduction to achieve the BPT limitations.

The second requirement to calculate mass limitations is the set of concentrations that are achievable by application of the BPT level of treatment technology. Section \overline{VII} discusses the various control and treatment technologies which are currently in place for each wastewater source. In most cases, the current control and treatment technologies consist of chemical precipitation and sedimentation (lime and settle) technology and a combination of reuse and recycle to reduce flow. Ammonia steam stripping is applied to streams with treatable concentrations of ammonia. Cyanide precipitation is applied to streams with treatable concentrations of free and complexed cyanide. Ion exchange is applied as an effluent polishing step to reduce combined metals (gold, platinum, and palladium), and palladium concentrations.

Using these regulatory flows and the achievable concentrations, the next step is to calculate mass loadings for each wastewater source or subdivision. This calculation was made on a stream-bystream basis, primarily because plants in this subcategory may perform one or more of the operations in various combinations.

The mass loadings (milligrams of pollutant per troy ounce of production -mg/T.0.) were calculated by multiplying the BPT regulatory flow (1/T.0.) by the concentration achievable by the BPT level of treatment technology (mg/1) for each pollutant parameter to be limited under BPT. These mass loadings are published in the <u>Federal Register</u> and in CFR Part 421 as the effluent limitations guidelines.

The mass loadings which are allowed under BPT for each plant will

be the sum of the individual mass loadings for the various wastewater sources which are found at particular plants. Accordingly, all the wastewater generated within a plant may be combined for treatment in a single or common treatment system, but the effluent limitations for these combined wastewaters are based on the various wastewater sources which actually contribute to the combined flow. This method accounts for the variety of combinations of wastewater sources and production processes which may be found at secondary precious metals plants.

The Agency usually establishes wastewater limitations in terms of mass rather than concentration. This approach prevents the use of dilution as a treatment method (except for controlling pH). The production normalized wastewater flow (1/T.0.) is a link between the production operations and the effluent limitations. The pollutant discharge attributable to each operation can be calculated from the normalized flow and effluent concentration achievable by the treatment technology and summed to derive an appropriate limitation for each plant.

INDUSTRY COST AND POLLUTANT REMOVAL ESTIMATES

In balancing costs in relation to pollutant removal estimates, EPA considers the volume and nature of existing discharges, the and nature of discharges expected after application of volume BPT, the general environmental effects of the pollutants, and the cost and economic impacts of the required pollution control level. The Act does not require or permit consideration of water quality problems attributable to particular point sources or industries, or water quality improvements in particular water quality bodies. Accordingly, water quality considerations were not the basis for selecting the proposed or promulgated BPT. See Weyerhaeuser Company v. Costle, 590 F.2d 1011 (D.C. Cir. 1978).

The methodology for calculating pollutant removal estimates and plant compliance costs is discussed in Section X. The pollutant removal estimates have been revised since proposal based on comments and on new data. Tables X-2 and XII-1 (pages 2574 and 2619) show the estimated pollutant removals for each treatment option for direct and indirect dischargers. Compliance costs are presented in Tables X-3 and XII-2 (pages 2575 and 2620).

BPT OPTION SELECTION

The technology basis for the BPT limitations is Option A, chemical precipitation and sedimentation technology to remove metals and solids from combined wastewaters and to control pH, ammonia steam stripping to remove ammonia, cyanide precipitation to remove free and complexed cyanide and ion exchange to remove gold, platinum and palladium. The promulgated technology is equivalent to the proposed technology with the addition of ion exchange. Chemical precipitation and sedimentation technology is already in-place at 20 of the plants in the subcategory including all four direct dischargers. One plant has cyanide precipitation in-place. Three plants currently practice ion exchange. The

technology bases for steam stripping and cyanide precipitation are discussed below. The pollutants specifically proposed for regulation at BPT are copper, cyanide, zinc, ammonia, combined platinum, and palladium), (gold, TSS, and pH. metals The Agency believes that it may be necessary for some facilities to use sulfide polishing in order to achieve the promulgated effluent limitations because of high zinc concentrations or complexing problems. Because the Agency believes that these situations will be the exception, rather than the rule, sulfide polishing is not specifically included as part of the model technology on which the BPT effluent limitations are based. The Agency has, however, evaluated the cost associated with the use of sulfide polishing at secondary precious metals plants. After performing this evaluation, the Agency has concluded that sulfide polishing will result in a very small (less than 5 percent) incremental increase in wastewater treatment costs at a typical secondary precious metals facility.

Implementation of the promulgated BPT limitations will remove annually an estimated 34,634 kg of priority pollutants (which includes 6.3 kg of cyanide), 490 kg of ammonia, and 18,420 kg of TSS.

The compliance costs for this subcategory are not presented here because the data on which they are based have been claimed to be confidential. The Agency has determined that the benefits justify the costs for this subcategory.

More stringent technology options were not selected for BPT since they require in-process changes end-of-pipe technologies less widely practiced in the subcategory, and, therefore, are more appropriately considered under BAT.

In response to a comment about high zinc concentrations in raw wastewater, EPA has included sulfide precipitation as a polishing step to treat high zinc concentrations in secondary precious metals wastewaters. The Agency concluded that the addition of sulfide precipitation would add only approximately 4 percent to the total annual costs for wastewater treatment at a typical secondary precious metals plant. Although not including it in the BPT treatment scheme, the Agency has no doubt that the CMDB concentrations could be achieved by secondary precious metals plants using sulfide polishing.

Ammonia steam stripping is demonstrated at six facilities in the nonferrous metals manufacturing category. These facilities are treating ammonia-bearing wastewaters associated with the production of primary tungsten, primary columbium and tantalum, primary molybdenum, secondary tungsten and cobalt, and primary zirconium and hafnium. EPA believes that performance data from the iron and steel manufacturing category provide a valid measure technology's performance on nonferr ng category wastewater because raw nonferrous metals of this manufacturing wastewater concentrations of ammonia are of the same order of magnitude in the respective raw wastewater matrices.

Chemical analysis data were collected of raw waste (treatment influent) and treated waste (treatment effluent) from one coke plant of the iron and steel manufacturing category. Using EPA sampling and chemical analysis protocols, six paired samples were collected in a two-month period. These data are the data base for determining the effectiveness of ammonia steam stripping technology. Ammonia treatment at this coke plant consisted of two steam stripping columns in series with steam injected countercurrently to the flow of the wastewater. A lime reactor for pH adjustment separated the two stripping columns.

The Agency has verified the proposed steam stripping performance values using steam stripping data collected at a primary zirconium and hafnium plant which has raw ammonia levels as high as any in the nonferrous metals manufacturing category. Data collected by the plant represent almost two years of daily operations, and support the long-term mean used to establish treatment effectiveness. Also, data from a bench scale treatability study which was submitted with comments on the proposed rulemaking show that the proposed steam stripping performance can be achieved with secondary precious metals wastewaters.

Cyanide precipitation is demonstrated in the secondary precious metals subcategory at one plant. Cyanide precipitation is required for the secondary precious technology metals subcategory because existing treatment within the subcategory does not effectively remove cyanide. Most secondary precious metals plant use alkaline oxidation to destroy free cyanide, but do not effectively remove complexed cyanide. Cyanide precipitation is directed at control of free and complexed cyanides in waste streams within the secondary precious metals subcategory collectively subcategory. This discharges approximately 557 kg/yr of cyanide. The achievable performance is transferred from three well-operated coil coating plants in the coil coating category, and are contained within the public record supporting this document. The Agency technology, and the achievable concentration The Agency believes this limits, are transferable to the secondary precious metals subcategory because raw wastewater cyanide concentrations (prior to dilution with waste streams without cyanide) are of the same order of magnitude in both categories. Further, no pollutants were identified in secondary precious metals wastewater that would interfere with the operation or performance of this technology.

Several discharging plants within the secondary precious metals subcategory use chlorine gas or hypochlorite solution to oxidize cyanide in their wastewater. EPA considered chemical oxidation using chlorine. Although the chlorine oxidation process can be used effectively for wastewater containing predominantly free cyanides and easily oxidizable cyanide complexes, the Agency determined that precipitation with ferrous sulfate is more effective than chlorine oxidation for the removal of iron-cyanide complexes which may be found in the secondary precious metals

wastewater.

WASTEWATER DISCHARGE RATES

A BPT discharge rate is calculated for each subdivision based on the average of the flows of the existing plants, as determined from analysis of the dcp. The discharge rate is used with the achievable treatment concentrations to determine BPT effluent limitations. Since the discharge rate may be different for each wastewater source, separate production normalized discharge rates for each of the 14 wastewater sources are discussed below and summarized in Table IX-1 (page 2546). The discharge rates are normalized on a production basis by relating the amount of wastewater generated to the mass of the intermediate product which is produced by the process associated with the waste stream in question. These production normalizing parameters, or PNPs, are also listed in Table IX-1.

Section V of this supplement further describes the discharge flow rates and presents the water use and discharge flow rates for each plant by subdivision.

FURNACE WET AIR POLLUTION CONTROL

The BPT wastewater discharge rate proposed and promulgated for furnace wet air pollution control is 71.8 liters per troy ounce of precious metals, including silver, incinerated or smelted, based on zero percent recycle. This rate is allocated only for plants practicing wet air pollution control for the furnace. Seven plants reported this wastewater, two of whom practice 100 percent recycle (plants 1094 and 1084). The BPT rate is based on the average water use rate of four of the remaining five plants. Plant 1105 was omitted because its water use rate was not reported, and its recycle rate was not quantified. The BPT rate is the average of 137, 116, 27.6, and 7.26 liters per troy ounce. The distribution of wastewater rates for this waste stream is presented in Section V (Table V-1, page 2360).

RAW MATERIAL GRANULATION

The BPT wastewater discharge rate promulgated for raw material granulation is 6.34 liters per troy ounce of precious metals in the granulated raw material. This rate is different than the proposed rate, which was 0 liters per troy ounce. Based on comments received following proposal, the Agency determined that a discharge is necessary for raw material granulation water. The promulgated BPT rate is based on the average of the rates reported by two of the three plants whose production normalized water use rates are quantified, plants 1008 and 1094. As shown in Table V-2 (page 2361), these rates are 8.67 and 4.0 liters per troy ounce, respectively. The production normalized flow rate for plant 1112 was not included in the calculation of the BPT discharge rate for this subdivision as discussed at the front of this section.

SPENT PLATING SOLUTIONS

The BPT wastewater discharge rate proposed and promulgated for spent plating solutions is 1.0 liter per liter of spent plating solution raw material. This rate is applicable to those plants which recover gold and other precious metals from spent or contaminated electroplaters solutions which they receive as a raw The discharge rate is given in terms of volume of raw material. because EPA believes plants cannot control material the concentration of precious metals in this raw material, and should be allowed to discharge the entire volume of solution coming into the plant, after recovering the precious metals. Only the volume of raw material solution should be allowed to be discharged, and this is why a discharge rate of 1.0 liter per liter was selected. The 12 plants with this subdivision are shown in Table V-3 (page 2362).

SPENT CYANIDE STRIPPING SOLUTIONS

The BPT wastewater discharge rate promulgated for spent cyanide stripping solutions is 3.7 liters per troy ounce of gold recovered by cyanide stripping. This rate applies to plants which recover gold by stripping it from a raw material such as electronic scrap, with a cyanide-based solution, and then recovering the gold from this solution. This rate is based on the average of the lower five of six water use rates reported for stream in Table V-4 (page 2363). Plant 1100 was omitted this because of its excessive water use. EPA proposed 1.1 liters per troy ounce for this waste stream, but stated that it was considering 4.7 liters per troy ounce and would select between the two based on a variety of factors, including public comment. EPA decided to revise the rate from 1.1 to 3.7 liters per troy ounce for promulgation based on comments received following proposal.

REFINERY WET AIR POLLUTION CONTROL

The BPT wastewater discharge rate proposed and promulgated for refinery wet air pollution control is 21.0 liters per troy ounce of precious metals, including silver, produced in the refinery, based on zero percent recycle. This rate applies to either acid alkaline scrubbers. If both acid and alkaline scrubbers are present in a particular facility, the same rate applies to each. This rate is allocated only for plants practicing wet air pollution control for acid or cyanide fumes in the refinery. Twenty-eight plants reported this waste stream, five of which practice zero percent recycle and discharge the wastewater (107, 32.8, 6.8, and 2.4 liters per troy ounce). The BPT rate is 42, based on the average of the lower four of these five dischargers. The highest flow rate (107 liters per troy ounce) was omitted from the BPT rate calculation because there is no reason to believe this much water is needed for this operation in light of rates from the other plants. Table V-5 (page 2364) shows the distribution of water use and discharge rates for refinery wet pollution control. The production normalized flow rate for air

plant 1112 was not included in the calculation of the BPT discharge rate for this subdivision as discussed at the front of this section.

GOLD SOLVENT EXTRACTION RAFFINATE AND WASH WATER

The BPT wastewater discharge rate proposed and promulgated for gold solvent extraction raffinate and wash water is 0.63 liters per troy ounce of gold recovered by solvent extraction. This discharge rate is allocated only to plants which refine gold by a solvent extraction process. The discharge rate is based on the rate reported by the only plant with this process (0.63 liters per troy ounce), as shown in Table V-6 (page 2366).

GOLD SPENT ELECTROLYTE

The BPT wastewater discharge rate proposed and promulgated for gold spent electrolyte is 0.0087 liters per troy ounce of gold recovered by electrolysis. This rate only applies to plants which refine gold by electrolysis. The discharge rate is based on the lower of the two rates reported for this wastewater stream (0.0087 liters per troy ounce), as shown in Table V-7 (page 2366). The other flow rate (0.294 liters per troy ounce) is more than 10 times higher than the selected BPT rate.

GOLD PRECIPITATION AND FILTRATION

The BPT wastewater discharge rate proposed and promulgated for gold precipitation and filtration is 4.4 liters per troy ounce of gold precipitated. This rate only applies to plants which refine gold by dissolving gold-containing raw material in acid, and then recovering gold by precipitation. This discharge rate is based on the final or net quantity of gold produced using this process, not the amount precipitated through each refining step. Of the 28 plants using this process, nine plants supplied insufficient information to calculate discharge rates, two plants report 100 percent reuse of this water, and six plants do not discharge this waste stream by means of contract hauling (these plants have water use rates of 560.5, 69.1, 3.34, 0.815, 0.63, and 0.05 liters per troy ounce). The BPT discharge rate is based on the average water use rate of 10 of the 11 discharging plants (24.3, 7.98, 4.1, 2.65, 2.5, 1.86, 0.341, 0.312. 0.27, and 0.144 liters per troy ounce). The plant reporting 404 liters per troy ounces rate was not considered in the average because this water use rate is almost 10 times that of the next highest plant. Eight of the discharging plants meet the BPT rate. Water use and discharge rates are presented in Table V-8 (page 2367).

PLATINUM PRECIPITATION AND FILTRATION

The BPT wastewater discharge rate proposed and promulgated for platinum precipitation and filtration is 5.2 liters per troy ounce of platinum precipitated. This rate only applies to plants which refine platinum by dissolving it in acid or base, and recover it by precipitation. This discharge rate is based on the

final or net quantity of metal produced using this process, not the amount precipitated through each refining step. 0f the 18 plants using this process, 13 supplied insufficient information to calculate discharge rates. Five plants reported sufficient data (354, 30.2, 10.4, 4.5, and 0.58 liters per troy ounce). Table V-9 (page 2369) presents the water use and discharge rates for this waste stream. The BPT discharge rate is based on thé average of the three lowest water use rates. The 354 and 30.2 liters per troy ounce water use rates were omitted from the average because there is no reason to believe this much water is needed for this operation in light of the rates from the other plants.

PALLADIUM PRECIPITATION AND FILTRATION

The BPT wastewater discharge rate proposed and promulgated for palladium precipitation and filtration was 3.5 liters per troy ounce of palladium precipitated. As a result of the settlement proposing to modify the flow rate from the agreement, EPA is palladium precipitation and filtration building block from 3.5 to liters per troy ounce of precious metals produced. 6.0 This reflects a recalculation of the average flow for change this building block and the incorporation of new data. This rate only applies to plants which refine palladium by dissolving it in acid or base, and then recovering it by precipitation. This discharge rate is based on the final or net quantity of metal produced using this process, not the amount precipitated through each refining step.

OTHER PLATINUM GROUP METALS PRECIPITATION AND FILTRATION

The BPT wastewater discharge rate proposed and promulgated for other platinum group metals (rhodium, iridium, osmium, and ruthenium) precipitation and filtration is 5.2 liters per troy ounce of platinum group metals precipitated. This rate only applies to plants which refine these metals by dissolving them in either acid or base, and then precipitating them. This discharge rate is based on the final or net quantity of metal produced using this process, not the amount precipitated through each refining step. Three plants use this process and none reported sufficient information to calculate water use or discharge rates. This is shown in Table V-11. The BPT discharge rate is therefore based on the platinum precipitation and filtration BPT discharge rate derived from Table V-9. These two subdivisions are expected to have similar flows because all five metals (platinum, rhodium, iridium, osmium, and ruthenium) are part of the platinum group, and all are refined in a similar manner.

SPENT SOLUTION FROM PGC SALT PRODUCTION

The BPT wastewater discharge rate proposed and promulgated for spent solution from the PGC salt production process is 0.9 liters per troy ounce of gold contained in PGC product. This rate applies only to plants which manufacture a potassium gold cyanide salt product by reacting pure gold with potassium cyanide.

There are four plants reporting this process, as shown solution. in Table V-12 (page 2371). The plant reporting 260 liters per troy ounce explained that part to that water is used in a scrubber above the reaction vessel, and the two flow rates (discharging excess solution and scrubber liquor) could not be separated. It is likely that most of that plant's water discharge is due to the scrubber. The plant reporting 0.9 liters per troy ounce confirmed that its water discharge was due entirely to the excess reaction solution. Because the data from the plant reporting 260 liters per troy ounce could not be apportioned between scrubber liquor and spent solution, the BPT wastewater discharge is based upon 0.9 liters per troy ounce. The production normalized flow rate for plant 1112 was not included in the calculation of the BPT discharge rate for this subdivision as discussed at the front of this section.

EQUIPMENT AND FLOOR WASH

The BPT wastewater discharge rate proposed and promulgated for equipment and floor wash is 0 liters per troy ounce of precious metals, including silver, produced in the refinery. The BPT discharge rate is based on recycle of treated effluent for use as raw water for equipment and floor wash. In precious metals refineries, EPA realizes there is a possibility of accidental leaks and spills, which may contain precious metals and silver, and need to be recovered by washing the equipment and the floor. We believe that wastewater treatment plant effluent can be recycled for this purpose, increasing the capacity of treatment but not the actual amount of water discharged.

PRELIMINARY TREATMENT

As discussed in Sections V and VII, EPA agreed to establish BPT limitations for a new building block, preliminary treatment, for the secondary precious metals subcategory. The BPT water discharge rate for this block is 50 liters per troy ounce of precious metal produced. The flow basis is based on information that is considered confidential.

REGULATED POLLUTANT PARAMETERS

The raw wastewater concentrations from individual operations and the subcategory as a whole were examined to select certain pollutant parameters for limitation. This examination and evaluation was presented in Section VI. Seven pollutant or pollutant parameters selected for limitation under BPT and are listed below:

120. copper
121. cyanide
128. zinc
ammonia (as N)
combined metals (gold, platinum, and palladium)
total suspended solids (TSS)

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EFFLUENT LIMITATIONS

The concentrations achievable by application of the promulgated BPT treatment are explained in Section VII of Vol. I and summarized there in Table VII-21 (page 248). The achievable treatment concentrations (both one-day maximum and monthly average values) are multiplied by the BPT normalized discharge flows summarized in Table IX-1 (page 2546) to calculate the mass of pollutants allowed to be discharged per mass of product. The results of these calculations in milligrams of pollutant per troy ounce of product represent the BPT effluent limitations and are presented in Table IX-2 (page 2548) for each individual building block.

TABLE IX-1

BPT WASTEWATER DISCHARGE RATES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

Wastewater Stream	BAT Normalized <u>Discharge</u> <u>Rates</u>	Production Normalizing Parameter
Furnace wet air pollution control	4.5 l/t.o.	Troy ounces of precious metals, including silver, incinerated or smelted
Raw material granulation	0.64 l/t.o.	Troy ounces of precious metals in the granulated raw material
Spent plating solutions	1.1 1/1	Liters of spent plating solution used as a raw material
Spent cyanide stripping solutions	3.7 l/t.o.	Troy ounces of gold stripped
Refinery wet air pollution control	1.0 l/t.o.	Troy ounces of precious metals produced in refinery, including silver
Gold solvent extraction raffinate	0.63 l/t.o.	Troy ounces of gold produced by solvent extraction
Gold spent electrolyte	0.0087 l/t.o.	Troy ounces of gold produced by electrolysis
Gold precipitation and filtration	4.4 l/t.o.	Troy ounces of gold precipitated
Platinum precipitation and filtration	5.2 l/t.o.	Troy ounces of platinum precipitated

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XI

TABLE IX-1 (Continued)

BPT WASTEWATER DISCHARGE RATES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

Wastewater Stream	BAT Normalized <u>Discharge</u> <u>Rates</u>	Production Normalizing Parameter
Palladium precipitation and filtration	3.5 l/t.o.	Troy ounces of palladium precipitated
Other platinum group metals precipitation and filtration	5.2 l/t.o.	Troy ounces of other platinum group metals precipitation
Spent solution from PGC salt production	0.9 1/t.o.	Troy ounces of gold contained in PGC product
Equipment and floor wash	0.0 l/t.o.	Troy ounces of precious metals metals including silver produced in refinery

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NOTES 1/t.o. = liters per troy ounce 1/1 = liters per liter

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TABLE IX-2

BPT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(a) Furnace Wet Air Pollution Control BPT

Pollutant or	Maximum for	Maximum for
pollutant property	any one day	monthly average
·····		
mg/troy ounce	of precious metals	incinerated or smelted
Antimony	206.100	91.900
Arsenic	150.100	66.770
Cadmium	24.410	10.770
Chromium	31.590	12.920
*Copper	136.400	71 .8 00
*Cyanide	20.820	8.616
Lead	30.160	14.360
Nickel	137.900	91.190
Selenium	88.310	39.490
Silver	29.440	12,210
Thallium	147.200	65.340
*Zinc	104.800	43.800
*Ammonia	9,571.000	4,207.000
*Combined metals	21.540	
*TSS	2,944.000	1,400.000
	ange of 7.5 to 10.0	

TABLE IX-2 (Continued)

BPT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(b) Raw Material Granulation BPT

Pollutant pollutant		Maximum fo any one da		
mg/troy	ounce of	precious metals	in the granulated	raw material
Antimony	a de la composición d Composición de la composición de la comp	18.200	8.11	.5
Arsenic		13.250	5.89	6
Cadmium	-	2.156	5.4 J	
Chromium		2.790	1.14	1
*Copper		12.050	6.34	0
*Cyanide		1.839	.76	1
Lead		2.663	1.26	8
Nickel		12.170	8.05	2
Selenium		7.798	3.48	7
Silver		2.599	1.07	- · · ·
Thallium		13.000		8 1 a s 1
*Zinc	· · · · · ·	9.256		
*Ammonia		845.100	371.50	
*Combined	metals	1.902		•
*TSS	· ·	259.900	123.60	0
*pH Wi	thingthe		10.0 at all times	

TABLE IX-2 (Continued)

BPT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(c) Spent Plating Solutions BPT

Pollutant or pollutant property	Maximum for any one day	Maximum for monthly average
mg/liter of spent	plating solutions	used as a raw material
Antimony	2.870	1.280
Arsenic	2.090	.930
Cadmium	.340	.150
Chromium	.440	.180
*Copper	1.900	1.000
*Cyanide	.290	.120
Lead	.420	.200
Nickel	1.920	1.270
Selenium	1.230	.550
Silver	.410	.170
Thallium	2.050	.910
*Zinc	1.460	.610
*Ammonia	133.300	58.6 00
*Combined metals	.300	
*TSS	41.000	19.500
*pH Within the ra	nge of 7.5 to 10.0	at all times

TABLE IX-2 (Continued)

BPT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(d) Spent Cyanide Stripping Solutions BPT

	any one day mont g/troy ounce of gold stri	
	g/troy ounce of gold stri	
	g/troy ounce of gold stri	
Antimony		pped
Anciaony	10.620	4.736
Arsenic	7.733	3.441
Cadmium	1.258	.555
Chromium	1.628	.666
*Copper	7.030	3.700
*Cyanide	1.073	.444
Lead	1.554	.740
Nickel	7.104	4.699
Selenium	4.551	2.035
Silver	1.517	.629
Thallium	7.585	3.367
*Zinc	5.402	2.257
*Ammonia	493.200	216.800
*Combined metals	1.110	
*TSS	151.700	72.150
*pH Within the	range of 7.5 to 10.0 at a	ll times

TABLE IX-2 (Continued)

BPT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(e) <u>Refinery Wet Air</u> <u>Pollution</u> <u>Control¹</u> BPT

Pollutant or	Maximum for	Maximum for	1
pollutant property	any one day	monthly average	

mg/troy ounce of precious metals produced in refinery

Antimony	60 270	26.880
Arsenic	43.890	19.530
Cadmium	7.140	3.150
Chromium	9.240	3.780
*Copper	39.900	21.000
*Cyanide	6.090	2.520
Lead	8.820	4.200
Nickel	40.320	26.670
Selenium	25.830	11.550
Silver	8.610	3.570
Thallium	43.050	19.110
*Zinc	30.660	12.810
*Ammonia	2,799.000	1,231.000
*Combined metals	6.300	بنيبة بنيد لنعد
*TSS	861.000	409.500
*pH Within the range	of 7.5 to 10.0 at	

*Regulated Pollutant

¹This allowance applies to either acid or alkaline wet air pollution control scrubbers. If both acid and alkaline wet air pollution control scrubbers are present in a particular facility the same allowance applies to each.

TABLE IX-2 (Continued)

BPT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(f) Gold Solvent Extraction Raffinate and Wash Water BPT

Pollutant or pollutant property			imum for one day		laximum onthly			
	mg/troy	ounce	of gol	d produc	ed by	solvent	extract	ion
Antimo	ny	x		1.808	·		.806	
Arseni				1.317	• •		.586	
Cadmiu	ım			.214			.095	1.1
Chromi	um			.277			.113	
*Coppe	er			1.197	н. н. 1		.630	
*Cyani				.183			.076	· · · ·
Lead		•	τ	.265			.126	
Nickel	Ľ			1.210	•		.800	· · ·
Seleni	um	.'		.775			.347	
Silver		•		.258			.107	•
Thalli	um			1.292			.573	
*Zinc				.920			.384	
*Ammor	nia			83.980			36.920	
	ned meta	ls		.189			`	
*TSS				25.830	1. S. 1.	•	12.290	
*pH	Within	the ra	nge of	7.5 to	10.0 a	t all t	imes	

SECONDARY PRECIOUS METALS SUBCATEGORY

TABLE IX-2 (Continued)

BPT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(g) Gold Spent Electrolyte BPT

Pollut pollut			pert	у		imum one			ximum nthly	for average	
	1	mg/t	roy	ounce	of	gold	produce	ed by	y ele	ctrolysis	
Antimo	ny					. ()25			.011	
Arseni	.c					. ()18			.008	
Cadmiu						. (03			.001	
Chromi						. (04			.002	
*Coppe						. ()17			.009	
*Cyani	de					. (03			.001	
Lead						. ()04			.002	4 6
Nickel						. ()17			.011	
Seleni						. ()11			.005	
Silver						. (04			.001	,
Thalli	um					. ()18			.008	
*Zinc							13			.005	
*Ammon			_			1.1	.60			.510	
*Combin	ned	meta	als			.0	03				
*TSS							57			.170	÷
*pH	Wit	hin	the	range	of	7.5	to 10.0	at	all	times	. •

TABLE IX-2 (Continued)

BPT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(h) Gold Precipitation and Filtration BPT

	and the second		1 · · · · · · · · · · · · · · · · · · ·	
Pollutant	or	Maximum for	Maximum for .	
pollutant	property	any one day	monthly average	
-		· · · · · · · · · · · · · · · · · · ·		
	•			
	mg/troy	ounce of gold	precipitated	
Antimony		12.630	5.632	
Arsenic		9.196	4.092	
Cadmium		1.496	.660	
Chromium		1.936	.792	
*Copper		8.360	4.400	
*Cyanide		1.276	.528	
Lead		1.848	.880	
Nickel		8.448	5.588	
Selenium	` x	5.412	2,420	
Silver		1.804	.748	
Thallium		9.020	4.004	* <u>'</u>
*Zinc		6.424	2.684	
*Ammonia		586.500	257.800	
*Combined	metals	1.320		
*TSS		180.400	85.800	
	thin the rang		.0 at all times	

TABLE IX-2 (Continued)

BPT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(i) Platinum Precipitation and Filtration BPT

Pollutant pollutant			imum fo one da		ximum onthly	for average	
	mg/t	roy ounce	of pla	tinum pr	ecipit	ated	
Antimony Arsenic Cadmium Chromium *Copper *Cyanide Lead Nickel Selenium Silver Thallium *Zinc *Ammonia *Combined *TSS		2	14.920 10.870 1.768 2.288 9.880 1.508 2.184 9.984 6.396 2.132 10.660 7.592 93.200 1.560 213.200	. 5	1	6.656 4.836 .780 .936 5.200 .624 1.040 6.604 2.860 .884 4.732 3.172 04.700 01.400	
*pH Wit	hin the	range of	7.5 to	10.0 at	all t	imes	

*Regulated Pollutant

.

TABLE IX-2 (Continued)

BPT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(j) Palladium Precipitation and Filtration BPT

Pollutant pollutant	· · · · · · · · · · · · · · · · · · ·	Maximum for any one day	Maximum for monthly average	
				1.5 B
	mg/troy ou	nce of palladium	precipitated	
Antimony		17.230	7.680	
Arsenic		12.540	5.580	•
Cadmium	(1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,1,	2.040	.900	
Chromium		2.640	1.08	
*Copper	i i i i i i i i i i i i i i i i i i i	11.400	6.000	
*Cyanide		1.740	.720	
Lead		2.520	1.200	
Nickel	1. 人名法法法	11.520	7.620	
Selenium		7.380	3.300	1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1
Silver	·	2.460	1.020	
Thallium		12.300	5.460	
*Zinc	A Charles of the second	8.760	3.660	
*Ammonia		799.800	351.600	
*Combined	metals	1.800		
*TSS		246.000	117.000	
*pH Wi	thin the range	of 7.5 to 10.0	at all times	9

TABLE IX-2 (Continued)

BPT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(k) Other Platinum Group Metals Precipitation and Filtration BPT

Pollutant or	Montinum			
	Maximum for	Maximum	for	
pollutant property	any one day	monthly	average	

mg/troy ounce of other platinum group metals precipitated

B		н. 1
Antimony	14.920	6. 656
Arsenic	10.870	4.836
Cadmium	1.768	.780
Chromium	2.288	
*Copper	9.880	.936
*Cyanide		5.200
—	1.508	.624
Lead	2.184	1.040
Nickel	9.984	6.604
Selenium	6.396	2.860
Silver	2.132	.884
Thallium	10.660	
*Zinc	7.592	4.732
*Ammonia		3.172
	693.200	304.700
*Combined metals	1.560	
*TSS	213.200	101.400
*pH Within the range	of 7.5 to 10.0 at	all times
•		

SECONDARY PRECIOUS METALS SUBCATEGORY

TABLE IX-2 (Continued)

BPT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(1) Spent Solution from PGC Salt Production BPT

Pollutant pollutant		У	Maximum any one		Maximum month1	m for y average	
n	ng/troy	ounce	of gold	containe	ed in P	GC product	
Antimony			2.	5 83		1.152	·
Arsenic			1.4	881		.837	· .
Cadmium			•	306		.135	
Chromium			•	396		.162	
*Copper			1.	710		.900	
*Cyanide			•	261		.108	
Lead				378		.180	
Nickel			1.	728		1.143	
Selenium	· · · ·		1.	107	. ••	.495	
Silver			•	369		.153	
Thallium			1.	845		.819	
*Zinc			1.	314		.549	
*Ammonia	· . *	· · ·	120.	000		52.740	
*Combined	metals	· .	•	270			
*TSS			36.		,	17.550	
*pH Wi	thin the	e range	e of 7.5	to 10.0	at all	times	

TABLE IX-2 (Continued)

BPT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(m) Equipment and Floor Wash BPT

		and the second
Pollutant or	Maximum for	Maximum for
pollutant property	any one day	monthly average
mg/troy ound	e of precious metal	s produced in refinery
Antimony	0.000	0.000
Arsenic	0.000	0.000
Cadmium	0.000	0.000
Chromium	0.000	0.000
*Copper	0.000	0.000
*Cyanide	0.000	0.000
Lead	0.000	0.000
Nickel	0.000	0.000
Selenium	0.000	0.000
Silver	0.000	0.000
Thallium 🤄	0.000	0.000
*Zinc	0.000	0.000
*Ammonia	0.000	0.000
Combined metals	0.000	0.000
*TSS	0.000	0.000
*pH Within the :	range of 7.5 to 10.0	0 at all times
2		

*Regulated Pollutant

1.72

TABLE IX-2 (Continued)

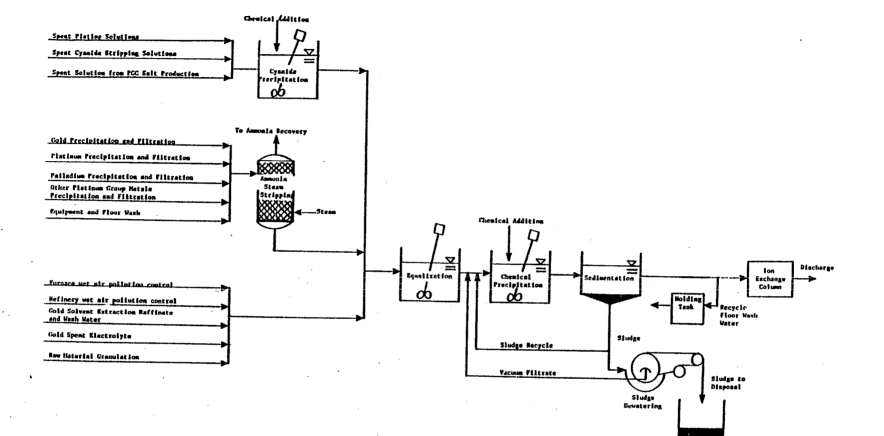
BPT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(n) Preliminary Treatment BPT

Pollutant	or	Maximum for	Maximum for	÷
Pollutant	Property	Any One Day	Monthly Average	

mg/troy ounce of total precious metals produced through this operation

Antimony	143.500 Jam 14	64.000
Arsenic	104.500	
Cadmium	17.000	
Chromium	22.000	
*Copper	95.000	50.000
*Cyanide (total)	14.500	
	21.000	
Lead	96.000	
Nickel	61.500	
Selenium	20.500	
Silver		45.5
Thallium	102.5	1
*Zinc	73.000	
*Ammonia (as N)	6665.000	2930.000
*Combined metals	15.000	a the second
*Total suspended	2050.000	975.000
solids	4 -	
*pH	Within the ran at all t	ge of 7.5 to 10.0 imes



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Figure IX-1

BPT TREATMENT SCHEME FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

SECTION X

BEST AVAILABLE TECHNOLOGY ECONOMICALLY ACHIEVABLE

These effluent limitations are based on the best control and treatment technology used by a specific point source within the industrial category or subcategory, or by another category where it is readily transferable. Emphasis is placed on additional treatment techniques applied at the end of the treatment systems currently used, as well as reduction of the amount of water used and discharged, process control, and treatment technology optimization.

The factors considered in assessing best available technology economically achievable (BAT) include the age of equipment and facilities involved, the process used, process changes, nonwater quality environmental impacts (including energy requirements), and the costs of application of such technology. BAT represents the best available technology economically achievable at plants of various ages, sizes, processes, or other characteristics. Where the Agency has found the existing performance to be uniformly inadequate, BAT may be transferred from a different subcategory or category. BAT may include feasible process changes or internal controls, even when not in common practice.

The statutory assessment of BAT considers costs, but does not require a balancing of costs against pollutant removals (see <u>Weyerhaeuser</u> v. <u>Costle</u>, 11 ERC 2149 (D.C. Cir. 1978)). However, in assessing the proposed and promulgated BAT, the Agency has given substantial weight to the economic achievability of the technology.

TECHNICAL APPROACH TO BAT

The Agency reviewed a wide range of technology options and evaluated the available possibilities to ensure that the most effective and beneficial technologies were used as the basis to BAT. To accomplish this, the Agency elected to examine three technology options which could be applied to the secondary precious metals subcategory as alternatives for the basis of BAT effluent limitations.

the development of BAT effluent limitations, mass loadings For were calculated for each wastewater source or subdivision in the subcategory using the same technical approach as described in Section IX for BPT limitations development. The differences in the mass loadings for BPT and BAT are due to increased treatment effectiveness achievable with the more sophisticate treatment technology and reductions in the effluent sophisticated BAT flows allocated to various waste streams.

In summary, the treatment technologies considered for the secondary precious metals subcategory are:

Option A (Figure X-1, page 2592):

- Cyanide precipitation preliminary treatment for streams containing cyanide at treatable concentrations
- Ammonia steam stripping preliminary treatment for streams containing ammonia at treatable concentrations
- o Chemical precipitation and sedimentation
- o Ion exchange end-of-pipe treatment

Option B (Figure X-2, page 2593) is based on

- In-process flow reduction of wet air pollution control and raw material granulation water
- Cyanide precipitation preliminary treatment for streams containing cyanide at treatable concentrations
- Ammonia steam stripping preliminary treatment for streams containing ammonia at treatable concentrations
- o Chemical precipitation and sedimentation
- o Ion exchange end-of-pipe treatment

Option C (Figure X-3, page 2594) is based on

- In-process flow reduction of wet air pollution control and raw material granulation water
- Cyanide precipitation preliminary treatment for streams containing cyanide at treatable concentrations
- Ammonia steam stripping preliminary treatment for streams containing ammonia at treatable concentrations
- o Chemical precipitation and sedimentation
- o Multimedia filtration
- o Ion exchange end-of-pipe treatment

The three options examined for BAT are discussed in greater detail below. The first option considered is the same as the BPT treatment technology which was presented in the previous section.

OPTION A

Option A for the secondary precious metals subcategory is equivalent to the control and treatment technologies which were analyzed for BPT in Section IX. The BPT end-of-pipe treatment scheme includes chemical precipitation, sedimentation (caustic or lime and settle), and ion exchange, with ammonia steam stripping preliminary treatment of wastewaters containing treatable concentrations of ammonia, and cyanide precipitation preliminary treatment of wastewaters containing treatable concentrations of cyanide (see Figure X-1, page 2592). The discharge rates for Option A are equal to the discharge rates allocated to each stream as a BPT discharge flow.

OPTION B

Option B for the secondary precious metals subcategory achieves lower pollutant discharge by building upon the Option A (ammonia

steam stripping, cyanide precipitation, chemical precipitation, sedimentation and ion exchange) treatment technology. Flow reduction measures are added to the Option A treatment scheme (see Figure X-2, page 2593). These flow reduction measures. including in-process changes, result in the concentration to pollutants in some wastewater streams. As explained in Section VII of the General Development Document, treatment of a more concentrated effluent allows achievement of a greater net pollutant removal and introduces the possible economic benefits associated with treating a lower volume of wastewater.

Option B flow reduction measures are reflected in the BAT wastewater discharge rates. Flow reduction has been included in determining the BAT discharge rates for furnace wet air pollution control, raw material granulation, and refinery wet air pollution control. Based on available data, the Agency did not feel that further flow reduction over BPT would be feasible for the remaining 10 waste streams in the secondary precious metals subcategory. These waste streams are:

- 1. Spent plating solutions,
- 2. Spent cyanide stripping solutions,
- 3. Gold solvent extraction raffinate and wash water,
- 4. Gold spent electrolyte,
- 5. Gold precipitation and filtration,
- 6. Platinum precipitation and filtration,
- 7. Palladium precipitation and filtration
- 8. Other platinum group metals precipitation and filtration,
- 9. Spent solution from PGC salt production, and
- 10. Equipment and floor wash.

Flow reduction measures used in Option B to reduce process wastewater generation or discharge rates include the following:

Recycle of Water Used in Wet Air Pollution Control

There are two wastewater sources associated with wet air pollution control which are regulated under these effluent limitations:

- 1. Furnace scrubber, and
- 2. Refinery scrubber.

Table X-1 (page 2592) presents the number of plants reporting wastewater from the wet air pollution control sources listed above, the number of plants practicing recycle, and the range of recycle values being listed. Recycle of both furnace scrubber water and refinery scrubber water are required for BAT. The recycle rate used for both sources is based on the average of all discharging plants which currently practice recycle of these waste streams (currently practicing greater than 90 percent recycle), as will be shown later.

Recycle of Water Used for Raw Material Granulation

As shown in Table V-2 (page 2361), three plants generate wastewater from their raw material granulation operation. One plant (1082) practices 100 percent recycle of this water, and two plants do not practice any recycle of this water. Flow reduction based on 90 percent recycle of raw material granulation water using holding tanks, is included under Option B. Recycle is demonstrated at one plant for this subdivision, and there is no reason to believe that 90 percent recycle cannot be practiced at the two other plants with this subdivision.

OPTION C

Option C for the secondary precious metals subcategory consists of all control and treatment requirements of Option B (in-process flow reduction, ammonia steam stripping, cyanide precipitation, and chemical precipitation, sedimentation and ion exchange) plus multimedia filtration technology added between the sedimentation and ion exchange operations at the end of the Option B treatment scheme (see Figure X-3 (page 2594). Multimedia filtration is used to remove suspended solids, including precipitates of priority metals, beyond the concentration attainable by gravity sedimentation. The filter suggested is of the gravity, mixed media type, although other filters, such as rapid sand filters or pressure filters, would perform satisfactorily.

INDUSTRY COST AND ENVIRONMENTAL BENEFITS

As one means of evaluating each technology option, EPA developed estimates to the pollutant removals and the compliance costs associated with each option. The methodologies are described below.

POLLUTANT REMOVAL ESTIMATES

A complete description of the methodology used to calculate the estimated pollutant removal, or benefit, achieved by the application of the various treatment options is presented in Section X of the General Development Document. The pollutant removal estimates have been revised from proposal based on comments and new data; however, the methodology for calculating pollutant removals was not changed. The data used for estimating removals are the same as those used to revise the compliance costs.

Sampling data collected during the field sampling program were used to characterize the major waste streams considered for regulation. At each sampled facility, the sampling data was production normalized for each unit operation (i.e., mass of pollutant generated per mass of product manufactured). This value, referred to as the raw waste, was used to estimate the mass of priority pollutants generated within the secondary precious metals subcategory. The pollutant removal estimates were calculated for each plant by first estimating the total mass

of each pollutant in the untreated wastewater. This was calculated by first multiplying the raw waste values by the corresponding production value for that stream and then summing these values for each pollutant for every stream generated by the plant.

The volume of wastewater discharged after the application of each treatment option was estimated for each operation at each plant by comparing the actual discharge to the regulatory flow. The smaller of the two values was selected and summed with the other plant flows. The mass of pollutant discharged was then estimated by multiplying the achievable concentration values attainable with the option (mq/1) by the estimated volume of process wastewater discharged by the subcategory. The mass of pollutant removed is the difference between the estimated mass of pollutant generated within the subcategory and the mass of pollutant discharged after application of the treatment option. The pollutant removal estimates for direct dischargers in the secondary precious metals subcategory are presented in Table X-2 (page 2574).

COMPLIANCE COSTS

In estimating subcategory-wide compliance costs, the first step was to develop a cost model, relating the total costs associated with installation and operation of wastewater treatment technologies to plant process wastewater discharge. EPA applied model on a per plant basis, a plant's costs (both capital, the and operating and maintenance) being determined by what treatment it has in place and by its individual process wastewater discharge (as discussed above, this flow is either the actual or the BAT regulatory flow, whichever is lesser). The final step to annualize the capital costs, and to sum the annualized was capital costs, and the operating and maintenance costs, yielding the cost of compliance for the subcategory. These costs were used in assessing economic achievability. Table X-3 (page 2575) shows the costs developed for promulgation for direct dischargers in the secondary precious metals subcategory. Compliance costs indirect dischargers are presented in Table XII-2 for (page 2620).

BAT OPTION SELECTION - PROPOSAL

EPA selected Option C for the proposed BAT, which included ammonia steam stripping and cyanide precipitation preliminary treatment, flow reduction for furnace and refinery wet air pollution control, and chemical precipitation, sedimentation and multimedia filtration end-of-pipe treatment. Although the compliance costs for the proposed BAT are not presented because the data on which they are based has been claimed confidential, it was determined that the pollutant removals justified the costs for this subcategory. Implementation of the proposed BAT was estimated to remove 34,580 kg of priority pollutants annually.

BAT OPTION SELECTION - PROMULGATION

Following proposal, EPA received comments and collected additional data showing a need for revising two regulatory flows (raw material granulation and spent cyanide stripping solutions). to consider additional treatment for high zinc concentrations, and to include compliance costs for two facilities not included in the proposal costs. Implementation of the promulgated BAT limitations would remove annually an estimated 34,650 kg of priority pollutants, which is 14 kg of priority pollutants greater than the estimated BPT removal. No additional ammonia or cyanide is removed at BAT.

EPA has added ion exchange end-of-pipe treatment to the BAT treatment scheme discussed at proposal. Ion exchange is an effective method for removing gold, platinum and palladium from wastewaters generated in the subcategory. EPA has determined that no additional costs will be generated in this subcategory by adding ion exchange because of the value of the precious metals recovered in the column.

EPA is promulgating BAT limitations for this subcategory based on ammonia steam stripping and cyanide precipitation preliminary treatment, flow reduction for furnace and refinery wet air pollution control, as well as for raw material granulation, and chemical precipitation, sedimentation, multimedia filtration and ion exchange end-of-pipe treatment. This treatment scheme is the same as that selected at proposal with the addition of flow reduction for one stream and ion exchange end-of-pipe treatment. The treatment performance concentrations, upon which the mass limitations are based, are equal to the values used to calculate the proposed mass limitations.

EPA is promulgating multimedia filtration as part of the BAT technology because this technology results in additional removal of priority metals. Filtration is also presently demonstrated at 25 plants throughout the nonferrous metals manufacturing category, including one plant in this subcategory. Filtration adds reliability to the treatment system by making it less susceptible to operator error and to sudden changes in raw wastewater flow and concentrations.

Based on comments received after proposal, the Agency believes it may be necessary for some facilities to use sulfide that order .to achieve the promulgated effluent polishing in limitations because of high zinc concentrations or complexing problems. Because the Agency believes that these situations will be the exception, rather than the rule, sulfide polishing is not specifically included as part of the model technology on which effluent limitations and performance standards are based. The Agency has, however, evaluated the cost associated with the use of sulfide polishing at secondary precious metals plants. After performing this evaluation, the Agency has concluded that sulfide polishing will result in a very small (less than 5 percent) incremental increase in wastewater treatment costs at a typical

secondary precious metals facility.

BAT treatment for the secondary precious metals subcategory is shown schematically in Figure X-3 (page 2594).

on information made available to the Agency after Based promulgation, the Agency has agreed to propose to make selected amendments to the regulation promulgated on September 20, 1985, FR 38276). These changes included: adding a new (50 (omitted) block for preliminary treatment; increasing building the production normalized flow for the palladium precipitation and filtration building block, reflecting the Agency's recalculation of the average flow for the building block and the incorporation new data; revising the manner of regulating gold, platinum, of and palladium to regulate the sum of these; and, expanding the refinery wet air pollution control building block to include separate flow allowances for acid and alkaline scrubbers at facilities with both types of scrubbers.

WASTEWATER DISCHARGE RATES

A BAT discharge rate was calculated for each subdivision based upon the flows of the existing plants, as determined from analysis of the data collection portfolios. The discharge rate is used with the achievable treatment concentration to determine BAT effluent limitations. Since the discharge rate may be different for each wastewater source, separate pro normalized discharge rates for each of the 14 wastewater production sources were determined and are summarized in Table X-4 (page 2576). The discharge rates are normalized on a production basis by relating amount of wastewater generated to the mass the of the intermediate product which is produced by the process associated with the waste stream in question. These production normalizing parameters (PNP) are also listed in Table X-4.

As discussed previously, the promulgated BAT wastewater discharge rate equals the promulgated BPT wastewater discharge rate for 11 of the 14 waste streams in the secondary precious metals subcategory. Based on the available data, the Agency determined that further flow reduction would not be feasible for these wastewater sources. Wastewater streams for which BAT discharge rates differ from BPT are discussed below.

FURNACE WET AIR POLLUTION CONTROL

The BAT wastewater discharge rate proposed and promulgated for furnace wet air pollution control is 4.5 liters per troy ounce of precious metal, including silver, incinerated or smelted. This rate is based on the value reported by the only discharging plant practicing recycle, and it is supported by the tact that two plants achieve zero discharge through 100 percent recycle.

RAW MATERIAL GRANULATION

The BAT wastewater discharge rate promulgated for raw material

granulation is 0.64 liters per troy ounce of precious metals in the granulated raw material. This rate is based on 90 percent recycle of the promulgated BPT rate of 6.34 liters per troy ounce. As discussed earlier in this section, 100 percent recycle is practiced by one plant for this subdivision, and recycle is considered feasible for the two non-recycling plants within this subdivision.

REFINERY WET AIR POLLUTION CONTROL

The BAT wastewater discharge rate proposed and promulgated for refinery wet air pollution control is 1.0 liter per troy ounce of precious metals, including silver, produced in the refinery. This rate is based on the average of all discharging plants which practice at least 90 percent recycle. These plants are shown in Table V-5 (page 2364) (1.75 liters per troy ounce, 0.19 liters per troy ounce, 0.234 liters per troy ounce, 14.2 liters per troy ounce, 0.072 liters per troy ounce, 0.67 liters per troy ounce, 0.7 liters per troy ounce, 2.3 liters per troy ounce, 0.04 liters per troy ounce, 0.174 liters per troy ounce, 0.036 liters per troy ounce, 1.665 liters per troy ounce, 1.41 liters per troy 1.1 liters per troy ounce, 0.06 liters per troy ounce, ounce, 4.64 liters per troy ounce, and 0.21 liters per troy ounce). Omitting the plant discharging 14.2 liters per troy ounce as being out to line with the water use at the majority of other dischargers, the average discharge rate equals 1.0 liter per troy This rate is supported by the average water use ounce. calculation presented in Section IX. That calculation shows average water use of 19.8 liters per troy ounce. Assuming 95 percent recycle is achievable (18 of 21 plants reported greater than 90 percent recycle for this stream), the BAT wastewater discharge rate of 1.0 liter per troy ounce is supported.

REGULATED POLLUTANT PARAMETERS

In implementing the terms of the Consent Agreement in <u>NRDC</u> v. <u>Train</u>, Op. Cit., and 33 U.S.C. \$1314(b)(2)(A and B) (1976), the Agency placed particular emphasis on the priority pollutants. The raw wastewater concentrations from individual operations and the subcategory as a whole were examined to select certain pollutants and pollutant parameters for consideration for limitation. This examination and evaluation was presented in Section VI. The Agency, however, has chosen not to regulate all 12 priority pollutants selected in this analysis.

The high cost associated with analysis for priority metal pollutants has prompted EPA to develop an alternative method for regulating and monitoring priority pollutant discharges from the nonferrous metals manufacturing category. Rather than developing specific effluent mass limitations and standards for each of the priority metals found in treatable concentrations in the raw wastewaters from a given subcategory, the Agency is promulgating effluent mass limitations only for those pollutants generated in the greatest quantities as shown by the pollutant removal analysis. The pollutants selected for specific limitation are listed below:

120. copper
121. cyanide
128. zinc
ammonia (as N)
combined metals (gold, platinum, and palladium)

By establishing limitations and standards for certain priority metal pollutants, dischargers will attain the same degree of control over priority metal pollutants as they would have been required to achieve had all the priority metal pollutants been directly limited.

This approach is technically justified since the treatable concentrations used for caustic precipitation and sedimentation technology are based on optimized treatment for concomitant multiple metals removal. Thus, even though metals have somewhat different theoretical solubilities, they will be removed at very nearly the same rate in a lime precipitation and sedimentation treatment system operated for multiple metals removal. Filtration as part of the technology basis is likewise justified because this technology removes metals non-preferentially.

The priority metal pollutants selected for specific limitation in the secondary precious metals subcategory to control the discharges of priority metal pollutants are copper and zinc. Cyanide, ammonia, and combined metals (gold, platinum, and palladium), are also selected for limitation since the methods used to control copper and zinc are not effective in the control of cyanide, ammonia, and combined metals (gold, platinum, and palladium).

The following toxic pollutants are excluded from limitation on the basis that they are effectively controlled by the limitations developed for copper and zinc:

114. antimony
115. arsenic
118. cadmium
119. chromium
122. lead
124. nickel
125. selenium
126. silver
127. thallium

EFFLUENT LIMITATIONS

The treatable concentrations achievable by application of the BAT technology (Option C) are summarized in Table VII-21 (page 248) of Vol. I. These treatable concentrations (both one-day maximum and monthly average) are multiplied by the BAT normalized discharge flows summarized in Table X-4 (page 2576) to calculate the mass of pollutants allowed to be discharged per mass of

product. The results of these calculations in milligrams of pollutant per troy ounce of product represent the promulgated BAT effluent limitations for the secondary precious metals subcategory. BAT effluent limitations based on Option C (ammonia steam stripping, cyanide precipitation, chemical precipitation, sedimentation, in-process flow reduction, multimedia filtration and ion exchange) are presented in Table X-5 (page 2578).

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	Number of Plants With Wastewater	Number of Plants Practicing Recycle	Range of Recycle Values (%)	
Furnace Scrubber	7	4	≥90 - 100	
Raw Material Granulation	4	1	.100	
Refinery Scrubber	29	21	75 - 100	

TABLE X-2

POLLUTANT REMOVAL ESTIMATES FOR DIRECT DISCHARGERS

		Option A	Option A	Option B	Option B	Option C	Option C
	Raw Waste	Discharged	Removed	Discharged		Discharged	
<u>Pollutant</u>	(kg/yr)	<u>(kg/yr)</u>	<u>(kg/yr)</u>	<u>(kg/yr)</u>	<u>(kg/yr)</u>	<u>(kg/yr)</u>	<u>(kg/yr)</u>
Antimony	1.23	0.90	0.33	0.90	0.33	0.90	0.33
Arsenic	0.40	0.30	0.10	0.30	0.11	0.30	0.11
Cadmium	3.22	0.34	2.88	0.28	2.94	0.18	3.05
Chromium (Tota	1) 3.88	0.29	3.59	0.29	3.59	0.24	3.63
Copper	293.11	6.49	286.62	4.78	288.33	3.22	289.89
Cyanide (Total	.) 17.76	11.48	6.29	11.46	6.31	11.46	6.31
Lead	23.27	1.34	21.93	0.99	22.28	0.66	22.61
Mercury	0.01	0.01	0.00	0.01	0.00	0.01	0.00
Nickel	89.31	8.25	81.06	6.09	83.22	1.81	87.50
Selenium	224.90	1.86	223.04	0.97	223.93	0.65	224.25
Silver	11.23	0.49	10.74	0.36	10.87	0.25	10.98
Thallium	0.38	0.28	0.10	0.28	0.10	0.28	0.10
Zinc 3	4,001.15	3.69	33,937.46	2.72	33,998.43	1.89 3	13,999.26
	4,669.85	35.72	34,634.14	29.43	34,639.81	21.85 3	4,648.02
Priority Pollu	tants						
Ammonia	615.38	120.98	494.39	120.80	494.57	120.80	494.57
Cobalt	0.93	0.33	0.61	0.18	0.75	0.12	0.81
Fluoride	3.01	2.28	0.73	2.26	0.74	2.26	0.74
Gold	7.86	0.08	7.78	0.05	7.81	0.05	7.81
Platinum	3.58	0.08	3.50	0.05	3.53	0.05	3.53
Palladium	3.58	0.08	3.50	0.05	3.53	0.05	3.53
TOTAL NONCONVENTIONA	634.34	123.83	510.51	123.40	510.94	123.34	511.00
				00.01	10 456 43	22.30	18,533.04
	.8,555.34		18,421.05		18,456.43	6.66	2.21
Oil & Grease	8.87	6.70	2.17	6.66	2.21	0.00	• 2•61
TOTAL 1 CONVENTIONALS	8,564.21	140.99	18,423.22	105.57	18,458.64	28.96	18,535.25
TOTAL 5 POLLUTANTS	53,868.40	300.54	53,567.87	258.40	53,609.39	174.15	53,694.27

OPTION A - Chemical precipitation, sedimentation, ion exchange, ammonia steam stripping, and cyanide precipitation.

Option B - Chemical precipitation, sedimentation, ion exchange, ammonia steam stripping, cyanide precipitation, and flow reduction

Option C - Chemical precipitation, sedimentation, ion exchange, ammonia steam stripping, cyanide precipitation, flow reduction and filtration

SECONDARY PRECIOUS METALS SUBCATEGORY

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Table X-3

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COST OF COMPLIANCE FOR DIRECT DISCHARGERS IN THE SECONDARY PRECIOUS METALS SUBCATEGORY

The costs for this subcategory cannot be presented here because the data on which they are based have been claimed to be confidential.

TABLE X-4

BAT WASTEWATER DISCHARGE RATES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

<u>Wastewater</u> <u>Stream</u>	BAT Normalized <u>Discharge</u> <u>Rates</u>	Production Normalizing Parameter
Furnace wet air pollution control	4.5 l/t.o.	Troy ounces of precious metals, including silver, incinerated or smelted
Raw material granulation	0.64 l/t.o.	Troy ounces of precious metals in the granulated raw material
Spent plating solutions	1.1 1/1	Liters of spent plating solution used as a raw material
Spent cyanide stripping solutions	3.7 l/t.o.	Troy ounces of gold stripped
Refinery wet air pollution control	1.0 l/t.o.	Troy ounces of precious metals produced in refinery, including silver
Gold solvent extraction raffinate	0.63 l/t.o.	Troy ounces of gold produced by solvent extraction
Gold spent electrolyte	0.0087 l/t.o.	Troy ounces of gold produced by electrolysis
Gold precipitation and filtration	4.4 l/t.o.	Troy ounces of gold precipitated
Platinum precipitation and filtration	5.2 l/t.o.	Troy ounces of platinum precipitated
	м. •	

SECONDARY PRECIOUS METALS SUBCATEGORY

SECT - X

TABLE X-4 (Continued)

BAT WASTEWATER DISCHARGE RATES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

Wastewater Stream	BAT Normalized Discharge Rates	Production Normalizing Parameter
Palladium precipitation and filtration	3.5 l/t.o.	Troy ounces of palladium precipitated
Other platinum group metals precipitation and filtration	5.2 l/t.o.	Troy ounces of other platinum group metals precipitation
Spent solution from PGC salt production	0.9 l/t.o.	Troy ounces of gold contained in PGC product
Equipment and floor wash	0.0 l/t.o.	Troy ounces of precious metals metals including silver produced
		in refinery

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NOTES 1/t.o. = liters per troy ounce 1/1 = liters per liter

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TABLE X-5

BAT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(a) <u>Furnace Wet</u> <u>Air</u> <u>Pollution</u> <u>Control</u> BAT

					•		
Pollutant pollutant		Maximum any one		Maximum for monthly average			
mg/t:	roy ounce	of precious	metals	incinerat	ted or smelted		
Antimony Arsenic Cadmium Chromium *Copper *Cyanide Lead Nickel Selenium		6.2 0.9 1.6 5.7 0.9 1.2 2.4	900 260 175	• • •	3.870 2.790 0.360 0.675 2.745 0.360 0.585 1.665		
Selenium Silver Thallium *Zinc *Ammonia *Combined Metals		1.3 6.3 4.5 599.9	3.690 1.305 6.300 4.590 599.900 1.350		1.665 0.540 2.745 1.890 263.700		

TABLE X-5 (Continued)

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BAT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(b) Raw Material Granulation BAT

	, •	11 A	· · ·
Pollutant or		Maximum for	
pollutant property	any one day	monthly average	
mg/troy ounce of p	recious metals in th	e granulated raw	material
Antimony	1.235	0.550	
Arsenic	0.890	0.397	
Cadmium	0.128	0.051	
Chromium	0.237	0.096	·, · · · ·
*Copper	0.819	0.390	the second
*Cyanide	0.128	0.051	·
Lead	0.179	0.083	н. 1814 -
Nickel	0.352	0.237	
Selenium	0.525	0.237	
Silver	0.186	0.077	and a start of the
Thallium	0.896	0.390	1
*Zinc	0.653	0.269	
*Ammonia	85.310	37.500	· · ·
*Combined metals	0.192		

TABLE X-5 (Continued)

BAT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

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(c) Spent Plating Solutions BAT

Pollutant or pollutant prop		operty	Max: any		imum thly	for average		
mg/l	of	spent	plating	solutions	used	as a	raw material	
Antimony Arsenic Cadmium Chromium *Copper *Cyanide Lead Nickel Selenium Silver Thallium *Zinc *Ammonia *Combined	met	als		1.930 1.390 0.200 0.370 1.280 0.200 0.280 0.550 0.820 0.290 1.400 1.020 L33.300 0.300	· ·		0.860 0.620 0.080 0.150 0.610 0.080 0.130 0.370 0.370 0.370 0.120 0.610 0.420 58.600	

TABLE X-5 (Continued)

BAT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(d) Spent Cyanide Stripping Solutions BAT

Pollutant		Maximum for	Maximum for	
pollutant	property	any one day	monthly average	
	mg	/troy ounce of gol	d tripped.	·
Antimony		7.141	3.182	·
Arsenic		5.143	2.294	
Cadmium		0.740	0.296	
Chromium		1.369	0.555	S. 14.
*Copper		4.736	2.257	· • ·
*Cyanide	÷	0.740	0.296	
Lead		1.036	0.481	
Nickel		2.035	1.369	
Selenium		3.034	1.369	۰.
Silver	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	1.073	0.444	
Thallium		5.180	2.257	
*Zinc	*	3.774	1.554	. ,
*Ammonia	1997 - 1997 -	493.200	216.800	5 A. 19
*Combined	metals	1.110		· •

TABLE X-5 (Continued)

BAT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(e) <u>Refinery Wet</u> <u>Air</u> <u>Pollution</u> <u>Control¹</u> BAT

Pollutant or pollutant property				Maximum for any one day	Maximum for monthly average		
mg/	troy	ounce	of	precious metals	s produced in refinery		
Antimony Arsenic Cadmium Chromium *Copper *Cyanide Lead Nickel Selenium Silver Thallium *Zinc				1.930 1.390 0.200 0.370 1.280 0.200 0.280 0.280 0.550 0.820 0.290 1.400 1.020	0.860 0.620 0.080 0.150 0.610 0.080 0.130 0.370 0.370 0.120 0.610 0.420		
*Ammonia *Combined	meta	als		133.300 0.300	58.600 		

*Regulated Pollutant

¹This allowance applies to either acid or alkaline wet air pollution control scrubbers. If both acid and alkaline wet air pollution control scrubbers are present in a particular facility the same allowance applies to each.

TABLE X-5 (Continued)

BAT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(f) Gold Solvent Extraction Raffinate and Wash Water BAT

									· · · · ·
Pollutant pollutant		pertv	;	Maximu any on			imum thly	for average	
<u> </u>	T 1	2 N. 19 4	· ·	1	-	•		-	
		r	·			. ,	-		·····
mg/	troy	ounce	of	gold p	roduced	by so	lvent	extrac	tion
Antimony				1	.216	1		0.542	
Arsenic				. 0	.876			0.391	
Cadmium				1	.126			0.050	
Chromium					.233		-	0.095	•
*Copper					.806			0.384	
*Cyanide					.126			0.050	-
Lead					.176	,	1	0.082	•
Nickel					.347		`	0.233	
Selenium					.517		· · · ·	0.233	
Silver				-	.183			0.076	
Thallium					.882			0.384	
*Zinc					.643			0.265	
*Ammonia				-	.980		÷	36.920	
*Combined	meta	als			.189		,		•
				-					

TABLE X-5 (Continued)

BAT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(g) Gold Spent Electrolyte BAT

Pollutant pollutant	or property	Maximum for any one day	Maximum for monthly average	
	mg/troy ounce	of gold produce	d by electrolysis	
Antimony Arsenic Cadmium Chromium *Copper *Cyanide Lead Nickel Selenium	. ,	0.017 0.012 0.002 0.003 0.011 0.002 0.002 0.005 0.007	0.007 0.005 0.000 0.001 0.005 0.000 0.001 0.003 0.003	
Silver Thallium *Zinc *Ammonia *Combined	metals	0.003 0.012 0.009 1.160 0.003	0.001 0.005 0.004 0.510	

SECONDARY PRECIOUS METALS SUBCATEGORY

TABLE X-5 (Continued)

BAT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(h) Gold Precipitation and Filtration BAT

Pollutant pollutant		Maximum for any one day	Maximum for monthly average
	mg/troy	ounce of gold	precipitated
Antimony Arsenic Cadmium Chromium *Copper *Cyanide Lead Nickel Selenium Silver Thallium *Zinc *Ammonia *Combined	motals	8.492 6.116 0.880 1.628 5.632 0.880 1.232 2.420 3.608 1.276 6.160 4.488 586.500 1.320	3.784 2.728 0.352 0.660 2.684 0.352 0.572 1.628 1.628 1.628 2.684 1.848 257.800

TABLE X-5 (Continued)

BAT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

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(i) Platinum Precipitation and Filtration BAT

						÷		
Pollutant	or	Maxi	imum	for	Maximum	for	,	<u>.</u>
pollutant	property	any	one	day		average	-	1. 1. 1.
	~ ~ _	• ·		·····				
	mg/troy	ounce	of g	platinum	precipit	tated		
Antimony			10.0	040		4.472		
Arsenic			7.2	228		3.224		+ 1
Cadmium			1.0	40		0.416		ана. Стала стала ста Стала стала стал
Chromium			1.9	24		0.780	. 1	ta a stational A stational stational
*Copper			6.6	556		3.172	,	1.5.1
*Cyanide		1	1.0	40		0.416		
Lead			1.4	56		0.676		
Nickel			2.8	60		1.924		1000
Selenium			4.2	264		1.924	•	Star De L
Silver			1.5	608		0.624		
Thallium			7.2	280	н.	3.172		i di pi
*Zinc			5.3	04	•	2.184		e na le
*Ammonia		6	93.2	00 .	3	304.700		·
*Combined	metals		1.5					

TABLE X-5 (Continued)

BAT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(j) Palladium Precipitation and Filtration BAT

Pollutant or pollutant propertyMaximum for any one dayMaximum for monthly averagemg/troy ounce of palladium precipitatedAntimony11.5805.160Arsenic8.3403.720Cadmium1.2000.480Chromium2.2000.900*Copper7.6803.660*Cyanide1.2000.480Lead1.6800.780Nickel3.3002.220Silver1.7400.720Thallium8.4003.660*Zinc6.1202.520*Ammonia799.800351.600*Combined metals1.800					
Antimony11.5805.160Arsenic8.3403.720Cadmium1.2000.480Chromium2.2000.900*Copper7.6803.660*Cyanide1.2000.480Lead1.6800.780Nickel3.3002.220Selenium4.4202.220Silver1.7400.720Thallium8.4003.660*Zinc6.1202.520*Ammonia799.800351.600			10 M M		<u>, ., ., .</u> ,
Arsenic8.3403.720Cadmium1.2000.480Chromium2.2000.900*Copper7.6803.660*Cyanide1.2000.480Lead1.6800.780Nickel3.3002.220Selenium4.4202.220Silver1.7400.720Thallium8.4003.660*Zinc6.1202.520*Ammonia799.800351.600		mg/troy	ounce of palladi	um precipitated	
Cadmium1.2000.480Chromium2.2000.900*Copper7.6803.660*Cyanide1.2000.480Lead1.6800.780Nickel3.3002.220Selenium4.4202.220Silver1.7400.720Thallium8.4003.660*Zinc6.1202.520*Ammonia799.800351.600		• •			
Chromium2.2000.900*Copper7.6803.660*Cyanide1.2000.480Lead1.6800.780Nickel3.3002.220Selenium4.4202.220Silver1.7400.720Thallium8.4003.660*Zinc6.1202.520*Ammonia799.800351.600	Arsenic		8.340	3.720	
*Copper7.6803.660*Cyanide1.2000.480Lead1.6800.780Nickel3.3002.220Selenium4.4202.220Silver1.7400.720Thallium8.4003.660*Zinc6.1202.520*Ammonia799.800351.600	Cadmium		1.200	0.480	1 - E
*Cyanide1.2000.480Lead1.6800.780Nickel3.3002.220Selenium4.4202.220Silver1.7400.720Thallium8.4003.660*Zinc6.1202.520*Ammonia799.800351.600	Chromium	·	2.200	0.900	
*Cyanide1.2000.480Lead1.6800.780Nickel3.3002.220Selenium4.4202.220Silver1.7400.720Thallium8.4003.660*Zinc6.1202.520*Ammonia799.800351.600	*Copper		7.680	3.660	· · ·
Lead1.6800.780Nickel3.3002.220Selenium4.4202.220Silver1.7400.720Thallium8.4003.660*Zinc6.1202.520*Ammonia799.800351.600			1.200	0.480	. * .
Nickel3.3002.220Selenium4.4202.220Silver1.7400.720Thallium8.4003.660*Zinc6.1202.520*Ammonia799.800351.600			1.680	0.780	
Selenium4.4202.220Silver1.7400.720Thallium8.4003.660*Zinc6.1202.520*Ammonia799.800351.600			3.300	2.220	
Silver1.7400.720Thallium8.4003.660*Zinc6.1202.520*Ammonia799.800351.600	/		4.420	2.220	
Thallium8.4003.660*Zinc6.1202.520*Ammonia799.800351.600				0.720	
*Zinc 6.120 2.520 *Ammonia 799.800 351.600			· · · ·		,
*Ammonia 799.800 351.600		1			
		metals		~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~~	· · · ·

SECONDARY PRECIOUS METALS SUBCATEGORY

TABLE X-5 (Continued)

BAT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(k) Other Platinum Group Metals Precipitation and Filtration BAT

Pollutant or pollutant property	Maximum for any one day	Maximum for monthly average
		• • • • • • • • • • • • • • • • • • •
mg/troy ounce of	other platinum	group metals precipitated
Antimony	10.040	4.472
Arsenic	7.228	3.224
Cadmium	1.040	0.416
Chromium	1.924	0,780
*Copper	6.656	3.172
*Cyanide	1.040	0.416
Lead	1.456	0.676
Nickel	2.860	1.924
Selenium	4.264	1.924
Silver	1.508	0.624
Thallium	7.280	3.172
*Zinc	5.304	2.184
*Ammonia	693.200	304.700
*Combined metals	1.560	

SECONDARY PRECIOUS METALS SUBCATEGORY

TABLE X-5 (Continued)

BAT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(1) Spent Solution from PGC Salt Production BAT

				· · · · ·				1 Mar 11	
Pollutant	or			ximum		Maxim			
pollutant	proper	сy	an	y one	day	month	ly	average	
	mg/troy	ounce	of	gold	containe	ed in	PGC	product	· · · · · ·
Antimony				1.	737			0.774	
Arsenic		•		1.	251			0.558	· · ·
Cadmium	p.			0.	180			0.072	
Chromium					333			0.135	
*Copper				1.	152		• .	0.549	· · · ·
*Cyanide		· •			180			0.072	• •
Lead					252			0.117	· • * .
Nickel				0.	495			0.333	
Selenium				0.	738			0.333	
Silver				0.	261			0.108	
Thallium	.5	•			260			0.549	
*Zinc		1 a.			918			0.378	
*Ammonia	· .			120.	-			52.740	
*Combined	metals				270				
									5 T. 1

TABLE X-5 (Continued)

BAT MASS LIMITATIONS FOR THE SECONDARY PRÉCIOUS METALS SUBCATEGORY

(n) Equipment and Floor Wash BAT

Pollutant or pollutant property	Maximum for any one day	Maximum for monthly average
mg/troy ounce of	precious metals	produced in refinery
Antimony Arsenic Cadmium Chromium *Copper *Cyanide Lead Nickel Selenium Silver Thallium *Zinc *Ammonia *Combined metals	0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000	0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000

SECONDARY, PRECIOUS METALS SUBCATEGORY

TABLE X-5 (Continued)

BAT MASS LIMITATIONS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(n) Preliminary Treatment BAT

Pollutant	or	Maximum for	Maximum for	1 - 4 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -
Pollutant		Any One Day	Monthly Average	· · · · · · · · · · · · · · · · · · ·

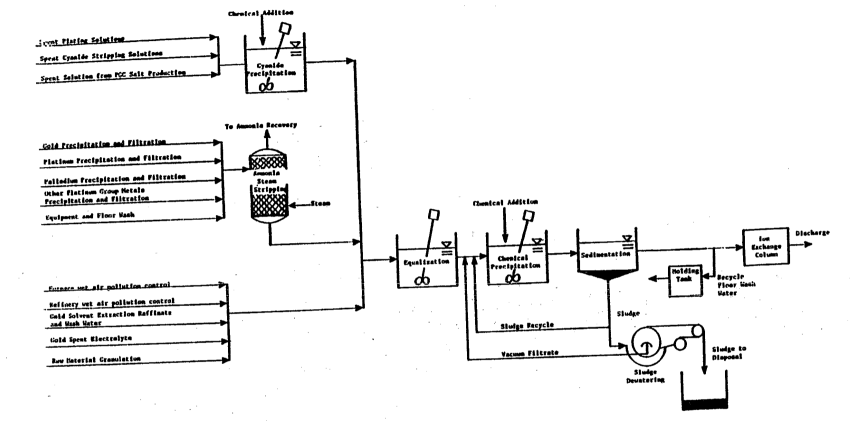
mg/troy ounce of total precious metals produced through this operation

Antimony	96.500	43.000
Arsenic	69.500	31.000
Cadmium	10.000	4.000
Chromium	18.500	7.500
Copper	64.000	30.500
Cyanide (total)	10.000	4.000
Lead	14.000	1.500
Nickel	27.500	18.500
Selenium	41.000	18.500
Silver	14.500	6.000
Thallium	70.000	30.500
Zinc	51.000	21.000
Combined metals	15.000	
Ammonia (as N	6665.000	2930.000
munonita (ab m		

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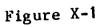
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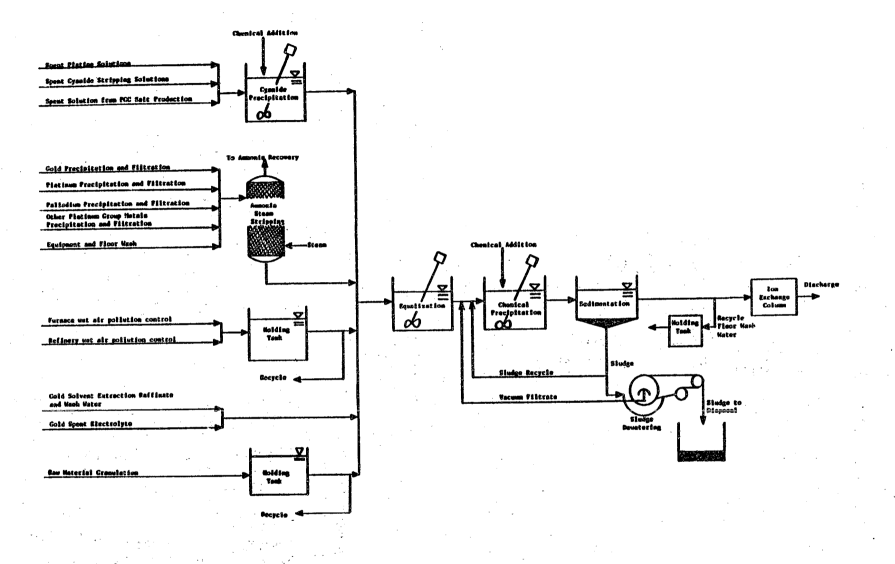




BAT TREATMENT SCHEME FOR OPTION A









BAT TREATMENT SCHEME FOR OPTION B

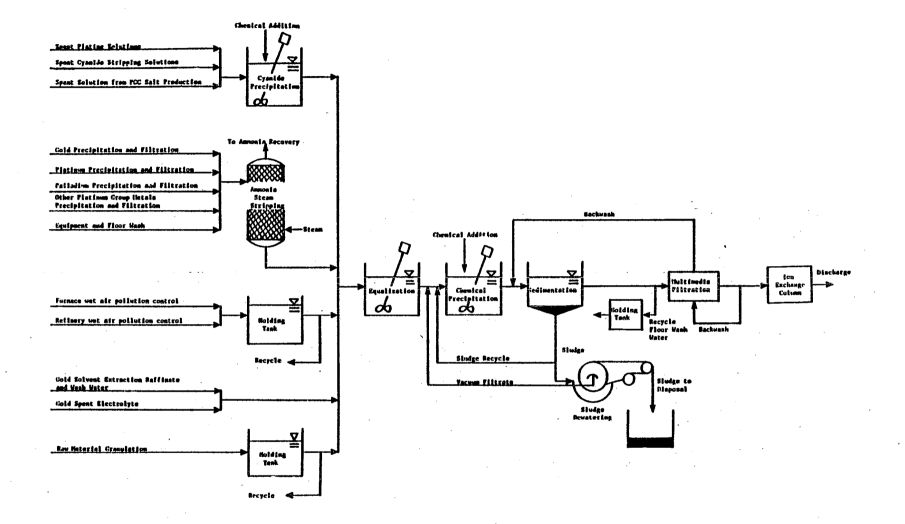
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SECONDARY PRECIOUS METALS SUBCATEGORY





BAT TREATMENT SCHEME FOR OPTION C

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SECONDARY PRECIOUS METALS SUBCATEGORY SECT н

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SECTION XI

NEW SOURCE PERFORMANCE STANDARDS

section describes the technologies for treatment of This wastewater from new sources and presents mass discharge standards regulated pollutants for NSPS in the secondary precious for subcategory, based on the selected treatment technology. metals plants have the opportunity to design the best and most New wastewater production processes and treatment efficient technologies without facing the added costs and restrictions encountered in retrofitting an existing plant. Therefore, EPA considered the best demonstrated process changes, in-plant controls, and end-of-pipe treatment technologies which reduce pollution to the maximum extent feasible.

TECHNICAL APPROACH TO NSPS

New source performance standards are equivalent to the best available technology (BAT) selected for currently existing secondary precious metals plants. This result is a consequence of careful review by the Agency of a wide range of technical options for new source treatment systems. Additionally, there was nothing found to indicate that the wastewater flows and characteristics of new plants would not be similar to those from existing plants since the processes used by new sources are not expected to differ from those used at existing sources. BAT production normalized discharge rates, which are based on the best existing practices of the subcategory, can also be applied to new sources. These rates are presented in Table XI-1 (page 2599).

EPA proposed that furnace wet air pollution control be given a zero discharge allowance for new sources, based on dry air pollution control techniques. After reviewing comments received on the proposal, EPA has decided to revise the allowance to be equal to the allowance for furnace wet air pollution control at BAT.

Treatment technologies considered for the NSPS options are identical to the treatment technologies considered for the BAT options. These options are:

OPTION A

- Preliminary treatment with cyanide precipitation (where required)
- Preliminary treatment with ammonia steam stripping (where required)
- o Chemical precipitation and sedimentation
- o Ion exchange end-of-pipe treatment

OPTION B

- Preliminary treatment with cyanide precipitation (where required)
- Preliminary treatment with ammonia steam stripping (where required)
- o Chemical precipitation and sedimentation
- o In-process flow reduction of refinery scrubber liquor and raw material granulation water
- o Ion exchange end-of-pipe treatment

OPTION C

- Preliminary treatment with cyanide precipitation (where required)
- Preliminary treatment with ammonia steam stripping (where required)
- o Chemical precipitation and sedimentation
- o In-process flow reduction to refinery scrubber liquor and raw material granulation water
- o Multimedia filtration
- o Ion exchange end-of-pipe treatment

NSPS OPTION SELECTION - PROPOSAL

EPA proposed that the best available demonstrated technology for secondary precious metals subcategory be equivalent to the the proposed Option C with additional flow reduction, which included ammonia steam stripping and cyanide precipitation preliminary treatment, dry air pollution control for furnace emissions, flow reduction for refinery wet air pollution control, and end-of-pipe treatment consisting of chemical precipitation, sedimentation, and multimedia filtration. Zero discharge for furnace air pollution control was based on dry scrubbing, which is demonstrated at 11 of 16 plants which control off-gases from the furnace. Except for furnace air pollution control, EPA proposed that NSPS wastewater discharge rates be equivalent to the proposed BAT rates. Flow reduction beyond BAT for the other 13 streams in this subcategory is not demonstrated.

NSPS OPTION SELECTION - PROMULGATION

EPA is promulgating best available demonstrated technology for secondary precious metals subcategory equivalent to Option C the which includes ammonia steam stripping and cyanide precipitation preliminary treatment, flow reduction for furnace and refinery wet air pollution control and raw material granulation, and endof-pipe treatment consisting of. chemical precipitation, multimedia filtration and ion exchange. sedimentation, As discussed under BAT option selection-promulgation, ion exchange end-of-pipe treatment is necessary to reduce gold, platinum and palladium concentrations. Regulatory flows for furnace wet air pollution control, raw material granulation and spent cyanide stripping solutions have been revised since proposal, based on comments received by the Agency on the proposed flow rates. The promulgated NSPS option is equivalent to the promulgated BAT option.

REGULATED POLLUTANT PARAMETERS

The Agency has no reason to believe that the pollutants that will be found in treatable concentrations in processes within new sources will be any different than with existing sources. Accordingly, pollutants and pollutant parameters selected for limitation under NSPS, in accordance with the rationale of Sections VI and X, are identical to those selected for BAT. The conventional pollutant parameters TSS and pH are also selected for limitation.

NEW SOURCE PERFORMANCE STANDARDS

The promulgated NSPS discharge flows for each wastewater source are the same as the promulgated discharge rates for BAT, except for furnace wet air pollution control, and are shown in Table XI-1 (page 2599). The mass of pollutant allowed to be discharged per mass of product is calculated by multiplying the appropriate treatable concentration (mg/l) by the production normalized wastewater discharge flows (liter/troy ounce). The treatable concentrations are listed in Table VII-21 (page 248) of Vol. I. The results of these calculations are the production-based new source performance standards. These standards are presented in Tables XI-2 (page 2600).

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TABLE XI-1

NSPS WASTEWATER DISCHARGE RATES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

<u>Wastewater</u> <u>Stream</u>	BAT Normalized <u>Discharge</u> <u>Rates</u>	Production Normalizing Parameter
Furnace wet air pollution control	4.5 l/t.o.	Troy ounces of precious metals, including silver, incinerated or smelted
Raw material granulation	0.64 l/t.o.	Troy ounces of precious metals in the granulated raw material
Spent plating solutions	1.1 1/1	Liters of spent plating solution used as a raw material
Spent cyanide stripping solutions	3.7 1/t.o.	Troy ounces of gold stripped
Refinery wet air pollution control	1.0 l/t.o.	Troy ounces of precious metals produced in refinery, including silver
Gold solvent extraction raffinate	0.63 l/t.o.	Troy ounces of gold produced by solvent extraction
Gold spent electrolyte	0.0087 l/t.o.	Troy ounces of gold produced by electrolysis
Gold precipitation and filtration	4.4 l/t.o.	Troy ounces of gold precipitated
Platinum precipitation and filtration	5.2 l/t.o.	Troy ounces of platinum precipitated

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NSPS WASTEWATER DISCHARGE RATES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

Wastewater Stream	BAT Normalized <u>Discharge</u> <u>Rates</u>	Production Normalizing Parameter
Palladium precipitation and filtration	3.5 l/t.o.	Troy ounces of palladium precipitated
Other platinum group metals precipitation and filtration	5.2 l/t.o.	Troy ounces of other platinum group metals precipitation
Spent solution from PGC salt production	0.9 l/t.o.	Troy ounces of gold contained in PGC product
Equipment and floor wash	0.0 l/t.o.	Troy ounces of precious metals metals including silver produced in refinery

NOTES 1/t.o. = liters per troy ounce 1/1 = liters per liter

IX

SECONDARY PRECIOUS METALS SUBCATEGORY

TABLE XI-2

NSPS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(a) <u>Furnace Wet Air</u> <u>Pollution</u> <u>Control</u> NSPS

Pollutant or	Maximum for	Maximum	for	• .
pollutant property	any one day	monthly	average	• .

mg/troy ound	e of precious metals	incinerated or smelted
Antimony	8 685	3.870
Arsenic	6.255	2.790
Cadmium	0.900	0.360
Chromium	1.665	0.675
*Copper	5.760	2.745
*Cyanide	0.900	0.360
Lead	1.260	0.585
Nickel	2.475	1.665
Selenium	3.690	1.665
Silver	1.305	1.665
Thallium	6.300	0.540
*Zinc	4.590	2.745
*Ammonia	599.900	1.890
*Combined metals	1.350	263.700
*TSS	67.500	54.000
*pH Within the	range of 7.5 to 10.0	at all times

SECT - XI

TABLE XI-2 (Continued)

NSPS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(b) Raw Material Granulation NSPS

Pollutant or pollutant property	Maximum for any one day	Maximum for monthly average	. <u></u>
			······
mg/troy ounce of	precious metals in	the granulated raw	material
Antimony	1.235	0.550	
Arsenic	0.890	0.397	
Cadmium	0.128	0.051	
Chromium	0.237	0.096	•
*Copper	0.819	0.390	•••
*Cyanide	0.128	0.051	
Lead	0.179	0.083	
Nickel	0.352	0.237	
Selenium	0.525	0.237	
Silver	0.186	0.077	
Thallium	0.896	0.390	
*Zinc	0.653	0.269	
*Ammonia	85.310	37.500	
*Combined metals	0.192		
*TSS	9.600	7.680	
	range of 7.5 to 10.	•	
-	· · ·		

NSPS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(c) Spent Plating Solutions NSPS

Pollutant or pollutant property	Maximum for any one day	Maximum monthly	for average
mg/liter of spent	plating solutions	used as	a raw material
Antimony	1.930		0.860
Arsenic	1.390		0.620
Cadmium	0.200		0.080
Chromium	0.370		0.150
*Copper	1.280		0.610
*Cyanide	0.200		0.080
Lead	0.280		0.130
Nickel	0.550		0.370
Selenium	0.820		0.370
Silver	0.290		0.120
Thallium	1.400		0.610
*Zinc	1.020		0.420

Ammon	La			_	L33.3	300	•			58.600	
Combined metals					0.3	300		•			
trss					15.0	000		•	•	12.000	
tpH	Within	the	range	of	7.5	to	10.0	at	all	times	

NSPS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(d) Spent Cyanide Stripping Solutions NSPS

Pollutant		pert		aximum ny one			imum thly	for average	3
		,		- · · ·				·····	· · ·
	•	1	mg/troy	ounce	of go	ld str	ippe	1	
Antimony			* .	7.	141	•		3.182	
Arsenic		,		5.	143			2.294	· · ·
Cadmium				0.	740			0.296	·
Chromium				1.	369 🖂 🗸	• •		0.555	
*Copper				4.	736			2.257	
*Cyanide		. •		0.	740			0.296	
Lead	6.5			1.0	036			0.481	
Nickel			а. 1	2.0	035			1.369	
Selenium				3.0	034			1.369	
Silver				1.0	073			0.444	
Thallium		1.51.5	ŧ	5.1	180			2.257	
*Zinc			1. A.	3.1	774	s		1.554	
*Ammonia				493.2	200			216.800	
*Combined	l meta	als		1.1	L10				. ;•
*TSS				55.5	- · · ·	¢		44.400	
*pH Wi	thin	the	range	of 7.5	to 10	.0 at	all t	imes	. · · ·

NSPS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(e) <u>Refinery Wet</u> <u>Air</u> <u>Pollution</u> <u>Control¹</u> NSPS

			•		1
Pollutant	or	Maximum fo	or Max	kimum .	for
pollutant	property	any one da	ay mor	nthly a	average

n	ng/troy	ounce	of	precious	metals	produced	in re	finery
Arsenic Cadmium Chromiu *Copper *Cyanid Lead Nickel Seleniu Silver Thalliu *Zinc *Ammoni *Combin *TSS	im le im im im la hed meta	als		1.3 0.2 0.3 1.2 0.2 0.2 0.2 0.2 0.2 1.4 1.0 133.3 0.3 15.0	390 200 370 280 200 280 350 320 290 300 300 300 300	- · ·	0.620 0.080 0.150 0.610 0.080 0.130 0.370 0.370 0.120 0.610 0.420 58.600 	
*pH	Within	the r	ange	e or 7.5	to 10.0	at all t	imes	

*Regulated Pollutant

¹This allowance applies to either acid or alkaline wet air pollution control scrubbers. If both acid and alkaline wet air pollution control scrubbers are present in a particular facility the same allowance applies to each.

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - XI

法法法法法法法法 TABLE XI-2 (Continued)

新学家 (1) 胡椒香 (1) A 小学会 (1) A 把 A 种 NSPS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

Υģ. j (f) Gold Solvent Extraction Raffinate and Wash Water NSPS

Pollutant	or	Maximum	for	Maximum	for	
pollutant	property	any one	day	monthly	average	,
· · ·	•					

		• •					1.12.2
,	mg/troy	ounce o	f gold	produced	by solver	nt extract	ion
Antimo	ony			1.216		0.542	۹
Arseni	.c •			0.876		0.391	
Cadmiu	ım			0.126	*	0.050	- 1
Chromi	.um			0.233	•	0.095	
*Coppe	er '	and a second second	1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -	0.806		0.384	÷
*Cyani	.de 🖓			0.126		0.050	
Lead				0.176		0.082	
Nickel	•			0.347		0.233	
Seleni	.um	2 8 2 - 6 8	•	0.517		0.233	
Silver	• • •			0.183		0.076	
Thalli	um 🚲	18.20		0882		0.384	
*Zinc				0.643		0.265	
*Ammon	ia		8	83.980		36.920	
*Combi	ned meta	als		0.189			
*TSS			2	9.450	· · ·	7.560	$L \in \mathbb{N}^{k}$
*pH	Within	the rand	ge of 🕻	7.5 to 10.			K
						5 6 8 F 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	· •••

TABLE XI-2 (Continued)

NSPS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(g) Gold Spent Electrolyte NSPS

Pollutant pollutant		Maximum for any one day	Maximu m for monthl y avera ge
<u></u>	mg/troy our	ce of gold produc	ced by electrolysis
Antimony		0.017	0.007
Arsenic		0.012	0.005
Cadmium		0.002	0.000
Chromium		0.003	0.001
*Copper		0.011	0.005
*Cyanide		0.002	0.000
Lead		0.002	0.001
Nickel		0.005	0.003
Selenium		0.007	0.003
Silver		0.003	0.001
Thallium		0.012	0.005
*Zinc		0.009	0.004
*Ammonia		1.160	0.510
*Combined	metals	0.003	
*TSS		0.131	0.104
	thin the ran	ge of 7.5 to 10.0) at all times

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - XI

TABLE XI-2 (Continued)

NSPS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(h) Gold Precipitation and Filtration NSPS

	Maximum	for
pollutant property any one day	monthly	average

mg/troy c	ounce of gold prec	ipitated
Antimony	8.492	3.784
Arsenic	6.116	2.728
Cadmium	0.880	0.352
Chromium	1.628	0.660
*Copper	5.632	2.684
*Cyanide	0.880	0.352
Lead Nickel	1.232	0.572
Selenium	2.420 3.608	1.628
Silver	1.276	1.628 0.528
Thallium	6.160	2.684
*Zinc	4.488	1.848
*Ammonia	586.500	257.800
*Combined metals	1.320	
*TSS	66.000	52.800
*pH Within the range	of 7.5 to 10.0 at	all times

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - XI

TABLE XI-2 (Continued)

NSPS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(i) Platinum Precipitation and Filtration NSPS

			· · · · · · · · · · · · · · · · · · ·
Pollutant	or	Maximum for	Maximum for
pollutant	property	any one day	monthly average
	mg/troy	ounce of platinum	precipitated
Antimony		10.040	4.472
Arsenic		7.228	3.224
Cadmium		1.040	0.416
Chromium		1.924	0.780
*Copper		6.656	3.172
*Cyanide		1.040	0.416
Lead		1.456	0.676
Nickel		2.860	1.924
Selenium		4.264	1.924
Silver		1.508	0.624
Thallium		7.280	3.172
*Zinc		5.304	2.184
*Ammonia		693,200	304.700
*Combined	motale	1.560	
*TSS	MCCATD	78.000	62,400
	thin the rat	nge of 7.5 to 10.0	

NSPS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(j) Palladium Precipitation and Filtration NSPS

Pollutant pollutant	or property	Maximum for any one day	Maximum for monthly average	······
<u></u>	mg/troy	ounce of palladiu	ım pre ci pitat e d	
Antimony Arsenic Cadmium Chromium *Copper *Cyanide Lead Nickel Selenium Silver Thallium *Zinc *Ammonia *Combined *TSS		11.580 8.340 1.200 2.220 7.680 1.200 1.680 3.300 4.920 1.740 8.400 6.120 799.800 1.800 90.000	5.160 3.720 0.480 0.900 3.660 0.480 0.780 2.220 2.220 2.220 0.720 3.660 2.520 351.600 72.000	
*pH Wi	thin the ra	nge of 7.5 to 10.0) at all times	

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - XI

TABLE XI-2 (Continued)

NSPS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(k) Other Platinum Group Metals Precipitation and Filtration NSPS

Pollutant or			Maximum for				Maximum for						
pollut	lutant property			any one day				monthly average			age		
			ı		········								·
mg	/troy	ounce	of	other	plat	inum	gro	oup	meta	als	pre	cipi	tated
Antimo					10.0	40	•			4	. 47	2	
Arseni	С				7.2	28				3	.22	4	
Cadmiu	m				1.0	40				0	.41	.6	
Chromi	um				1.9	24				0	.78	0	
Copper					6.6	56		•		3	.17	2	
*Cyani	de				1.0	40				0	.41	.6	
Lead					1.4	56				0	.67	6	
Nickel					2.8	60				1	.92	4	
Seleni	um				4.2	64				· 1	.92	4	
Silver					1.5	08				0	.62	4	
Thalli	um				7.2	80				3	.17	2	
*Zinc					5.3	04				2	.18	4	1
*Ammon	ia	•			693.2	00				30,4	.70	0	
*Combi	ned me	etals			1.5	60						-	
*TSS					78.0	00				62	.40	0	
*pH	Withi	.n the	rar	nge of	7.5	to 10	.0	at	all	tim	es		
				-									

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - XI

TABLE XI-2 (Continued)

NSPS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(1) Spent Solution from PGC Salt Production NSPS

Pollutant			Maximum		Maximu	
pollutant	proper	cy a	any one	aay	monthl	y average
	mg/troy	ounce o	of gold	contain	ed in P	GC product
Antimony	· .		1.7	737		0.774
Arsenic			1.2	251	5 	0.558
Cadmium			0.1	180		0.072
Chromium			0.3	333	_	0.135
*Copper			1.1	152		0.549
*Cyanide			0.3	L80 .		0.072
Lead	· .	•	0.2	252		0.117
Nickel	•	•	0.4			0.333
Selenium	× .		0.7	738	۰. ۱	0.333
Silver			0.2	261		0.108
Thallium			1.2	260		0.549
*Zinc		•	. 0.9	918		0.378
*Ammonia			120.0	000		52.740
*Combined	metals	н. 1	0.2	270		
*TSS			13.5			10.800
*pH Wi	thin the	e range	of 7.5	to 10.0	at all	times

TABLE XI-2 (Continued)

NSPS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(m) Equipment and Floor Wash NSPS

Pollutant or pollutant property	Maximum for any one day	Maximum for monthly average
mg/troy ounce of	precious metals	produced in refinery
Antimony	0.000	0.000
Arsenic	0.000	0.000
Cadmium	0.000	0.000
Cl romium	0.000	0.000
*Copper	0.000	0.000
*Cyanide	0.000	0.000
Lead	0.000	0.000
Nickel	0.000	0.000
Selenium	0.000	0.000
Silver	0.000	0.000
Thallium	0.000	0.000
*Zinc	0000	0.000
*Ammonia	0.000	0.000
*Combined metals	0.000	
*TSS	0.000	0.000
*pH W_thin the range	of 7.5 to 10.0	at all times

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - XI

TABLE XI-2 (Continued)

NSPS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(n) Preliminary Treatment NSPS

Pollutant or	Maximum for	Maximum for	
Pollutant Property	Any One Day	Monthly Average	а. •

mg/troy, ounce of total precious metals produced through this operation

Antimony	96.500	43.000
Arsenic	69.500	31.000
Cadmium	10.000	4.000
Chromium	18.500	7.500
Copper	64.000	30.500
Cyanide (total)	10.000	4.000
Lead	14.000	6.500
Nickel	27.500	18.500
Selenium	41.000	18.500
Silver	14.500	6.000
Thallium	70.000	30.500
Zinc	51.000	21.000
Combined metals	15.000	
Ammonia (as N)	6665.000	2930.000
Total suspended	750.000	600.000
solids		
рН	Within the rang	e of 7.5 to 10.0
		at all times

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SECTION XII

PRETREATMENT STANDARDS

This section describes the control and treatment technologies for pretreatment of process wastewaters from existing sources and new sources in the secondary precious metals subcategory. Pretreatment standards for regulated pollutants are presented based on the selected control and treatment technology.

TECHNICAL APPROACH TO PRETREATMENT

Before proposing and promulgating pretreatment standards, the Agency examines whether the pollutants discharged by the industry pass through the POTW or interfere with the POTW operation or its sludge disposal practices. In determining whether chosen pollutants pass through a well-operated POTW achieving secondary treatment, the Agency compares the percentage of a pollutant removed by POTW with the percentage removed by direct dischargers applying the best available technology economically achievable. A pollutant is deemed to pass through the POTW when the average percentage removed nationwide by well-operated POTW meeting secondary treatment requirements, is less than the percentage removed by direct dischargers complying with BAT effluent limitations guidelines for that pollutant. (See generally, 46 FR at 9414-16 (January 28, 1981)).

This definition of pass through satisfies two competing objectives set by Congress: (1) that standards for indirect dischargers be equivalent to standards for direct dischargers while at the same time, (2) that the treatment capability and performance of the POTW be recognized and taken into account in regulating the discharge of pollutants from indirect dischargers.

The Agency compares percentage removal rather than the mass or concentration of pollutants discharged because the latter would not take into account the mass of pollutants discharged to the POTW from non-industrial sources or the dilution of the pollutants in the POTW effluent to lower concentrations due to the addition of large amounts of non-industrial wastewater.

INDUSTRY COST AND POLLUTANT REMOVAL ESTIMATES

The industry cost and pollutant removal estimates of each treatment option were used to determine the most cost-effective option. The methodology applied in calculating pollutant removal estimates and plant compliance costs is discussed in Section X. Table XII-1 (page 2619) shows the estimated pollutant removal benefits for indirect dischargers. A comparison of proposal and promulgation compliance costs for indirect dischargers are presented in Table XII-2 (page 2620).

PRETREATMENT STANDARDS FOR EXISTING AND NEW SOURCES

Options for pretreatment of wastewaters from both existing and new sources are based on increasing the effectiveness of end-ofpipe treatment technologies. All in-plant changes and applicable end-of-pipe treatment processes have been discussed previously in Sections X and XI. The options for PSES and PSNS, therefore, are the same as the BAT and NSPS options discussed in Sections X and XI, respective|y.

A description of each option is presented in Sections X and XI, while a more detailed discussion, including pollutants controlled by each treatment process is presented in Section VII of Vol. I.

Treatment technologies considered for the PSES and PSNS options are:

OPTION A

- Preliminary treatment with cyanide precipitation (where required)
- Preliminary treatment with ammonia steam stripping (where required)
- o Chemical precipitation and sedimentation
- o Ion exchange end-of-pipe treatment

OPTION B

- Preliminary treatment with cyanide precipitation (where required)
- Preliminary treatment with ammonia steam stripping (where required)
- o Chemical precipitation and sedimentation
- In-process flow reduction of furnace and refinery scrubber liquor as well as for raw material granulation water
- o Ion exchange end-of-pipe treatment

OPTION C

- Preliminary treatment with cyanide precipitation (where required)
- Preliminary treatment with ammonia steam stripping (where required)
- o Chemical precipitation and sedimentation
- In-process flow reduction of furnace and refinery scrubber liquor as well as for raw material granulation water
- o Multimedia filtration
- o Ion exchange end-of-pipe treatment

PSES OPTION SELECTION

SECT - XII

EPA promulgated PSES equal to BAT (Option C) for this subcategory. It is necessary to promulgate this PSES to prevent pass-through of copper, cyanide, zinc, ammonia, gold, platinum and palladium. These priority pollutants are removed by a well operated POTW achieving secondary treatment at an average of 32 percent, while BAT level technology removes approximately 99 percent.

The technology basis for PSES thus is chemical precipitation and sedimentation, ammonia steam stripping, cyanide precipitation, flow reduction, multimedia filtration and ion wastewater The achievable concentration for ammonia steam exchange. stripping is based on iron and steel manufacturing category data, explained in the discussion of BPT and BAT for this as subcategory. Flow reduction is based on the same recycle of scrubber effluent and raw material granulation water that is the flow basis of BAT. Recycle is practiced by 22 of the 31 existing plants in the subcategory.

Implementation of the promulgated PSES limitations would remove annually an estimated 110,000 kg of priority pollutants including 866 kg of cyanide, and an estimated 10,534 kg of ammonia. Capital cost for achieving promulgated PSES is \$1,809,400 and annualized cost of \$1,100,500. The promulgated PSES will not result in adverse economic impacts.

Based on comments received after proposal, the Agency believes that it may be necessary for some facilities to use sulfide order to achieve the promulgated effluent polishing in limitations because of high zinc concentrations or complexing problems. Because the Agency believes that these situations will be the exception, rather than the rule, sulfide polishing is not specifically included as part of the model technology on which pretreatment standards for existing sources are based. The Agency has, however, evaluated the cost associated with the use of sulfide polishing at secondary precious metals plants. After performing this evaluation, the Agency has concluded that sulfide polishing will result in a very small (less than 5 percent) incremental increase in wastewater treatment costs at a typical secondary precious metals facility.

PSNS OPTION SELECTION

We are promulgating PSNS equivalent to NSPS (Option C). The technology basis for promulgated PSNS is identical to NSPS. This is equivalent to PSES and BAT. The same pollutants pass through PSNS as at PSES, for the same reasons. We know of no at economically feasible, demonstrated technology that is better than NSPS technology. The NSPS flow allowances are based on minimization of process wastewater wherever possible through the use of holding tanks to recycle wet scrubbing wastewater and raw material granulation water. The discharges are based on recycle of these waste streams (see Section X - Recycle of Water Used in Wet Air Pollution Control).

SECT - XII

EPA believes that the promulgated PSNS are achievable, and that they are not a barrier to entry of new plants into this subcategory.

REGULATED POLLUTANT PARAMETERS

Pollutants selected for limitation, in accordance with the rationale of Sections VI and X, are identical to those selected for limitation for BAT. It is necessary to promulgate PSES and PSNS to prevent the pass-through of copper, cyanide, zinc, ammonia, and combined metals (gold, platinum, and palladium), which are the limited pollutants.

PRETREATMENT STANDARDS

Pretreatment standards are based on the treatable concentrations from the selected treatment technology, (Option C), and the discharge rates determined in Sections X and XI for BAT and NSPS, respectively. A mass of pollutant per mass of product (mg/troy ounce) allocation is given for each subdivision within the sub-This pollutant allocation is based on the product of category. the treatable concentration from the promulgated treatment (mg/l) and the production normalized wastewater discharge rate (liter/ troy ounce). The achievable treatment concentrations for BAT are identical to those for PSES and PSNS. These concentrations are listed in Table VII-21 (page 248) of Vol. I. PSES and PSNS are presented in Tables XII-4 (page 2623) and XII-5 (page 2637).

Table XII-1

POLLUTANT REMOVAL ESTIMATES FOR INDIRECT DISCHARGERS

<u>Pollutant</u>	Raw Waste (kg/yr)	Option A Discharge (kg/yr)	Option A Removed (kg/yr)	Option B Discharge (kg/yr)	Qption B Removed (kg/yr)	Option C Discharge (kg/yr)	Option C Removed (kg/yr)	
Artinony	34.87	9.85	25.01	8.15	26.72	6.31	28.56	
Antimony	12.16	5.73	6.42	4.61	7.55	3.40	8.75	
Arsenic Cadmium	17.02	2.96	14.06	1.13	15.88	0.70	16.31	
'Chromium (Total)	187.43	2.96	184.47	1.20	186.22	1.00	186.42	
	2,603.86	21.71	2.582.15	8.31	2,595.55	5.59	2,598.27	
Copper Copper	881.57	19.53		15.59	865.98	15.34	866.28	
Cyanide (Total)	112.79		108.29	1.72	111.07	1.15	111.64	
Lead	0.10	0.09	0.00	0.09	0.01	0.09	0.01	
Mercury	1,161.39	27.70	1,133.69	10.60	1,150.79	3.15	1,158.24	
Nickel	2,630.50	11.12	2,619.43	4.19	2.626.36	2.79	2,627.76	
Selenium	42.41	3.74	38.67	1.43	40.98	1.00	41.41	
Silver	11.62	5.47	6.15	4.37	7.25	3.23	8.38	
Thallium		12.35	102,665.53	4.73	102,673.15	3.30	102,674.59	
Zinc	102,677.88	12.35	102,003133		100,075755			
TOTAL TOXICS	110,373.62	127.70	110,245.92	66.13	110,307,49	47.06	110,326.56	
Amnonia	10,962.04	884.29	10,077.75	427.92	10,534.12	427.92	10,534.12	
Cobalt	5.04	1.68	3.36	0.71	4.33	0.48	4.57	
Fluoride	12.12	10.57	1.54	9.84	2.27	9.84	2.27	
Gold	39.43	0.39	39.04	0.17	39.26	0.17	39.26	•
Platinum	17.96	0.39	17.57	0.17	17.79	0.17	17.79	
Palladium	17.96	0.39	17.57	0.17	17.79	0.17	17.79	
TOTAL NONCONVENTIONALS	11,054.55	896.71	10,156.84	438.99	10,615.56	438.76	10,615.79	
TSS	61,714.13	449.16	61,264.96	171.94	61,542.19	37.25	61,676.87	
Oil and Grease	37.66	32.87	4.78	30.51	7.15	30.51	7.15	•
TOTAL CONVENTIONALS	61,751.78	482.04	61,269.75	202.44	61,549.34	67.76	61,684.03	
TOTAL POLLUTANTS	183,179.95	1,507.45	181,672.50	707.55	182,472.40	553.58	182,626.37	

Option A - Chemical precipitation, sedimentation, ion exchange, ammonia steam stripping, and cyanide precipitation.

Option B - Chemical precipitation, sedimentation, ion exchange, ammonia steam stripping, cyanide precipitation, and flow reduction.

Option C - Chemical precipitation, sedimentation, ion exchange, ammonia steam stripping, cyanide precipitation, flow reduction, and filtration.

SECONDARY PRECIOUS METALS SUBCATEGORY SECT

XII

TABLE XII-2

COST OF COMPLIANCE FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY INDIRECT DISCHARGERS

Option	Total Required Capital Cost	Total Annual Cost
Α	l,774,400	1,078,000
В	1,707,000	1,034,000
С	1,809,400	1,100,500

TABLE XII-3

PSES AND PSNS WASTEWATER DISCHARGE RATES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

Wastewater Stream	BAT Normalized <u>Discharge</u> <u>Rates</u>	Production Normalizing Parameter	
Furnace wet air pollution control	4.5 l/t.o.	Troy ounces of precious metals, including silver, incinerated or smelted	
Raw material granulation	0.64 l/t.o.	Troy ounces of precious metals in the granulated raw material	
Spent plating solutions	1.1 1/1	Liters of spent plating solution used as a raw material	
Spent cyanide stripping solutions	3.7 l/t.o.	Troy ounces of gold stripped	
Refinery wet air pollution control	1.0 l/t.o.	Troy ounces of precious metals produced in refinery, including silver	
Gold solvent extraction raffinate	0.63 l/t.o.	Troy ounces of gold produced by solvent extraction	
Gold spent electrolyte	0.0087 l/t.o.	Troy ounces of gold produced by electrolysis	
Gold precipitation and filtration	4.4 l/t.o.	Troy ounces of gold precipitated	
Platinum precipitation and filtration	5.2 l/t.o.	Troy ounces of platinum precipitated	
	•		

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PSES AND PSNS WASTEWATER DISCHARGE RATES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

Wastewater Stream	BAT Normalized <u>Discharge</u> <u>Rates</u>	Production Normalizing Parameter
Palladium precipitation and filtration	3.5 1/t.o.	Troy ounces of palladium precipitated
Other platinum group metals precipitation and filtration	5.2 l/t.o.	Troy ounces of other platinum group metals precipitation
Spent solution from PGC salt production	0.9 l/t.o.	Troy ounces of gold contained in PGC product
Equipment and floor wash	0.0 l/t.o.	Troy ounces of precious metals metals including silver produced in refinery

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NOTES 1/t.o. = liters per troy ounce 1/1 = liters per liter

TABLE XII-4

PSES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(a) Furnace Wet Air Pollution Control PSES

Pollutant or	Maximum for	Maximum for
pollutant property	any one day	monthly average
mg/troy ounce o	f precious metals	incinerated or smelted
Antimony	8.685	3.870
Arsenic	6.255	2.790
Cadmium	0.900	0.360
Chromium	1.665	0.675
*Copper	5.760	2.745
*Cyanide	0.900	0.360
Lead	1.260	0.585
Nickel	2.475	1.665
Selenium	3.690	1.665
Silver	1.305	1.665
Thallium	6.300	0.540
*Zinc	4.590	2.745
*Ammonia	599.900	1.890
*Combined metals	1.350	263.700

PSES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(b) Raw Material Granulation PSES

Pollutan pollutan		ert		imum one			Maximum monthly	for average	······································
mg/troy	ounce	of	precious	meta	al in	the	granula	ted raw	material
Antimony Arsenic Cadmium Chromium *Copper *Cyanide Lead Nickel Selenium Silver Thallium *Zinc *Ammonia *Combined	metal	- 5	х	1.2 0.8 0.1 0.2 0.8 0.1 0.3 0.5 0.1 0.8 0.6 85.3 0.1	90 28 37 19 28 79 52 25 86 96 53 10	•		0.550 0.397 0.051 0.096 0.390 0.051 0.083 0.237 0.237 0.237 0.077 0.390 0.269 37.500	

PSES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(c) Spent Plating Solutions PSES

Pollutant o pollutant p		erty	Maximum for any one day	Maximum monthly	for average	
mg/liter	c of	spent	plating solution	s used as	a raw ma	terial
Antimony			1.930	· · ·	0.860	, <u>,</u>
Arsenic			1.390		0.620	. ,
Cadmium			0.200		0.080	
Chromium			0.370		0.150	
*Copper		. ,	1.280		0.610	
*Cyanide			0.200		0.080	• • • •
Lead			0.280		0.130	
Nickel	•		0.550		0.370	
Selenium			0.820		0.370	
Silver			0.290		0.120	•
Thallium			1.400		0.610	
*Zinc			1.020		0.420	
*Ammonia	,		133.300		58.600	
*Combined 1	netal	ls	0.300			•

TABLE XII-4 (Continued)

PSES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(d) Spent Cyanide Stripping Solutions PSES

Pollutant		Maximum for	Maximum for
pollutant		any one day	monthly average
	mg/	troy ounce of gold	d stripped
Antimony	metals	7.141	3.182
Arsenic		5.143	2.294
Cadmium		0.740	0.296
Chromium		1.369	0.555
*Copper		4.736	2.257
*Cyanide		0.740	0.296
Lead		1.036	0.481
Nickel		2.035	1.369
Selenium		3.034	1.369
Silver		1.073	1.369
Thallium		5.180	0.444
*Zinc		3.774	2.257
*Ammonia		493.200	1.554
*Combined		1.110	216.800

SECONDARY PRECIOUS METALS SUBCATEGORY SECT XII

TABLE XII-4 (Continued)

PSES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(e) <u>Refinery Wet Air Pollution Control¹ PSES</u>

· · · · · · · · · · · · · · · · · · ·		· · · · · ·			
Pollutant or	r M	aximum	for	Maximum	for
pollutant pr	roperty a	nyone	day	monthly	average
•			and the second second		· · · · · · · · · · · · · · · · · · ·

mg/troy ounce of precious metals produced in refinery

Antimony	1.930	0.860
Arsenic	1.390	0.620
Cadmium	0.200	0.080
Chromium	0.370	0.150
*Copper	1.280	0.610
*Cyanide	0.200	0.080
Lead	0.280	0.130
Nickel	0.550	0.370
Selenium	0.820	0.370
Silver	0.290	0.120
Thallium	1.400	0.610
*Zinc	1.020	0.420
*Ammonia	133.300	58.600
*Combined metals	0.300	

*Regulated Pollutant

¹This allowance applies to either acid or alkaline wet air pollution control scrubbers. If both acid and alkaline wet air pollution control scrubbers are present in a particular facility the same allowance applies to each.

PSES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(f) Gold Solvent Extraction Raffinate and Wash Water NSPS

Pollutant or pollutant property	Maximum for any one day	Maximum for monthly average	
mg/troy ounce of	of gold produced	by solvent extraction	
Antimony	1.216	0.542	
Arsenic	0.876	0.391	
Cadmium	0.126	0.050	
Chromium	0.233	0.095	
*Copper	0.806	0.384	
*Cyanide	0.126	0.050	
Lead	0.176	0.082	
Nickel	0.347	0.233	
Selenium	0.517	0.233	
Silver	0.183	0.076	
Thallium	0.882	0.384	
*Zinc	0.643	0.265	
*Ammonia	83.980	36.920	
*Combined metals	0.189		

PSES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(g) Gold Spent Electrolyte PSES

		and the second	
Pollutant pollutant			imum for thly average
1	1	-	and the second
· · .]	mg/troy ounce	of gold produced by	electrolysis
Antimony	, e	0.017	0.007
Arsenic		0.012	0.005
Cadmium		0.002	0.000
Chromium		0.003	0.001
*Copper	25 ji	0.011	0.005
*Cyanide	, p	0.002	0.000
Lead		0.002	0.001
Nickel	a di unita a	0.005	0.003
Selenium		0.007	0.003
Silver		0.003	0.001
Thallium		0.012	0.005
*Zinc		0.009	0.004
*Ammonia	2 4 4 6	1.160	0.510
*Combined	metals	0.003	
*Combined	metals	0.003	

PSES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(h) Gold Precipitation and Filtration PSES

Pollutant pollutant		Maximum for any one day	Maximum for monthly average	
	mg/troy	ounce of gold	precipitated	
Antimony Arsenic Cadmium Chromium *Copper *Cyanide Lead Nickel Selenium Silver Thallium *Zinc *Ammonia *Combined	metals	8.492 6.116 0.880 1.628 5.632 0.880 1.232 2.420 3.608 1.276 6.160 4.488 586.500 1.320	3.784 2.728 0.352 0.660 2.684 0.352 0.572 1.628 1.628 1.628 2.684 1.848 257.800	

PSES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(i) Platinum Precipitation and Filtration PSES

Pollutant		Maximum for	Maximum for	<u> </u>
pollutant	property	any one day	monthly average	
	mg/troy	ounce of platinum	precipitated	
Antimony		10.040	4.472	-
Arsenic		7.228	3.224	
Cadmium	ç	1.040	0.416	
Chromium		1.924	0.780	
*Copper		6.656	3.172	
*Cyanide		1.040	0.416	•
Lead	*	1.456	0.676	
Nickel	· · · · · · · · · · · · · · · · · · ·	2.860	1.924	
Selenium		4.264	1.924	
Silver	· .	1.508	0.624	
Thallium	· · ·	7.280	3.172	
*Zinc	· ·	5.304	2.184	
*Ammonia	.	693.200	304.700	
*Combined	metals	1.560		, ¹

SECONDARY PRECIOUS METALS SUBCATEGORY SECT - XII

TABLE XII-4 (Continued)

PSES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(j) Palladium Precipitation and Filtration PSES

		· · ·	
Pollutant	or	Maximum for	Maximum for
pollutant	property	any one day	monthly average
-	_	. – –	
	mg/troy	ounce of <u>p</u> alladiu	m precipitated
Antimony		11.580	5.160
Arsenic		8.340	3.720
Cadmium		1.200	0.480
Chromium		2.200	0.900
*Copper		7.680	3.660
*Cyanide		1.200	0.480
Lead	ł	1.680	0.780
Nickel		3.300	2.220
Selenium		4.920	2.220
Silver	· ·	1.740	0.720
Thallium	• .	8.400	3.660
*Zinc		6.120	2.520
*Ammonia		799.800	351.600
*Combined	metals	1.800	
		•	

SECONDARY PRECIOUS METALS SUBCATEGORY

TABLE XII-4 (Continued)

PSES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(k) Other Platinum Group Metals Precipitation and Filtration PSES

Pollutant or pollutant property	Maximum for any one day	Maximum for monthly average
mg/troy ounce of	f other platinum	group metals precipitated
Antimony	10.040	4.472
Arsenic	7.228	3.224
Cadmium	1.040	0.416
Chromium	1.924	0.780
*Copper	6.656	3.172
*Cyanide	1.040	0.416
Lead	1.456	0.676
Nickel	2.860	1.924
Selenium	4.264	1.924
Silver	1.508	0.624
Thallium	7.280	3.172
*Zinc	5.304	2.184
*Ammonia	693.200	304.700
*Combined metals	1.560	9700 9700 9700

TABLE XII-4 (Continued)

PSES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(1) <u>Solution from PGC Salt Production</u> PSES

Pollutant or	Maximum for	Maximum for
pollutant property	any one day	monthly average
mg/troy ounce of gold of	contained in PGC	product
Antimony	1.737	0.774
Arsenic	1.251	0.558
Cadmium	0.180	0.072
Chromium	0.333	0.135
*Copper	1.152	0.549
*Cyanide	0.180	0.072
Lead	0.252	0.117
Nickel	0.495	0.333
Selenium	0.738	0.333
Silver	0.261	0.333
Thallium	1.260	0.108
*Zinc	0.918	0.549
*Ammonia	120.000	0.378
*Combined metals	0.270	52.740

PSES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(m) Equipment and Floor Wash PSES

Pollutant pollutant		Maximum for any one day	Maximum for monthly average
		• • • • • • • • • • • • • • • • • • •	
mg/t	roy ounce	of precious metals	produced in refinery
Antimony		0.000	0.000
Arsenic	· · ·	0.000	0.000
Cadmium		0.000	0.000
Chromium	× .	0.000	0.000
*Copper	· •	0.000	0.000
-*Cyanide		0.000	0.000
Lead	. ,	0.000	0.000
Nickel	:	0.000	0.000
Selenium	4 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -	0.000	0.000
Silver	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	0.000	0.000
Thallium		0.000	0.000
*Zinc	1 - E - L	0.000	0.000
*Ammonia		0.000	0.000
*Combined	metals	0.000	

PSES FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(n) Preliminary Treatment PSES

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average

mg/troy ounce of total precious metals produced through this operation

Antimony	96.500	43.000
Arsenic	69.500	31.000
Cadmium	10.000	4.000
Chromium	18.500	7.500
Copper	64.000	30.500
Cyanide (total)	10.000	4.000
Lead	14.000	6.500
Nickel	27.500	18.500
Selenium	41.000	18.500
Silver	14.500	6.000
Thallium	70.000	30.500
Zinc	51.000	21.000
Combined metals	15.000	
Ammonia (as N)	6665.000	2930.000

TABLE XII-5

PSNS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

.....

(a) <u>Furnace Wet Air Pollution Control</u> PSNS

		•. 1		in an ann an a	<u> </u>		· · · · · · · · · · · · · · · · · · ·
Pollutant	or	Ma	ximum fo:	r. *	Maximum	for	
pollutant	property	an	y one dag	y n	nonthly	average	
·		· · · ·		4			
	, · · · · ·	.					
mg/ti	oy ounce	of pre	cious me	tals in	ncineral	ced or sr	nelted
Antimony			8.685			3.870	
-		. •		, i	•		· · · ·
Arsenic	· · · · ·	••	6.255			2.790	
Cadmium			0.900			0.3 60	4.00
Chromium	4		1.665			0.675	
*Copper		-	5.760			2.745	
*Cyanide		•	0.900			0.360	
Lead	\$	a di Cara	1.260		-	0.585	1 e g - 97
	· · · · ·						
lickel	· .		2.475	1 · · · · · · · · · · · · · · · · · · ·		1.665	
Selenium	•	54 - L	3.690		`	1.665	
Silver			1.305	s. 		0.540	
hallium			6.300		s	2.745	- 4782
Zinc			4.590			1.890	
Ammonia	2	and the second sec	599.900	· · · ,	•	263.700	
Combined	motola		1.350	÷ •			
.comprised	metals	· . ·	T.220			· · · · · · · · · · · · · · · · · · ·	

TABLE XII-5 (Continued)

PSNS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(b) Raw Material Granulation PSNS

Pollutant or pollutant property	Maximum for any one day	Maximum for monthly average
mg/troy ounce of pr	ecious metals in	the granulated raw materia
Antimony	1.235	0.550
Arsenic	0.890	0.397
Cadmium	0.128	0.051
Chromium	0.237	0.096
*Copper	0.819	0.390
*Cyanide	0.128	0.051
Lead	0.179	0.083
Nickel	0.352	0.237
Selenium	0.525	0.237
Silver	0.186	0.077
Thallium	0.896	0.390
*Zinc	0.653	0.269
*Ammonia	85.310	37.500
*Combined metals	0.192	-

PSNS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(c) Spent Plating Solutions PSNS

Pollutant or pollutant property	Maximum for any one day	Maximum for monthly average
mg/liter of spent	plating solutions	used as a raw materia
Antimony	1.930	0.860
Arsenic	1.390	0.620
Cadmium	0.200	0.080
Chromium	0.370	0.150
*Copper	1.280	0.610
*Cyanide	0.200	0.080
Lead	0.280	0.130
Nickel	0.550	0.370
Selenium	0.820	0.370
Silver	0.290	0.120
Thallium	1.400	0.610
*Zinc	1.020	0.420
*Ammonia	133.300	58.600
*Combined metals	0.300	

PSNS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(d) Spent Cyanide Stripping Solutions PSNS

Pollutant pollutant		Maximum for any one day	Maximum monthly		
		· ·			<u></u>
mg/troy ou stripped	unce of gold	· · · · · · · · · · · · · · · · · · ·			
Antimony		7.141		3.182	
Arsenic	1	5.143		2.294	
Cadmium		0.740		0.296	
Chromium		1.369		0.555	
*Copper		4.736		2.257	
*Cyanide		0.740		0.296	
Lead	1	1.036	>	0.481	
Nickel		2.035		1.369	
Selenium		3.034		1.369	
Silver		1.073		0.444	• •
Thallium		5.180		2.257	
*Zinc		3.774		1.554	
*Ammonia		493.200	2	216.800	
*Combined	metals	1.110			

PSNS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(e) <u>Refinery Wet Air Pollution Control¹ PSNS</u>

Pollutant or	Maximum for	Maximum for		
pollutant property	any one day	monthly average		
mg/troy ounce	of precious metals	produced in refinery		
Antimony	1.930	0.860		
Arsenic	1.390	0.620		
Cadmium	0.200	0.080		
Chromium	0.370	0.150		
*Copper	1.280	0.610		
*Cyanide	0.200	0.080		
Lead	0.280	0.130		
Nickel	0.550	0.370		
Selenium	0.820	0.370		
Silver	0.290	0.370		
Thallium	1.400	0.120		
*Zinc	1.020	0.610		
*Ammonia	133.300	0.420		
*Combined metals	0.300	58.600		

*Regulated Pollutant

¹This allowance applies to either acid or alkaline wet air pollution control scrubbers. If both acid and alkaline wet air pollution control scrubbers are present in a particular facility the same allowance applies to each.

TABLE XII-5 (Continued)

PSNS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(f) Gold Solvent Extraction Raffinate and Wash Water PSNS

Pollutant or	Maximum for	Maximum for
pollutant property	any one day	monthly average
mg/troy ounce of	gold produced	by solvent extraction
Antimony	1.216	0.542
Arsenic	0.876	0.391
Cadmium	0.126	0.050
Chromium	0.233	0.095
*Copper	0.806	0.384
*Cyanide	0.126	0.050
Lead	0.176	0.082
Nickel	0.347	0.233
Selenium	0.517	0.233
Silver	0.183	0.233
Thallium	0.882	0.233
*Zinc	0.643	0.233
*Ammonia	83.980	0.265
*Combined metals	0.189	36.920

PSNS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(g) Gold Spent Electrolyte PSNS

Pollutant pollutant		су	Maximum for Maximum for any one day monthly average					е	
	mg/troy	ounce	of gold	prod	uced by	/ elec	ctrolysis		
Antimony			0.0)17		•	0.007		
Arsenic			0.0)12			0.005		
Cadmium			0.0)02	3.		0.000		
Chromium	e.		0.0	03		,	0.001		
*Copper		•	0.0	011		1 a	0.005		
*Cyanide			0.0	002			0.000		
Lead		ъ.	0.0	002		£	0.001		
Nickel			0.0	05			0.003		
Selenium			0.0)07			0.003		
Silver			0.0)03			0.001		
Thallium			0.0)12			0.005		
*Zinc			0.0)09			0.004		
*Ammonia			1.]	L 60		ŕ	0.510		
*Combined	metals			003					

PSNS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(h) Gold Precipitation and Filtration PSNS

Pollutant		Maximum	for	Maximum	
pollutant	property	any one	day	monthly	average
<u></u>		· · · · · · · · · · · · · · · · · · ·			

mg/troy ounce of gold precipitated 8.492 Antimony 3.784 Arsenic 6.116 2.728 Cadmium 0.880 0.352 Chromium 1.628 0.660 5.632 *Copper 2.684 *Cyanide 0.880 0.352 Lead 1.232 0.572 Nickel 2.420 1.628 Selenium 3.608 1.628 Silver 1.276 0.528 Thallium 6.160 2.684 *Zinc 4.488 1.848 *Ammonia 586.500 257.800 *Combined metals 1.320 ----

*Regulated Pollutant

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PSNS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(i) Platinum Precipitation and Filtration PSNS

			a
Pollutant pollutant		Maximum for any one day	Maximum for monthly average
	mg/troy	ounce of platinum	n precipitated
Antimony Arsenic Cadmium Chromium *Copper *Cyanide Lead Nickel Selenium Silver Thallium *Zinc		10.040 7.228 1.040 1.924 6.656 1.040 1.456 2.860 4.264 1.508 7.280 5.304	4.472 3.224 0.416 0.780 3.172 0.416 0.676 1.924 1.924 0.624 3.172 2.184
*Ammonia *Combined	metals	693.200 1.560	304.700

PSNS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(j) <u>Palladium</u> <u>Precipitation</u> and <u>Filtration</u> PSNS

Pollutant pollutant	Maximum any one	_	Maximum monthly	

mg/troy ounce of palladium precipitated Antimony 11.580 5.160 3.720 Arsenic 8.340 Cadmium 1.200 0.480 Chromium 2.200 0.900 *Copper 7.680 3.660 1.200 *Cyanide 0.480 Lead 1.680 0.780 Nickel 3.300 2.220 Selenium 4.920 2.220 Silver 1.740 0.720 Thallium 8.400 3.660 *Zinc 6.120 2.520 *Ammonia 799.800 351.600

1.800

*Regulated Pollutant

*Combined metals

PSNS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(k) Other Platinum Group Metals Precipitation and Filtration PSNS

Pollutant or pollutant property	Maximum for any one day	Maximum for monthly average
mg/troy ounce of	other platinum	group metals precipitated
Antimony	10.040	4.472
Arsenic	7.228	3.224
Cadmium	1.040	0.416
Chromium	1.924	0.780
*Copper	6.656	3.172
*Cyanide	1.040	0.416
Lead	1.456	0.676
Nickel	2.860	1.924
Selenium	4.264	1.924
Silver	1.508	0.624
Thallium	7.280	3.172
*Zinc	5.304	2.184
*Ammonia	693.200	304.700
*Combined metals	1.560	

TABLE XII-5 (Continued)

PSNS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(1) Spent Solution from PGC Salt Production PSNS

Pollutant pollutant			imum one		Maxir month		for average
n	ng/troy o	unce of	gold	containe	ed in	PGC	product
Antimony Arsenic Cadmium Chromium *Copper *Cyanide Lead Nickel Selenium Silver Thallium *Zinc *Ammonia *Combined	metals		1.7 1.2 0.1 0.3 1.1 0.2 0.4 0.2 1.2 0.9 120.0 0.2	251 80 333 52 80 252 95 38 61 60 18 00			0.774 0.558 0.072 0.135 0.549 0.072 0.117 0.333 0.333 0.333 0.108 0.549 0.378 52.740

TABLE XII-5 (Continued)

PSNS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(m) Equipment and Floor Wash PSNS

or prop	berty	· .	Maximum for any one day	Maximum monthly	
roy	ounce	of	precious metals	produced	in refinery
			0.000		0.000
			0.000		0.000
			0.000		0.000
			0.000		0.000
			0.000		0.000
			0.000	-	0.000
			0.000		0.000
			0.000		0.000
			0.000		0.000
			0.000		0.000
			0.000		0.000
			000		0.000
			0.000		0.000
meta	als		0.000	•	~ ~ ~
	roy	property	property roy ounce of	property any one day roy ounce of precious metals 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000	property any one day monthly roy ounce of precious metals produced 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000 0.000

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TABLE XII-5 (Continued)

PSNS FOR THE SECONDARY PRECIOUS METALS SUBCATEGORY

(n) Preliminary Treatment PSNS

Pollutant	Maximum for	Maximum for
Pollutant	Any One Day	Monthly Average

Antimony	96.500	43.000
Arsenic	69.500	31.000
Cadmium	10.000	4.000
Chromium	18.500	7.500
Copper	64.000	30.500
Cyanide (total)	10.000	4.000
Lead	14.000	6.500
Nickel	27.500	18.500
Selenium	41.000	18.500
Silver	14.500	6.000
Thallium	70.000	30.500
Zinc	51.000	21.000
Combined metals	15.000	
Ammonia (as N)	6665.000	293,0.000
1		4

SECONDARY PRECIOUS METALS SUBCATEGORY

SECTION XIII

BEST CONVENTIONAL POLLUTANT CONTROL TECHNOLOGY

EPA is not promulgating best conventional pollutant control technology (BCT) for the secondary precious metals subcategory at this time.

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NONFERROUS METALS MANUFACTURING POINT SOURCE CATEGORY

DEVELOPMENT DOCUMENT SUPPLEMENT

for the

Secondary Silver Subcategory

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May 1989

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SECTION I

SUMMARY

This document provides the technical basis for promulgating effluent limitations based on best practicable technology (BPT) and best available technology economically achievable (BAT) for existing direct dischargers, pretreatment standards for existing indirect dischargers (PSES), pretreatment standards for new indirect dischargers (PSNS), and standards of performance for new source direct dischargers (NSPS) for plants in the Secondary Silver Subcategory.

The secondary silver subcategory is comprised of 61 plants. Of the 61 plants, seven discharge directly to rivers, lakes, or streams; 26 discharge to publicly owned treatment works (POTW); and 28 achieve zero discharge of process wastewater.

EPA first studied the secondary silver subcategory to determine materials, final whether differences in raw products, manufacturing processes, equipment, age and size of plants, or water usage, required the development of separate effluent and standards for different segments of the limitations subcategory. This involved a detailed analysis of wastewater discharge and treated effluent characteristics, including the sources and volume of water used, the processes used, the sources of pollutants and wastewaters in the plant, and the toxic and other pollutant constituents of wastewater.

As a result ll subdivisions or building blocks have been identified in this subcategory that warrant separate effluent limitations. These include:

- 1. Film stripping
- 2. Film stripping wet air pollution control and precipitation and filtration of film stripping solutions wet air pollution control,
- 3. Precipitation and filtration of film stripping solutions,
- 4. Precipitation and filtration of photographic solutions,
- 5. Precipitation and filtration of photographic solutions wet air pollution control,
- 6. Electrolytic refining,
- 7. Furnace wet air pollution control,
- 8. Leaching,
- 9. Leaching wet air pollution control and precipitation of nonphotographic solutions wet air pollution control,
- 10. Precipitation and filtration of nonphotographic solutions, and
- 11. Floor and equipment washdown.

Several distinct control and treatment technologies (both inplant and end-of-pipe) applicable to the secondary silver subcategory were identified. The Agency analyzed both historical

SECONDARY SILVER SUBCATEGORY SECT - I

and newly generated data on the performance of these technologies, including their nonwater quality environmental impacts (such as air quality impacts and solid waste generation), and energy requirements. EPA also studied various flow reduction techniques reported in the data collection portfolios (dcp) and plant visits.

Engineering costs were prepared for each of the control and treatment options considered for the category. These costs were then used by the Agency to estimate the impact of implementing the various options on the subcategory. For each control and treatment option that the Agency found to be most effective and technically feasible in controlling the discharge of pollutants, the number of potential closures, number of employees affected, and impact on price were estimated. These results are reported in a separate document entitled "The Economic Impact Analysis of Effluent Limitations and Standards for the Nonferrous Metals Manufacturing Industry."

Based on consideration of the above factors, EPA identified various control and treatment technologies which formed the basis for BPT and selected control and treatment appropriate for each set of standards and limitations. The mass limitations and standards for BPT, BAT, NSPS, PSES, and PSNS are presented in Section II.

After examining the various treatment technologies, the Agency has identified BPT to represent the average of the best existing technology. Metals removal based on lime precipitation and sedimentation technology is the basis for the BPT limitations. Steam stripping was selected as the technology basis for ammonia limitations. Complete recycle of treated flow and equipment washdown water is also included. To meet the BPT effluent limitations, the secondary silver subcategory will incur an estimated capital cost of \$0.110 million (1982 dollars) and an annual cost of \$0.309 million (1982 dollars).

For BAT, the Agency has built upon the BPT basis of lime precipitation and sedimentation by adding in-process control technology and multimedia filtration. In-process control technology includes recycle or reuse of process water from wet air pollution control. Filtration is added as a polishing step to further reduce metals and suspended solids concentrations. To meet BAT effluent limitations based on this technology, the secondary silver subcategory is estimated to incur a capital cost of \$0.278 million (1982 dollars) and an annual cost of \$0.390 million (1982 dollars).

The best demonstrated technology, BDT, which is the technical basis of NSPS, is equivalent to BAT. In selecting BDT, EPA recognizes that new plants have the opportunity to implement the best and most efficient manufacturing processes and treatment technology. However, the technology basis of BAT has been determined as the best demonstrated technology because no additional process modifications or treatment technologies have been identified that substantially improve BAT performance.

Pretreatment standards for existing sources are based on the same technology as BAT. The technology basis is in-process flow reduction, ammonia steam stripping preliminary treatment, lime precipitation, sedimentation, complete recycle of treated floor and equipment washdown water, and multimedia filtration. To meet PSES, the secondary silver subcategory is estimated to incur a capital cost of \$0.634 million (1982 dollars) and an annual cost of \$0.422 million (1982 dollars).

For pretreatment standards for new sources (PSNS), the Agency selected preliminary treatment, end-of-pipe treatment, and inprocess flow reduction control techniques equivalent to BDT. As such, the PSNS are identical to the NSPS for all wastewater streams.

SECONDARY SILVER SUBCATEGORY SECT - I

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SECTION II

CONCLUSIONS

EPA has divided the secondary silver subcategory into ll subdivisions or building blocks for the purpose of effluent limitations and standards. These subdivisions are:

- (a) Film stripping,
- (b) Film stripping wet air pollution control and precipitation and filtration of film stripping solutions wet air pollution control,
- (c) Precipitation and filtration of film stripping solutions,
- (d) Precipitation and filtration of photographic solutions,
- (e) Precipitation and filtration of photographic solutions
- wet air pollution control,
- (f) Electrolytic refining,
- (g) Furnace wet air pollution control,
- (h) Leaching,
- (i) Leaching wet air pollution control and precipitation of nonphotographic solutions wet air pollution control,
- (j) Precipitation and filtration of nonphotographic solutions, and
- (k) Floor and equipment washdown.

Promulgated BPT is based on the performance achievable by the application of chemical precipitation and sedimentation (lime and settle) technology, along with preliminary treatment consisting of ammonia steam stripping for selected waste streams. Complete recycle of treated floor and equipment washdown is also included. The following \mathbf{BPT} effluent limitations are promulgated for existing sources:

(a) Film Stripping BPT

Pollutant or	Maximum	for	Maximum for
Pollutant Property	 Any One	Day	Monthly Average

mg/troy ounce of silver from film stripping

Copper	95.670	50.350
Zinc	73.510	30.720
Ammonia (as N)	6,712.000	2,951.000
Total Suspended Solids	2,065.000	981.800
рн	Within the range o at all ti	

SECONDARY SILVER SUBCATEGORY SECT - II

(b) Film Stripping Wet Air Pollution Control and Precipitation and Filtration of Film Stripping Solutions Wet Air Pollution Control BPT

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average

mg/troy ounce of silver from precipitation and filtration of film stripping solutions

Copper	1.843	0.970
Zinc	1.416	0.592
Ammonia (as N)	129.300	56.840
Total Suspended Solids	39.770	18.920
pH	Within the range of 7.0	to 1 0.0
	at all times	

(c) <u>Precipitation and Filtration of Film Stripping</u> Solutions BPT

Pollutant or	Maximum	for	Maximum for
Pollutant Property	Any One	Day	Monthly Average

mg/troy ounce of silver precipitated

Copper	109.400	57.570
Zinc	84.050	35.120
Ammonia (as N)	7,674.000	3,374.000
Total Suspended Solids	2,361.000	1,123.000
pH	Within the range of	E 7.0 to 10.0
	at all times	

(d) <u>Precipitation and Filtration of Photographic</u> Solutions BPT

Pollutant or	Maximum for	Maximum for
Pollutant Property		onthly Average
mg/troy ounce o	of silver precipitated	
Copper	50.540	26.600
Zinc	38.836	16,226
Ammonia (as N)	3,545.000	1,559.000
Total Suspended Solids	1,090.600	51 8.700
pH	Within the range	of 7.0 to 10.0
E	at all t	

(e) <u>Precipitation</u> and <u>Filtration</u> <u>Solutions Wet</u> <u>Air</u> <u>Pollution</u> (•	
Pollutant or	Maxim	uum f	or	Maxim	um for	
Pollutant Property				Monthly		ae
	7		· · ·	7		
mg/troy ounce of silver from prec of photographic solutions	ipitatio	on ar	nd fil	ltration	,	
Copper	· · · · ·	23.0	170		12.14	0
Zinc		17.7			7.40	-
Ammonia (as N)		18.0			711.40	
		97.8			236.80	
Total Suspended Solids						
pH	WICHTH			e of 7.0 times	10 IU	• 0
	,	a	all	CIMES		•
				· · · · ·		
(f) Electrolytic Refining BPT	•		•			
(1) <u>Riectiolytic</u> <u>Relining</u> Bil	•					
Pollutant or	Maxim				um for	
Pollutant Property	Any O	ne I	Day 🐇	Monthly	Avera	ge
mg/troy ounce of silver fro	om elect	roly	ytic 1	cefining	1 N J 	
Copper		1.4	444		0.76	0
Zinc		1.1	110		0.46	4
Ammonia (as N)	1	.01.3	300		44.54	0
Total Suspended Solids		31.1			14.82	0
pH	Within			e of 7.0		
				times		
· · · · · · · · · · · · · · · · · · ·					· · ·	
(g) <u>Furnace Wet Air</u> Pollution Con	<u>ntrol</u> B	BPT	۰.,			
Pollutant or	Maxim	ium 1	Eor	Maxim	um for	
Pollutant Property	Any C)ne I	Day	Monthly	Avera	ge
mg/troy ounce of silver roasted,	smelted	ord	dried	۰ ۲		
Copper	• • • • •	1.2	273		0.67	0
Zinc			978	11	0.40	
Ammonia (as N)		89.3			39.26	
Total Suspended Solids	1	27.4		• •	13.07	
pH	Within			e of 7.0		
E			all			
· · · · · ·					. · .	
	······································	·····		· · · ·	÷ ' •	

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(h) Leaching BPT

Pollutant or Pollutant Property		lmum for Ly Average
mg/troy ounce of silver	r produced from leaching	<u> </u>
Copper	0.164	0.086
Zinc	0.126	0.053
Ammonia (as N)	11.470	5.040
Total Suspended Solids	3.526	1.677
- Hq	Within the range of 7.	0 to 10.0
-	at all times	

(i) <u>Leaching Wet Air Pollution Control and Precipitation of</u> Nonphotographic Solutions Wet Air Pollution Control BPT

Pollutant or	Maximum fo		r Maximum for		
Pollutant Property	Any One	Day	Monthly Average		

mg/troy ounce of silver produced from leaching or silver precipitated

Copper	8.417	4.430
Zinc	6.468	2.703
Ammonia (as N)	590.500	259.600
Total Suspended Solids	181.700	86.390
pH	Within the range of	7.0 to 10.0
-	at all time	S

(j) <u>Precipitation and Filtration of Nonphotographic</u> Solutions BPT

Pollutant or	 Maximum fo	r Maxi	mum for
Pollutant Property	Any One Da	y Monthl	y Average

mg/troy ounce of silver precipitated

Copper Zinc Ammonia (as N) Total Suspended Solids pH	5.833 4.482 409.300 125.900 Within the range of at all time	
	at all time	es

Pollutant or Pollutant Property			imum for One Day	Maximum for Monthly Average		
	mg/troy	ounce of	silver	production		······································
Copper				0.000		0.000
Zinc				0.000		0.000
Ammonia (as N)				0.000		0.000
Total Suspended	d Solids			0.000		0.000 .
рН			Withi	n the range	e of 7.0	to 10.0
				at all		1 ₈
and the second			4 C	· · · · ·		· · ·

(k) Floor and Equipment Washdown BPT

EPA is promulgating BAT based on the performance achievable by the application of in-process flow reduction, chemical precipitation, sedimentation, complete recycle of treated floor and facility washdown water, and multimedia filtration, along with ammonia steam stripping for selected wastewater streams. The following BAT effluent limitations are promulgated for existing sources:

(a) Film Stripping BAT

Pollutant :			·······		lmum for One Day		um for Average
	mg/troy	ounce of	silver	from	film str	ipping	
Copper Zinc Ammonia(as	N)	n an an Angelan An Angelan Angelan Angelan Angelan Angelan Angelan		6,	64.450 51.360 712.000	2,	30.720 21.150 951.000

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(b) <u>Film Stripping Wet Air Pollution Control and</u> <u>Precipitation and Filtration of Film Stripping</u> <u>Solutions Wet Air Pollution Control BAT</u>

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
		e

mg/troy ounce of silver from precipitation and filtration of film stripping solutions

Copper	1	1.242	0.592
Zinc		0.990	0.408
Ammonia (as N)		129.300	56.840
			e

(c) <u>Precipitation and Filtration of Film Stripping</u> Solutions BAT

Pollu Pollutar	itant nt Pro			······································		ximum y One			um for Average	•
		mg/troy	ounce	of sil	lver	preci	pitate	đ		-
Copper Zinc Ammonia	(as N)			i		690 720 000	3,	35.120 24.180 374.000	,

(d) Precipitation and Filtration of Photographic Solutions BAT

Pollutant or Pollutant Property			Maximum for Any One Day M			Maximum for Monthly Average		
	mg/troy	ounce	of	silver	preci	pitated	9 5	
Copper Zinc Ammonia (as 1	N)			•		.048 .132 .000	1,	16.226 11.172 559.000

<u>(e)</u>	Precipitation and		Photographic	Solutions
	Wet Air Pollution	Control BAT		

		• •		
Pollutant or		Maximum	for	Maximum for
Pollutant Property		Any One	Day	Monthly Average
·				
	·			

mg/troy ounce of silver from precipitation and filtration of photographic solutions

Copper Zinc	15.540	7.406
	12.380	5.099
Ammonia (as N)	1,618.000	711.400

(f) Electrolytic Refining BAT

Pollutant or	Maximum for Maximum for
Pollutant Property	Any One Day Monthly Average
mg/troy ounce of silver	from electrolytic refining
Copper	0.973 0.464
Zinc	0.775 0.319
Ammonia (as N)	101.300 44.540

(g) Furnace Wet Air Pollution Control BAT

Pollutant or Pollutant Property			Maximum for Any One Day		Maximum for Monthly Average		
	mg/troy	ounce	of silver	roasted,	smelted,	or dried	
Copper Zinc Ammoni					0.000 0.000 0.000		0.000 0.000 0.000

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(h) Leaching BAT

Pollutant or Pollutant Property			Maximum Any One			um for Average	
<u></u>	mg/troy o	ounce of	silver	produced fr	om le	eaching	
Copper					110		0.053
Zinc Ammonia	(as N)				088 470		0.036 5.040

(i) <u>Leaching Wet Air Pollution Control and Precipitation of</u> Nonphotographic Solutions Wet Air Pollution Control BAT

Pollutant or	Maximum for Maximum for			
Pollutant Property	Any One Day Monthly Average			
mg/troy ounce of silver p pre	produced from leachin cipitated	g or silver		
Copper	5.671	2.703		
Zinc	4.519	1.861		
Ammonia (as N)	590.500	259.600		

Ammonia	(as N)	4.519	259.600
			, <u>, ,</u>

(j) <u>Precipitation and Filtration of Nonphotographic</u> Solutions BAT

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average

		mg/troy	ounce	of	silver	precipitated		
Copper Zinc Ammonia	(as l	N)	¥ K			3.930 3.132 409.300	,	1.873 1.290 179.900

(k) Floor and Equipment Washdown BAT

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
mg/troy ounce of	silver production	Na 1999 (1999) (1999) (1999) (1999) (1999) (1999) (1999) (1999) (1999) (1999) (1999) (1999) (1999) (1999) (199
Copper	0.000	0.000
Zinc	0.000	0.000
Ammonia (as N)	0.000	0.000

NSPS are promulgated based on the performance achievable by the application of chemical precipitation, sedimentation, and multimedia filtration (lime, settle, and filter) technology and in-process flow reduction control methods, along with complete recycle of treated floor and equipment washdown water and preliminary treatment consisting of ammonia steam stripping for selected waste streams. The following effluent standards are promulgated for new sources:

(a) Film Stripping NSPS

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
	·	

mg/troy ounce of silver from film stripping

Copper	64.450 30.720
Zinc	51.360 21.150
Ammonia(as N)	6,712.000 2,951.000
Total Suspended Solids	755.300 604.200
pH	Within the range of 7.0 to 10.0
- ·	at all times

(b) <u>Film Stripping Wet Air Pollution Control and</u> <u>Precipitation and Filtration of Film Stripping</u> <u>Solutions Wet Air Pollution Control</u> NSPS

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
mg/troy ounce of silver of film s	from precipitation an stripping solutions	nd filtration
Copper	1.242	0.592
Zinc	0.990	0.408
Ammonia (as N)	129.300	56.840
Total Suspended Solids	14.550	11.640
pH Within	the range of 7.0 to	10.0 at all times

(c) <u>Precipitation and Filtration of Film Stripping</u> Solutions NSPS

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average

mg/troy ounce of silver precipitated

Copper Zinc	73.690 58.720 7,674.000	35.120 24.180 3,374.000
Ammonia (as N) Total Suspended Solids pH	863.600 Within the range of at all times	690.900 7.0 to 10.0

(d) Precipitation and Filtration of Photographic Solutions NSPS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of silver preci	pitated	
Copper Zinc Ammonia (as N) Total Suspended Solids pH	34.048 27.132 3,545.000 399.000 Within the range at all t	

(e) <u>Precipitation</u> and <u>Filtrati</u> <u>Wet</u> <u>Air</u> <u>Pollution</u> <u>Control</u>		otogr	aphic	<u>Solutic</u>	ons
Pollutant or	Maxi	Lmum	for	Maxin	num for
		One			Average
Pollutant Property	Any	one	Day	Montury	Average
·					
mg/troy ounce of silver from photograph			and f	filtrati	on of
Copper	· ·	15.	540		7.406
Zinc			380		5.099
	1	,618.		· .	711.400
Ammonia (as N)	<u>т</u> ,				
Total Suspended Solids		182.			145.700
pH	Within				to 10.0
		at	all ti	lmes	
	o.				
		, i		1 · · · · · · · · · · · · · · · · · · ·	·
(f) <u>Electrolytic</u> <u>Refining</u> NSE	?S				
Pollutant or	Max	imum	for	Maxin	num for
Pollutant Property	Anv	One	Dav	Monthly	Average
Follucane riopercy	7	U.LC	201		merage
mg/troy ounce of silver Copper Zinc Ammonia (as N) Total Suspended Solids pH		0. 0. 101. 11. the	973 775 300 400	of 7.0	0.464 0.319 44.540 9.120 to 10.0
<u></u>			`	1	
(g) Furnace Wet Air Pollution	Control	NSF	, S		
$(g) \underline{IuInacc} \underline{ncc} \underline{n11} \underline{IoIIucIon}$	<u></u>		- -		
Pollutant or	Max	imum	for	Maxin	num for
Pollutant Property		One			v Average
mg/troy ounce of silver	roasted,	smel	.ted, d	or dried	1
Copper	1	0			0
Zinc		0			0
Ammonia (as N)		Õ			Õ
		Ő			ň
Total Suspended Solids	747 L L L L		*****	0 5 7 0	Fo 10 0
pH	Within				to 10.0
		at	all t	Imes	
					- <u>-</u>

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(h) <u>Leaching</u> NSPS

Pollutant or	Maximum f	for	Maxim	um for
Pollutant Property	Any One I	Day M	onthly	Average
mg/troy ounce of silver	produced fro	om leac	hing	
Copper	0.1	L10		0.053
Zinc	0.0	88		0.036
Ammonia (as N)	11.4	170		5.040
Total Suspended Solids	1.2	290		1.032
pH	Within the r			to 10.0
	at a	all tim	es	
Nonphotographic Solutions We	et Air Pollut	ion Co	ntrol	NSPS
(i) <u>Leaching Wet Air Pollution C</u> <u>Nonphotographic Solutions We</u> Pollutant or Pollutant Property	Control and F t Air Pollut Maximum f Any One D	<u>ion Co</u> for	ntrol Maximu	NSPS
Nonphotographic Solutions We Pollutant or Pollutant Property mg/troy ounce of silver produ precipi Copper Zinc Ammonia (as N) Total Suspended Solids	Air Pollut Maximum f Any One D Iced from lea tated 5.6	cion Co For Day M Aching 571 519 500 150	ntrol Maximu onthly or silv	NSPS m for Average ver 2.703 1.861 259.600 53.160

(j) <u>Precipitation and Filtration of Nonphotographic</u> Solutions NSPS

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average

mg/troy ounce of silver precipitated

Copper	3.930	1.873
Zinc	3.132	1.290
Ammonia (as N)	409.300	179.900
Total Suspended Solids	46.050	36.840
pH	Within the range of 7.	0 to 10.0
	at all times	

(k)	Floor	and	Equipment	Washdown	NSPS
• •	the second se				

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce	of silver production	
Copper Zinc Ammonia (as N) Total Suspended Solids pH	0.000 0.000 0.000 0.000 Within the range at all t	

EPA is promulgating PSES based on the performance achievable by the application of in-process flow reduction, chemical precipitation, sedimentation, complete recycle of treated floor and facility washdown water, and multimedia filtration, along with ammonia steam stripping for selected waste streams. The following pretreatment standards are promulgated for existing sources:

(a) Film Stripping PSES

Pollutan Pollutant I			lmum for One Day	Maximum for Monthly Average
	mg/troy ounce of s	ilver from	film stri	pping
Copper Zinc Ammonia(as	N)	6,	64.450 51.360 712.000	30.720 21.150 2,951.000

(b) Film Stripping Wet Air Pollution Control and Precipitation and Filtration of Film Stripping Solutions Wet Air Pollution Control PSES

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average

mg/troy ounce of silver from precipitation and filtration of film stripping solutions

Copper	1.242	0.592
Zinc	0.990	0.408
Ammonia (as N)	129.300	56.840

Pollutant or Pollutant Property			Maximum for	Maximum for
	Property	• .	Any One Day	Monthly Average
	mg/troy	ounce of silv	er precipitat	eđ
Copper		•	73.690	35.120
Zinc		14 - 14 - 14 - 14 - 14 - 14 - 14 - 14 -	58.720	24.180
Ammonia (a	s N)		7,674.000	3,374.000

(c) <u>Precipitation and Filtration of Film Stripping</u> Solutions PSES

(d) Precipitation and Filtration of Photographic Solutions PSES

Pollutant or	Maximum for	Maximum for	
Pollutant Property	Any One Day	Monthly Average	
mg/troy ounce of	silver precipitated		
Copper	34.048	16.226	
Zinc	27.132	11.172	
Ammonia (as N)	3,545.000	1,559.000	

(e) <u>Precipitation and</u> <u>Filtration of</u> <u>Photographic</u> <u>Solutions</u> <u>Wet Air Pollution</u> <u>Control</u> <u>PSES</u>

Pollutant or	Maximum	for	Maximum for
Pollutant Property	Any One	Day	Monthly Average

mg/troy ounce of silver from precipitation and filtration of photographic solutions

Copper	15.540	7.406
Zinc	12.380	5.099
Ammonia (as N)	1,618.000	711.400

(f) <u>Electrolytic</u> <u>Refining</u> PSES

		Maximu	m for	Mavim	um for
Pollutant or Pollutant Property		Any On		Monthly	
mg/troy ounce of silver	from	electr	olytic	refining	• •
Copper		· ·	0.973		0.464
Zinc	· . ·		0.775		0.319
Ammonia (as N)		10	1.300		44.540
				• • • • • • • • • • • • • • • • • • •	. <u>.</u>
(g) <u>Furnace Wet</u> <u>Air</u> <u>Pollutior</u>	n <u>Cont</u>	<u>rol</u> I	SES	:	
Pollutant or Pollutant Property		Maximu Any Or	im for le Day	Maxim Monthly	um for Averag
mg/troy ounce of silver	r roas	ted, sn	elted,	or dried	
Copper			0		0
Zinc	÷		0	ан сайта (с. 1916) 1917 — Прила Прила (с. 1917) 1917 — Прила (с. 1917)	0
Ammonia (as N)			0		0
		· · · · · · · · · · · · · · · · · · ·			
(h) <u>Leaching</u> PSES					
Pollutant or	· · · · · · · · · · · · · · · · · · ·		im for	Maxim Monthly	um for

Pollutar	nt Propert	з у		•	Any Or	ne Day	Monthly	Average
	mg/troy	ounce	of	silver	produced	from	leaching	
Copper Zinc Ammonia	(as N)		×	•	· · · · · · · · · · · · · · · · · · ·	0.110 0.088 1.470		0.053 0.036 5.040

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(i) <u>Leaching Wet Air Pollution Control and Precipitation of</u> <u>Nonphotographic</u> <u>Solutions Wet Air Pollution</u> <u>Control</u> PSES

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average

mg/troy ounce of silver produced from leaching or silver precipitated

Copper Zinc	5.671	2.703
Ammonia (as N)	4.519 590.500	1.861 259.600

(j) <u>Precipitation and Filtration of Nonphotographic</u> Solutions PSES

Pollutant or		Maximum for	Maximum for	
Pollutant Property		Any One Day	Monthly Average	
mg/troy	ounce of silver precipi	tated	······	
Copper	(as N)	3.930	1.873	
Zinc		3.132	1.290	
Ammonia		409.300	179.900	

(k) Floor and Equipment Washdown PSES

	Pollutant or Pollutant Property			Maximum for Any One Day			Maximum for Monthly Average	
		mg/troy	ounce	of silver	prod	uction		
Copper Zinc Ammonia	(as N)				0.000)	(0.000 0.000 0.000

PSNS are promulgated based on the performance achievable by the application of chemical precipitation, sedimentation, and multimedia filtration (lime, settle, and filter) technology and in-process flow reduction control methods, along with complete recycle of treated floor and equipment washdown water and preliminary treatment consisting of ammonia steam stripping for selected waste streams. The following pretreatment standards are promulgated for new sources:

(a) Film Stripping PSNS

	tant or t Property				imum One	for Day	Maxim Monthly	um for Average
	mg/troy	ounce o	f silver	from	filı	n str	ipping	
Cannor					64	450		30.720

Copper		64.450		30.720
Zinc	к. В	51.360		21.150
Ammonia(as N)		6,712.000	2,9	51.000
• •				

(b) Film Stripping Wet Air Pollution Control and Precipitation and Filtration of Film Stripping Solutions Wet Air Pollution Control PSNS

Pollutant or ,	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
mg/troy ounce of silver from p of film strippi	precipitation an ing solutions	nd filtration

1.242	0.592
0.990	0.408
129.300	56.840
	0.990

(c) <u>Precipitation and Filtration of Film Stripping</u> Solutions PSNS

Pollut		Maximum for	Maximum for	
Pollutant		Any One Day M	onthly Average	
· · ·	mg/troy ounce of	silver precipitated		
Copper	as N)	73.690	35.120	
Zinc		58.720	24.180	
Ammonia (7,674.000	3,374.000	

(d) <u>Precipitation and Filtration of Photographic Solutions</u> PSNS

Pollutant or Pollutant Property			Maximum for Any One Day N			Maximum for Monthly Average			
		mg/troy	ounce	of	silver	preci	pitated		
Copper Zinc Ammonia	(as	N)					048 132 000	1,!	16.226 11.172 559.000

(e) <u>Precipitation and Filtration of Photographic</u> Solutions <u>Wet Air Pollution</u> Control PSNS

Pollutant or	Maximum fo	r Maximum for
Pollutant Property	Any One Da	y Monthly Average

mg/troy ounce of silver from precipitation and filtration of photographic solutions

Copper	15.540	7.406
Zinc	12.380	5.099
Ammonia (as N)	1,618.000	711.400

(f) Electrolytic Refining PSNS

Pollutant or Pollutant Property		· · · · · · · · · · · · · · · · · · ·		Maximum for Any One Day		Maximum for Monthly Average			
<u> </u>	mg/troy	ounce	of	silver	from	electrol	lytic	refining	
Copper Zinc Ammonia	a (as N)			•			973 775 300		0.464 0.319 44.540

		Max	mum for	Maximu	m for
Pollutant or	an a		One Day	Monthly	
ollutant Property	۰	Any	one buj		
mg/troy ounce	of gilver	roasted.	smelted.	or dried	`
mg/troy bunce	Of SITAGE	LOADECU	D	•• •• ••	. •
Copper			0.000		. 0.000
linc			0.000	•	0.000
Ammonia (as N)			0.000	·	0.000
				د ر م	-
(h) Leaching PSNS				5 1	
(ii) <u>Leaching</u> Fond			<u>.</u>	· ·	
Pollutant or	· ·		imum for		um for
Pollutant Property	a set and a set a	Any	One Day	Monthly	Average
	······			aachina	
mg/troy ound	e of silve	r produc	ed from f	eaching	
Copper			0.110		0.053
Zinc	,		0.088		0.036
Ammonia (as N)			11.470		5.040
				· · · · ·	· · ·
(i) Leaching Wet Ai	Pollution	Control	and Prec	ipitation	of
(i) <u>Leaching Wet Air</u> Nonphotographic	Pollution Solutions	<u>Control</u> Wet <u>Air</u>	and Prec Pollution	ipitation Control	of PSNS
Nonphotographic	Pollution Solutions	Wet Air	Pollution	CONTROL	POND
Nonphotographic Pollutant or	Pollution Solutions	<u>Wet Air</u> Max	imum for	Maxim	um for
Nonphotographic	Pollution Solutions	<u>Wet Air</u> Max	Pollution	Maxim	um for
Nonphotographic Pollutant or Pollutant Property	Solutions	Wet Air Max Any	imum for One Day	<u>Maxim</u> Monthly	um for Averag
Nonphotographic Pollutant or	<u>Solutions</u> silver pro	Wet Air Max Any	imum for One Day	<u>Maxim</u> Monthly	um for Averag
Nonphotographic Pollutant or Pollutant Property mg/troy ounce of	<u>Solutions</u> silver pro	Wet Air Max Any oduced fr	imum for One Day om leachi	<u>Maxim</u> Monthly	um for Averag ver
Nonphotographic Pollutant or Pollutant Property mg/troy ounce of Copper	<u>Solutions</u> silver pro	Wet Air Max Any oduced fr	imum for One Day om leachi 5.671	<u>Maxim</u> Monthly	um for Averag ver 2.703
Nonphotographic Pollutant or Pollutant Property mg/troy ounce of Copper Zinc	<u>Solutions</u> silver pro	Wet Air Max Any oduced fr	imum for One Day om leachi	<u>Maxim</u> Monthly	um for Averag
Nonphotographic Pollutant or Pollutant Property mg/troy ounce of Copper Zinc	<u>Solutions</u> silver pro	Wet Air Max Any oduced fr	imum for One Day om leachi 5.671 4.519	<u>Maxim</u> Monthly	ver 2.703
Nonphotographic Pollutant or Pollutant Property mg/troy ounce of Copper	<u>Solutions</u> silver pro	Wet Air Max Any oduced fr	imum for One Day om leachi 5.671 4.519	<u>Maxim</u> Monthly	ver 2.703
Nonphotographic Pollutant or Pollutant Property mg/troy ounce of Copper Zinc	<u>Solutions</u> silver pro	Wet Air Max Any oduced fr	imum for One Day om leachi 5.671 4.519	<u>Maxim</u> Monthly	ver 2.703

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
mg/troy ounce	of silver precipitate	đ
Copper	3.930	1.873
Zinc	3.132	1.290
Ammonia (as N)	409.300	179.900

(j) <u>Precipitation and Filtration of Nonphotographic</u> Solutions PSNS

(k) Floor and Equipment Washdown PSNS

Pollutant or Pollutant Property			Maximum for Any One Day	Maximum for Monthly Average	
		mg/troy ounce of	silver production		
Copper Zinc Ammonia	(as N)		0.000 0.000 0.000	0.000 0.000 0.000	

SECTION III

SUBCATEGORY PROFILE

This section of the secondary silver supplement describes the raw materials and processes used in refining secondary silver and presents a profile of the secondary silver plants identified in this study.

The production of secondary silver can be divided into two processing types based on the source of raw materials: photographic and nonphotographic. Photographic processes for recovering silver include film stripping and precipitation, film incineration, chemical precipitation from solution, metallic replacement in solution, and direct electrolytic refining. Nonphotographic manufacturing involves precipitation of silver from waste plating solutions, melting and casting of sterling silver scrap, and processing electrical component scrap.

RAW MATERIALS

The principal raw materials used by plants recovering silver from photographic materials are discarded photographic film (both color and black and white), scrap photographic papers, and silver-rich sludges and solutions from photographic processing. Waste plating solutions, sterling ware scrap, and electrical component scrap are the principal raw materials used in the nonphotographic category.

PHOTOGRAPHIC MATERIALS

Photographic raw materials silver recovery can be divided into two primary sources, discarded film and photographic papers, and film processing solutions. There are many different processes for recovering silver from photographic materials. The most common methods are discussed below.

The silver in the emulsion on discarded film can be recovered by stripping and precipitation, or incineration. Figure III-1 (page 2697) represents a general flow diagram of photographic film scrap processes. The primary steps are:

- 1. Granulation,
- 2. Stripping,
- 3. Sedimentation and filtration,
- 4. Precipitation,
- 5. Roasting,
- 6. Casting,
- 7. Purification, and
- 8. Melting and casting.

Photographic film can be stripped directly or first shredded and granulated. Dust generated by granulation is collected with a

baghouse and recycled to the precipitation step further along in the process. The film can be stripped of the silver-bearing emulsion in a number of ways. In one method, the film is stripped using nitric acid, resulting in a silver nitrate The reaction of emulsion with nitric acid produces solution. nitrogen-containing air emissions (NO_X), which are removed with a scrubber, resulting in a wastewater stream. Another method uses wet oxidation with a catalyst at high temperature and pressure to produce a silver liquor. A third stripping process converts silver in the film to silver chloride using ferric chloride solution containing hydrochloric acid.

A silver-rich solution is usually separated from the granulated base (residue) by sedimentation, decantation, film and filtration. The residue is discarded as solid waste, usually in a landfill. Some plants salvage the film base with washing, dewatering, and drying steps. The film base can be reused as a raw material for new photographic film. Wastewater can be generated during the washing of the film base.

Silver in solution can be precipitated by various precipitating agents. Caustic soda, soda ash (Na2CO3), and proteolytic enzymes are commonly used. Alum is used as a flocculating agent in some processes. The addition of chloride ion will precipitate silver chloride which can be reduced to silver by direct hydrogen reduction with gaseous hydrogen under high temperature and Thiosulfate solution also converts silver chloride to pressure. a soluble silver complex, silver thiosulfate, from which the silver may be electrodeposited. Recovered baghouse dust from the granulation step may also be added during the precipitation step.

silver-free supernatant is decanted and is a source of The wastewater. Silver sludge is dewatered by gravity or filter thickening, vacuum filtration, centrifuging, or drying. The water removed is sent to waste treatment. Alkaline or acidic fumes emitted from the precipitation step are scrubbed, resulting in a wastewater stream. Silver sludge filtration and washing produces another silver-free wastewater stream.

The dried cake is roasted in a reverberatory furnace. The impure silver is then cast into ingots or dore plates. The furnace slag is crushed and classified, and the silver concentrate recycled as furnace feed, while the tailings are landfilled. Most processes have baghouses for control of particulates in furnace off-gases. Some use scrubbers and electrostatic precipitators. Contact cooling water is used at some plants during casting. Other plants air cool the ingots on dore plates.

Dore plates are electrolytically refined on-site or shipped to other facilities. The electrolytic purification is carried out The electrolytic purification is carried out in either Balbach-Thum cells (horizontal electrodes) or Moebius cells (vertical electrodes). A current is passed between an anode and a cathode which are suspended in solution. Silver collects on the cathode. A typical electrolyte solution consists of silver nitrate and a small amount of nitric acid. The

electrolyte is kept slightly to mildly acidic, a pH range of approximately 2 to 6. The refined silver is periodically scraped from the cathode and washed to remove residual electrolyte. In addition to refined silver, electrolysis produces a waste stream of spent electrolyte and wash water, and a slime containing precious metals such as gold and platinum. The slime may be further refined for precious metals recovery.

The refined silver is dried, melted in a melting furnace, and cast as ingots. Pollution control of furnace off-gases is handled with a baghouse, scrubber, or electrostatic precipitator. Contact cooling water is used in the casting steps, as well as casting scrubbers which produce wastewater streams.

Photographic film and photographic papers may be incinerated, rather than processed by granulation, stripping, and The temperature and rate of burning must be precipitation. carefully controlled if high efficiency is to be maintained. Air emissions include organic vapors from the volatilization and decomposition of organic scrap contaminants, as well as sometimes The emissions are combustion gases and dust. controlled by afterburners; at some plants, afterburners are used in series with a baghouse or scrubber. Wet scrubbing techniques produce a wastewater discharge. Silver-bearing ash is then fed directly to roasting and the impure silver is cast and electrolytically refined as described above. Some refineries buy silver-bearing ash from scrap dealers.

There are three basic methods for recovering silver from photographic processing solutions: chemical precipitation, metallic replacement, and direct electrolytic refining. Silver recovery from baths has also been successful by adsorption from solution by ion exchange. Reverse osmosis has been used on dilute solutions.

Silver-rich solutions from photographic film developing and manufacturing undergo precipitation and purification as described above. One alternate method uses sulfide compounds, particularly sodium sulfide as the precipitating agent. Gaseous emissions, such as hydrogen sulfide, are controlled with a wet scrubber, resulting in a wastewater stream. The subsequent process for silver recovery is identical to other precipitation methods.

Silver ions can be effectively reduced from solution to a solid state by a replacement reaction. Any metal more active than silver will go into solution as an ion, while the silver ion becomes solid metal. Zinc, aluminum, copper, and iron are commonly used to recover silver by replacement from photographic fixing solutions. The silver sludge produced can be filtered, roasted, and cast as described previously.

Although used as a purification step in other recovery processes, electrolytic refining is also a direct means of silver recovery. In the electrolytic method, a current is passed between an anode and a cathode which are suspended in a solution which contains

greater than 1.0 mg/l of silver. Solutions containing silver below this concentration are difficult to refine electrolytically. Silver, about 99 percent pure, collects on the cathode. The cathode is periodically stripped to recover the If the current density is too high for the amount of silver. silver in the solution, thiosulfate in solution will decompose, forming silver sulfide. This reduces current efficiency and will render the regenerated solution unsuitable for reuse. Spent electrolyte solution is discarded or further refined for other precious metals. If the thiosulfate in solution is allowed to decompose, gaseous sulfur emissions (SO_x), must be controlled with a scrubber. Wastewater may also be generated from washing the silver scraped off the cathode.

NONPHOTOGRAPHIC MATERIALS

Based on the source of raw materials, the nonphotographic materials category can be divided into three basic processes for recovery of silver: precipitation of waste plating the solutions, melting of silverware, dental scrap, coins, and sterling-silver industry scrap (e.g., turnings, sweeps), and refining of electrical components scrap.

Silver-plated tableware is produced by electroplating silver from cyanide solutions onto preformed shapes made of tin, iron, zinc, or copper. Silver wastes generated are spills of silver-rich electrolyte, dilute wash solutions, and spent electrolyte. Cyanide plating solutions are treated to precipitate the silver and oxidize the cyanide. As shown in Figure III-2 (page 2698), the process consists of precipitation, filtration and washing, drying or roasting, casting, refining, and recasting. Some processors cast the silver before refining and sell the ingots to other refiners.

Precipitation is usually accomplished by addition of sodium hypochlorite, resulting in silver chloride. After settling, the silver chloride is washed, filtered, and dried to be sold as product or further processed with methods similar to those used for photographic silver precipitates. The cyanide left in solution may be oxidized with sodium hypochlorite and lime and discarded as wastewater. Wastewater streams also result from waste washing water and the filtrate and dewatering wastes. Wet scrubbers are used to control fumes from the precipitation and filtration steps. Roasting and melting furnaces also require air pollution control to remove particulates. Baghouses are usually used.

An alternate silver recovery method is precipitation of silver as the metal, using zinc metal with sodium chloride solution. The subsequent steps are identical to other precipitation processes.

The solid waste products from the sterling silver industry include defective tableware, trimmings, turnings, punchings, fumes, spillage, drosses from melting and casting, and dusts. The different wastes vary in impurity and the relatively pure

materials are melted, assayed, and reused. Lower quality wastes are combined, melted and cast, and the bullions are electrolytically refined as described above.

Silver scrap from electrical components includes electrical contacts, wire, silver-bearing batteries, condensers and solders. Figure III-3 (page 2699) shows typical production processes followed if electrical scrap is not suitable for electrolytic refining. Silver is recovered from electrical component scrap using one of three methods.

In the first method, the base metals are leached from the silver residue after smelting. After careful sorting and sampling, the scrap is smelted in a reverberatory furnace to produce lead bullion, copper matte, and slag. The slag is smelted in a blast furnace to separate the lead and copper portions, which are recycled. Blast furnace slag is discarded. Dust and fumes from both the reverberatory and blast furnaces are collected and The lead bullion from the reverberatory smelting recycled. lead from the blast furnace is fed to а furnace and reverberatory-type cupola furnace. The cupellation produces litharge and precious metal layers. The litharge is sent to a lead refinery or reduced for recycle to the reverberatory The cupola furnace requires a baghouse or smelting unit. scrubber to remove emission gas pollutants. The precious metal layer is cast into anodes (dore plates) for electrolytic The silver collects on the cathodes, which are melted refining. and cast as refined ingots. The slime residue, containing gold and platinum, is further refined. The spent electrolyte solution may be discarded as waste. Wastewater streams are also generated if contact cooling water is used in casting, and if melting furnace and casting scrubbers, which remove particulates emitted from these operations, are used. The copper matte from the initial smelting is crushed, ground, roasted, and leached. A wet scrubber may be used to control particulate air emissions from the roasting furnace, producing a wastewater stream. Leaching may be effected with sulfuric or hydrochloric acid. The leaching agent dissolves the base metals, leaving silver as a residue which can be filtered and washed for further processing. This leaching operation usually produces two wastewater streams: а silver-free leachate, which may be discharged or recycled, and a scrubber discharge stream.

In the second leaching process, silver is dissolved or stripped directly from the electrical component parts and later precipitated from solution. Nitric acid is the most common stripping agent. This leaching also results in two wastewater streams: a lead-iron residue and a scrubber discharge stream, resulting from the control of acid fumes.

In the third method, the base metals are dissolved directly using hydrochloric or sulfuric acid. The silver residue is then roasted, cast, and electrolytically refined. A scrubber stream, from control of acid fumes, and the lead-iron residue stream are sources of wastewater.

Silver in solution from leaching or direct stripping is precipitated by metallic replacement (usually with copper or zinc) and then filtered. Copper sulfate or zinc sulfate is usually the principal constituent of the supernatant and filtrate and is either purified for copper recovery or discarded. Water from washing the silver precipitate is also discharged. Wet scrubbers may provide control of acidic fumes emitted during the precipitation step, producing an additional wastewater stream.

The recovered silver is melted in a furnace and cast as ingots. Silver of insufficient purity is electrolytically refined. Particulate emissions from the melting furnace are controlled with a baghouse or scrubber. Venturi scrubbers are commonly used and a wastewater stream is discharged.

Silver-rich sludges from waste plating solutions, stripping solutions, and photographic solutions are leached and the silver recovered, resulting in a silver-rich solution. The leaching agent used is hydrochloric acid, sulfuric acid, or nitric acid. The silver-rich solution is processed through precipitation, filtration, roasting, melting, and casting steps to produce refined silver ingots.

PROCESS WASTEWATER SOURCES

The principal uses of water in secondary silver plants are:

- 1. Film stripping,
- 2. Film stripping wet air pollution control and precipitation of film stripping solutions wet air pollution control,
- 3. Precipitation and filtration of film stripping solutions,
- 4. Precipitation and filtration of photographic solutions,
- 5. Precipitation and filtration of photographic solutions wet air pollution control,
- 6. Electrolytic refining,
- 7. Furnace wet air pollution control,
- 8. Leaching,
- 9. Leaching wet air pollution control and precipitation of nonphotographic solutions wet air pollution control,
- 10. Precipitation and filtration of nonphotographic solutions, and
- 11. Floor and equipment washdown.

OTHER WASTEWATER SOURCES

There may be other wastewater streams associated with the production of secondary silver. These wastewater streams may include maintenance and cleanup water, and direct electrolytic refining wet air pollution control wastewater. These wastewater streams are not considered as part of this rulemaking. EPA believes that the flows and pollutant loadings associated with these streams are insignificant relative to the waste streams

SECONDARY SILVER SUBCATEGORY

selected and are best handled by the appropriate permit authority on a case-by-case basis under the authority of Section 402 of the Clean Water Act.

Casting contact cooling water is not considered as part of this rulemaking because, although several plants do discharge this stream, sampling data from an integrated secondary silver secondary precious metals plant indicate that the pollutant loadings are insignificant compared with the other waste streams selected.

AGE, PRODUCTION, AND PROCESS PROFILE

Of the 61 plants recovering silver from photographic and nonphotographic materials, Figure III-4 (page 2700) shows that the plants are concentrated in the Northeast and California, with plants also located in Idaho, Utah, Louisiana, Florida, and Texas.

Table III-1 (page 2694) summarizes the general type and shows the relative ages of the secondary silver plants. Seven plants discharge directly, 26 are indirect dischargers, and 28 are zero dischargers. Of the discharging plants, five process only photographic materials, 26 process only nonphotographic materials, and two plants process both types of materials. The average plant age is between 10 and 25 years.

Table III-2 (page 2695) shows the production ranges for the 61 secondary silver plants. Over half of the plants that reported production data produce less than 50,000 troy ounces of silver per year. Eight of these plants produce over 1,000,000 troy ounces of silver per year.

Table III-3 (page 2696) provides a summary of the plants having the various secondary silver processes. The number of plants generating wastewater from the processes is also shown.

Table III-1

INITIAL OPERATING YEAR (RANGE) SUMMARY OF PLANTS IN THE SECONDARY SILVER SUBCATEGORY BY DISCHARGE TYPE

Type of Plant <u>Discharge</u>	1983- 1974 <u>0-10</u>	1973- 1969 <u>10-15</u>	1968- 1959 <u>15-25</u>	1958- 1949 <u>25-35</u>	1948- 1939 <u>35-45</u>	1938- 1929 <u>45-55</u>	1928- 1919 <u>55-65</u>	1918- 1904 <u>65-80</u>	Before 1904 <u>80+</u>	Not Reported in dcp	<u>Total</u>
Direct	0	1	0	2	0	1	1	0	0	2	7
Indirect	8	6	4	1	1	·· 0	1	3	. 1	1	26
Zero	_8_	_3	_7	<u>1</u>	<u>1</u>	<u>0</u>	<u>0</u>	<u>0</u>	<u>0</u>	_8	<u>28</u>
Total	16	10	- 11	4	2	1	2	3	1	11	61

SECONDARY SILVER SUBCATEGORY SECT TTT

Table III-2

PRODUCTION RANGES FOR THE SECONDARY SILVER SUBCATEGORY

	<u></u>	Silver	Production R	anges (troy o	unces/year)	
Type of Plant	0 - 50,000	50,001- 100,000	100,001- 500,000	500,001- 1,000,000	1,000,000+	Not Reported in dcp
Direct Discharger	3	0	2	0	1	1
Indirect Discharger	21	0	3	1	1	0
Zero Discharger	_5	<u>1</u>	_7	<u>1</u>	<u>6</u>	8
Total	29	1	12	2	8	9

2695

SECONDARY SILVER SUBCATEGORY SECT -III

Table III-3

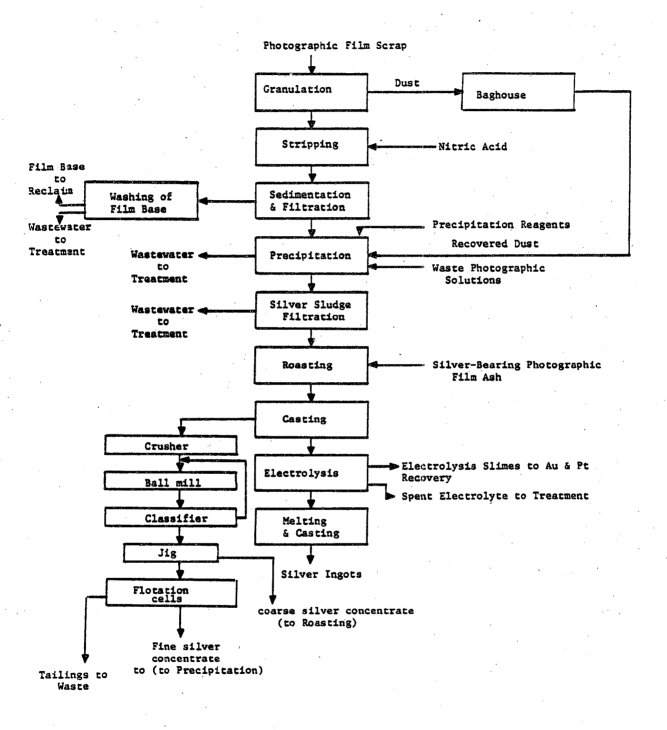
SUMMARY OF SECONDARY SILVER SUBCATEGORY PROCESSES AND ASSOCIATED WASTE STREAMS

Process	P1	umber of ants With Process	Number of Plants Generating <u>Wastewater</u> *
File Stripping o Film Base Processing o Precipitation and Filtration o Air Pollution Control		5 3 5 1	1 4 1
Photographic Solution Processing o Precipitation and Filtration o Air Pollution Control		7 7 3	- 7 3
Electrolytic Refining		15	13
Furnace Air Pollution Control	- ;	27	12
Casting		34	16
Leaching o Air Pollution Control		13 13†	11 13
Nonphotographic Solution Processing o Precipitation and Filtration		27 27	27

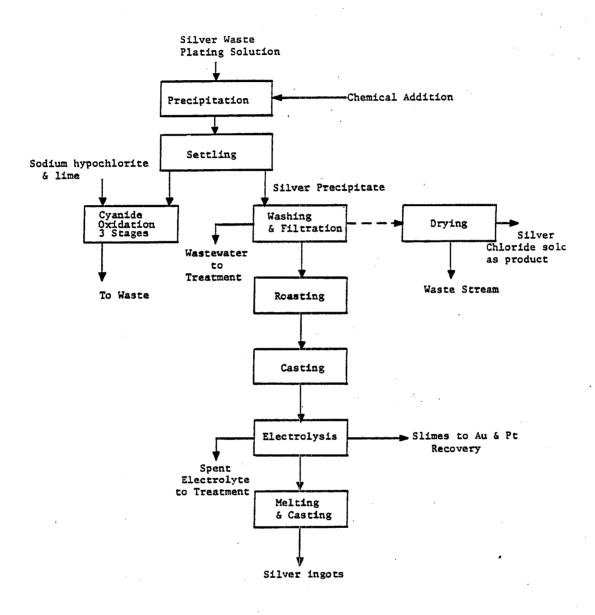
* Through reuse or evaporation practices, a plant may "generate" a wastewater from a particular process but not discharge it.

† Includes two scrubbers over leaching and precipitation of nonphotographic solutions
processes

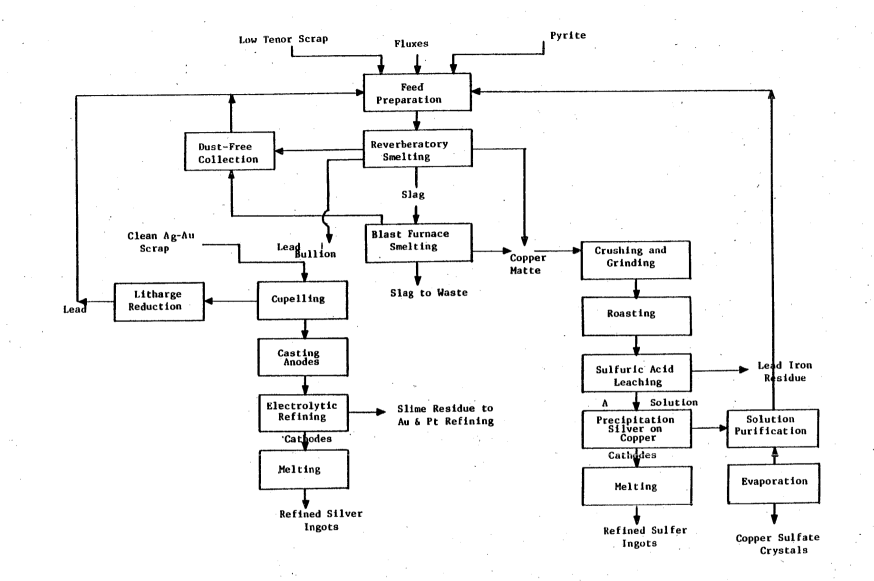
SECONDARY SILVER SUBCATEGORY SECT I. III



SILVER REFINING FROM PHOTOGRAPHIC MATERIALS



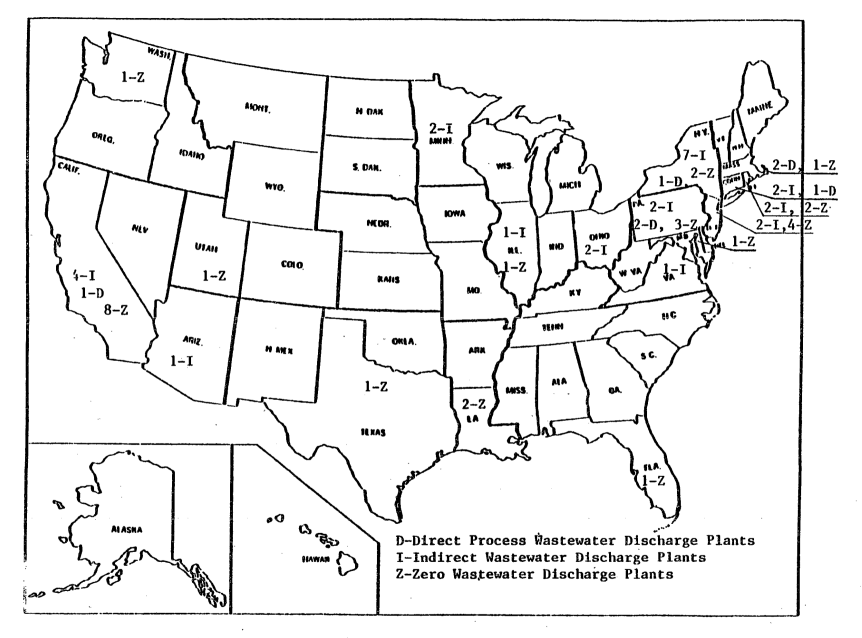
SILVER REFINING FROM WASTE PLATING SOLUTIONS



SECONDARY SILVER PRODUCTION PROCESS FROM NONPHOTOGRAPHIC SCRAP

2699

SECONDARY SILVER SUBCATEGORY SECT - III



GEOGRAPHIC LOCATIONS OF SECONDARY SILVER PLANTS

SECONDARY SILVER SUBCATEGORY SECT T H H H H

2700

SECTION IV

SUBCATEGORIZATION

This section summarizes the factors considered during the designation of the secondary silver subcategory and its related subdivisions.

The factors listed for general subcategorization were each evaluated when considering subdivision of the secondary silver subcategory. In the discussion that follows, the factors will be described as they pertain to this particular subcategory.

The rationale for considering segmentation of the secondary silver subcategory is based primarily on the production processes Within the subcategory, a number of different operations used. which may or may not have a water use or are performed, discharge, and which may require the establishment of separate effluent limitations and standards. While the secondary silver industry is considered a single subcategory, a more thorough examination of the production processes, water use and discharge practices, and pollutant generation rates has illustrated the need for limitations and standards based on a specific set of streams. Limitations and standards will be based wastewater on specific flow allowances for the following subdivisions:

- 1. Film stripping,
- 2. Film stripping wet air pollution control and precipitation of film stripping solutions wet air pollution control,
- 3. Precipitation and filtration of film stripping solutions,
- 4. Precipitation and filtration of photographic solutions,
- 5. Precipitation and filtration of photographic solutions wet air pollution control,
- 6. Electrolytic refining,
- 7. Furnace wet air pollution control,
- 8. Leaching,
- 9. Leaching wet air pollution control and precipitation of nonphotographic solutions wet air pollution control,
- 10. Precipitation and filtration of nonphotographic solutions, and
- 11. Floor and equipment washdown.

A number of other factors considered in this evaluation were shown to be inappropriate bases for further segmentation. Air pollution control methods, treatment costs, nonwater quality aspects, and total energy requirements are functions of the selected subcategorization factors -- raw materials and production processes. As such, they support the method of subcategorization which has been developed. Factors determined to be inappropriate for use as bases for subcategorization are discussed briefly below.

PLANT SIZE

It is difficult to categorize secondary silver plants on the basis of size. The individual processes involved in silver production often process different amounts of silver-bearing material. Therefore, it is more appropriate to categorize silver plants on the basis of process production, e.g., precipitation production.

PLANT AGE

Plants within the secondary silver subcategory differ in age, in terms of initial operating year. However, to remain competitive, plants are constantly modernized. Modifications to process operations have been made, resulting in greater production efficiency and reduced air pollution emissions. As a result, neither the concentration of constituents in wastewater nor the capability to meet the limitations is related to plant age.

PRODUCTION NORMALIZING PARAMETERS

The effluent limitations and standards developed in this document establish mass limitations for the discharge of specific pollutant parameters. To allow these limitations to be applied to plants with various production capacities, the mass of pollutant discharged must be related to a unit of production. This factor is known as the production normalizing parameter (PNP). In general, the actual silver production from the respective manufacturing process is used as the PNP. This is based on the principle that the amount of water generated is proportional to the amount of product made. Therefore, the PNPs for the ll secondary silver subdivisions are as follows:

Building Block

PNP

1.	Film stripping	troy ounces of silver produced from film stripping
2.	Film stripping wet air pollu- tion control and precipita- tion of film stripping solu- tions wet air pollution control	troy ounces of silver produced from precipitation and filtra- tion of film stripping solutions
3.	Precipitation and filtration of film stripping solutions	troy ounces of silver precipitated
4.	Precipitation and filtration of photographic solutions	troy ounces of silver precipitated

5. Precipitation and filtration troy ounces of silver of photographic solutions precipitated wet air pollution control

- 6. Electrolytic refining
- 7. Furnace wet air pollution control
- 8. Leaching
- 9. Leaching wet air pollutio control and precipitation of nonphotographic solutions wet air pollution control
- 10. Precipitation and filtration of non-photographic solutions
- 11. Floor and equipment washdown

troy ounces of silver from electrolytic refining

troy ounces of silver smelted, roasted, or dried

troy ounces of silver produced from leaching

troy ounces of silver produced from leaching or silver precipitated

troy ounces of silver precipitated

troy ounces of silver produced

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SECTION V

WATER USE AND WASTEWATER CHARACTERISTICS

This section describes the characteristics of wastewater associated with the secondary silver subcategory. Data used to quantify wastewater flow and pollutant concentrations are presented, summarized, and discussed. The contribution of specific production processes to the overall wastewater discharge from secondary silver plants is identified whenever possible.

Two principal data sources were used in the development of effluent limitations and standards for this subcategory; data collection portfolios (dcp) and field sampling results. Data collection portfolios contain information regarding wastewater flows and production levels. Data were also collected through comments and a special request.

In order to quantify the pollutant discharge from secondary silver plants, a field sampling program was conducted. A complete list of the pollutants considered and a summary of the techniques used in sampling and laboratory analyses are included in Section V of Vol. I. Wastewater samples were collected in two screening and verification. The first phase, screen phases: sampling, was to identify which toxic pollutants were present in the wastewaters from production of the various metals. Screening samples were analyzed for 125 of the 126 toxic pollutants and other pollutants deemed appropriate. (Because the analytical standard for TCDD was judged to be too hazardous to be made generally available, samples were never analyzed for this There is no reason to expect that TCDD would be pollutant. present in secondary silver wastewater). A total of 10 plants were selected for screen sampling in the nonferrous metals manufacturing category, one of these being a secondary silver Of the 36 plants selected for verification sampling, plant. three were from the secondary silver subcategory. In general, the samples were analyzed for three classes of pollutants: toxic pollutants, toxic metal pollutants, and criteria organic pollutants (which includes both conventional and nonconventional pollutants).

As described in Section IV of this supplement, the secondary silver subcategory has been further segmented into 11 subdivisions or building blocks, so that the promulgated regulation contains mass discharge limitations and standards for manufacturing processes discharging process wastewater. 11 Differences in the wastewater characteristics associated with these subdivisions are to be expected. For this reason, streams corresponding to each subdivision are wastewater addressed separately in the discussions that follow.

The wastewater data presented in this section were evaluated in light of production process information compiled during this

study. As a result, it was possible to identify the principal wastewater sources in the secondary silver subcategory. They are:

- 1. Film stripping,
- 2. Film stripping wet air pollution control and precipi-
- tation of film stripping solutions wet air pollution control,
- 3. Precipitation and filtration of film stripping solutions,
- Precipitation and filtration of photographic solutions,
 Precipitation and filtration of photographic solutions
- 5. Precipitation and filtration of photographic solutions wet air pollution control,
- 6. Electrolytic refining,
- 7. Furnace wet air pollution control,
- 8. Leaching,
- 9. Leaching wet air pollution control and precipitation of nonphotographic solutions wet air pollution control,
- 10. Precipitation and filtration of nonphotographic solutions, and
- 11. Floor and equipment washdown.

In the proposed development document, separate subdivisions were identified for precipitation and filtration of film stripping solutions wet air pollution control, casting contact cooling water, casting wet air pollution control, and precipitation and filtration of nonphotographic solutions wet air pollution control. Based on new data gathered since proposal from secondary precious metals plants' dcp, sampling at two integrated secondary silver and secondary precious metals plants, and re-evaluation of existing data, these subdivisions were either combined with other subdivisions or deleted. A subdivision for floor and equipment washdown also was added. The reasons for these changes are discussed in detail in Section IX - Wastewater Discharge Rates.

Although flow and production data were collected in secondary precious metals dcp in the nonferrous metals manufacturing category, these data were not used to modify the proposed regulatory flow allowances. The new data support the proposed flow allowance and the Agency did not receive any comments suggesting that the allowances should be revised. The new flow data are included in the water use and discharge tables at the end of this section.

Additionally, since the dcp were collected, the Agency has learned that one secondary silver plant has closed and six plants no longer process secondary silver. Flow and production data (when available) for these plants are presented in this section and in the remainder of the development document. These data are an integral part of the BPT and BAT effluent limitations because the processes remain representative in determining BPT and BAT and flow rates. Therefore, it is necessary to present this information so that the effluent limitations are documented.

WASTEWATER FLOW RATES

Data supplied by dcp responses were evaluated, and two flow-to-production ratios were calculated for each stream. The two The two flow, and wastewater discharge are water use ratios, differentiated by the flow value used in calculation. Water use is defined as the volume of water or other fluid (e.g., leachate, spent electrolyte) required for a given process per mass of silver product and is therefore based on the sum of recycle and make-up flows to a given process. Wastewater flow discharged after pretreatment or recycle (if these are present) is used in calculating the production normalized flow--the volume of wastewater discharged from a given process to further treatment, disposal, or discharge per mass of silver produced. Differences between the water use and wastewater flows associated with a given stream result from recycle, evaporation, and carryover on the product. The production values used in calculation correspond to the production normalizing parameter, PNP, assigned to each stream, as outlined in Section IV. The production normalized flows were compiled and statistically analyzed by stream type. Where appropriate, an attempt was made to identify factors that could account for variations in water use. This information is summarized in this section. A similar analysis of factors affecting the wastewater values is presented in Sections IX, Χ, XI, and XII where representative BPT, BAT, BDT, and pretreatment discharge flows are selected for use in calculating the effluent limitations and standards. As an example, silver precipitation scrubbing wastewater flow is related to filtration and precipitate production. As such, the discharge rate is expressed in liters of scrubber wastewater discharged per troy ounce of silver produced by precipitation.

WASTEWATER CHARACTERIZATION DATA

In order to quantify the concentrations of pollutants present in wastewater from secondary silver plants, wastewater samples were collected at four plants. Diagrams indicating the sampling sites and contributing production processes are shown in Figures V-1 through V-4 (pages 2714 - 2723).

The raw wastewater sampling data for the secondary silver subcategory are presented in Tables V-2, V-5, and V-8 (pages 2715, 2724, and 2728. Treated wastewater sampling data are shown in Tables V-13 through V-16 (pages 2736 - 2740). The stream codes presented in the tables may be used to identify the location of each of the samples on the process flow diagrams in Figures V-1 through V-4. Where no data are listed for a specific day of sampling, the wastewater samples for the stream were not collected. If the analysis did not detect a pollutant in a waste stream, the pollutant was omitted from the table.

The data tables include some samples measured at concentrations considered not quantifiable. The base-neutral extractable, acid fraction extractable, and volatile organics are generally considered not quantifiable at concentrations equal to or less

than 0.010 mg/l. Below this concentration, organic analytical results are not quantitatively accurate; however, the analyses are useful to indicate the presence of a particular pollutant. The pesticide fraction is considered not quantifiable at concentrations equal to or less than 0.005 mg/l. Nonquantifiable results are designated in the tables with an asterisk (double asterisk for pesticides).

These detection limits shown on the data tables are not the same in all cases as the published detection limits for these pollutants by the same analytical methods. The detection limits used were reported with the analytical data and hence are the appropriate limits to apply to the data. Detection limit variation can occur as a result of a number of laboratoryspecific, equipment-specific, and daily operator-specific factors. These factors can include day-to-day differences in machine calibration, variation in stock solutions, and variation in operators.

The statistical analysis of data includes some samples measured at concentrations considered not quantifiable. Data reported as an asterisk are considered as detected but below quantifiable concentrations, and a value of zero is used for averaging. Toxic organic, nonconventional, and conventional pollutant data reported with a "less than" sign are considered as detected, but not further quantifiable. A value of zero is also used for averaging. If a pollutant is reported as not detected, it is excluded in calculating the average. Finally, toxic metal values reported as less than a certain value were considered as not detected and a value of zero is used in the calculation of the average. For example, three samples reported as ND, *, and 0.021 mg/l have an average value of 0.010 mg/l.

The method by which each sample was collected is indicated by number, as follows:

1	one-time grab
2	24-hour manual composite
3	24-hour automatic composite
4	48-hour manual composite
5	48-hour automatic composite
6	72-hour manual composite
7	72-hour automatic composite

In the 1977 data collection portfolios, the secondary silver plants which discharge wastewater were asked to specify the presence or absence of the toxic pollutants in their effluent. Of the 44 secondary silver plants submitting a 1977 dcp, 19 did not respond to this portion of the questionnaire. All plants responding to the organic compounds portion of the questionnaire reported that all toxic organic pollutants were known to be absent or believed to be absent from their wastewater.

The responses for the toxic metals and cyanide are summarized below:

•	Known	Believed	Believed	Known
Antimony	2	4	14	5
Arsenic	1	2	16	6
Beryllium	0	2	16	7
Cadmium	4	5	10	6
Chromium	5	4	10	6
Copper	10	4	б	5
Cyanide	4	1	13	7
Lead	7	4	8	6
Mercury	1	2	16	6
Nickel	8	3	9	5
Selenium	1	2	15	· 7
Silver	13	5	3	· 4
Thallium	0	1	16	8
Zinc	10	4	7	. 4

WASTEWATER CHARACTERISTICS AND FLOWS BY SUBDIVISION

Since secondary silver production involves 11 principal sources of wastewater and each has potentially different characteristics and flows, the wastewater characteristics and discharge rates corresponding to each subdivision will be described separately. A brief description of why the associated production processes generate a wastewater and explanations for variations of water use within each subdivision will also be discussed.

FILM STRIPPING

Photographic film may be stripped of emulsion and the silver The film base can be screened and rinsed, precipitated. producing wastewater. Water discharge rates are presented in Table V-1 (page 2714) in liters per troy ounce of silver produced from film stripping. Table V-2 (page 2715) (stream 14) shows combined raw wastewater data from film stripping and wet air on film stripping and control film pollution stripping precipitation. Data are not available for separate waste streams because discrete points in each stream were not accessible. However, based on the combined wastewater data and the raw materials and process used, film stripping wastewater should contain toxic organics and metals, cyanide, and total suspended solids above treatable concentrations, as well as phenols.

FILM STRIPPING WET AIR POLLUTION CONTROL AND PRECIPITATION OF FILM STRIPPING SOLUTIONS WET AIR POLLUTION CONTROL

One plant engaged in film stripping uses a wet scrubber to control air emissions. This plant uses the same scrubber to control emissions from film stripping and film stripping precipitation. A 99+ percent recycle of the scrubber water is maintained and the discharge rate is 0.034 liters per troy ounce of silver produced from precipitation and filtration of film stripping solutions. Table V-2 (stream 14) shows combined raw wastewater data from film stripping and wet air pollution control on film stripping and film stripping precipitation. Data are not available for separate waste streams because discrete points in each stream were not accessible. However, based on the combined wastewater data and the raw materials and process used, film stripping wet air pollution control wastewater should contain toxic organics and metals, cyanide, phenolics, and total suspended solids.

PRECIPITATION AND FILTRATION OF FILM STRIPPING SOLUTIONS

The solution resulting from stripping granulated film is treated to precipitate the silver. After settling or filtration, the silver-free solution may be discarded as wastewater. Four of the five photographic plants that use this process discharge a waste stream. The water discharge rates, reported in liters per troy ounce of silver precipitated, are shown in Table V-3 (page 2722). Sampling data for film stripping solutions precipitation are summarized in Table V-2 (Stream 12). Raw wastewater from this process contains toxic organics and metals, cyanide, phenolics, and total suspended solids at treatable concentrations.

PRECIPITATION AND FILTRATION OF PHOTOGRAPHIC SOLUTIONS

Silver can be precipitated from discarded photographic hypo solutions. After filtration, the silver-free solution constitutes waste stream. Solutions resulting from a photographic sludges are also included in this subdivision. A11 seven plants which have precipitation processes discharge process The discharge rates from these plants, presented in wastewater. liters per troy ounce of silver precipitated, are shown in Table (page 2723). The Agency did not sample the raw wastewater V-4 from silver solution precipitation directly; however, wastewater samples were collected after filtering with sawdust (which is This part of the process). wastewater contains 1,2dichloroethane, chloroform, phthalates, and tetrachloroethylene, all above treatable concentrations (0.025 to 0.132 mg/l). Toxic metals are also found, including a high concentration of zinc (200 mg/l). Ammonia (4,630 mg/l), and chloride (734 mg/l) are Total suspended solids are evident, but most also present. solids in the raw wastewater were probably removed by the filter. Raw wastewater sampling data are given in Table V-5 (page 22724).

PRECIPITATION AND FILTRATION OF PHOTOGRAPHIC SOLUTIONS WET AIR POLLUTION CONTROL

Of the seven photographic silver plants precipitating silver solutions, three use wet air pollution control; two of these discharge wastewater from wet scrubbers. The water discharge flow rates are shown in Table V-6. Although wastewater samples were not collected from precipitation of photographic solutions wet air pollution control, raw wastewater data are available from a film stripping precipitation scrubber. The wastewater characteristics for the two scrubbers are expected to be similar

because of the similarities in the raw materials and processes used. Wastewater samples collected from the analogous wet scrubber stream contain toxic organics and metals, cyanide, and total suspended solids above treatable concentrations.

ELECTROLYTIC REFINING

Fifteen plants use electrolytic refining as a purification step in secondary silver processing. Thirteen plants generate waste streams consisting of spent electrolyte and wash water; 12 of those discharge the wastewater. Table V-7 (page 2725) shows the water discharge rates in liters per troy ounce of silver refined.

Electrolytic refining is similar for photographic and nonphotographic plants, therefore wastewater from each may have similar characteristics. Table V-8 (page 2728) summarizes the raw wastewater sampling data for the toxic and selected conventional and nonconventional pollutants.

The samples were collected at a nonphotographic plant from a combined waste stream comprised of raw wastewater from electrolytic refining, as well as metal-depleted solutions. This raw wastewater contains toxic organics and metals, ammonia, fluoride, cyanide, and total suspended solids above treatable concentrations.

FURNACE WET AIR POLLUTION CONTROL

Of the secondary silver plants with reverberatory furnaces, incinerators, or casting furnaces, 27 control off-gas emissions. Twelve plants use wet scrubbers, and four of these discharge wastewater, as shown in Table V-9 (page 2732). The Agency did not collect samples from furnace scrubber wastewater at a secondary silver plant before proposal. However, after proposal, samples of this wastewater were collected at an integrated secondary silver and secondary precious metals facility. This plant processes nonphotographic materials and the scrubber controls off-gas emissions from a furnace used to smelt silver, gold, platinum, and palladium. These data are presented in the administrative record supporting this regulation. Furnace scrubber wastewater contains a treatable concentration of total suspended solids (5,600 mg/l) and has a pH of approximately 7.

LEACHING

In nonphotographic materials plants, leaching is used to recover silver from silver sludges and copper matte associated with the melting of electrical component parts. Of the 13 nonphotographic plants that leach, 11 produce wastewater, consisting of either silver-free leachate or lead-iron residue. Water discharge rates are given in Table V-12 (page 2735) in liters per troy ounce of silver produced from leaching.

Table V-8 (page 2728) (stream 40) shows combined raw wastewater data from nonphotographic solutions precipitation and

electrolytic refining. Leaching wastewaters have similar characteristics as precipitation wastewater because of the nature of the nonphotographic materials processed. Data are not available for separate waste streams because discrete points in each stream were not accessible. However, based on the combined wastewater data and the raw materials and process used, raw wastewater from leaching should contain toxic organics and metals, ammonia, fluoride, cyanide, phenolics, and total suspended solids above treatable concentrations.

LEACHING WET AIR POLLUTION CONTROL AND PRECIPITATION OF NONPHOTOGRAPHIC SOLUTIONS WET AIR POLLUTION CONTROL

For leaching emissions, discharge rates are shown in Table V-13 (page 2736). All 13 of the plants with leaching and precipitation emissions control discharge wastewater. For proposal, the Agency did not have discrete samples of this waste stream. After proposal, EPA collected samples of scrubbers over leaching and precipitation processes at two integrated secondary silver and secondary precious metals plants. These plants also process gold, platinum, and palladium from nonphotographic materials. The scrubbers control emissions from the secondary silver and the secondary precious metals leaching and precipitation processes. The scrubber wastewater contains treatable concentrations of toxic metals and total suspended solids. The pH is approximately 8 when a caustic solution is used as a scrubber liquor. At plants scrubbing with water only, this wastewater is expected to be acidic. The wastewater data for this stream are presented in the administrative record supporting this regulation.

PRECIPITATION AND FILTRATION OF NONPHOTOGRAPHIC SOLUTIONS

Silver may be recovered by precipitation from leachates, waste silver-plating solutions or melted silver scrap. Twenty-seven nonphotographic plants report this process, and all 27 discharge wastewater. Depleted solutions may be discarded as wastewater, along with wash water and silver-free filtrates. Discharge water rates are presented in Table V-15 (page 2740).

Table V-8 (stream 40) shows combined raw wastewater data from nonphotographic solutions precipitation and electrolytic refining. Data are not available for separate waste streams because discrete points in each stream were not accessible. However, based on the combined wastewater data and the raw materials and process used, precipitation of nonphotographic solutions wastewater should be characterized by the presence of toxic organics and metals, ammonia, cyanide, chloride, fluoride, phenolics, and suspended solids above treatable concentrations.

FLOOR AND EQUIPMENT WASHDOWN

Many plants wash equipment and floors to recover silver values that may be contained in accidental leaks and spills of process solutions. Flow data on the floor wash were not generally available in the data collection portfolios. However, data from

a sampling effort at one secondary silver-secondary precious metals integrated plant and one secondary precious metals plant show that one liter per troy ounce of washdown water is typically generated. Sampling data collected at the two plants show this wastewater to contain treatable concentrations of toxic metals and total suspended solids. The flow and sampling data for this waste stream are included in the administrative record supporting this regulation.

TABLE V-1

WATER USE AND DISCHARGE RATES FOR FILM STRIPPING

(1/troy ounce of silver produced from film stripping)

Plant <u>Code</u>	Percent Recycle	Production Normalized Water <u>Use</u>	Production Normalized Discharge Flow
30927	0	50.35	50.35
596	NR	NR	NR
74	NR	NR	NR

NR = data not reported in dcp.

Table V-2

SECONDARY SILVER SAMPLING DATA PHOTOGRAPHIC - MISCELLANEOUS RAW WASTEWATER

	Stream	Conc Sample	entrations	(mg/l, ex	cept as no	oted)	
Pollutant(a)	_Code_	Typet	Source(b)	Day 1	Day 2	Day 3	Average
4. benzene	12 14 16	1 1 1	•	0.084 0.17 0.149	0.132	2.05	0.084 0.784 0.149
6. carbon tetra- chloride	12 14 16	1 1 1		ND ND ND	ND	0.07	0.07
10. 1,2-dichloro- ethane	12 14 16	1 1 1	۷.	0.061 0.58 0.156	ND	ND	0.061 0.58 0.156
11. 1,1,1-trichloro- ethane	12 14 16	1	•	ND ND 0.022	ND	ND	0.022
23. c loroform	12 14 16	1 1 1		0.244 1.31 0.36	ND	ND	0.244 1.31 0.36
29. 1,1-dichloro- ethylene	12 14 16	1 1 1		ND 0.33 6.1	ND	ND	0.33 6.1
38. ethylbenzene	12 14 16	1 1 1		0.017 0.016 *	*	ND	0.017 0.008 *

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SECONDARY SILVER SUBCATEGORY

SECONDARY SILVER SAMPLING DATA PHOTOGRAPHIC - MISCELLANEOUS RAW WASTEWATER

	Stream	Con Sample	centrations	(mg/l, e	xcept as no	oted)	v
<u>Pollutant(a)</u>	<u>Code</u>	Typet	Source(b)	Day 1	Day 2	Day 3	Average
44. methylene	12	1		0.67		ND	0.67
chloride	14	1		3.23	ND	ND	3.23
	16	1		3.1			3.1
66. bis(2-ethylhexy)		1		*			*
phthalate	14	1		0.034			0.034
	16	1		0.011			0.011
68. di-n-butyl	12	1		0.015			0.015
phthalate	14	1		0.014			0.015
· · · ·	16	1		0.047			0.047
69. di-n-octyl	12	1		0.033			0.033
phthalate	14	1		0.058		·	0.058
	16	1	`	ND			
. diethyl	12	1		ND			• •
phthalate	14	1		0.038			
	16	1		ND			
85. tetrachloro-	12	1		*			*
ethylene	14	1		0.087	ND	ND	0.087
	16	1		0.042			0.041
86. toluene	12	1		0.029			0.029
	14	1		0.027	ND	0.032	0.03
	16	· 1 ,		0.013			0.013

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SECONDARY SILVER SAMPLING DATA PHOTOGRAPHIC - MISCELLANEOUS RAW WASTEWATER

			ncentrations	(mg/1, e	xcept as n	oted)	, `` ,
<u>Pollutant(a)</u>	Stream <u>Code</u>	Sample Typet	<u>Source</u> (b)	Day 1	<u>Day 2</u>	Day 3	Average
87. trichloro- ethylene	12 14 16	1 1 1		0.473 0.93 0.832	ND	ND	0.473 0.93 0.832
91. chlordane	12 16	1 · · 1	· · · ·	** <0.01	· ·		** <0.01
106. PCB-1242 (c) 107. PCB-1254 (c) 108. PCB-1221 (c)	12 16 230	1 1 6		<0.014 <0.007 0.012		•	<0.0 <0.0 0.012
109. PCB-1232 (d) 110. PCB-1248 (d) 111. PCB-1260 (d)	12 16 230	1 1 6		<0.017 <0.015 0.012		• .	<0.017 <0.015 0.012
113. toxaphene	12 16 230	1 1 1		** <0.01 ND			** <0.01
114. antimony	12 14 16	1 1 1	, 	12 0.7 1.5			12.0 0.7 1.5
115. arsenic	12 14 16	1 1 1		2.2 0.2 1.9			2.2 0.2 1.9

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SECONDARY SILVER SUBCATEGORY

SECONDARY SILVER SAMPLING DATA PHOTOGRAPHIC - MISCELLANEOUS RAW WASTEWATER

			Stream	Con Sample	centrations	(mg/1,	except as	noted)	
	<u>Poll</u>	utant(a)	Code	Typet	Source(b)	<u>Day 1</u>	<u>Day 2</u>	Day 3	Average
	117.	beryllium	12 14 16	1 1 1		<0.02 <0.02 <0.02			<0.02 <0.02 <0.02
N	118.	cadmium	12 14 16	1 1 1		0.37 5 0.65			0.37 5.0 0.65
2718	119.	chromium	12 14 16	1 1 1		100 9 7			100.0 9.0 7.0
	120.	copper	12 14 16	1 3 1 3 1 3 3 1 3 3 3 1 3		30 2 0.72			30.0 2.0 0.72
	121.	cyanide	12 14 16	1 1 1		5.95 1.83 0.311	1.13	1.29	5.95 1.416 0.311
-	122.	lead	12 14 16	1 1 1	6	9 2 6			9.0 2.0 6.0
	123.	mercury	12 16	1 · · · · · · · · · · · · · · · · · · ·		0.017 0.0008			0.017 0.0008

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SECONDARY SILVER SAMPLING DATA PHOTOGRAPHIC - MISCELLANEOUS RAW WASTEWATER

			Con	centrations (mg/l, e	xcept as no	ted)	
•	Pollutant(a)	Stream Code	Sample Typet	Source(b) Day 1	Day 2	<u>Day 3</u> <u>Aver</u>	age
	124. nickel	12 14 16	1 1 1	<0.5 1 2		<0.5 1.0 2.0)
	125. selenium	12 14 16	1 1 1	0.9 0.6 0.25		0.9 0.6 0.2	
2719	126. silver	12 16	1 1	5 3		5.0 3.0	
	127. thallium	12 16	1 1	0.4 0.2		0.4 0.2	
2	128. zinc	12 14 16	1 1 1	20 4 10		20.0 4.0 10.0)
	Nchconventionals	•	· ·				
	chemical oxygen demand (COD)	12 16 230	1 1 6	10,100 6,460 14,800	a da antara da antar Antara da antara da an	10,100 6,460 14,800	
	phenols (total; by 4-AAP method)	12 14 16	1 1 1	0.197 32 62.5	28.8	0. 16.7 25.8 62.5	

SECONDARY SILVER SAMPLING DATA PHOTOGRAPHIC - MISCELLANEOUS RAW WASTEWATER

Pollutert (.)	Stream	Sample	centrations (mg/l, e	xcept as n	oted)	
<u>Pollutant(a)</u>	Code	Typet	<u>Source(b)</u> Day 1	Day 2	Day 3	Average
total organic carbon (TOC)	12 16 230	1 1 6	4,040 2,410 13,040			4,040 2,410 13,040
Conventionals						13,040
oil and grease	12 14 16	1	111 130 20			111 130 20
total suspended solids (TSS)	12 16 230	1 1 6	3,66 4 162 484			3,664 162.0 484.0
pH (standard units		1 1 1	2.95 8.4 1.1	6.05	5.88	

(a) Three samples were analyzed for the acid extractable pollutants; none were detected. The samples were also analyzed for pesticides, but none were detected, except as noted.

- (b) No source water samples were analyzed.
- (c), (d) Reported together

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SECONDARY SILVER SUBCATEGORY SECT - V

SECONDARY SILVER SAMPLING DATA PHOTOGRAPHIC - MISCELLANEOUS RAW WASTEWATER

SECONDARY SILVER SUBCATEGORY

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tSample type. Note: These numbers also apply to subsequent data tables.

one-time grab
 24-hour manual composite
 24-hour automatic composite
 48 hour manual composite
 48 hour automatic composite
 72 hour manual composite
 72 hour automatic composite

* Less than or equal to 0.01 mg/l

** Less than or equal to 0.005 mg/l

TABLE V-3

WATER USE AND DISCHARGE RATES FOR PRECIPITATION AND FILTRATION OF FILM STRIPPING SOLUTIONS

(1/troy ounce of silver produced from film stripping)

Plant <u>Code</u>	Percent Recycle	Production Normalized Water <u>Use</u>	Production Normalized <u>Discharge</u> Flow
30927	0	112.7	112.7
596	0	2.31	2.31
74	0	0.74	0.74
566	NR	NR	NR
602	No	Wastewater Produce	đ

NR = data not reported in dcp.

TABLE V-4

WATER USE AND DISCHARGE RATES FOR PRECIPITATION AND FILTRATION OF PHOTOGRAPHIC SOLUTIONS

(1/troy ounce of silver precipitated)

Plant <u>Code</u>	Percent Recycle	Production Normalized <u>Water Use</u>	Production Normalized Discharge Flow
30927	0	89.9	89.9
538	0	21.1	21.1
9022(a)	0	13.5	13.5
437	0	6.75	6.75
615(a)	0	1.6	1.6
74	0	1.35	1.35
563	0	NR	NR
567	0	NR	NR
4301	0	NR	NR

NR = data not reported in dcp.

(a) - Plant closed or no longer processing secondary silver.

Table V-5

SECONDARY SILVER SAMPLING DATA PHOTOGRAPHIC SOLUTIONS RAW WASTEWATER

SECONDARY SILVER SUBCATEGORY

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	Stream	Con	centrations (mg/l	, except as n	oted)
<u>Pollutant(a)</u>	<u>Code</u>	Sample <u>Typet</u>	Source(b) Day	<u>1 Day 2</u>	<u>Day 3 Average</u>
Toxic Pollutants					
10. 1,2-dichloro- ethane	61	1	0.06	B	0.060
15. 1,1,2,3-tetra- chloroethane	61	1	<0.02	9	<0.029
23. chloroform	61	1	0.13	2	0.132
29. 1,1-dichloro- ethylene	61	1	0.04		0.049
30. 1,2-trans- dichloroethylend	61	1	0.04	9	0.049
66. bis(2-ethyl- hexyl)phthalate	61	1	0.119	95	0.1195
67. butyl benzyl phthalate	61	1	0.052	2	0.052
85. tetrachloro- methylene	61	1	<0.025	5	<9.025
115. arsenic	61	1	0.03		0.03
118. cadmium	61	1	6		6.0
119. chromium	61	1	0.3		0.3
120. copper	61	1	1		1.0
122. lead	.61	1	0.5		0.5
123. mercury	61	-1	1		1.0
124. nickel	61	1	0.4		0.4
125. selenium	61	1	<0.04		<0.04
126. silver	61	1	<9.2		<9.2
127. thallium	61	1	<0.2		<0.2
128. zinc	61	1	200		200

SECONDARY SILVER SAMPLING DATA PHOTOGRAPHIC SOLUTIONS RAW WASTEWATER

	0 +		centrations	(mg/l, ex	cept as no	oted)	in a start and a
<u>Pollutant(a)</u>	Stream Code	Sample <u>Typet</u>	Source(b)	Day 1	Day 2	Day 3	Average
Nonconventionals	• •	· •	• •				ж. т.
ammonia chemical oxygen demand (COD)	61 61	1 ³ 1		4,630 40,700			4,630 40,700
chloride total organic carbon (TOC)	61 61	1 1	•	734 3,085	· · ·	•	734 3,085
<u>Conventionals</u>		• • • • • •					
oil and grease total suspended solids (TSS)	61 61	1 1 1		3 92			3 92

SECONDARY SILVER SUBCATEGORY

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- (a) One sample was analyzed for pesticides: dieldrin, chlordane, 4,4'DDT, 4,4'DDE, endrin, endrin aldehyde, heptachlor, alpha-BHC, beta-BHC and gamma-BHC were detected but below the quantification limit. One sample was analyzed for PCB fractions. PCB-1242, 1254, 1221, 1232, 1248, 1260 and 1016 were detected, but below the quantification limit.
- (b) No source water samples were analyzed.

TABLE V-6

WATER USE AND DISCHARGE RATES FOR PRECIPITATION AND FILTRATION OF PHOTOGRAPHIC SOLUTIONS WET AIR POLLUTION CONTROL

(1/troy ounce of silver precipitated)

Plant Code	Percent Recycle	Production Normalized Water Use	Production Normalized Discharge Flow
553	99+	1214.	0.58
74(a)	99+	96,5	0.068
459	100	NR	0
567	68	NR	NR

NR = data not reported in dcp.

- (a) Same scrubber used for film stripping and precipitation of film stripping solutions
- (b) Plant closed

TABLE V-7

WATER USE AND DISCHARGE RATES FOR PRECIPITATION AND FILTRATION OF FILM STRIPPING SOLUTIONS

(1/troy ounce of silver from electrolytic refining)

Plant <u>Code</u>	Percent <u>Recycle</u>	Production Normalized Water Use	Production Normalized Discharge Flow
567(a)	0	1.97	1.97
448(b)	0	1.77	1.77
457	0	1.64	1.64
553	0	0.63	0.63
615(a)	0	0.49	0.49
1104(b)	0	0.44	0.44
460	0	0.31	0.31
460(b)	0	0.06	0.06
65	0	0.28	0.28
1053(b)	0	0.16	0.16
4301	0	0.068	0.068
1084(b)	0	0.035	0.035
578(b)	0	0.024	0.024
11 38(b)	0	NR	NR
1071(b)	0	NR	NR
1088(b)	0	NR	NR

NR = data not reported in dcp.

(a) - Plant closed or no longer processes secondary silver

(b) - Date from nonferrous metals manufacturing phase II 1983 dcp

Table V-8

SECONDARY SILVER SAMPLING DATA NONPHOTOGRAPHIC - MISCELLANEOUS RAW WASTEWATER

	Stream	Con	centration	s (mg/l, e	xcept as n	oted)	
<u>Pollutant(a)</u>	<u>Code</u>	Sample <u>Type</u>	Source	Day 1	Day 2	Day 3	Average
<u>Toxic Pollutants</u>							ĸ
1. acenaphthne	40	6	ND	0.010	ND	ND	0.01
4. benzene	40	2	ND	0.054	*	0.038	0.031
	44	1	ND	*		0.050	*
5. carbon tetra-	40	2	ND	ND	2.3	1.66	1.98
chloride	44	1	ND	ND			
7. chlorobenzene	40	2	ND	*	*	<0.022	*
	44	1	ND	*		(0.022	*
11. 1,1,1-trichlo-	40	2	ND	ND	0.022	ND	0.022
roethane	44	1	ND	ND			0.011
15. 1,1,2,3-tetra-	40	2	ND	ND	ND	ND	
chloroethane	44	1	*	<0.038			<0.038
23. chloroform	40	2	0.021	ND	ND	0.312	0.312
	» 44	1	*	0.109		••••	0.109
ී8. ethylbenzene	40	2	ND	0.021	*	ND	0.011
- , -	44	1.	ND	ND			0.011
47. bromoform	40	2	ND	0.065	ND	ND	0.065
÷	44	1	ND	ND			00003
51. chlorodibro-	40	2	ND	ND	<0.064	ND	<0.064
momethane	44	1 👘	ND .	ND		÷	
66. is(2-ethyl-	40	2	0.016	0.047		u en u	0.047
hexyl)phthalate	e 44	1	*	0.011			0.011
67. butyl benzyl	40 [°]	2	ND	0.054			0.054
pithalate	4 4 [~]	1	*	ND			
68. di-n-butyl	40	· 2	*	0.3			0.3
phthalate	44	1	*	*			*

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SECT - V

SECONDARY SILVER SUBCATEGORY

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SECONDARY SILVER SAMPLING DATA NONPHOTOGRAPHIC - MISCELLANEOUS RAW WASTEWATER

			centration	as (mg/l, e	except as n	oted)	
Pollutant(a)	Stream <u>Code</u>	Sample Type	Source	Day 1	Day 2	Day 3	Average
69. di-n-octyl	40	2	ND	0.055		•	0.055
phthalate	44	1	*	*			*
78. anthracene (b)	40	2	*	<0.014			<0.014
81. phenanthrene (b) 44	1	ND	ND			
84. pyrene	40	2	*	2.15	-		2.150
	44	1	ND	ND			
85. tetrachloro-	40	2	0.011	0.123	0.017	ND	0.07
ethylene	44	1	*	<0.046	· · · · · · · · · · · · · · · · · · ·		<0.046
86. toluene	40	2	*	0.057	*	<0.014	0.019
	- 44	1 .	ND	0.013			0.013
87. trichloro-	40	2	ND	ND	<0.019	ND	<0.019
ethylene	44	1	ND	<0.015			<0.015
90. dieldrin	44	1	**	**	· · ·		**
91. chlordane	44	1	**	**			**
92. 4,4'-DDT	44	1	**	**	,		**
93. 4.4'-DDE	44	1	ND	**			**
3. endrin	44	1	**	**			**
100. heptechlor	44	1	**	**			**
103. b-BHC-Beta	44	· 1	**	**	•	,	**
107. PCB-1254	44	1	**	**		· ·	**
• PCB-1248	44	1	**	**			**
. antimony	40	2	<0.1	<0.1			<0.1
115. arsenic	40	2	<0.01	0.05			0.05
	44	1	<0.01	0.05			0.05
117. beryllium	40	2	<0.001	<0.001			<0.001
· · · · · · · · · · · · · · · · · · ·	44	1	<0.001	0.02	· .		0.02
118. cadmium	.40	2	<0.002	1.0	and the second second		1.0
	44	1	<0.002	80.0	к		80.0

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SECT

SECONDARY SILVER SUBCATEGORY

SECONDARY SILVER SAMPLING DATA NONPHOTOGRAPHIC -- MISCELLANEOUS RAW WASTEWATER

SECONDARY SILVER SUBCATEGORY

SECT

4

Concentrations (mg/l, except as noted)							
Pollutant(a)	Stream _Code_	Sample <u>Type</u>	Source	Day 1	Day 2	Day 3	Average
119. chromium	40	2	<0.005	2.0			2.0
	44	1	<0.005	20.0			20.0
120. copper	40	2	0.2	70.0			70.0
1201 00pp01	44	1	0.04	60.0	_		60.0
121. cyanide	40	2		.0.018	0.132	0.019	0.056
	44	1		0.001		•	0.001
122. lead	40	2	<0.02	4.0	1 N.		4.0
	44	1	<0.02	50.0			50.0
123. mercury	40	2	<0.0001	<0.0001			<0.0001
	44	1	0.0002	0.0001	~		0.0001
124. nickel	40	2	<0.005	30.0			30.0
	-44	1	<0.005	800	•		800
125. selenium	40	2	<0.01	<0.01			<0.01
	44	, 1	<0.01	<0.01			<0.01
126. silver	40	2	<0.02	0.43			0.43
	44	1	<0.02	4.7			4.7
127. thallium	40	2	<0.1	<0.1			<0.1
	44	1	<0.1	<0.1		. •	<0.1
128. zinc	40	2	<0.06	20.0			20.0
	44	1	<0.06	2,000			2,000
Nonconventionals		е 14					
ommoni o	40	· 1		2,180	750	675	1,202
ammonia	40	i		9.5	-		9.5
aborteal average	40	2	<5		,040		3,040
chemical oxygen demand (COD)	40	1.	<5	231	•		231

SECONDARY SILVER SAMPLING DATA NONPHOTOGRAPHIC - MISCELLANEOUS RAW WASTEWATER

	Stream	Con Sample	centratio	ns (mg/l, ex	cept as no	oted)	
Pollutant(a)	<u>Code</u>		Source	Day 1	Day 2	Day 3	Average
chloride fluoride phenols (total; by	44 40 40	1	41.0 1.3	32,300 0.017	1.2 0.012	32 0 . 014	2,300 1.2 0.014
4-AAP method) total organic carbon (TOC)	44 40 44	1 2 1	4.0 5.0	0.044 24.0	435.0		0.044 435.0 24.0
<u>Conventionals</u>			•		• •	·	
oil and grease	40 44	1	- · · ·	11 8	27	13	17.0 8.0
total suspended solids (TSS)	40 44	2 1	14 <1	112	118		118 112
pH (standard units)	40 44	1 1		1.9	2.2	2.3	

(a) Stream 40 was analyzed for the pesticide fraction, but none were detected above its quantification limit.

(b) Reported together.

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SECONDARY SILVER SUBCATEGORY SECT

TABLE V-9

WATER USE AND DISCHARGE RATES FOR FURNACE WET AIR POLLUTION CONTROL

(1/troy ounce of silver roasted, smelted or dried)

Plant <u>Code</u>	Percent Recycle	Production Normalized Water Use	Production Normalized Discharge Flow
78	99.9	143.7	0.14
553	99.7	49.21	0
65	100	19.85	0
549	100(A)	11.6	0
437	0	9.4	0
9020	0	7.87	7.87
1138(C)	0	0.37	0.37
596	100	NR	NR
441	100	NR	NR
1084	100	NR	NR
459	0	NR	NR
4567	NR	NR	NR

NR = data not reported in dcp.

(a) - Partial evaporation.

(b) - 100 % evaporation.

(c) - Data from nonferrous metals manufacturing phase II 1983 dcp.

TABLE V-10

WATER USE AND DISCHARGE RATES FOR LEACHING

(1/troy ounce of silver produced from leaching)

Plant <u>Code</u>	Percent Recycle	Production Normalized <u>Water</u> <u>Use</u>	- 1.4.5. € ∦ - 4. 1.5.5. € - 1.4. 1.5.5.	Production Normalized Discharge Flow
9022	0	635.2		635.2
9020	0 0	98.31		98 .3 1
9020(b)	0	2.97	· · · · ·	2.97
1145	0	1,56		1.56
1150(b)	0	NR		NR
615(a)	0	0.11	۱ ۱	0.11
78	0	0.079		0.079
55 3	0	0.068		0.068
25	NR	NR	* *	NR
82(a)	NR	NR		NR
567(a)	0	NR		NR
459	NR	NR		NR

NR = data not reported in dcp.

(a) - Partial evaporation.

(b) - 100 % evaporation.

(c) - Data from nonferrous metals manufacturing phase II 1983 dcp.

SECT - V

TABLE V-11

WATER USE AND DISCHARGE RATES FOR LEACHING AND PRECIPITATION OF NONPHOTOGRAPHIC SOLUTIONS WET AIR POLLUTION CONTROL

Plant <u>Code</u>	Percent Recycle	Production Normalized Water Use	Production Normalized Discharge Flow
9020	99	984	9.84
74	99+	218	0.12
549	99	90	0.90
83(a)	79.2	54.56	11.3
553	99+	7.0	0.014
78	100	0.078	0
82(a)	97.4	NR	NR
459	100	NR	0
664	100	NR	0
448	, NR	NR	NR
567(a)	65	NR	NR
1204(b)	100	13.5	0
5 78	NR	NR	NR

NR = data not reported in dcp.

(a) - Plant closed or no longer processing secondary silver.

(b) - Data from nonferrous metals manufacturing phase II 1983 dcp.

TABLE V-12

WATER USE AND DISCHARGE RATES FOR PRECIPITATION AND FILTRATION OF NONPHOTOGRAPHIC SOLUTIONS

(1/troy ounce of silver precipitated)

Plant <u>Code</u>	Percent Recycle	Production Normalized Water <u>Use</u>	Production Normalized Discharge Flow
1100	0	109.5	109.5
1092	0	103.3	103.3
9020*	0	78.6	78.6
9020	0	19.9	19.9
1018	0	47.5	47.5
1104	0	17.7	17.7
18	0	12.96	12.96
615(a)	0	7.87	7.87
1128	0	7.53	7.53
1165	0	5.16	5.16
549	0	2.7	2.7
1164	0	2.1	2.1
448	0	1.77	1.77
1029	0	1.23	1.23
1023	0	1.0	1.0
74	0	0.90	0.90
1053(a)	0	0.65	0.65
1167(a)	0	0.52	0.52
4640(a)	0	0.5	0.5
460	0	0.42	0.42
1072	0	0.37	0.37
57 8	0	0.19	0.19
1204	0	0.183	0.183
1063	0	0.13	0.13
1117	0	0.098	0.098

NR = data not reported in dcp.

(a)	-		from	nonferrous	metals	manufacturing	phase	II	1983
		dcp.			·	2			

(b) - Plant colsed or no longer processes secondary silver.

Table V-13

SECONDARY SILVER SAMPLING DATA NONPHOTOGRAPHIC - TREATMENT PLANT SAMPLES - PLANT A

			Concentrations (mg/l, except as noted)					
<u>Pollutant</u>	Stream <u>Code</u>	Sample <u>Type</u>	Source	Day 1	Day 2	<u>Day 3</u>	Average	
<u>Toxic Pollutants</u>		×						
4. benzene	· 41	2	ND	0.014	ND	ND	0.014	
6. carbon tetra- chloride	41	2	ND	0.394	0.305	0.401	0.367	
15. 1,1,2,3-tetra- chloroethane	41	2	ND	ND	0.025	ND	0.025	
23. chloroform	41	2	0.021	0.04	0 - 305	ND	0.173	
38. ethylbenzene	41	2 2 7	ND	0.012	ND	*	0.006	
39. fluoranthene	41		*	0.198			0.198	
47. bromoform	41	2	ND	ND	0.013	ND	0.013	
48. dichlorobromo- methane	41	2	ND	2.8	2.4	1.58	2.26	
51. chlorodibro-	41	2	ND	<0.047	ND .	ND	<0.047	
momethane 66. bis(2-ethyl- hexyl)phthalate	41	7	0.016	0.022			0.022	
67. butyl benzyl	41	7	ND	0.038		е п Н	0.038	
phthalate 68. di-n-butyl	41	7	*	0.082			0.082	
phthalate					•	- ÷		
69. di-n-octyl phthalate	41	7	ND	0.069			0.069	
84. pyrene	41	7	*	0.179	×.		0.179	
85. tetrachloro- ethylene	41	2	0.011	0.017	*	*	0.006	
87. trichloro- ethylene	41	2	ND	<0.014	ND	ND	<0.014	
114. antimony	41	7	<0.1	1.5			1.5	

SECONDARY SILVER SUBCATEGORY SECT - V

SECONDARY SILVER SAMPLING DATA NONPHOTOGRAPHIC - TREATMENT PLANT SAMPLES - PLANT A

	-		oncentrat	ions (mg/l	, except a	s noted)	• .
Pollutant	Stream <u>Code</u>	Sample Type	Source	Day 1	Day 2	Day 3	Average
15. arsenic 21. cyanide 26. silver	41 41 41	7 7 7	<0.01 <0.02	1.26 0.020 6.9	0.075	0.053	1.26 0.049 6.9
Nonconventionals		· ·					
amm tia	41	1		2,200	2,080	1,600	1,960
chem_c:l oxygen demand (COD)	41	7	<5		556		556
fluoride	41	7	1.3		2.4		2.4
phenols (total; by 4-AAP method)	41	2		0.008	0.023	0.018	0.016
total organic carbon (TOC)	41	7	4	• •	97		c 97
Conventionals							
oil and grease	41	1		82	5	10	32
total suspended solids (TSS)	41	7	14		3,140		3,140
pH (standard units)	41	1		7.3	8.1	8.7	

Table V-14

SECONDARY SILVER SAMPLING DATA NONPHOTOGRAPHIC - TREATMENT PLANT SAMPLES - PLANT B

			centration	s (mg/l, ex	cept as no	oted)	
	Stream	Sample			_		
<u>Pollutant</u>	Code	_Туре_	Source	<u>Day 1</u>	<u>Day 2</u>	<u>Day 3</u>	Average
Toxic Pollutants			•				
6. carbon tetra- chloride	45	1	ND	0.019			0.019
66. bis(2-ethyl-	45	2	*	0.03			0.03
hexyl) phthalat		· .	· ·				
118. cadmium	45	2	<0.002	0.1			0.1
119. chromium	45	2 2 2 2 2	<0.005	<0.05			<0.05
120. copper	45	2	0.04	0.6			0.6
121. cyanide	45	2		0.001			0.001
122. lead	45	2	<0.02	<0.2			<0.2
123. mercury	45	2 2	<0.0002	0.0001			0.0001
124. nickel	45	. 2	<0.005	<0.05 °			<0.05
128. zinc	45	2	<0.06	1			1
Nonconventionals							
ammonia	45	1		0.49	-		0.49
chemical oxygen demand (COD)	45	1	<5	<5			<5.0
chloride	45	1	41	669	х.		6 69
phenols (total; by by 4-AAP method)	45	1		0.011			0.011
total organic carbon (TOC)	45	1	5	<1	•		<1.0

SECONDARY SILVER SUBCATEGORY SECT -

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SECONDARY SILVER SAMPLING DATA NONPHOTOGRAPHIC - TREATMENT PLANT SAMPLES - PLANT B

	Concentrations (mg/l, except as noted)								
Pollutant	Stream <u>Code</u>	Sample Type	Source	Day 1	<u>Day 2</u>	Day 3	Average		
Conventionals			• • •		•				
oil and grease total suspended	45 45	1 1	<1	10 10			10 10		
solids (TSS) pH (standard units)	45	1	•	9.9					

SECONDARY SILVER SUBCATEGORY

SECT

Table V-15

SECONDARY SILVER SAMPLING DATA PHOTOGRAPHIC - TREATMENT PLANT SAMPLE - PLANT C

	Stream	Concentrations (mg/l, except as noted) Sample						
<u>Pollutant</u>	_Code_	Туре	Source	Day 1	Day 2	Day 3	Average	
Taxic Pollutants								
4. benzene	13	1		0.06	2 9	ND	0.06	
	15 17	1		0.03	3.2	ND	0.03	
0. 1,2-dichloro-	13	1		0.126 0.044	ND	0.05	0.126 0.047	
ethane	15 17	1		0.044	1412	0.05	0.26	
23. chloroform	13 15 17	1 1 1		0.404 0.076 3.18	0.07	0.032	0.404 0.0593 3.18	
29. 1,1-dichloro- ethylene	13 15 17	1 1 1		0.101 0.013 3.418	ND	ND	0.101 0.013 3.418	
38. ethylbenzene	13 15 17	1 1 1		0.014 ND ND	0.036	0.05	0.014 0.043	
44. methylene chloride	13 15 17	1 1 1		0.876 0.086 0.89	ND	ND	0.876 0.086 0.89	
85. tetrachloro-	13	1		0.012	ND		0.012	
methylene	15 17	1 1		ND 0.041	ND	ND	0.041	

SECONDARY SILVER SUBCATEGORY

SECT

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Table V-15 (Continued)

SECONDARY SILVER SAMPLING DATA PHOTOGRAPHIC - TREATMENT PLANT SAMPLE - PLANT C

SECONDARY SILVER SUBCATEGORY

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		Co	ncentrations	; (mg/1, e	except as n	oted)	
Pollutant	Stream <u>Code</u>	Sample Type	Source	Day 1	Day 2	Day 3	Average
86. toluene	13 15 17	1 1 1		0.019 * ND	ND	ND	0.019 *
87. trichloro- ethylene	13 15 17	1 1 1 1		0.334 0.047 0.19	ND	ŇD	0.334 0.047 0.19
114. antimony	15 17	1 1		0.45 ND			0.45
115. arsenic	15 17	1		0.7 ND		н Т	0.7
117. beryllium	15 17	1		<0.02 ND			<0.02
118. cadmium	15 17	1		3 ND			3
119. chromium	15 17	1		8 ND			8
120. copper	15 17	. 1		1 ND			1

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Table V-15 (Continued).

SECONDARY SILVER SAMPLING DATA PHOTOGRAPHIC - TREATMENT PLANT SAMPLE - PLANT C

		Con	centrations	(mg/1, ex	cept as no	oted)	
Pollutant	Stream Code	Sample Type	Source	Day 1	Day 2	Day 3	Average
121 cyanide	13 15 17	1 1 1		2.19 1.29 0.098	1.62	2.04	2.19 1.65 0.098
122. lead	15 17	1 1		3 ND			3
N 123 mercury	13 15 17	1 1 1		0.0032 0.0016 ND			0.0032 0.0016
i aickel	15 17	1		4 ND			4
12. selemium	15 17	1 1		0.4 ND		· · ·	0.4
126. si. e	13 15	1 . 		1	· · · · · ·		1
127. thallium	13 15	1		0.65 0.2			0.65 0.2
128. zinc	15	1		5 ND		. • · ·	5

SECT -

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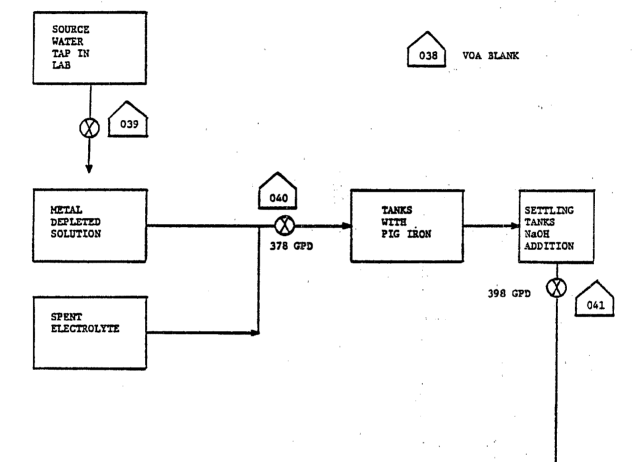
Table V-15 (Continued)

SECONDARY SILVER SAMPLING DATA PHOTOGRAPHIC - TREATMENT PLANT SAMPLE - PLANT C

		ncentration	ns (mg/l, e	xcept as	noted)	- 	
Pollutant	Stream Code	Sample <u>Type</u>	Source	Day 1	Day 2	Day 3	Average
Nonconventionals		: •			· · ·		
phenols (total; by 4-AAP method)	13 15 17	1 1 1		0.421 26.5 51.3	26.8	20.3	0.421 24.5 51.3
CONVENTIONALS	N	·• · · · · · · · · · · · · · · · · · ·	•				
oil and grease	15 17	1 1 1	· · · · · · · · · · · · · · · · · · ·	50 21	51	195	99 21
pH (standard units)	13 15 17	1 1 1		7.14 8.55 6.68	7.16	6.97	

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SECONDARY SILVER SUBCATEGORY SECT



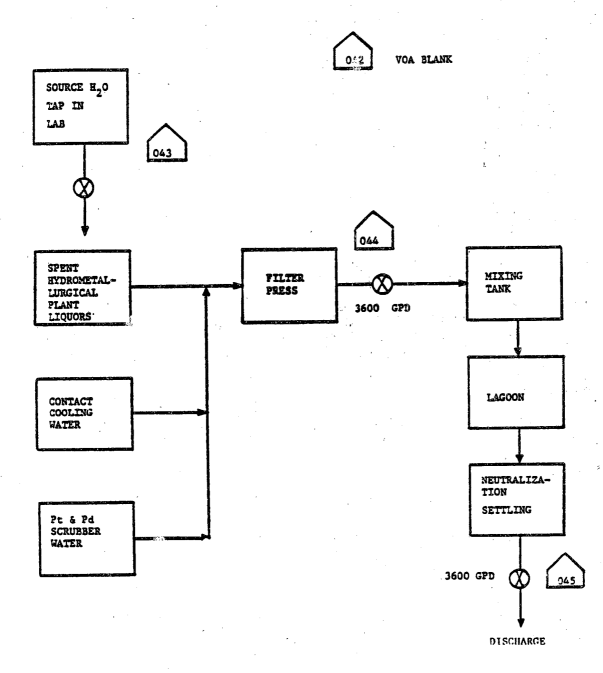
SECT

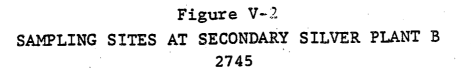
V

DISCHARGE

igure V-1 SAMPLING SITES AT SECONDARY SILVER PLANT A

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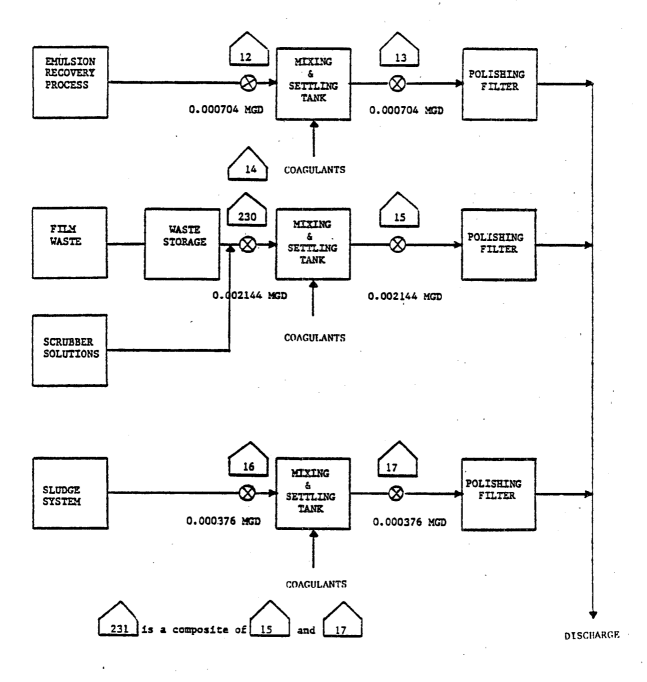
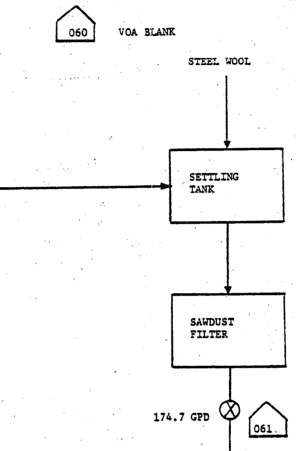


Figure V-3 SAMPLING SITES AT SECONDARY SILVER PLANT C

SECONDARY SILVER SUBCATEGORY SECT - V

SPENT

HYPO SOLUTION



DISCHARGE

Figure V-4 SAMPLING SITES AT SECONDARY SILVER PLANT D

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SECONDARY SILVER SUBCATEGORY SECT - VI

SECTION VI

SELECTION OF POLLUTANT PARAMETERS

This section examines chemical analysis data presented in Section V and discusses the selection or exclusion of pollutants for potential limitation.

The discussion that follows describes the analysis that was or exclude pollutants for performed to select further consideration for limitations and standards. Pollutants will be considered for limitation if they are present in concentrations treatable by the technologies considered in this analysis. The treatable concentrations used for the toxic metals were the long-term performance values achievable by lime precipitation, sedimentation, and filtration. The treatable concentrations used for the toxic organics were the long-term values achievable by carbon adsorption.

After proposal, the Agency re-evaluated the treatment performance activated carbon adsorption to control toxic organic of pollutants. The treatment performance for the acid extractable, base-neutral extractable, and volatile organic pollutants has been set equal to the analytical quantification limit of 0.010 The analytical quantification limit for pesticides and mq/l.total phenols (by 4-AAP method) is 0.005 mg/l, which is below the 0.010 mg/l accepted for the other toxic organics. However, to be consistent, the treatment performance of 0.010 mg/l is used for pesticides and total phenols. The 0.010 mg/l concentration is achievable, assuming enough carbon is used in the column and a suitable contact time is allowed. The frequency of occurrence for 36 of the toxic pollutants has been redetermined based on the revised treatment performance value. As a result, the following pollutants, which were not selected at proposal, have been selected for further consideration for limitation:

- 11. 1,1,1-trichloroethane
- 30. 1,2-trans-dichloroethylene
- 38. ethylbenzene
- 84. pyrene
- 85. tetrachloroethylene
- 86. toluene

The selection of these pollutants is discussed in greater detail below.

This study examined samples from the secondary silver subcategory for three conventional pollutant parameters (oil and grease, total suspended solids, and pH) and six nonconventional pollutant parameters (ammonia, chemical oxygen demand, chloride, fluoride, total organic carbon, and total phenols).

CONVENTIONAL AND NONCONVENTIONAL POLLUTANT PARAMETERS SELECTED

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The conventional and nonconventional pollutants and pollutant parameters selected for consideration for limitation in this subcategory are:

ammonia phenols (total; by 4-AAP method) total suspended solids (TSS) pH

Ammonia was found in all four samples analyzed in concentrations ranging from 675 to 4,630 mg/l. All of the values recorded are well above the treatable concentration of 32.2 mg/l, attainable by the available treatment technology. Therefore, ammonia is selected for consideration for limitation.

Total phenols are detected in all eight samples analyzed. All eight samples contained phenols in concentrations above the treatable concentration of 0.010 mg/l. Concentrations for the samples ranged from 0.012 to 62.5 mg/l. Therefore, total phenols are also selected for consideration for limitation.

Total suspended solids (TSS) concentrations ranging from 92 to 3,664 mg/l were observed in the five samples analyzed for this study. All five samples exhibited concentrations above the treatable concentration attainable by the identified treatment technology. Furthermore, most of the specific methods for removing toxic metals do so by precipitation, and the resulting toxic metals precipitates should not be discharged. Meeting a limitation on TSS also aids in removal of precipitated toxic metals. For these reasons, total suspended solids is considered for limitation in this subcategory.

The pH values observed in four of seven samples were outside the 6.0 to 10.0 range considered desirable for discharge to receiving waters. Four pH values ranged from 1.1 to 2.95. The remaining three samples ranged from 5.9 to 8.4. Effective removal of toxic metals by chemical precipitation requires careful control of pH. Therefore, pH is considered for limitation in this subcategory.

TOXIC PRIORITY POLLUTANTS

The frequency of occurrence of the toxic pollutants in the wastewater samples taken is presented in Table VI-1 (page 2758). These data provide the basis for the categorization of specific pollutants, as discussed below. Table VI-1 is based on the raw wastewater data from streams 12, 14, 16, 40, 61, and 230 (see Section V). Treatment plant samples were not considered in the frequency count. Raw waste stream 44 was not used in the count because it contained gold, platinum, and palladium processing wastewater in addition to silver processing wastewater.

TOXIC POLLUTANTS. NEVER DETECTED

The toxic pollutants listed in Table VI-2 (page 2762) were not

detected in any wastewater samples from this subcategory; therefore, they are not selected for consideration in establishing limitations.

TOXIC POLLUTANTS NEVER FOUND ABOVE THEIR ANALYTICAL QUANTIFICA-TION LIMIT

The toxic pollutants listed below were never found above their analytical quantification concentration in any wastewater samples from this subcategory; therefore, they are not selected for consideration in establishing limitations.

- chlorobenzene 7. 1,1,2,2-tetrachloroethane 15. chlorodibromomethane 51. 78. anthracene (a) 81. phenanthrene (a) 90. dieldrin 91. chlordane 92. 4.4' - DDT93. 4,4'-DDE 98. endrin 99. endrin aldehyde 100. heptachlor 102. alpha-BHC 103. beta-BHC 104. gamma-BHC 113. toxaphene
- 116. asbestos

(a) Reported together.

TOXIC POLLUTANTS PRESENT BELOW CONCENTRATIONS ACHIEVABLE BY TREATMENT

The pollutant listed below is not selected for consideration in establishing limitations because it was not found in any wastewater samples from this subcategory above concentrations considered achievable by existing or available treatment technologies.

Acenaphthene was detected in only one of nine samples analyzed. That sample contained 0.010 mg/l, which is the treatable concentration. Since the pollutant was not detected above the concentration attainable by identified treatment technology, acenaphthene is not considered for limitation.

TOXIC POLLUTANTS DETECTED IN A SMALL NUMBER OF SOURCES

The following pollutants were not selected for limitation on the basis that they were detectable in the effluent from only a small number of sources within the subcategory and are uniquely related to only those sources:

23. chloroform

44. methylene chloride

47. bromoform

66. bis(2-ethylhexyl) phthalate

67. butyl benzyl phthalate

68. di-n-butyl phthalate

- 69. di-n-octyl phthalate
- 70. diethyl phthalate
- 106. PCB-1242 (b)
- 107. PCB-1254 (b)
- 108. PCB-1221 (b)
- 109. PCB-1232 (C) 110. PCB-1248 (c)
- 111. PCB-1260
- (c) 112. PCB-1016 (C)

123. mercury

(b),(c) Reported together, as a combined value

Although these pollutants were not selected for consideration in establishing nationwide limitations, it may be appropriate, on a case-by-case basis, for the local permitter to specify effluent limitations.

Chloroform was found at concentrations ranging from 0.109 to 1.31 mg/l in five of nine samples. The achievable concentration treatment for chloroform is 0.010 mg/1. Chloroform cannot be traced to specific materials or processes associated with the secondary silver subcategory; however, it is a common laboratory solvent and the high concentrations found could be attributed to sample contamination. The presence of chloroform in the blank samples taken attest to this possibility, particularly since the pollutant was not detected in four samples. The results cannot be generalized as characteristic of the subcategory. All 25 of the secondary silver plants reporting the presence or absence of toxic pollutants indicated in the dcp that this pollutant was either known or believed to be absent from their wastewater. Therefore, chloroform is not considered for limitation.

Methylene chloride was measured at a concentration above its treatable concentration in three of nine samples, with values of 3.10, and 3.32 mg/1. - The treatable concentration is 0.010 0.67, All three treatable samples were from the same plant. This mq/l. pollutant is not attributable to specific materials or processes associated with the secondary silver subcategory, but is a common solvent used in analytical laboratories. All 25 of the secondary silver plants reporting the presence or absence of toxic pollutants indicated in the dcp that this pollutant was either known or believed to be absent from their wastewater. Because methylene chloride was not detected in six of nine samples, as well as the high probability of sample contamination, this pollutant is not considered for limitation.

Bromoform was not detected in eight of nine samples, but was found above its treatable concentration in one sample. The 0.065 mg/l found is greater than the 0.01 mg/l treatable concentration.

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All 25 of the secondary silver plants reporting the presence or absence of toxic pollutants indicated in the dcp that this pollutant was either known or believed to be absent from their wastewater. Since bromoform is present at only one source, bromoform is assumed to be unique to that source and not considered for limitation.

Bis(2-ethylhexyl) phthalate was found above its trea concentration of 0.010 mg/l in four of five samples. treatable The concentrations ranged from 0.011 to 0.119 mg/1. This pollutant is not associated with specific processes used in the secondary silver subcategory, but is commonly used as a plasticizer in laboratory and field sampling equipment. All 25 of the secondary silver plants reporting the presence or absence of pollutants indicated in the dcp that this pollutant was toxic either known or believed to be absent from their wastewater. Since the presence of this pollutant may be attributed to sample contamination, bis(2-ethylhexyl) phthalate is not considered for limitation.

Butyl benzyl phthalate was measured in two of five samples at concentrations of 0.052 and 0.054 mg/l. The treatable concentration for this pollutant is 0.010 mg/l. This pollutant is used as a plasticizer in laboratory and field sampling equipment. Since it was not detected in three of five samples, the measurements may be regarded as specific to the site and not characteristic of the subcategory as a whole. All 25 of the secondary silver plants reporting the presence or absence of toxic pollutants indicated in the dcp that this pollutant was either known or believed to be absent from their wastewater. Therefore, butyl benzyl phthalate is not considered for limitation.

Di-n-butyl phthalate was found above its treatable concentration (0.010 mg/l) in four of five samples analyzed. However, this compound is a plasticizer used in many products found in manufacturing plants; it is not associated with specific processes used in this subcategory. All 25 of the secondary silver plants reporting the presence or absence of toxic pollutants indicated in the dcp that this pollutant was either known or believed to be absent from their wastewater. Therefore, di-n-butyl phthalate is not considered for limitation.

Di-n-octyl phthalate was measured above its treatable concentration (0.010 mg/l) in three of five samples analyzed. However, this compound is a plasticizer used in many products found in manufacturing plants; it is not associated with specific processes in this subcategory. All 25 of the secondary silver plants reporting the presence or absence of toxic pollutants indicated in the dcp that this pollutant was either known or believed to be absent from their wastewater. Therefore, di-noctyl phthalate is not considered for limitation.

Diethyl phthalate was detected above its treatable concentration (0.010 mg/l) in one of five samples analyzed. However, this

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compound is a plasticizer used in many products found in manufacturing plants; it is not associated with specific processes in this subcategory. All 25 of the secondary silver plants reporting the presence or absence of toxic pollutants indicated in the dcp that this pollutant was either known or believed to be absent from their wastewater. Because of the site-specificity of the one result, diethyl phthalate is not considered for limitation.

The seven toxic pollutant PCBs (polychlorinated biphenyls) are not clearly separated by the analytical protocol used in this study; thus, they are reported in two groups. The first group contains PCB-1242, PCB-1254, and PCB-1221; the second contains PCB-1232, PCB-1248, PCB-1260, and PCB-1016. Both groups were found in one of five samples at the same plant. The concentration for each group was 0.012 mg/1, which slightly exceeds the treatable concentration of 0.010 mg/1. All 25 of the secondary silver plants reporting the presence or absence of toxic pollutants indicated in the dcp that this pollutant was either known or believed to be absent from their wastewater. Since these pollutants were found in only one plant, they are assumed to be unique to that source and are not considered for limitation.

Mercury was measured above its treatable concentration (0.036 mg/l) in one of four samples. Even though found at 1.0 mg/l, this pollutant is not attributable to specific materials and processes in this subcategory. Also, 22 of the 25 secondary silver plants reporting the presence or absence of toxic pollutants indicated in the dcp that mercury was known to be absent or believed to be absent from their wastewater. Since it was found in only one plant, mercury is not considered for limitation.

TOXIC POLLUTANTS SELECTED FOR CONSIDERATION IN ESTABLISHING LIMITATIONS

The pollutants listed below are selected for further consideration in establishing limitations and standards for this subcategory.

-		
4.	benzene	
6.	carbon tetrachloride	
10.	1,2-dichloroethane	
11.	1,1,1-trichloroethane	
29.	1,1-dichloroethylene	
30.	1,2-trans-dichloroethylene	
38.	ethylbenzene	
84.	pyrene	
85.	tetrachloroethylene	
86.	toluene	
87.	trichloroethylene	
114.	antimony	

- 115. arsenic
- 118. cadmium

119.	chromium
120.	copper
121.	cyanide
122.	lead
124.	nickel
125.	selenium
126.	silver
127.	thallium
128.	zinc

Benzene was detected above its treatable concentration (0.010 mg/l) in seven of nine samples. The concentrations ranged from 0.017 to 2.05 mg/l. Since benzene was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

Carbon tetrachloride was found above its treatable concentration (0.010 mg/l) in three of nine samples. Concentrations ranged from 0.07 to 2.3 mg/l. Since carbon tetrachloride was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

1,1,1-Trichloroethane was detected at two plants in two of nine samples, both at concentrations of 0.022 mg/l. The treatable concentration is 0.010 mg/l for this pollutant. Therefore, 1,1,1-trichlorethane is selected for consideration for limitation.

1,2-Dichloroethane was detected above its quantification limit in four of nine samples in two plants. Two samples, with concentrations of 0.58 and 0.156 mg/l, were above the concentration considered attainable by treatment (0.010 mg/l). Since 1,2-dichloroethane was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

1,1-Dichloroethylene was measured above its treatable concentration (0.010 mg/l) in three of nine samples in two plants, with concentrations of 0.049, 0.33, and 6.1 mg/l. Since 1,1-dichloroethylene was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

1,2-<u>Trans</u>-dichloroethylene was detected in one of nine samples with a concentration of 0.049 mg/l. This value is above the treatable concentration (0.010 mg/l). The raw wastewater stream containing this pollutant was sampled at only one plant. There fore, the presence of this pollutant cannot be regarded as sitespecific. For these reasons, $1,2-\underline{trans}$ -dichloroethylene is selected for consideration for limitation.

Ethylbenzene was found in six of nine samples. Only three samples contained this pollutant above its concentration considered attainable by treatment (0.010 mg/l). These values

ranged from 0.016 to 0.021 mg/l. Also, it was detected at two of three plants. Therefore, ethylbenzene is selected for consideration for limitation.

Pyrene was found in one of five samples at a concentration of 2.15 mg/l, which is much greater than its treatable concentration (0.010 mg/l). The raw wastewater stream containing this pollutant was sampled at only one plant. Therefore, the presence of this pollutant cannot be regarded as site-specific and is selected for consideration for limitation.

Tetrachloroethylene was measured above concentrations regarded achievable by identified treatment technology (0.010 mg/l) in four of nine streams. The values ranged from 0.017 mg/l to 0.123 mg/l. Tetrachloroethylene was detected below its quantification limit in two other samples. These samples represent five different vastewater streams at three plants. For these reasons, this pollutant is selected for consideration for limitation.

Toluene was detected in seven of nine samples representing four raw wastewater streams from three planes. Five of its measured concentrations ranged from 0.013 mg/l to 0.05 mg/l, which is above the concentration considered attainable by available treatment for this pollutant (0.010 mg/l). Therefore, toluene is selected for consideration for limitation.

Trichloroethylene was detected above its treatable concentration (0.010 mg/l) in three of nine samples. The concentrations ranged from 0.473 to 0.93 mg/l. Since trichloroethylene was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

Antimony was found above its treatable concentration (0.47 mg/l) in three of five samples. The concentrations ranged from 0.7 to 12.0 mg/l. Since antimony was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

Arsenic was measured above its quantification limit in all five samples analyzed. Two of the five samples contained this pollutant above the treatable concentration (0.34 mg/l), with concentrations of 1.9 and 2.2 mg/l. Since arsenic was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

Chromium was found above its treatable concentration (0.07 mg/l) in all five samples analyzed. The concentrations ranged from 0.3 to 100 mg/l. Since chromium was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

Copper was detected above its treatable concentration (0.39 mg/l) in all five samples analyzed. The concentrations ranged from

0.72 to 70.0 mg/l. Since copper was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

Cyanide was measured above its treatable concentration (0.047 mg/l) in six of nine samples from four of the five waste streams. The concentrations ranged from 0.132 to 5.95 mg/l, in two plants (one photographic and one nonphotographic). Since cyanide was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

Lead was found above its treatable concentration (0.08 mg/l) in all five samples analyzed. The concentrations ranged from 0.5 to 9.0 mg/l. Since lead was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

Nickel was measured above its treatable concentration (0.22 mg/l) in four of five samples. The concentrations ranged from 0.4 to 30.0 mg/l. Since nickel was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

Selenium was found above its treatable concentration (0.20 mg/l) in three of five samples. The concentrations ranged from 0.25 to 0.9 mg/l. Since selenium was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

Silver was detected above its quantification limit in three of five samples analyzed. Concentrations ranged from 0.07 to 5.0 mg/l. Three samples contained silver at concentrations above the concentration considered attainable by treatment (0.07 mg/l). Since silver was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

Thallium was found above its quantification limit in two of the five samples analyzed for this pollutant. One of the five samples contained thallium at a concentration of 0.4 mg/l, above the treatable concentration (0.34 mg/l) for this pollutant. Since thallium was present in concentrations exceeding the concentration achievable by identified treatment technology, it is selected for consideration for limitation.

Zinc was measured above its treatable concentration (0.23 mg/l) in all five samples analyzed. The concentrations ranged from 4.0 to 2,000 mg/l. Since zinc was present in concentrations exceeding the concentration attainable by identified treatment technology, it is selected for consideration for limitation.

Table VI-1

FREQUENCY OF OCCURRENCE OF TOXIC POLLUTANTS SECONDARY SILVER RAW WASTEWATER

Pollutant	Analytical Quantification Concentration (mg/l)(a)	Treatable Concentra- <u>tion (mg/1)</u> (b)	Number of Streams Analyzed	Number of Samples Analyzed	ND	Detected Below Quantification Concentration	Detected Below Treat- able Concen- tration	Detected Above Treat- able concen- tration
1. acenaphthene	0.010	0.010	5.	5	4		1	
2. acrolein	0.010	0.010	5	9	9			
3. acrylonitrile	0.010	0.010	5	9	9			
4. benzene	0.010	0.010	5	9	î t	1		7
5. benzidine	0.010	0.010	5	5	5			
6. carbon tetrachloride	0.010	0.010	5	9	6			3
7. chlorobenzene	0.010	0.010	5	9	6	3		•
8. 1,2,4-trichlorobenzene	0.010	0.010	5	5	5			
9. hexachlorobenzene	0.010	0.010	5	5	5			
10. 1,2-dichloroethane	0.010	0.010	. 5	9	5			4
11. 1,1,1-trichloroethane 12. hexachloroethane	0.010	0.010	2	9	1			2
13. 1,1-dichloroethane	0.010 0.010	0.010	2	5	5			
14. 1,1,2-trichloroethane	0.010	0.010 0.010	Ş	9.	9			
5. 1,1,2,2-tetrachloroethane	0.010		2	9	9			
	0.010	0.010 0.010	ž	9	8	1		
17. bis(chloromethyl) ether	0.010	0.010	2	9	9			
15. bis(2-chloroethyl) ether	0.010	0.010)	9	9		•	
1. 2-chloroethyl vinyl ether	0.010	0.010)	2	2			
2: 2-chloronaphthalene	0.010	0.010) E	9	9 c			
21. 2,4,6-trichlorophenol	0.010	0.010	5	2	2			
22. parachlorometa cresol	0.010	0.010	5	2	2			
23. chloroform	0.010	0.010	- 5	5	ر ۲			-
24. 2-chlorophenol	0.010	0.010	· J	2	4			5
25. 1,2-dichlorobenzene	0.010	0.010	2 ·	2	5			•
26. 1,3-dichlorobenzene	0.010	0.010	ś	Š	· č			
27. 1,4-dichlorobenzene	0.010	0.010	รั	ś	ś			
28. 3,3'-dichlorobenzidine	0.010	0.010	ŝ	Ś	Ś			
29. 1,1-dichloroethylene	0.010	0.010	ร้	ş	6			3
30. 1,2-trans-dichloroethylene	0.010	0.010	Ś	9	Ř			J 1
31. 2,4-dichlorophenol	0.010	0.010	· 5	í	ž			1
32. 1,2-dichloropropane	0.010	0.010	S	9	. <u>9</u>			
33. 1,3-dichloropropylene	0.010	0.010	5	ģ	ģ			
34. 2,4-dimethylphenol	0.010	0.010	5	3	3			
35. 2,4-dinitrotoluene	0.010	0.010	5	Š	5	· . *		
36. 2,6-dinitrotoluene	0.010	0.010	5	5	5			
37. 1,2-diphenylhydrazine	0.010	0.010	5	5	5		-	

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Table VI-1 (Continued)

FREQUENCY OF OCCURRENCE OF TOXIC POLLUTANTS SECONDARY SILVER RAW WASTEWATER

Pollutant	Analytical Quantification Concentration (mg/l)(a)	Treatable Concentra- <u>tion (mg/1)</u> (b)	Number of Streams Analyzed	Number of Samples Analyzed	ND	Detected Below Quantification Concentration	Detected Below Treat- able Concen- tration	Detected Above Treat- able Concen- tration
38. ethylbenzene	0.010	0.010	5	9	3	3		3
39. fluoranthene	0.010	0.010	5	5	5		•	
40. 4-chlorophenyl phenyl ether	0.010	0.010	. 5	5	5		1	
41. 4-bromophenyl phenyl ether	0.010	0.010	5	5	5			
42. bls(2-chloroisopropyl) ether	0.010	0.010	2	2	2			
43. bis(2-chloroethoxy) methane 44. methylene chloride	0.010	0.010	2		Ş			-
44. methyl chloride	0.010 0.010	0.010 0.010	2	9	0			3
45. methyl bromide	0.010	0.010	2	9	9			
47. bromoform	0.010	0.010	5	, 9	2			4
38. dichlorobromomethane	0.010	0.010		9.	ĝ			1
49. trichlorofluoromethane	0.010	0.010	Ś	9	ģ			•
50. dichlorodifluoromethane	0.010	0.010	5	9	9			
51. chlorodibromomethane	0.010	0.010	5	9	8	1		
52. hexachlorobutadiene	0.010	0.010	5	5	5			
53. hexachlorocyclopentadiene	0.010	0.010	5	5	5			
54. Isophorone	0.010	0.010	5	5	5			
55. naphthalene	0.010	0,010	5	5 ·	5			
56. nitrobenzene	0.010	0.010	5	5	5			
57. 2-nitrophenol	0.010	0.010	5	3	3		•	
58. 4-nitrophenol	0.010	0.010	5	3	3			-
59. 2,4-dinitrophenol	0.010	0.010	5	3	3			
60. 4,6-dinitro-o-cresol 61. N-nitrosodimethylamine	0.010 0.010	0.010		3	3			
62. N-nitrosodiphenylamine	0.010	0.010 0.010	2	2	2			
63. N-nitrosodi-n-propylamine	0.010	0.010	5		2	· .	•	
64. pentachlorophenol	0.010	0.010	5) 1				
65. phenol	0.010	0.010	- 5	ว้	2			
66. bis(2-ethylhexyl) phthalate	0.010	0.010	· 5	· 5	5	. 1		4
67. butyl benzyl phthalate	0.010	0.010	. 5	Ś	3	•		2
68. di-n-butyl phthalate	0.010	0.010	5	Š	ī			Ā
69. di-n-octyl phthalate	0.010	0.010	5	5	2			3
70. diethyl phthalate	0.010	0.010	5	5	4			i i
71. dimethyl phthalate	0.010	0.010	5	5	5		· ·	
72. benzo(a)anthracene	0.010	0.010	5	- 5	5			
73. benzo(a)pyrene	0.010	0.010	5	5	5		•	
74. 3,4-benzofluoranthene	0.010	0.010	5	5	5	· · · ·		

SECONDARY SILVER SUBCATEGORY

SECT - VI

Table VI-1 (Continued)

FREQUENCY OF OCCURRENCE OF TOXIC POLLUTANTS SECONDARY SILVER RAW WASTEWATER

<u>Pollutant</u>	Analytical Quantification Goncentration (mg/l)(a)	Treatable Concentra- <u>tion (mg/1)</u> (b)	Number of Streams <u>Analyzed</u>	Number of Samples Analyzed	ND	Detected Below Quantification Concentration	Detected Below Treat- able Concen- tration	Detected Above Treat- able Concen- tration
75. benzo(k)fluoranthene	0.010	0.010	5	5	5			
76. chrysene	0.010	0.010	5 5	5	5			
77. acenaphthylene	0.010	0.010		5	5			
78. anthracene (c)	0.010	0.010	5	5	4	1		· · · · · ·
79. benzo(ghi)perylene	0.010	0.010	5	5	5			
80. fluorene	0.010	0.010	5 -	5	-5			
81. phenanthrene (c)	0.010	0.010	5	5	4	1		
82. dibenzo(a,h)anthracene	0.010	0.010	5	5	5			
	0.010	0.010	5	5	5			
84. pyrene	0.010	0.010	5	5	4			1
85. tetrachloroethylene	0.010	0.010	5	9	3	2		4
86. toluene	0.010	0.010	5	9	2	2		5
87. trichloroethylene	0.010	0.010	5	9	5	I		3
88. vinyl chloride	0.010	0.010	5	9	9			
89. aidrin	0.005	0.010	5	5	5			
90. dieldrin	0.005	0.010	5	5	3	2		
91. chlordane	0.005	0.010	5	5	2	3		
92. 4,4'-DDT	0.005	0.010	5	5	3	2		
93. 4,4'-DDE	0.005	0.010	5	5	3	2		
94. 4.4'-DDD	0.005	0.010	5	5	5			
95. alpha-endosulfan	0.005	0.010	5	5	:5		4	
96. beta-endosulfan	0.005	0.010	5	5	5		*	
97. endosulfan sulfate	0.005	0.010	5	5.	5		•	
98. endrin	0.005	0.010	5	5	3	- 2		
99. endrin aldehyde	0.005	0.010	5	5	. 4	1		
100. heptachlor	0.005	0.010	5	5	3	2		•
101. heptachlor epoxide	0.005	0.010	5	5	5			r
102. alpha-BHC	0.005	0.010	5	5	4	1		
103. beta-BHC	0.005	0.010	5	5	3	2		
104. gamma-BHC	0.005	0.010	5	5	4	1		
105. delta-BHC	0.005	0.010	5	5	5			
106. PCB-1242 (d)	0.005	0.010	5	5	4			1
107. PCB-1254 (d)	0.005		-					
108. PCB-1221 (d)	0.005			_		<u>.</u>		
109. PCB-1232 (e)	0.005	0.010	5 -	5		4		1
110. PCB-1248 (e)	0.005							
111. PCB-1260 (e)	0.005							
112. PCB-1016 (e)	0.005			. `				

SECONDARY SILVER SUBCATEGORY SECT

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Table VI-1 (Continued)

FREQUENCY OF OCCURRENCE OF TOXIC POLLUTANTS SECONDARY SILVER RAW WASTEWATER

Pollutant	Analytical Quantification Concentration (mg/l)(a)	Treatable Concentra- <u>tion (mg/l)</u> (b)	Number of Streams Analyzed	Number of Samples Analyzed	ND	Detected Below Quantification Concentration	Detected Below Treat- able Concen- tration	Detected Above Treat- able Concen- tration
113. toxapliene	0.005	0.010	5	5	3	2		
114. artimo y	0.100	0.47	5	5	2			3
115. a. senic	0.010	0.34	5	5			. 3	2
116. asbestos	10 MFL	10 MFL	2	2	1	1		
117. beryllium	0.010	0.20	5	5	5	<u>۱</u> ۰		•
118. cadmium	0.002	0.49	5	5			•	5
119. chromium	0.005	0.07	5	5				5
120. copper	0.009	0.39	- 5	5				5
121. cyanide	0.02 (f)	0.047	5	9	1	2		6
122. lead	0.020	0.08	5	5				5
123. mercury	0.0001	0.036	4	4	1 1		2	1
124. nickel	0.005	0.22	5	5	1 1			4
125. selenium	0.01	0.20	5	5	2			3
126. silver	0.02	0.07	5	5	2			3
127. thallium	0.100	0.34	5	5	3		. 1	1
128. zinc	0.050	0.23	- 5	, 5				5
129. 2,3,7,8-tetrachlorodibenzo- p-dioxin (TCDD)		Not Analyzed	•				ι	

(a) Analytical quantification concentration was reported with the data (see Section V).

) Treatable concentrations are based on performance of lime precipitation, sedimentation, and filtration for toxic metal pollutants and activated carbon adsorption for toxic organic pollutants.

,(d),(e) Reported together.

(1, Analytical quantification concentration for EPA Method 335.2, Total Cyanide Methods for Chemical Analysis of Water and Wastes, EPA-600/4-79-020, March 1979.

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TABLE VI-2

TOXIC POLLUTANTS NEVER DETECTED

2. acrolein 3. acrylonitrile 5. benzidine 8. 1,2,4-trichlorobenzene hexachlorobenzene 9. 12. hexachloroethane 1,1-dichloroethane 13. 14. 1,1,2-trichloroethane chloroethane 16. 17. DELETED 18. bis(2-chloroethyl) ether 19. 2-chloroethyl vinyl ether 20. 2-chloronaphthalene 21. 2,4,6-trichlorophenol parachlorometa cresol 22. 24. 2-chlorophenol 25. 1,2-dichlorobenzene 26. 1,3-dichlorobenzene 27. 1,4-dichlorobenzene 3,3'-dichlorobenzidine 28. 2,4-dichlorophenol 31. 32. 1,2-dichloropropane 1,3-dichloropropylene 33. 34. 2,4-dimethylphenol 2,4-dinitrotoluene 35. 36. 2,6-dinitrotoluene 37. 1,2-diphenylhydrazine 39. fluoranthene 40. 4-chlorophenyl phenyl ether 41. 4-bromophenyl phenyl ether 42. bis(2-chloroisopropyl) ether 43: bis(2-chloroethoxy) methane 45. methyl chloride 46. methyl bromide 48. dichlorobromomethane 49. DELETED 50. DELETED 52. hexachlorobutadiene 53. hexachlorocyclopentadiene 54. isophorone 55. naphthalene 56. nitrobenzene 57. 2-nitrophenol 58. 4-nitrophenol 59. 2,4-dinitrophenol

TABLE VI-2 (Continued)

TOXIC POLLUTANTS NEVER DETECTED

61. N-nitrosodimethylamine 62. N-nitrosodiphenylamine 63. N-nitrosodi-n-propylamine 64. pentachlorophenol 65. phenol dimethyl phthalate 71. 72. benzo(a)anthracene 73. benzo(a)pyrene 74. 3,4-benzofluoranthene 75. benzo(k)fluoranthene 76. chrysene 77. acenaphthylene 79. benzo(ghi)perylene 80. fluorene 82. dibenzo(a,h)anthracene 83. indeno(1,2,3-cd)pyrene vinyl chloride 88. 89. aldrin 4, 4' - DDD94. alpha-endosulfan 95. 96. beta-endosulfan

2,3,7,8-tetrachlorodibenzo-p-dioxin

97. endosulfan sulfate 101. heptachlor epoxide

delta-BHC

beryllium

105.

117.

129.

2763

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SECTION VII

CONTROL AND TREATMENT TECHNOLOGIES

The preceding sections of this supplement discussed the sources, flows, and characteristics of the wastewaters from secondary silver plants. This section summarizes the description of these wastewaters and indicates the level of treatment which is currently practiced by the secondary silver subcategory for each waste stream.

CURRENT CONTROL AND TREATMENT PRACTICES

This section presents a summary of the control and treatment technologies that are currently being applied to each of the sources generating wastewater in this subcategory. As discussed Section V, wastewater associated with the secondary silver in subcategory is characterized by the presence of the toxic metal pollutants and suspended solids. (The raw (untreated) wastewater data for specific sources as well as combined waste streams are presented in Section V.) Generally, these pollutants are present in each of the waste streams at concentrations above treatable concentrations, so these waste streams are commonly combined for treatment to reduce the concentrations of these pollutants. Construction of one wastewater treatment system for combined treatment allows plants to take advantage of economies of scale in some instances, to combine streams of differing and, alkalinity to reduce treatment chemical requirements. Twenty-two plants in this subcategory currently have combined wastewater precipitation five treatment systems, have lime and sedimentation, two have lime precipitation, sedimentation and filtration, and nine have lime precipitation and filtration. As such, three options have been selected for consideration for BPT, BAT, BDT, and pretreatment in this subcategory, based on combined treatment of these compatible waste streams.

FILM STRIPPING

The film base (residue) resulting from the stripping of photographic film can be screened and rinsed, producing wastewater. One of the three plants with this process reported an effluent, none of which is recycled. As discussed in Section V, this wastewater should contain treatable concentrations of toxic metals, oil and grease, cyanide, phenolics, and total suspended solids. The one plant treats film stripping wastewater with sedimentation, pH adjustment, a trickling filter, followed by an activated sludge system.

FILM STRIPPING WET AIR POLLUTION CONTROL AND PRECIPITATION AND FILTRATION OF FILM STRIPPING SOLUTIONS WET AIR POLLUTION CONTROL

One of the three plants engaged in film stripping and precipitation of film stripping solutions uses a wet scrubber to

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control air emissions. Toxic organics, toxic metals, phenolics, total suspended solids, and cyanide should be present at treatable concentrations. This plant practices 99+ percent recycle of film stripping scrubber water. Treatment of the wastewater consists of neutralization, flocculation, and sedimentation, followed by polishing filtration.

PRECIPITATION AND FILTRATION OF FILM STRIPPING SOLUTIONS

Depleted silver solutions from film stripping must be discarded after precipitation. Four of five plants discharge this wastewater. Toxic organics, toxic metals, total suspended solids, phenolics, and cyanide should be present at treatable concentrations. No plants reported recycling this wastewater. Treatment at one plant consists of settling, pH adjustment, trickling filtration, and an activated sludge system. Another plant treats by neutralization with caustic soda or acid, flocculation by polymer addition, and settling followed by polishing filtration. Two plants discharge into municipal sewer lines without treatment.

PRECIPITATION AND FILTRATION OF PHOTOGRAPHIC SOLUTIONS

Silver-free solutions are usually discarded after precipitation. All seven plants precipitating photographic solutions produce wastewater from this process. Treatable concentrations of ammonia and toxic metals characterize this wastewater. Most suspended solids will have been removed with the silver precipitate during filtration. There are no plants that recycle this wastewater. A number of treatment methods are applied before this wastewater is discharged. They are:

- 1. Neutralization with limestone one plant,
- 2. Neutralization with caustic and filtration one plant,
- 3. Neutralization with caustic, sedimentation, and filtration one plant,
- 4. Settling, pH adjustment, trickling filter, and activated sludge system one plant,
- 5. No treatment two plants, and
- 6. Contractor disposal one plant.

PRECIPITATION AND FILTRATION OF PHOTOGRAPHIC SOLUTIONS WET AIR POLLUTION CONTROL

Three plants use wet scrubbers on precipitation and filtration processes. The wastewater characteristics are similar to scrubber wastewater from film stripping precipitation because of the similar materials and processes used. Toxic organics, metals, cyanide, and total suspended solids should be present in this wastewater at treatable concentrations. One plant practices complete recycle of silver solution scrubber water. The three others practice partial recycle of the scrubber liquor (99+ percent). The following treatment schemes are currently in use in the subcategory:

- 1. 100 percent evaporation one plant,
- 2. Neutralization with caustic, polymer addition,
- sedimentation, and polishing filtration one plant,
- 3. No treatment one plant.

ELECTROLYTIC REFINING

Wastewater discharges from electrolytic refining consist of spent electrolyte solution and water from washing the recovered silver. Of the 15 plants having an electrolytic refining process, 13 discharge wastewater. This wastewater should contain treatable concentrations of carbon tetrachloride, pyrene, bromoform, benzene, and tetrachloroethylene. Toxic metals, ammonia, cyanide, and total suspended solids are present above treatable concentrations. One plant reported recycling the spent electrolyte to a precipitation process. The following treatment methods are currently practiced:

- 1. No treatment three plants,
- 2. Neutralization with caustic one plant,
- 3. Precipitation with sodium bicarbonate and sedimentation - one plant,
- 4. Contractor disposal two plants,
- 5. Neutralization with caustic and sedimentation one plant,
- 6. Neutralization with caustic, flocculation with alum, and sedimentation one plant,
- Zinc cementation to recover precious metals, neutralization with caustic, polymer addition, and pressure filtration - one plant,
- 8. Iron cementation to recover precious metals and pH adjustment (chemical unspecified) one plant,
- 9. Neutralization with caustic and filtration one plant, 10. Iron cementation to recover precious metals.
- Iron cementation to recover precious metals, neutralization with caustic and sedimentation - one plant.

FURNACE WET AIR POLLUTION CONTROL

Air emission sources in secondary silver furnace operations are incinerators, roasting and drying furnaces, and melting furnaces. Nineteen secondary silver producers control air emissions, using various methods. These are:

- 1. Baghouse seven plants,
- 2. Dry electrostatic precipitator three plants,
- 3. Afterburner four plants,
- 4. Wet scrubber eight plants (includes Venturi, wet electrostatic precipitator, and spray type scrubbers),
- 5. Wet scrubber and baghouse two plants,
- 6. Wet scrubber, afterburner and baghouse two plants,
- 7. Afterburner and baghouse one plant.

Total suspended solids should be present at treatable concentrations in the wastewater produced by wet air pollution

control. Six plants producing this wastewater practice complete recycle or evaporation. Two others practice partial recycle (>99 percent). Treatment methods used are:

- 1. No treatment five plants,
- 2. 100 percent evaporation one plant,
- Neutralization and precipitation with caustic, sodium sulfide or calcium chloride, flocculation with polymer, and sedimentation - one plant,
- 4. Contractor disposal one plant.
- 5. Neutralization with caustic, filtration, evaporation and complete recycle one plant.
- 6. Complete recycle-mechanism not reported two plants.

LEACHING

Of the 13 nonphotographic silver plants that leach, 11 produce wastewater. This wastewater should contain treatable concentrations of toxic organics and metals, ammonia, cyanide, phenolics, and suspended solids. One plant recovers precious metals from the waste by electrolysis. Wastewater treatment methods used are:

- 1. Neutralization with caustic, filtration, and ion exchange one plant,
- 2. Neutralization with lime, polymer addition, and sedimentation - one plant,
- 3. Contractor disposal two plants,
- Neutralization and precipitation with caustic, sodium sulfide, or calcium chloride, followed by flocculation with polymer and sedimentation - one plant,
- 5. Evaporation one plant,
- 6. No treatment one plant.

LEACHING WET AIR POLLUTION CONTROL AND PRECIPITATION OF NONPHOTOGRAPHIC SOLUTIONS WET AIR POLLUTION CONTROL

All 13 plants that leach nonphotographic materials reported air emissions controls. Devices commonly used are packed bed, spray tower, and venturi scrubbers. This wastewater contains treatable concentrations of toxic metals and total suspended solids. Three plants practice complete recycle of the scrubber water. Seven other plants recycle from 65 to 99+ percent. Treatment methods used consist of:

- Neutralization and precipitation with caustic, sodium sulfide, or calcium chloride, followed by flocculation with polymer and sedimentation - one plant,
- 2. Neutralization with caustic, polymer addition, sedimentation, and polishing filtration one plant,
- 3. Neutralization with caustic, polymer addition, and sedimentation one plant,
- 4. Neutralization in a limestone bed one plant,
- Iron cementation and pH adjustment (chemical unspecified) - one plant,

- 6. Neutralization with caustic one plant,
- 7. Neutralization with caustic and filtration one plant,
- 8. Evaporation one plant,
- 9. Contractor disposal one plant,
- 10. No treatment three plants.

PRECIPITATION AND FILTRATION OF NONPHOTOGRAPHIC SOLUTIONS

All 27 of the silver plants with this process produce wastewater. This wastewater should contain toxic organics, toxic metals, ammonia, cyanide, phenolics, and total suspended solids. No plants reported recycling this waste stream. Treatment methods for this wastewater consist of:

- 1. Neutralization and precipitation with caustic, sodium sulfide, or calcium chloride, followed by flocculation with polymer and sedimentation one plant,
- Neutralization with caustic, polymer addition, sedimentation, and polishing filtration - one plant,
- 3. Neutralization with caustic and sedimentation two plants,
- Neutralization with caustic, polymer addition, and sedimentation - one plant,

5. Neutralization with caustic and filtration - one plant,

- 6. Contractor disposal three plants,
- 7. Neutralization with caustic four plants,
- 8. Line precipitation, polymer addition, and sedimentation - two plants,
- 9. Zinc cementation, caustic neutralization, polymer addition, sedimentation, and pressure filtration - one plant,
- 10. Chlorine addition and neutralization with caustic one plant,
- 11. pH adjustment (chemical unspecified) one plant,
- 12. Neutralization with limestone two plants,
- 13. Chlorination, lime precipitation, sodium sulfide precipitation, and sedimentation one plant,
- 14. Caustic and ammonia addition, sedimentation, and filtration one plant,
- 15. Sodium hydrosulfite and caustic addition, recycle of sludge to process, ion exchange, pH adjustment with sulfuric acid one plant,
- 16. Neutralization with caustic, flocculation with alum, and sedimentation one plant,
- 17. Iron cementation and pH adjustment (chemical unspecified) one plant,
- Iron cementation, neutralization with caustic, and sedimentation - one plant,
- 19. No treatment two plants.

FLOOR AND EQUIPMENT WASHDOWN

Many plants wash equipment and floors to recover silver values that may be contained in accidental leaks and spills of process solutions. Data on treatment of floor wash water were not generally available in the data collection portfolios. However, some plants practice cementation to recover the precious metals before discharging the wastewater to POTW or central treatment. This wastewater contains treatable concentration of toxic metals and total suspended solids. As described above, central treatment usually consists of neutralization with lime or caustic and sedimentation. Eleven plants use filtration either after sedimentation or as a solids removal step to replace sedimentation.

CONTROL AND TREATMENT OPTIONS

Based on an examination of the wastewater sampling data, three control and treatment technologies that effectively control the pollutants found in secondary silver wastewaters were evaluated after proposal. These technology options are discussed below. The effectiveness of these technologies is presented in Section VII of the General Development Document.

Other treatment technologies included activated alumina adsorption (Option D) and reverse osmosis (Option F). Although these technologies are theoretically applicable to wastewaters generated in the secondary silver subcategory, they were not selected for evaluation because they are not demonstrated in the nonferrous metals manufacturing category, nor are they clearly transferable.

OPTION A

Option A for the secondary silver subcategory requires treatment technologies to reduce pollutant mass. The Option A treatment scheme consists of ammonia steam stripping preliminary treatment applied to precipitation and filtration of photographic Preliminary treatment is followed by lime and settle solutions. (chemical precipitation and sedimentation) applied to the combined stream steam stripper effluent and the combined stream of all other wastewater. Complete recycle of treated floor and included. equipment washdown wastewater is also Chemical precipitation is used to remove metals by the addition of lime followed by gravity sedimentation. Suspended solids are also removed from the process.

OPTION B

Option B for the secondary silver subcategory consists of the ammonia steam stripping, lime precipitation, and sedimentation technology considered in Option A plus control technologies to reduce the discharge of wastewater volume. Complete recycle of treated floor and equipment washdown wastewater is also included. Water recycle and reuse of scrubber water is the principal control mechanism for additional flow reduction.

OPTION C

Option C for the secondary silver subcategory consists of the

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in-process flow reduction, lime ammonia steam stripping, precipitation, and sedimentation technology considered in Option B plus multimedia filtration technology added at the end of the Option B treatment scheme. Complete recycle of treated floor and equipment washdown wastewater is also included. Multimedia is used to remove suspended solids, including filtration precipitates of metals, beyond the concentration attainable by gravity sedimentation. The filter suggested is of the gravity, mixed media type, although other forms of filters such as rapid sand filters or pressure filters would perform satisfactorily. The addition of filters also provides consistent removal during periods in which there are rapid increases in flows or loadings of pollutants to the treatment system.

additional treatment technology was considered prior to An proposing effluent limitations for this subcategory as discussed below. Activated carbon adsorption was rejected because it is not necessary since toxic organic pollutants are not selected for limitation in this subcategory. (Refer to discussion of regulated pollutant parameters in Section X.)

OPTION E

Option E for the secondary silver subcategory consisted of the in-process flow lime reduction, ammonia steam stripping, filtration sedimentation, and multimedia precipitation, technology considered in Option C with the addition of granular activated carbon technology at the end of the Option C treatment The activated carbon process is utilized to control scheme. discharge of toxic organics.

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SECTION VIII

COSTS, ENERGY, AND NONWATER QUALITY ASPECTS

This section describes the method used to develop the costs associated with the control and treatment technologies discussed in Section VII for wastewaters from secondary silver plants. The energy requirements of the considered options as well as solid waste and air pollution aspects are also discussed.

As discussed in Section VII, three control and treatment options have been developed for the secondary silver subcategory. The options are summarized below and schematically presented in Figures X-1 through X-3 (pages 2818 - 2820).

OPTION A

Option A requires preliminary ammonia steam stripping treatment, and end-of-pipe technology consisting of lime precipitation, sedimentation, and complete recycle of treated floor and equipment washdown wastewater. The stream that will require ammonia steam stripping preliminary treatment is precipitation and filtration of photographic solutions wastewater.

OPTION B

Option B requires in-process flow reduction measures, preliminary ammonia steam stripping treatment, and end-of-pipe treatment technology consisting of lime precipitation, sedimentation, and complete recycle of floor and equipment washdown wastewater. The in-process flow reduction measures consist of the recycle of wet air pollution control water through holding tanks.

OPTION C

Option C requires the in-process flow reduction measures of Option B, preliminary ammonia steam stripping treatment, and endof-pipe treatment technology consisting of lime precipitation, sedimentation, complete recycle of treated floor and equipment washdown wastewater, and multimedia filtration.

A detailed discussion of the methodology used to develop the compliance costs is presented in Section VIII of the General Development Document. Plant-by-plant compliance costs have been estimated for the nonferrous metals manufacturing category and are presented in the administrative record supporting this regulation. A comparison of the costs developed for proposal and the revised costs for the final regulation are presented in Tables VIII-1 and VIII-2 (page 2776) for the direct and indirect dischargers, respectively.

Each of the major assumptions used to develop compliance costs is presented in Section VIII of Vol. I. However, each subcategory

contains a unique set of waste streams requiring certain subcategory-specific assumptions to develop compliance costs. Six major assumptions are discussed briefly below.

- (1) Since 23 of the plants whose compliance costs were estimated overlap with other nonferrous manufacturing subcategories or categories, costs are apportioned to each subcategory on a flow-weighted basis.
- (2) Although a discharge allowance for floor wash is not necessary, a flow of 1 liter of floor wash per troy ounce is used for cost estimation purposes for each plant on the basis of total production of all precious metals (including silver) that results in precipitation and filtration wastewater. Since acceptable floor wash water may be obtained from recycling treated wastewater, costs are estimated for a holding tank after chemical precipitation and settling to recycle water for floor wash use under all options.
- (3) Sodium hydroxide addition was used throughout the secondary silver subcategory in estimating costs for chemical precipitation since it is likely that most plants will recycle treatment plant sludges for additional metal recovery.
- (4) When a plant reported recycle of treatment plant sludges, capital and annual costs for sludge handling (vacuum filtration and contract hauling) are not included. Where the sludge disposal method is reported as contract hauling, or is unknown, contract hauling costs are included assuming nonhazardous disposal.
- (5) Recycle of air pollution control scrubber liquor is based on recycle through holding tanks. Annual costs associated with maintenance and sludge disposal are included in the estimated compliance costs. If a plant currently recycles scrubber liquor, capital costs of the recycle equipment (piping, pumps, and holding tanks) were not included in the compliance costs.
- (6) Costs for ammonia removal for streams with flow rates below 50 liters per hour (none of which are air pollution streams) are estimated using an air stripping system. Ammonia steam stripping is not considered feasible due to insufficient hydraulic loading in the stripping column (given the minimum column diameter of 2 feet used in cost estimation).

The chemical precipitation tank is used for the air stripping operation. Chemical precipitation is always operated in the "low flow" batch treatment mode with a five day holdup due to the low flow rate (see the discussion on chemical precipitation in Section VIII of the General Development Document for a description of

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the "low flow" batch treatment mode). An air sparger is incorporated into the reactor tank. The influent is sparged while the tank fills with wastewater, i.e., over the entire five day holdup period. A hood is placed over the tank to capture any ammonia-laden vapors.

Direct capital costs for the ammonia air stripping system include a blower, a sparger system, and a ventilation hood. Direct annual costs are assumed to consist solely of blower operation and maintenance costs. These are assumed to be 5 percent of the blower capital cost.

NONWATER QUALITY ASPECTS

A general discussion of the nonwater quality aspects of the control and treatment options considered for the nonferrous metals category is contained in Section VIII of the General Development Document. Nonwater quality impacts specific to the secondary silver subcategory including energy requirements, solid waste, and air pollution are discussed below:

ENERGY REQUIREMENTS

The methodology used for determining the energy requirements for the various options is discussed in Section VIII of Vol. I. Energy requirements for the three options considered are estimated at 0.88 mwh/yr, 0.88 mwh/yr, and 0.93 mwh/yr for Options A, B, and C respectively. Option C represents roughly eight percent of a typical plant's electrical usage. It is therefore concluded that the energy requirements of the treatment options considered will have no significant impact on total plant energy consumption.

SOLID WASTES

Sludges associated with the secondary silver subcategory will necessarily contain additional quantities (and concentrations) of toxic metal pollutants. Wastes generated by secondary metals industries can be regulated as hazardous. However, the Agency examined the solid wastes that would be generated at secondary nonferrous metals manufacturing plants by the suggested treatment technologies and believes they are not hazardous wastes under the Agency's regulations implementing Section 3001 of the Resource Conservation and Recovery Act. None of these wastes are listed specifically as hazardous. Nor are they likely to exhibit a characteristic of hazardous waste. This judgment is made based the recommended technology of lime precipitation, on sedimentation, and filtration. By the addition of excess lime during treatment, similar sludges, specifically toxic metal bearing sludges, generated by other industries such as the iron and steel industry, passed the Extraction Procedure (EP) toxicity test. See 40 CFR 8261.24. Thus, the Agency believes that the

waste water sludges will similarly not be EP toxic if the recommended technology is applied.

Although it is the Agency's view that solid wastes generated as a result of these guidelines are not expected to be hazardous, generators of these wastes must test the waste to determine if the wastes meet any of the characteristics of hazardous waste (see 40 CFR 262.11).

these wastes should be identified or are listed as hazardous, If they will come within the scope of RCRA's "cradle to grave" hazardous waste management program, requiring regulation from the point of generation to point of final disposition. EPA's generator standards would require generators of hazardous nonferrous metals manufacturing wastes to meet containerization, labeling, recordkeeping, and reporting requirements; if plants dispose of hazardous wastes off-site, they would have to prepare a manifest which would track the movement of the wastes from the generator's premises to a permitted off-site treatment, storage, or disposal facility. See 40 CFR 262.20 45 FR 33142 (May 19, 1980), as amended at 45 FR 86973 (December 31, 1980). The transporter regulations require transporters of hazardous wastes to comply with the manifest system to assure that the wastes are delivered to a permitted facility. See 40 CFR 263.20 45 FR 33151 (May 19, 1980), as amended at 45 FR 86973 (December 31, 1980).

Finally, RCRA regulations establish standards for hazardous waste treatment, storage, and disposal facilities allowed to receive such wastes. See 40 CFR Part 464 46 FR 2802 (January 12, 1981), 47 FR 32274 (July 26, 1982).

Even if these wastes were not identified as hazardous, they still must be disposed of in compliance with the Subtitle \bar{D} open dumping standards, implementing 4004 of RCRA. See 44 FR 53438 (September 13, 1979). The Agency has calculated as part of the costs for wastewater treatment the cost of hauling and disposing of these wastes. The Agency estimates implementation of lime and settle technology will generate approximately 2,900 tons per year of wastewater treatment sludge. Multimedia filtration technology will not generate any significant amount of sludge over that resulting from lime precipitation.

AIR POLLUTION

There is no reason to believe that any substantial air pollution problem will result from implementation of ammonia steam stripping, chemical precipitation, sedimentation, and multimedia filtration. These technologies transfer pollutants to solid waste and are not likely to transfer pollutants to air.

At three secondary silver plants, streams with treatable concentrations of ammonia having flows less than 50 l/hr were treated with air stripping for design and cost determination. None of the waste streams were air pollution control streams. The air stripping is accomplished by aeration and agitation in

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the chemical precipitation batch tank, which includes a ventilation hood. Air stripping is not a model treatment technology because it simply transfers the ammonia from one medium to another, whereas steam stripping allows for ammonia recovery, and if desired, reuse. Air stripping was used in costing instead of steam stripping because at such low flow, continuous operation of steam strippers is not possible. Therefore, the treatable concentration for ammonia would be difficult to attain. The Agency does not believe that under these circumstances (low flow, non-air pollution control streams) that air stripping will create an air quality problem.

TABLE VIII-1

COST OF COMPLIANCE FOR THE SECONDARY SILVER SUBCATEGORY DIRECT DISCHARGERS

Option		osal	Promulgation		
operon	capital Cost	Annual Cost	<u>Capital</u> Cost	<u>Annual</u> Cost	
А	169000	357000	110000	309000	
В	250000	379000	110000	309000	
С	280000	469000	278000	390000	

TABLE VIII-2

COST OF COMPLIANCE FOR THE SECONDARY SILVER SUBCATEGORY INDIRECT DISCHARGERS

Option	Prop <u>Capit</u> al Cost	osal Annual Cost	Promulgation Capital Cost Annual Cost		
A	1066000	1233000	596000	<u>381000</u>	
В	1400000	1302000	577000	385000	
C .	1549000	1454000	534000	422000	

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SECTION IX

BEST PRACTICABLE CONTROL TECHNOLOGY CURRENTLY AVAILABLE

This section defines the effluent characteristics attainable through the application of best practicable control technology currently available (BPT). BPT reflects the existing performance by plants of various sizes, ages, and manufacturing processes within the secondary silver subcategory, as well as the established performance of the recommended BPT systems. Particular consideration is given to the treatment already in place at plants within the data base.

The factors considered in identifying BPT include the total cost of applying the technology in relation to the effluent reduction benefits from such application, the age of equipment and the manufacturing processes facilities involved, employed, environmental impacts (including energy nonwater quality requirements), and other factors the Administrator considers appropriate. In general, the BPT level represents the average of the existing performances of plants of various ages, sizes, processes, or other common characteristics. Where existing performance is uniformly inadequate, BPT may be transferred from different subcategory or category. Limitations based on а transfer of technology are supported by a rationale concluding that the technology is, indeed, transferable, and a reasonable prediction that it will be capable of achieving the prescribed effluent limits. BPT focuses on end-of-pipe treatment rather than process changes or internal controls, except where such practices are common within the subcategory.

TECHNICAL APPROACH TO BPT

The Agency studied the nonferrous metals manufacturing category to identify the processes used, the wastewaters generated, and the treatment processes installed. Information was collected from industry using data collection portfolios, and specific plants were sampled and the wastewaters analyzed. Some of the factors which must be considered in establishing effluent limitations based on BPT have already been discussed. The age of equipment and facilities, processes used, and raw materials were taken into account in subcategorization and subdivision and are discussed fully in Section IV. Nonwater quality impacts and energy requirements are considered in Section VIII.

As explained in Section IV, the secondary silver subcategory has been subdivided into 11 potential wastewater sources. Since the water use, discharge rates, and pollutant characteristics of each of these wastewaters is potentially unique, effluent limitations will be developed for each of the 11 subdivisions.

For each of the subdivisions, a specific approach was followed the development of BPT mass limitations. To account for for production and flow variability from plant to plant, a unit of production or production normalizing parameter (PNP) was determined for each waste stream which could then be related to the flow from the process to determine a production normalized flow. Selection of the PNP for each process element is discussed in Section IV. Each process within the subcategory was then analyzed to determine (1) whether or not operations included generated wastewater, (2) specific flow rates generated, and (3) the specific production normalized flows for each process. This analysis is discussed in detail in Section \bar{v} . Nonprocess wastewater, such as rainfall runoff and noncontact cooling water, is not considered in the analysis.

Normalized flows were analyzed to determine which flow was to be used as part of the basis for BPT mass limitations. The selected flow (sometimes referred to as a BPT regulatory flow or BPT discharge rate) reflects the water use controls which are common practices within the subcategory. The BPT normalized flow is based on the average of all applicable data. Plants with normalized flows above the average may have to implement some method of flow reduction to achieve the BPT limitations. In most cases, this will involve improving housekeeping practices, better maintenance to limit water leakage, or reducing excess flow by turning down a flow valve. It is not believed that these modifications would incur any costs for the plants.

For the development of effluent limitations, mass limitations were calculated for each wastewater source or subdivision. This calculation was made on a stream-by-stream basis, primarily because plants in this category may perform one or more of the operations in various combinations. The mass limitations (milligrams of pollutant per troy ounce of production unit mg/troy ounce) were calculated by multiplying the BPT normalized flow (l/troy ounce) by the achievable treatment concentrations using the BPT treatment system (mg/l) for each pollutant parameter to be limited under BPT.

The mass limitations which are allowed under BPT for each plant will be the sum of the individual mass loadings for the various wastewater sources which are found at particular plants. Accordingly, all the wastewater generated within a plant may be combined for treatment in a single or common treatment system, but the effluent limitations for these combined wastewaters are based on the various wastewater sources which actually contribute to the combined flow. This method accounts for the variety of combinations of wastewater sources and production processes which may be found at secondary silver plants.

The Agency usually establishes wastewater limitations in terms of mass rather than concentration. This approach prevents the use of dilution as a treatment method (except for controlling pH). The production normalized wastewater flow (l/troy ounce) is a link between the production operations and the effluent

limitations. The pollutant discharge attributable to each operation can be calculated from the normalized flow and effluent concentration achievable by the treatment technology and summed to derive an appropriate limitation for each subcategory.

BPT effluent limitations are based on the average of the discharge flow rates for each source; consequently, the treatment technologies which are currently used by the lowest dischargers will be the treatment technologies most likely required to meet BPT guidelines. Section VII discusses the various treatment technologies which are currently in place for each wastewater source. In most cases, the current treatment technologies consist of chemical precipitation and sedimentation (lime and settle technology) and a combination of reuse and recycle to reduce flow. Ammonia steam stripping is added to streams containing treatable concentrations of ammonia.

The overall effectiveness of end-of-pipe treatment for the removal of wastewater pollutants is improved by the application of water flow controls within the process to limit the volume of wastewater requiring treatment. The controls or in-process technologies recommended under BPT include only those measures which are commonly practiced within the subcategory and which reduce flows to meet the production normalized flow for each operation.

INDUSTRY COST AND POLLUTANT REMOVAL ESTIMATES

In balancing costs in relation to effluent removal estimates, EPA considers the volume and nature of existing discharges, the volume and nature of discharges expected after application of BPT, the general environmental effects of the pollutants, and the cost and economic impacts of the required pollution control level. The Act does not require or permit consideration of water quality problems attributable to particular point sources or industries, or water quality improvements in particular water quality bodies. Accordingly, water quality considerations were not the basis for selecting the proposed BPT. See <u>Weyerhaeuser</u> Company v. Costle, 590 F.2d 1011 (D.C. Cir. 1978).

The methodology for calculating pollutant removal estimates and plant compliance costs is discussed in Section X. Table X-2 (page 2819) shows the estimated pollutant removal estimates for each treatment option for direct dischargers. Compliance costs are presented in Table VIII-1 (page 2818) for direct dischargers.

BPT OPTION SELECTION - PROPOSAL

The proposed best practicable technology consisted of chemical precipitation and sedimentation (lime and settle technology) with ammonia steam stripping preliminary treatment of wastewaters containing treatable concentrations of ammonia. The best practicable technology is presented schematically in Figure IX-1 (page 2797) of this supplement.

SECONDARY SILVER SUBCATEGORY SECT - IX

Ammonia steam stripping is demonstrated in the nonferrous metals manufacturing category. Two plants in the primary columbiumtantalum subcategory and three plants in the primary tungsten subcategory reported steam stripping in-place.

EPA believes that performance data from the iron and steel manufacturing category provide a valid measure of this technology's performance on nonferrous metals manufacturing category wastewater because raw wastewater concentrations of ammonia are of the same order of magnitude in the respective raw wastewater matrices. A detailed discussion of this technology transfer is found in Section VII of Vol. I.

Chemical analysis data were collected of raw waste (treatment influent) and treated waste (treatment effluent) from one coke plant of the iron and steel manufacturing category. A contractor for EPA, using EPA sampling and chemical analysis protocols, collected data paired samples in a two-month period. These data are the data base for determining the effectiveness of ammonia steam stripping technology and are contained within the public record supporting this document. Ammonia treatment at this coke plant consisted of two steam stripping columns in series with steam injected countercurrently to the flow of the wastewater. A lime reactor for pH adjustment separated the two stripping columns.

The raw untreated wastewater samples from the coke facility contained ammonia concentrations of 599, 226, 819, 502, 984, and 797 mg/l. Raw untreated wastewater samples from the secondary silver subcategory contained ammonia concentrations of 1,202 and 4,630 mg/l.

BPT OPTION SELECTION - PROMULGATION

EPA is promulgating BPT limitations for the secondary silver subcategory based on lime precipitation and sedimentation to remove toxic metals, control pH, and remove TSS and pretreatment with steam stripping to reduce ammonia concentrations. Complete recycle of treated floor and equipment washdown wastewater is also included. The end-of-pipe treatment technology basis for the BPT limitations being promulgated is the same as that for the proposed limitations. Lime and settle treatment technology is currently in place at five direct discharging facilities.

The Agency has collected data on secondary precious metals facilities through data collection portfolios (dcp) so that it may propose mass limitations for the secondary precious metals subcategory. Many of the plants in the subcategory overlap with the secondary silver subcategory. Review of these dcp, and the dcp collected only for the secondary silver subcategory, has led the Agency to revise the regulatory flows. Accordingly, the wastewater streams from film stripping wet air pollution control and precipitation and filtration of film stripping solutions wet air pollution control have been combined into one building block. Leaching wet air pollution control and precipitation of

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nonphotographic solutions wet air pollution control wastewater sources have also been combined into one building block. In addition, the mass limitations proposed for casting contact cooling water and casting wet air pollution control have been eliminated. Analytical data collected at a secondary precious metals plant demonstrate casting contact cooling water is not sufficiently contaminated to warrant treatment. Casting wet air pollution control limitations have been eliminated because the Agency believes this limitation is duplicated by the furnace wet pollution control limitations (these operations are air identical). A flow allowance is not provided for floor and equipment washdown based on reuse of recycled treatment effluent as facility washdown water. In developing compliance cost estimates, the Agency sized treatment equipment to allow for this flow.

The Agency has verified the proposed steam stripping performance values using steam stripping data collected at a zirconiumhafnium plant. Data collected by the plant represent almost two years of daily operations and support the long-term mean and variability used to establish treatment effectiveness.

Several comments were received (although none were from secondary silver plants) stating that ammonia steam stripping performance data transferred from the iron and steel category are not appropriate for the nonferrous metals manufacturing category. Many of the commenters believe plugging of the column due to precipitates will severely affect their ability to achieve the promulgated steam stripping performance values. In developing compliance costs, the Agency designed the steam stripping module to allow for a weekly acid cleaning to reduce plugging problems. Through special information requests, the Agency attempted to gather data at plants which stated they could not achieve the proposed limits. However, very little data were submitted to support their claims or document column performance. Therefore, the Agency has retained the proposed performance, which has been validated with steam stripping data from a zirconium-hafnium facility.

The promulgated BPT will result in the removal of an estimated 30,870 kg of toxic pollutants, 664,000 kg of ammonia, and 7,750 kg of TSS per year from raw discharge levels. The estimated capital investment cost of BPT is \$110,000 (March, 1982 dollars) and the estimated annual cost is \$309,000 (March, 1982 dollars). These costs represent wastewater treatment equipment not currently in place.

WASTEWATER DISCHARGE RATES

A BPT discharge rate is calculated for each subdivision based on the average of the flows of the existing plants, as determined from analysis of the dcp. The discharge rate is used with the achievable treatment concentration to determine BPT effluent limitations. Since the discharge rate may be different for each wastewater source, separate production normalized discharge rates

for each of the 11 wastewater sources are discussed below and summarized in Table IX-1 (page 2789). The discharge rates are normalized on a production basis by relating the amount of wastewater generated to the mass of the intermediate product which is produced by the process associated with the waste stream in question. These production normalizing parameters, or PNPs, are also listed in Table IX-1.

Section V of this supplement further describes the discharge flow rates and presents the water use and discharge flow rates for each plant by subdivision.

COLLECTION OF NEW DATA

In the proposed development document, separate subdivisions were identified for precipitation and filtration of film stripping solutions wet air pollution control, casting contact cooling water, casting wet air pollution control, and precipitation and filtration of nonphotographic solutions wet air pollution control. Based on new data gathered from secondary precious metals dcp and sampling efforts and re-evaluation of existing data, these subdivisions were either combined with other subdivisions or deleted. A subdivision for floor and equipment washdown was added.

Although flow and production data were collected from secondary precious metals dcp in the nonferrous metals manufacturing category, these data were not used to modify the proposed regulatory flow allowances. The new data support the proposed flow allowances and the Agency did not receive any comments suggesting that the allowances should be revised. The new flow data are included in the water use and discharge tables in Section V. Wastewater discharge allowances for the 11 subdivisions of the secondary silver subcategory are discussed below.

The regulatory flow allowance at proposal for casting contact cooling water has been eliminated. Analytical data collected at a secondary precious metals plant demonstrate casting contact cooling water is not sufficiently contaminated to require treatment. However, it is possible that toxic pollutants may be present in larger concentrations at any individual plant than the Agency's sampling data indicate. Therefore, the permitting or controlling authority should check for the presence of toxic pollutants on a case-by-case basis and determine if they require treatment.

FILM STRIPPING

The BPT wastewater discharge rate at proposal for film stripping was 50.35 l/troy ounce of silver produced from film stripping. At proposal, three plants reported wastewater discharges from film stripping, but the dcp data provided by two plants were insufficient to calculate discharge rates. Therefore, the discharge rate from one plant was used.

The BPT wastewater discharge rate is 50.35 l/troy ounce of silver produced from film stripping. The Agency received no new data for this waste stream after proposal. Therefore, the flow allowance is promulgated as proposed.

FILM STRIPPING WET AIR POLLUTION CONTROL AND PRECIPITATION AND FILTRATION OF FILM STRIPPING SOLUTIONS WET AIR POLLUTION CONTROL

The BPT wastewater discharge rate at proposal for film stripping wet air pollution control was 0.485 l/troy ounce of silver produced from film stripping, based on 99 percent recycle. This rate was allocated only for plants practicing wet air pollution control for film stripping. One plant reported this wastewater, recycling 99+ percent. This plant used the same scrubber to control air emissions from film stripping and film stripping precipitation.

The BPT wastewater discharge allowance for film stripping and precipitation of film stripping solutions wet air pollution control is 0.97 l/troy ounce of silver produced from precipitation and filtration of film stripping solutions. Because the one plant that uses a wet scrubber on film stripping also uses the same scrubber on the precipitation of film stripping solutions, the Agency believes only one allowance is necessary. This allowance is based on 99 percent recycle of the water use at the one plant.

PRECIPITATION AND FILTRATION OF FILM STRIPPING SOLUTIONS

The BPT wastewater discharge rate at proposal for film stripping precipitation and filtration waste streams was 50.57 l/troy ounce of silver precipitated. Of the six plants with this process, four reported producing wastewater. The proposed BPT rate was based on the average discharge rate of two plants, which generated 112.7 and 2.31 l/troy ounce. A third plant reported insufficient data to calculate the discharge rate. Another plant reported this waste stream as a combination of photographic and nonphotographic wastewater; therefore, this plant also was omitted from the calculation.

The BPT wastewater discharge rate at promulgation for precipitation and filtration of film stripping solutions is 57.57 l/troy ounce of silver precipitated. Since proposal, data from plant 74 were clarified and a flow was attributed to precipitation of photographic film. The discharge rate at this plant is below the proposed allowance. The Agency received no new data or comments suggesting that the proposed allowance should be changed. Water use and wastewater discharge rates are presented in Table V-3 (page 2718).

PRECIPITATION AND FILTRATION OF PHOTOGRAPHIC SOLUTIONS

The BPT wastewater discharge rate at proposal for the precipitation and filtration of photographic solutions was 26.6 l/troy ounce of silver precipitated. Of the 15 plants reporting this process at proposal, nine discharged wastewater. Four plants did not provide sufficient data to calculate discharge rates. The discharge rates for the five other plants ranged from 1.6 to 89.9 l/troy ounce. The proposed BPT rate was based on the average of the discharge rates of these five plants.

The BPT wastewater discharge rate is 26.6 l/troy ounce of silver precipitated. This is equivalent to the proposed allowance. Data from plant 74, which precipitates silver from solutions resulting from photographic sludges, were added since proposal. However, these data support the proposed allowance. The Agency received no new data demonstrating that the proposed allowance should be changed. The distribution of wastewater discharge rates is presented in Table V-4 (page 2719).

PRECIPITATION AND FILTRATION OF PHOTOGRAPHIC SOLUTIONS WET AIR POLLUTION CONTROL

The BPT wastewater discharge rate at proposal for precipitation and filtration of photographic solutions wet air pollution control was 12.14 1/troy ounce of silver precipitated. This rate was allocated only to plants having wet air pollution control for precipitation and filtration of photographic solutions. Of the 15 plants that reported this process at proposal, four used wet air pollution control devices. Three of the four plants did not report sufficient production data to calculate a discharge rate for this waste stream, although sufficient data were reported to determine recycle practices. One of the four plants achieved zero discharge of this waste stream through complete recycle, while two plants practiced 99 percent recycle or greater. The fourth plant recycled 68 percent of its precipitation and filtration of photographic solutions wet air pollution control water. Thus, extensive recycle is possible for this wastewater stream. However, zero discharge may not be technically feasible unless a recycle system controls dissolved solids buildup, the wastewater is evaporated, or this wastewater can be reused in another production operation that can accept water of this quality. Some of these zero-discharge possibilities are site-specific and, therefore, are not applicable to all secondary silver pollutants that generate this wastewater. Therefore, a BPT wastewater discharge rate was allocated for precipitation and filtration of photographic solutions wet air pollution control. This discharge rate was based on 99 percent recycle of the water used for precipitation and filtration of photographic solutions wet air pollution control at the only plant for which a discharge rate could be determined. In the absence of other data, the Agency normally bases limits on 90 percent recycle of scrubber discharges; however, the plant that the discharge rate was based on recycled 99.9 percent of this wastewater, and two other plants practiced 99 and 100 percent recycle. Thus 99 percent recycle represented current subcategory practices for precipitation and filtration of photographic solutions wet air pollution control water.

The BPT wastewater discharge rate for precipitation and filtration of photographic solutions wet air pollution control is 12.14 l/troy ounce of silver precipitated. This is equivalent to the proposed allowance. Data from plant 74 were added since proposal. However, the scrubber at this plant is used over the film stripping and film stripping precipitation operations as well as the precipitation of photographic solutions process. For this reason, it was not considered representative of this subdivision and was not used to revise the regulatory allowance. The Agency received no new data demonstrating that the proposed flow allowance should be revised. Water use and wastewater discharge rates are presented in Table V-6 (page 2722).

ELECTROLYTIC REFINING

The BPT wastewater discharge rate at proposal for electrolytic refining was 0.76 l/troy ounce of silver refined. Of the 20 plants reporting electrolytic refining operations at proposal, 12 produced wastewater. Four plants reported insufficient data to calculate discharge rates. Data from seven plants, with discharge rates ranging from 0.068 to 1.97 l/troy ounce were used to calculate the BPT rate. Only one plant practiced recycle of this wastewater and achieved zero discharge by 100 percent reuse.

The promulgated BPT wastewater discharge rate for electrolytic refining is 0.76 l/troy ounce of silver refined. This is equivalent to the proposed allowance. New data received by the Agency after proposal support this allowance. The Agency believes there is no reason to change the proposed allowance based on the data received. The water use and discharge rates are presented in Table V-7 (page 2723).

FURNACE WET AIR POLLUTION CONTROL

The BPT wastewater discharge rate at proposal for the furnace air wet scrubbing stream was 0.67 1/troy ounce of silver smelted, roasted, or dried. This rate was allocated only for plants practicing wet air pollution control for furnace emissions. Emissions from furnace operations are controlled by dry or wet Common dry methods involve baghouses or control devices. dry electrostatic precipitators. Wet devices include packed bed, spray, and Venturi scrubbers, and wet electrostatic precipitators. Of the 19 plants reporting furnace air pollution control at proposal, 11 produced waste streams. Seven of the eleven plants achieved zero discharge through 100 percent Two of the four plants that discharged this waste recycle. stream practiced 99 percent recycle or greater, while one plant used a once-through operation. The remaining plant did not report production or wastewater flow data for this waste stream. The proposed BPT discharge rate was based on 99 percent recycle of the average water use at the three plants for which discharge rates were determined. The 99 percent recycle basis represented current subcategory practices since nine of the eleven plants that produced this waste stream recycled 99 percent or greater.

The BPT wastewater discharge rate for furnace wet air pollution control is 0.67 l/troy ounce of silver smelted, roasted, or dried. This is equivalent to the proposed BPT allowance. The Agency received new data from one plant with this waste stream. These data support the proposed allowance. The water use and wastewater discharge rates are shown in Table V-9 (page 2732). There are no new data demonstrating that proposed allowances should be changed. This allowance also includes the casting wet air pollution control allowance which was proposed. These two operations were identical and two allowances are not justified. Plants having smelting furnaces and casting furnaces use the same scrubber on both operations (e.g., plant 553).

LEACHING

The BPT discharge rate at proposal for plants with nonphotographic leaching processes was 0.086 1/troy ounce of silver produced from leaching. Of the 15 plants using this process at proposal, 12 discharged wastewater. Six plants supplied sufficient information to calculate discharge rates. Three plants with once-through discharge had rates ranging from 0.068 to 0.11 1/troy ounce. The proposed BPT rate was an average of the discharge from these three plants. Three other oncethrough dischargers reported rates ranging from 2.7 to 635.2 1/troy ounce. The rates from these three plants were omitted from the BPT rate calculation because there was no reason to believe that water is needed in these amounts, in light of rates from the other plants.

The BPT wastewater discharge rate for leaching is 0.086 1/troy ounce of silver produced from leaching. This is equivalent to the proposed allowance. Since proposal it was determined that the wastewater reported at plant 549 for leaching was actually precipitation of nonphotographic solutions wastewater. This did not affect the regulatory flow because this plant was not used to calculate the proposed allowance. Data from a plant received after proposal support the proposed allowance. The distribution of wastewater discharge rates is shown in Table V-10 (page 2733). The Agency received no data demonstrating that the proposed allowances should be revised.

LEACHING WET AIR POLLUTION CONTROL AND PRECIPITATION OF NONPHOTOGRAPHIC SOLUTIONS WET AIR POLLUTION CONTROL

The BPT wastewater discharge rate at proposal for nonphotographic leaching wet scrubbing was 4.43 l/troy ounce of silver produced from leaching. This rate was allocated only for plants using wet air pollution control on leaching processes. At proposal, three plants achieved zero discharge through 100 percent recycle or reuse. The recycle in seven additional plants ranged from 65 to 99+ percent, four of those using at least 99 percent. Some of

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the zero discharge possibilities were site-specific and are not applicable on a nationwide basis. The proposed BPT discharge rate was based on the average of five plants with discharge rates ranging from 0.014 to 11.3 1/troy ounce. Insufficient data to calculate a discharge rate were reported from three of the eight discharging plants.

The BPT wastewater discharge allowance is 4.43 l/troy ounce silver produced from leaching or silver precipitated. This of is This allowance also equivalent to the proposed allowance. precipitation of proposed allowance for the replaces nonphotographic solutions wet air pollution control. The Agency determined that the same scrubbers were used over both processes and precipitation) and two allowances were not (leaching Water use and wastewater discharge rates are justified. presented in Table V-11 (page 2734). The Agency received no new data demonstrating that the proposed allowance should be revised. The one plant submitting data for this scrubber subsequent to proposal practices 100 percent recycle.

PRECIPITATION AND FILTRATION OF NONPHOTOGRAPHIC SOLUTIONS

The BPT wastewater discharge rate at proposal for nonphotographic precipitation and filtration was 3.07 l/troy ounce of silver precipitated. Of the nine plants using this process at proposal, two produced no wastewater. Three plants supplied insufficient information to calculate discharge rates. Four plants were oncethrough dischargers with rates ranging from 0.42 to 78.6 l/troy ounce. The proposed BPT discharge rate was based on the average discharge rate of three of these plants. The plant with the 78.6 l/troy ounce rate was not considered in the average because this discharge rate was nearly ten times that of the next highest plant.

The BPT wastewater discharge allowance at promulgation is 3.07 l/troy ounce of silver precipitated. This is equivalent to the proposed allowance. The Agency received revised or new data from 18 plants. The water use and wastewater discharge rates are presented in Table V-12 (page 2735). After excessive water users are discarded, the data support the proposed allowance. The Agency believes there is no reason to revise the proposed allowance.

FLOOR AND EQUIPMENT WASHDOWN

No BPT wastewater discharge allowance will be provided for floor and equipment washdown. Many plants generate this wastewater while recovering silver contained in spills and leaks of process solutions. Plants usually precipitate the silver by cementation before discharging the wastewater. The Agency believes that this wastewater can be reused as washdown water after toxic metals and total suspended solids removal during treatment. The compliance costs for the treatment system reflect the additional capacity and equipment needed to achieve complete recycle of this waste stream.

REGULATED POLLUTANT PARAMETERS

The raw wastewater concentrations from individual operations and the subcategory as a whole were examined to select certain pollutant parameters for limitation. This examination and evaluation was presented in Section VI. Five pollutant parameters are selected for limitation under BPT and are listed below:

120. copper
128. zinc
ammonia (N)
total suspended solids (TSS)
pH

The concentrations achievable by application of the proposed BPT treatment are explained in Section VII of this supplement. The achievable treatment concentrations (both one-day maximum and monthly average values) are multiplied by the BPT normalized discharge flows summarized in Table IX-1 (page 2791) to calculate the mass of pollutants allowed to be discharged per mass of product. The results of these calculations in milligrams of pollutant per troy ounce of product represent the BPT effluent limitations and are presented in Table IX-2 (page 2793) for each individual waste stream.

Table IX-1

BPT WASTEWATER DISCHARGE RATES FOR THE SECONDARY SILVER SUBCATEGORY

	BPT Normalized Discharge Rate	Production Normalizing Parameter
Wastewater Stream	l/troy ounce	Parameter
Film stripping	50.35	troy ounces of silver produced from film stripping
Film stripping wet air pollution control and precipitation and filtration of film stripping solutions wet air pollution control	0.97	troy ounces of silver produced from precipitation and filtration of film stripping solutions
Precipitation and filtration of film stripping solutions	57.57	troy ounces of silver precipitated
Precipitation and filtration of photo- graphic solutions	26.6	troy ounces of H Silver precipitated H
Precipitation and filtration of photo- graphic solutions wet air pollution control	12.14	troy ounces of H silver precipitated H
Electrolytic refining	0.76	troy ounces of silver refined
Furnace wet air pollution control	0.67	troy ounces of silver smelted, roasted, or dried

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Table IX-1 (Continued)

BPT WASTEWATER DISCHARGE RATES FOR THE SECONDARY SILVER SUBCATEGORY

	BPT Normalized <u>Discharge Rate</u>	Production Normalizing
Wastewater Stream	<u>l/troy_ounce</u>	Parameter
Leaching	0.086	troy ounces of silver produced from leaching
Leaching wet air pollution control and precipitation of nonphoto- graphic solutions wet air pollu- tion control	4.43	troy ounces of silver produced from leaching or precipitation
Precipitation and filtration of non- photographic solutions	3.07	troy ounces of silver precipitated
Floor and equipment washdown wastewater	0	troy ounces of silver produced

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TABLE IX-2

BPT EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(a) Film Stripping BPT

Pollutant	A#	Nouimum	6	Marshaun G
h		Maximum	ror	Maximum for
Pollutant	Property	Any One	Day	Monthly Average
. 042 u cuito	- rober ol	imy one	Duy	Monthly A

	mg/troy	ounce of	f silver	from film	stripping	
Antimony				144.500	64.450	
Arsenic				105.200	46.830	
Cadmium				17.120	7.553	T
Chromium				22.150	9.063	1 ÷
*Copper			_	95.660	50.350	
Lead				21.150	10.070	
Nickel				96.670	63.950	
Selenium				61.930	27.690	9 .
Silver				20.640	8.560	
Thallium			£	103.200	45.209	-
*Zinc			÷.,	73.510	30.710	
*Ammonia (a	as N)			6712.000	2951.000	
*TSS				2064.000	981.800	
*pH		Within	the rar		to 10.0 at all	times

(b) Film Stripping Wet Air Pollution Control and Precipitation and Filtration of Film Stripping Solutions Wet Air Pollution Control BPT

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average

mg/troy ounce of silver from precipitation and filtration of film stripping solutions

Antimony Arsenic Cadmium Chromium	2.784 2.027 0.330 0.427	1.242 0.902 0.146
*Copper		0.175
	1.843	0.970
Lead	0.407	0.194
Nickel	1.862	1.232
Selenium	1.193	0.534
Silver	0.398	0.165
Thallium	1.988	0.883
*Zinc	1.416	0.592
*Ammonia (as N)	129.300	56.840
*TSS	39.770	18.920
*pH	Within the range of 7.5 to	10.0 at all times
		· · · · · · · · · · · · · · · · · · ·

*Regulated Pollutant

SECONDARY SILVER SUBCATEGORY SECT - IX

TABLE IX-2 (Continued)

BPT EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(c) Precipitation and Filtration of Film Stripping Solutions BPT

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average

...

	mg/troy ou	nce of	silver	precip	itated	
Antimony			165	.230	73.690	
Arsenic			120	.320	53.540	
Cadmium			19	.570	8.636	
Chromium			25	.330	10.360	
*Copper			109	.400	57.570	
Lead			. 24	.180	11.510	
Nickel			110	.500	73.110	
Selenium			70	.810	31.660	
Silver			23	.600	9.787	
Thallium			118	.000	52.390	
*Zinc			84	.050	35.120	
*Ammonia (as	N)		7674	.000	3374.000	
*TSS	•		2360	.000	1123.000	
*pH	Withi	n the r			o 10.0 at all	times

(d) Precipitation and Filtration of Photographic Solutions BPT

Pollutant or		Maximum fo	r Maximum	for
Pollutant P	roperty	Any One Day	y Monthly	Average
	mg/troy oun	ce of silver preci	pitated	
Antimony	ĩ	76.340	34.050	
Arsenic		55.590	24.740	
Cadmium		0.044	3.990	
Chromium		11.700	4.788	1
*Copper		50.540	26.600	
Lead		11.170	5.320	
Nickel		51,070	33.780	
Selenium		32.720	14.630	
Silver		10,910	4.522	
Thallium		54.530	24.210	
*Zinc		38.840	16.230	
*Ammonia (as	N)	3546.000	1559.000	
*TSS	•	1091.000	518.700	
*pH	Within	the range of 7.5		times

*Regulated Pollutant

BPT EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(e) <u>Precipitation and</u> <u>Filtration of</u> <u>Photographic</u> <u>Solutions</u> <u>Wet Air Pollution</u> <u>Control</u> BPT

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average

mg/troy ounce of silver from precipitation and filtration of photographic solutions

Antimony				34.	840	15.	540		
Arsenic				25.	370	11.	290		
Cadmium		14		4.	128	1.	821		
Chromium				5.	342		185		
*Copper			• `	23.	070	12.			
Lead					099	-	428		ć
Nickel		· •		23.	310	15.	-		
Selenium			,	14.			677	;	
Silver					977	,	064	· ·	
Thallium				24.	•••	11.			
*Zinc				17.	-		405		
*Ammonia (as	sN)			1618.0	-	711.	-		
*TSS				497.		236.		· · · · ·	
*pH		Within	the	range of				+ imaa	
· · · · ·	1		CITC	range or		IV.V al	arr	times	

(f) Electrolytic Refining BPT

						1. to
Pollutant	or			Maximum fo	or Maximum	for
Pollutant	Proj	perty		Any One Da	-	
mg/	'troy	ounce c	of silve	r from electro	lytic refining	
Antimony				2.181	0.973	
Arsenic			-	1.588	0.707	
Cadmium				0.258	0.114	
Chromium				0.334	0.137	
*Copper				1.444	0.760	
Lead			•	0.319	0.152	
Nickel			x	1.459	0.965	
Selenium				0.935	0.418	
Silver				0.312	0.129	
Thallium			-	1.558	0.692	
*Zinc				1.110	0.646	
*Ammonia (as N)			101.310	44.540	
*TSS		· .		31.160	14.820	• • •
*pH		With	in the		o 10.0 at all t	imes

*Regulated Pollutant

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BPT EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(g) Furnace Wet Air Pollution Control BPT

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of si	ilver roasted, smelted	or dried
Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc *Ammonia (as N) *TSS *pH Within t	$\begin{array}{c} 1.923\\ 1.400\\ 0.228\\ 0.295\\ 1.273\\ 0.281\\ 1.286\\ 0.824\\ 0.275\\ 1.374\\ 0.978\\ 89.310\\ 27.470\\ he \ range \ of \ 7.5 \ to \ 10.0 \end{array}$	<pre></pre>

(h) Leaching BPT

Pollutant or Pollutant Prope	erty	Maximum for Any One Day	Maximum for Monthly Average
mg/troj	y ounce of silver	produced from	leaching
Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc *Ammonia (as N) *TSS *pH	Within the range	0.247 1.180 0.029 0.038 0.163 0.163 0.165 0.106 0.035 0.176 0.126 11.460 3.526 2 of 7.5 to 10.	0.110 0.080 0.013 0.015 0.086 0.017 0.109 0.047 0.015 0.047 0.015 0.078 0.052 5.040 1.677 0 at all times

*Regulated Pollutant

BPT EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(i) <u>Leaching Wet Air Pollution Control and Precipitation of</u> <u>Nonphotographic Solutions Wet Air Pollution Control</u> BPT

Pollutant or Pollutant Prop	perty	Maximum for Any One Day	Maximum for Monthly Average
mg/troy oz. of	silver produced	from leaching o	r silver precipitated
Antimony		12.710	5.670
Arsenic		9.259	4.120
Cadmium		1.506	0.665
Chromium		1.949	0.797
*Copper		8.417	4.430
Lead		1.861	0.886
Nickel		8.506	5.626
Selenium		5.449	2.437
Silver		1.816	0.753
Thallium	-	9.082	4.031
*Zinc	•	6.468	2.702
*Ammonia (as N)	590.500	259.600
*TSS	•	181.600	86.390
*pH	Within the ra	ange of 7.5 to 1	

(j) <u>Precipitation and Filtration of Nonphotographic</u> Solutions BPT

Pollutant or	Noui mum for	
Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce o	f silver precipita	ted
Antimony	8.811	3.930
Arsenic	6.416	2.855
Cadmium	1.044	0.461
Chromium	1.351	0.553
*Copper	5.833	3.070
Lead	1.289	0.614
Nickel	5.894	3.899
Selenium	3.776	1.689
Silver	1.259	0.522
Thallium	6.293	2.794
*Zinc	4.482	1.873
*Ammonia (as N)	409.200	170.900
*TSS	125.900	59.870
*pH Within the	range of 7.5 to 10	.0 at all times

*Regulated Pollutant

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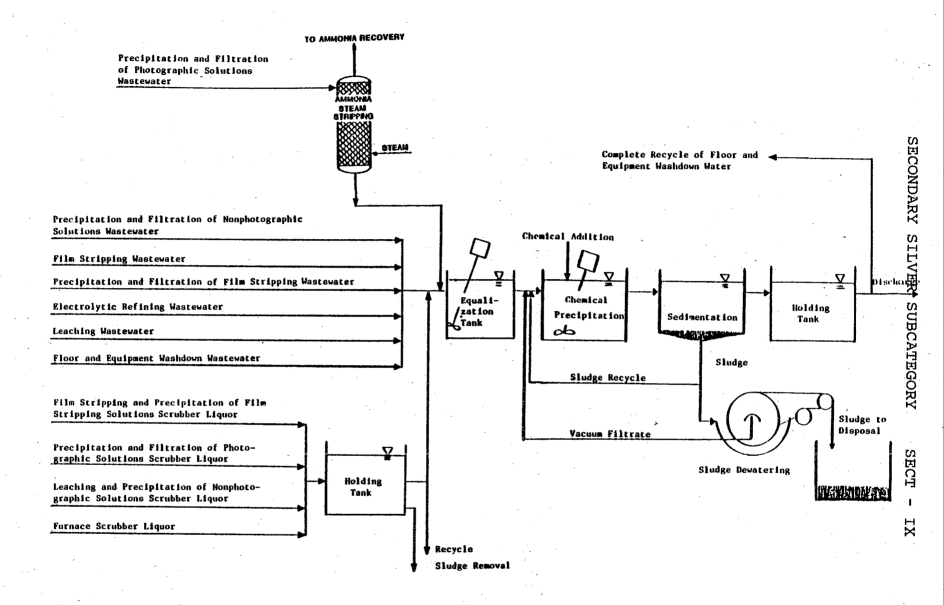
BPT EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(k) Floor and Equipment Washdown Water BPT

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
Politicant Property	Any One Day	Monthly Average

mg	/troy ounce	of silver produc	ction
Antimony		0.000	0.000
Arsenic		0.000	0.000
Cadmium		0.000	0.000
Chromium		0.000	0.000
*Copper		0.000	0.000
Leãd		0.000	0.000
Nickel		0.000	0.000
Selenium		0.000	0.000
Silver		0.000	0.000
Thallium		0.000	0.000
*Zinc		0.000	0.000
*Ammonia (as N)		0.000	0.000
*TSS		0.000	0.000
*pH	Within the	range of 7.5 to	10.0 at all times

*Regulated Pollutant



BPT TREATMENT SCHEME SECONDARY SILVER SUBCATEGORY

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SECONDARY SILVER SUBCATEGORY

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SECTION X

BEST AVAILABLE TECHNOLOGY ECONOMICALLY ACHIEVABLE

These effluent limitations are based on the best control and treatment technology used by a specific point source within the industrial category or subcategory, or by another category from which the technology is transferable. Emphasis is placed on additional treatment techniques applied at the end of the treatment systems currently used, as well as reduction of the amount of water used and discharged, process control, and treatment technology optimization.

The factors considered in assessing best available technology economically achievable (BAT) include the age of equipment and facilities involved, the process used, process changes, nonwater quality environmental impacts (including energy requirements), and the costs of application of such technology. BAT may include feasible process changes or internal controls, even when not in common practice.

The statutory assessment of BAT considers costs, but does not require a balancing car costs against effluent reduction benefits. However, in assessing BAT, the Agency has given substantial weight to the economic achievability of the technology.

TECHNICAL APPROACH TO BAT

The Agency reviewed a wide range of technology options and evaluated the available possibilities to ensure that the most effective and beneficial technologies were used as the basis of BAT. To accomplish this, the Agency elected to examine three technology options which could be applied to the secondary silver subcategory as alternatives for the basis of BAT effluent limitations.

For the development of BAT effluent limitations, mass loadings were calculated for each wastewater source or subdivision in the subcategory using the same technical approach as described in Section IX for BPT limitations development. The differences in the mass loadings for BPT and BAT are due to increased treatment effectiveness achievable with the more sophisticated BAT treatment technology and reductions in the effluent flows allocated to various waste streams. In summary, the treatment technologies considered for the secondary silver subcategory are:

Option A (Figure X-1, page 2818) is based on

- o Preliminary treatment of precipitation and filtration of photographic solutions wastewater with ammonia steam stripping
- o Chemical precipitation and sedimentation
- Complete recycle of floor and equipment washdown wastewater after treatment

Option B (Figure X-2, page 2819) is based on

- o In-process flow reduction of wet air pollution control water
- Preliminary treatment of precipitation and filtration of photographic solutions wastewater with ammonia steam stripping
- o Chemical precipitation and sedimentation
- Complete recycle of floor and equipment washdown wastewater after treatment

Option C (Figure X-3, page 2820) is based on

- o In-process flow reduction of wet air pollution control water
- Preliminary treatment of precipitation and filtration of photographic solutions wastewater with ammonia steam stripping
- o Chemical precipitation and sedimentation
- o Complete recycle of floor and equipment washdown wastewater after treatment
- o Multimedia filtration

The three options examined for BAT are discussed in greater detail below. The first option considered is the same as the BPT treatment technology which was presented in the previous section.

OPTION A

Option A for the secondary silver subcategory is equivalent to the control and treatment technologies which were analyzed for BPT in Section IX. The BPT end-of-pipe treatment scheme includes chemical precipitation, and sedimentation (lime and settle), with preliminary treatment of precipitation and filtration of photographic solutions wastewater with ammonia steam stripping (see Figure X-1). Complete recycle of treated floor and equipment washdown wastewater is also included. The discharge rates for Option A are equal to the discharge rates allocated to each stream as a BPT discharge flow.

OPTION B

Option B for the secondary silver subcategory achieves lower pollutant discharge by building upon the Option A (ammonia steam stripping, chemical precipitation, sedimentation, and recycle of treated floor and equipment washdown wastewater) treatment technology. Flow reduction measures are added to the Option A scheme (see Figure X-2). treatment These flow reduction measures, including in-process changes, result in the elimination of some wastewater streams and the concentration of pollutants in As explained in Section VII of the General effluents. other Development Document, treatment of a more concentrated effluent allows achievement of a greater net pollutant removal and introduces the possible economic benefits associated with treating a lower volume of wastewater.

Option B flow reduction measures are reflected in the BAT wastewater discharge rates. Flow reduction has been included in determining the BAT discharge rates for furnace wet air pollution control. Based on available data, the Agency did not feel that further flow reduction over BPT would be feasible for the remaining 10 waste streams in the secondary silver subcategory.

Flow reduction measures used in Option B to reduce process wastewater generation or discharge rates include the following:

There are four wastewater sources associated with wet air pollution control which are regulated under these effluent limitations:

- 1. Film stripping scrubber and precipitation of film stripping solutions scrubber,
- 2. Precipitation and filtration of photographic solutions scrubber,
- 3. Furnace scrubber, and
- 4. Leaching and precipitation of nonphotographic solutions scrubber.

Table X-1 (page 2818) presents the number of plants reporting wastewater with the wet air pollution control sources listed above, the number of plants practicing recycle, and the range of recycle values being listed. Complete recycle of furnace scrubber water will be required for BAT. The Agency is not requiring further flow reduction at BAT for the remaining wet air pollution control waste streams.

OPTION C

Option C for the secondary silver subcategory consists of all control and treatment requirements of Option B (in-process flow reduction, ammonia steam stripping, chemical precipitation, sedimentation, and recycle of treated floor and equipment washdown wastewater) plus multimedia filtration technology added at the end of the Option B treatment scheme (see Figure X-3). Multimedia filtration is used to remove suspended solids, including precipitates of toxic metals, beyond the concentration attainable by gravity sedimentation. The filter suggested is of the gravity, mixed media type, although other filters, such as rapid sand filters or pressure filters, would perform satisfactorily.

As one means of evaluating each technology option, EPA developed estimates of the pollutant removal estimates and the compliance costs associated with each option. The methodologies are described below.

POLLUTANT REMOVAL ESTIMATES

A complete description of the methodology used to calculate the estimated pollutant removals achieved by the application of the various treatment options is presented in Section X of Vol. I. The pollutant removal estimates have been revised from proposal based on comments and new data. However, the methodology for calculating pollutant removals was not changed. The data used for estimating removals are the same as those used to revise the compliance costs.

Sampling data collected during the field sampling program were used to characterize the major waste streams considered for regulation. At each sampled facility, the sampling data were production normalized for each unit operation (i.e., mass of pollutant generated per mass of product manufactured). This value, referred to as the raw waste, was used to estimate the mass of toxic pollutants generated within the secondary silver subcategory. By multiplying the total subcategory production for a unit operation by the corresponding raw waste value, the mass of pollutant generated for that unit operation was estimated.

The volume of wastewater discharged after the application of each treatment option was estimated for each operation at each plant by comparing the actual discharge to the regulatory flow. The smaller of the two values was selected and summed with the other plant flows. The mass of pollutant discharged was then estimated by multiplying the achievable concentration values attainable by the option (mg/l) by the estimated volume of process wastewater discharged by the subcategory. The mass of pollutant removed is simply the difference between the estimated mass of pollutant generated within the subcategory and the mass of pollutant disclarged after application of the treatment option. pollutant removal estimates for direct dischargers in The the secondary silver subcategory are presented in Table X-2 (page 2819).

COMPLIANCE COSTS

Compliance costs presented at proposal were estimated using cost curves, which related the total costs associated with installation and operation of wastewater treatment technologies to plant process wastewater discharge. EPA applied these curves on a per plant basis, a plant's costs -- both capital, and

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operating and maintenance -- being determined by what treatment has in place and by its individual process wastewater it discharge (from dcp). The final step was to annualize the capital costs, and to sum the annualized capital costs, and the operating and maintenance costs, yielding the cost of compliance for the subcategory. Since proposal, the cost estimation methodology has been changed as discussed in Section VIII of this document and in Section VIII of Vol. I. A design model and plant-specific information were used to size a wastewater treatment system for each discharging facility. After completion of the design, capital and annual costs were estimated for each unit of the wastewater treatment system. Capital costs were developed from vendor quotes and annual costs were developed from literature. The revised compliance costs are presented in Table VIII-1 (page 2776).

BAT OPTION SELECTION - PROPOSAL

At proposal, EPA selected both Option B and Option C as the basis for alternative BAT effluent limitations for the secondary silver subcategory due to adverse structural economic changes that the Agency suspected were not reflected in the Agency's proposed economic analysis. These alternative limitations were based on ammonia steam stripping preliminary treatment, lime precipitation and sedimentation, end-of-pipe technology, and in-process control technologies to reduce the volume of process wastewater discharged for Option B, and the addition of multimedia filtration to the end-of-pipe technology for Option C.

Option E was eliminated because the addition of activated carbon technology is not necessary since toxic organic pollutants are not selected for limitation in this subcategory.

BAT OPTION SELECTION - PROMULGATION

EPA is promulgating BAT effluent mass limitations based on lime precipitation and sedimentation and ammonia steam stripping with additional reduction in pollutant discharge with the use of filtration as an effluent polishing step. The end-of-pipe and The end-of-pipe and pretreatment technology basis for BAT limitations being promulgated is the same as that proposed for Alternative B. The expressed concerns at proposal about this option's Agency economic achievability, but after revising the compliance costs and the economic analysis, the Agency believes that filtration as an end-of-pipe treatment technology is economically achievable. The treatment performance concentrations upon which the mass limitations are based are equal to values used to calculate the proposed mass limitations.

Application of the promulgated BAT would remove 31,000 kg/yr of toxic metals and 664,154 kg/yr of ammonia over the estimated raw discharge. The BAT effluent mass limitations will remove 132 kg/yr of toxic pollutants above the estimated BPT discharge. The option is economically achievable. EPA believes that incremental removal justifies selection of filtration as part of BAT model technology. In addition, filtration is demonstrated at 11 secondary silver facilities. The estimated capital investment cost of the promulgated BAT is \$278,000 (March, 1982 dollars) and the annual cost is \$390,000 (March, 1982 dollars).

WASTEWATER DISCHARGE RATES

A BAT discharge rate was calculated for each subdivision based upon the flows of the existing plants, as determined from analysis of the data collection portfolios. The discharge rate is used with the achievable treatment concentration to determine BAT effluent limitations. Since the discharge rate may be different for each wastewater source, separate production normalized discharge rates for each of the 11 wastewater sources were determined and are summarized in Table X-3 (page 2810). The discharge rates are normalized on a production basis by relating amount of wastewater generated to the mass the of the intermediate product which is produced by the process associated with the waste stream in question. These production normalizing parameters (PNP) are also listed in Table X-3.

As discussed previously, the BAT wastewater discharge rate equals the BPT wastewater discharge rate for 10 of the 11 waste streams in the secondary silver subcategory. Based on the available data, the Agency did not feel that further flow reduction would be feasible for these wastewater sources. Wastewater streams for which BAT discharge rates differ from BPT are discussed below.

FURNACE WET AIR POLLUTION CONTROL

No BAT wastewater discharge allowance was provided for furnace wet air pollution control at proposal. This rate applies to all air pollution control of furnace operations and was based on complete recycle of wastewater. Since 15 of the 19 plants with furnace air pollution control did not discharge water, including plants with wet scrubber systems, the Agency believed that zero discharge is feasible for all secondary silver furnace air pollution control.

No BAT wastewater discharge allowance is provided for furnace wet air pollution control for promulgation. Only five of the 19 plants with furnace air pollution control currently discharge wastewater. The Agency believes, as it did at proposal, that zero discharge is feasible for all secondary silver furnace air pollution control. No comments were received challenging the Agency's conclusion.

REGULATED POLLUTANT PARAMETERS

In implementing the terms of the Consent Agreement in <u>NRDC</u> v. <u>Train</u>, Op. Cit., and 33 U.S.C. {1314(b)(2)(A and B) (1976), the Agency placed particular emphasis on the toxic pollutants. The raw wastewater concentrations from individual operations and the subcategory as a whole were examined to select certain pollutant parameters for consideration for limitation. This examination

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and evaluation, presented in Section VI, concluded that 26 pollutants and pollutant parameters are present in secondary silver wastewaters at concentrations than can be effectively reduced by identified treatment technologies. (Refer to Section VI.)

However, the cost associated with analysis for toxic metal pollutants has prompted EPA to develop an alternative method for regulating and monitoring toxic pollutant discharges from the nonferrous metals manufacturing category. Rather than developing specific effluent mass limitations and standards for each of the toxic metals found in treatable concentrations in the raw wastewaters from a given subcategory, the Agency is promulgating effluent mass limitations only for those pollutants generated in the greatest quantities as shown by the pollutant removal estimates analysis. The pollutants selected for specific limitation are listed below:

120. copper 128. zinc ammonia

By establishing limitations and standards for certain toxic metal pollutants, dischargers will attain the same degree of control over toxic metal pollutants as they would have been required to achieve had all the toxic metal pollutants been directly limited.

This approach is justified technically since the treatable concentrations used for lime precipitation and sedimentation technology are based on optimized treatment for concomitant multiple metals removal. Thus, even though metals have somewhat different theoretical solubilities, they will be removed at very nearly the same rate in a lime precipitation and sedimentation treatment system operated for multiple metals removal. Filtration as part of the technology basis is likewise justified because this technology removes metals non-preferentially.

The toxic metal pollutants selected for specific limitation in the secondary silver subcategory to control the discharges of toxic metal pollutants are copper and zinc. Ammonia is also selected for limitation since the methods used to control copper and zinc are not effective in the control of ammonia.

The following toxic pollutants are excluded from limitation on the basis that they are effectively controlled by the limitations developed for copper and zinc:

114. antimony
115. arsenic
118. cadmium
119. chromium
122. lead
124. nickel
125. selenium
126. silver

127. thallium

The Agency believes that the organic pollutants in this subcategory are present only in trace (deminimus quantities) and are neither causing nor likely to cause toxic effects. Therefore, the following organic pollutants are excluded from limitation:

- 4. benzene
- 6. carbon tetrachloride
- 10. 1,2-dichloroethane
- 11. 1,1,1-trichloroethane
- 29. 1,1-dichloroethylene
- 30. 1,2-trans-dichloroethylene
- 38. ethylbenzene
- 84. pyrene
- 85. tetrachloroethylene
- 86. toluene
- 87. trichloroethylene, and total phenols (by 4-AAP method)

Cyanide was present in the secondary silver subcategory in certain waste streams at concentrations that can be effectively reduced by identified treatment technologies. Treatable concentrations of cyanide were found in one photographic materials plant and one nonphotographic materials plant. different process waste streams were sampled; four contained cyanide at treatable concentrations, in six of nine samples. Five However, at proposal, when waste streams were combined for treatment, cyanide was found at a concentration below that achievable by identified treatment technology. This determination was made by comparing the raw (untreated) wasteload and treated discharge estimates presented in the pollutant removal estimates. Cyanide was thus excluded from limitation.

For promulgation, cyanide is not chosen as a regulated pollutant parameter on a subcategory-wide basis for the secondary silver mass limitations. However, secondary silver plants process plating solutions, which may contain cyanide, to recover silver contained in the solution. Cyanide is present due to its use as a process chemical in plating operations. The permitting authority should check for the presence of cyanide in this waste stream and develop discharge limitations if necessary. Α discharge allowance can be developed by locating the flow allowance for precipitation and filtration of nonphotographic solutions at BPT and BAT in Sections IX and X, respectively, of the silver supplemental development secondary document. Treatment performance for cyanide precipitation is presented in Section VII of the General Development Document. The discharge allowance (or mass limitation) is the product of the flow allowance and the treatment performance.

EFFLUENT LIMITATIONS

The treatable concentrations, achievable by application of the

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BAT technology (Option C) are summarized in Section VII of this supplement. These treatable concentrations (both one day maximum and monthly average) are multiplied by the BAT normalized discharge flows summarized in Table X-3 (page 2810) to calculate the mass of pollutants allowed to be discharged per mass of product. The results of these calculations in milligrams of pollutant per troy ounce of product represent the BAT effluent limitations for the secondary silver subcategory. The BAT effluent limitations are presented in Table X-4 (page 2811).

Table X-1

CURRENT RECYCLE PRACTICES WITHIN THE SECONDARY SILVER SUBCATEGORY

	Number of Plants with Wastewater	Number of Plants Practicing Recycle	Range of Recycle <u>Values (%)</u>
Film stripping and precipitation of film stripping solutions scrubber	1	. 1	99+
Photographic solution precipi- tation and filtration scrubber	3	3	99.93 - 100
Furnace scrubber	. 11	7	99.7 - 100
Leaching and precipitation of nonphotographic solutions scrubber	13	8	99 - 100

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Table X-2

POLLUTANT REMOVAL ESTIMATES FOR SECONDARY SILVER DIRECT DISCHARGERS

POLLUTANT	TOTAL RAW WASTE (kg/yr)	OPTION A Discharged (kg/yr)	OPTION A Removed (kg/yr)	OPTION B DISCHARGED (kg/yr)	OPTION B REMOVED (kg/yr)	OPTION C DISCHARGED (kg/yr)	OPTION C REMOVED (kg/yr)
Arsenic	6.2	6.2	0.0	6.2	0.0	6.2	0.0
Ant Lmony	7.0	7.0 7.6	0.0	7.0	0.0	7.0 7.6	5 0.0
Cadmium	7.6		15.4	71.7	15.4		27.3
Chromium	87.1 112.8	71.7 102.5	10.3	102.5	10.3	59.8 68.3	44.4
Lead	126.6	126.6	0.0	126.6	0.0	126.6	0.0
Nickel Selenium	9.0	9.0	0.0	9.0	0.0	9.0	0.0
Silver	6.9	6.9	0.0	6.9	0.0	6.9	0.0
Thallium	0.5	0.5	b 0.0	0.5	0.0	0.5	0.0
Copper	224.2	224.2	0.0	224.2 281.8	0.0	224.2 196.4	0.0 30,928.9
Zinc	31,125.3	281.8	30,843.5	281.8	30,843.5	190.4	30,920.9
TOTAL TOXIC METALS	31,713.3	844.1	30,869.2	844.1	30,869.2	712.6	31,000.7
Ammonia	691,477.2	27,323.2	664,154.0	27,323.2	664,154.0	27,323.2	664,154.0
TOTAL NONCONVENTIONALS	691,477.2	27,323.2	664,154.0	27,323.2	664,154.0	27,323.2	664,154.0
TSS	17,999.2	10,246.2	7,753.0	10,246.2	7,753.0	2,220.0	15,779.2
TOTAL CONVENTIONALS	17,999.2	10,246.2	7,753.0	10,246.2	7,753.0	2.220.0	15,779.2
TOTAL POLLUTANTS	741,189.7	38,413.5	702,776.1	38,413.5	702,776.1	30,255.8	710,933.8
FLOW (1/yr)		853,850,000		853,850,000		853,850,000	

NOTE: TOTAL TOXIC METALS = Arsenic + Antimony + Cadmium + Chromium + Lead + Nickel + Selenium + Silver + Thaiiium + Copper + Zinc

TOTAL CONVENTIONALS = TSS

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TOTAL POLLUTANTS = Total Toxic Metals + Total Nonconventionals + Total Conventionals

OPTION A = Ammonia Steam Stripping, Lime Precipitation, and Sedimentation OPTION B = Option A, plus In-Process Flow Reduction OPTION C = Option B, plus Multimedia Filtration

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Table X-3

BAT WASTEWATER DISCHARGE RATES FOR THE SECONDARY SILVER SUBCATEGORY

	BAT Normalized Discharge Rate	Production
Wastewater Stream	l/troy ounce	Normalizing <u>Parameter</u>
Film stripping	50.35	troy ounces of silver produced from film stripping
Film stripping wet air pollution control and precipitation and fitration of film stripping solutions wet air pollution control	0.97	troy ounces of silver produced from precipitation and filtration of film stripping solutions
Precipitation and filtration of film stripping solutions	57.57	troy ounces of silver precipitated
Precipitation and filtration of photo- graphic solutions	26.6	troy ounces of silver precipitated
Precipitation and filtration of photo- graphic solutions wet air pollution control	12.14	troy ounces of silver precipitated
Electrolytic refining	0.76	troy ounces of silver refined
Furnace wet air pollution control	0	troy ounces of silver smelted, roasted, or dried

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SECONDARY SILVER SUBCATEGORY SECT -

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Table X-3 (Continued)

BAT WASTEWATER DISCHARGE RATES FOR THE SECONDARY SILVER SUBCATEGORY

	BAT Normalized Discharge Rate	· *	Production
Wastewater Stream	l/troy ounce	• •	Normalizing Parameter
Leaching	0.086		troy ounces of silver produced from leaching
Leaching wet air pollution control and precipitation of nonphoto- raphic solutions wet air pollu- ion control	4.43	2	troy ounces of silver produced from leaching or precipitation
Precipitation and filtration of non- photographic solutions	3.07	• •	troy ounces of silver precipitated
Floor and equipment washdown wastewater	0		troy ounces of silver produced

SECONDARY SILVER SUBCATEGORY

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SECONDARY SILVER SUBCATEGORY SECT - X

TABLE X-4

BAT EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(a) Film Stripping BAT

Pollutant Pollutant	or Property	?	Maximum for Any One Day	Maximum Monthly	
	mg/troy	ounce of	silver from film	stripping	
Antimony			97.180	43.300	
Arsenic			69.990	31.220	
Cadmium			10.070	4.028	
Chromium			18.630	7.553	
*Copper			64.450	30.710	
Lead			14.100	6.546	
Nickel			27.690	18.630	
Selenium			41.290	18.630	
Silver			14.600	6.042	
Thallium			70.490	30.710	
*Zinc			51.360	21,150	
*Ammonia (as N)		6712.000	2951.000	

(b) <u>Film Stripping Wet Air Pollution</u> <u>Control and Precipitation</u> <u>and Filtration of Film Stripping</u> <u>Solutions Wet Air Pollution</u> <u>Control BAT</u>

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average

mg/troy ounce of silver from precipitation and filtration of film stripping solutions

	· · · · · · · · · · · · · · · · · · ·		
Antimony	1.872	0.834	
Arsenic	1.348	0.601	
Cadmium	0.194	0.078	
Chromium	0.359	0.146	
*Copper	1.242	0.592	
Lead	0.272	0.126	
Nickel	0.534	0.359	
Selenium	0.795	0.359	
Silver	0.281	0.116	
Thallium	1.358	0.592	
*Zinc	0.989	0.407	
*Ammonia (as N)	129.300	56.840	

BAT EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(c) <u>Precipitation</u> and <u>Filtration</u> of <u>Film</u> <u>Stripping</u> <u>Solutions</u> BAT

Pollutant or Pollutant Pr	operty		Maximum for Any One Day	Maximum for Monthly Aver	age
	mg/troy ound	e of	silver precipita	ated	
Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc *Ammonia (as	N)		111.100 80.020 11.510 21.300 73.690 16.120 31.660 47.210 16.700 80.600 58.720 7674.000	49.510 35.690 4.606 8.636 35.120 7.484 21.300 21.300 6.908 35.120 24.180 3374.000	

(d) <u>Precipitation</u> and <u>Filtration</u> of <u>Photographic</u> Solutions BAT

Pollutant or	coperty	Maximum for	Maximum for
Pollutant Pr		Any One Day	Monthly Average
	mg/troy ounce	of silver precipita	ated
Antimony	N)	51.340	22.880
Arsenic		36.970	16.490
Cadmium		5.320	2.128
Chromium		9.842	3.990
*Copper		34.050	16.230
Lead		7.448	3.458
Nickel		14.630	9.842
Selenium		21.810	9.842
Silver		7.714	3.192
Thallium		37.240	16.230
*Zinc		27.130	11.170
*Ammonia (as		3546.000	1559.000

*Regulated Pollutant

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TABLE X-4 (Continued)

BAT EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(e) <u>Precipitation and Filtration of Photographic Solutions</u> <u>Wet Air Pollution Control BAT</u>

Pollutant or Pollutant Prope	rty	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce	of silver f photog	rom precipitation and raphic solutions	filtration of
Antimony		23.430	10.440
Arsenic	· .	16.880	- 7.527
Cadmium		2.428	0.971
Chromium		4.492	1.821
*Copper		15.540	7.405
Lead		3.399	1.578
Nickel		6.677	4.492
Selenium		9.955	4.492
Silver		3.521	1.457
Thallium		17.000	7.405
*Zinc	ı	12.380	5.099
*Ammonia (as N)		1618.000	711.400

(f) Electrolytic Refining BAT

Pollutant or Pollutant Pr	operty	Maximum for Any One Day	Maximum for Monthly Average
mg/tro	y ounce of silv	er from electrolytic	refining
Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc *Ammonia (as 1	N)	1.467 1.056 0.152 0.281 0.973 0.213 0.418 0.623 0.220 1.064 0.775 101.300	0.654 0.471 0.061 0.114 0.464 0.099 0.281 0.281 0.281 0.091 0.464 0.319 44.549

*Regulated Pollutant

BAT EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(g) Furnace Wet Air Pollution Control BAT

ollutant or Collutant P		•	· • •		mum for One Day	Maximum Monthly	
mg/troy our	ce of si	lver f	rom	silver	roasted,	smelted, or	dried
Antimony	•		ю.		0.000	0.000	
Arsenic					0.000	0.000	÷.
Cadmium					0.000	0.000	
Chromium				•	0.000	0.000	
Copper					0.000	0.000	
Lead					0.000	0.000	
Nickel					0.000	0.000	
Selenium					0.000	0.000	
Silver					0.000	0.000	· .
Thallium					0.000	0.000	
*Zinc					0.000	0.000	· · ·
*Ammonia (as	- NI)				0.000	0.000	
		3		,			
(h) <u>Leachi</u>	ng BAT	,					
(h) <u>Leachin</u> Pollutant o Pollutant 1	r				imum for One Day	Maximum Monthly	
Pollutant of Pollutant	r Property	silve	r fro	Any	One Day		Average
Pollutant o Pollutant 1 mg/troy	r Property	silve	r fro	Any	One Day er produc	Monthly	Average hing
Pollutant o Pollutant i mg/troy Antimony	r Property	silve	r fro	Any	One Day	Monthly ced from leac	Average hing
Pollutant o Pollutant i mg/troy Antimony Arsenic	r Property	silve	r fro	Any	One Day er produc 0.166 0.120	Monthly ced from leac 0.074	Average hing
Pollutant o Pollutant i mg/troy Antimony Arsenic Cadmium	r Property	silve	r fro	Any	One Day er produc 0.166 0.120 0.017	Monthly ced from leac 0.074 0.053	Average hing
Pollutant o Pollutant i mg/troy Antimony Arsenic Cadmium Chromium	r Property	silve	r fro	Any	One Day er produc 0.166 0.120	Monthly ced from leac 0.074 0.053 0.007	Average hing
Pollutant o Pollutant i mg/troy Antimony Arsenic Cadmium Chromium *Copper	r Property	silve	r fro	Any	One Day er produc 0.166 0.120 0.017 0.032	Monthly ced from leac 0.074 0.053 0.007 0.013	Average hing
Pollutant o Pollutant i mg/troy Antimony Arsenic Cadmium Chromium *Copper Lead	r Property	silve	r fro	Any	One Day er produce 0.166 0.120 0.017 0.032 0.110 0.024	Monthly ced from leac 0.074 0.053 0.007 0.013 0.052	Average
Pollutant of Pollutant of mg/troy Antimony Arsenic Cadmium Chromium *Copper Lead Nickel	r Property	silve	r fro	Any	One Day er product 0.166 0.120 0.017 0.032 0.110 0.024 0.047	Monthly ced from leac 0.074 0.053 0.007 0.013 0.052 0.011	Average
Pollutant of Pollutant of mg/troy Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium	r Property	silve	r fro	Any	One Day er product 0.166 0.120 0.017 0.032 0.110 0.024 0.047 0.071	Monthly ced from leac 0.074 0.053 0.007 0.013 0.052 0.011 0.032	Average
Pollutant o Pollutant o Mg/troy Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver	r Property	silve	r fro	Any	One Day er product 0.166 0.120 0.017 0.032 0.110 0.024 0.047 0.071 0.025	Monthly ced from leac 0.074 0.053 0.007 0.013 0.052 0.011 0.032 0.032	Average
Pollutant of Pollutant of mg/troy Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium	r Property	silve	r fro	Any	One Day er product 0.166 0.120 0.017 0.032 0.110 0.024 0.047 0.071	Monthly ced from leac 0.074 0.053 0.007 0.013 0.052 0.011 0.032 0.032 0.010	Average

BAT EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(i) <u>Leaching Wet Air Pollution Control and Precipitation of</u> Nonphotographic Solutions Wet Air Pollution Control BAT

Pollutant of Pollutant	or Property		Maximum for Any One Day	Maximum Monthly	
mg/troy	ounce of	silver	from silver produced of silver	from leach	ing
Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc *Ammonia (a	ISN)	, ,, ,	8.550 6.158 0.886 1.639 5.670 1.240 2.437 3.633 1.285 6.202 4.519 590.500	3.810 2.747 0.354 0.665 2.702 0.576 1.639 1.639 0.532 2.702 1.861 259.600	•

(j) <u>Precipitation and Filtration of Nonphotographic Solutions</u> BAT

Pollutant or Pollutant I	r Property	Maximum for Any One Day	Maximum Monthly	for Average
	mg/troy ounce of	E silver precipitated	1	······································
Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc *Ammonia (as	; N)	5.925 4.267 0.614 1.136 3.930 0.860 1.689 2.517 0.890 4.298 3.131 409.200	2.640 1.903 0.246 0.461 1.873 0.399 1.136 1.136 0.399 1.873 1.289 179.900	

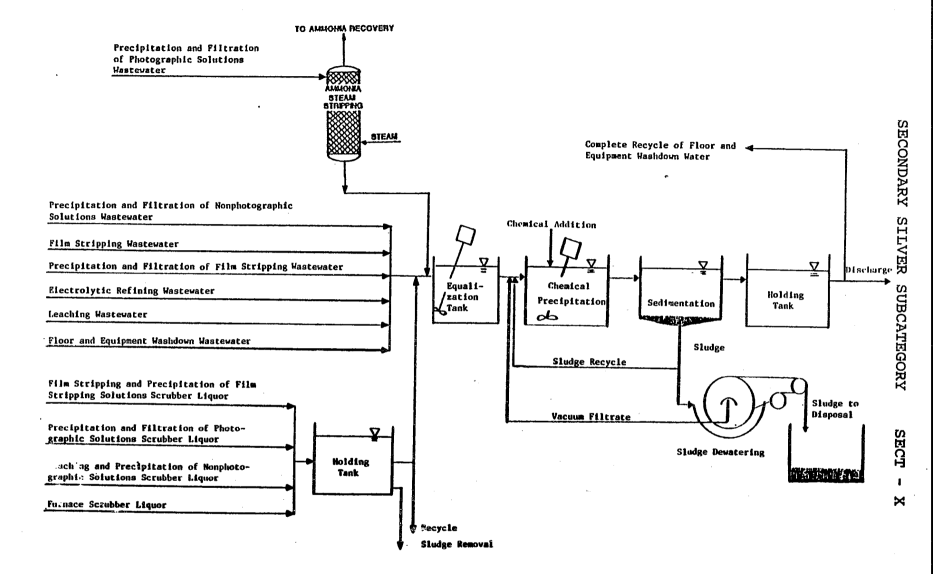
SECONDARY SILVER SUBCATEGORY SECT - X

TABLE X-4 (Continued)

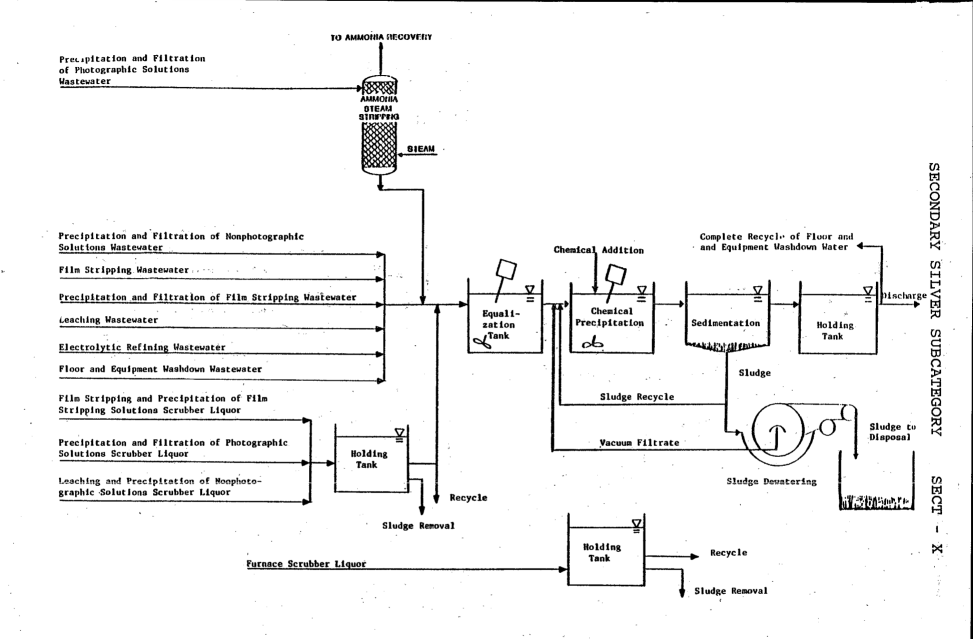
BAT EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(k) Floor and Equipment Washdown Water BAT

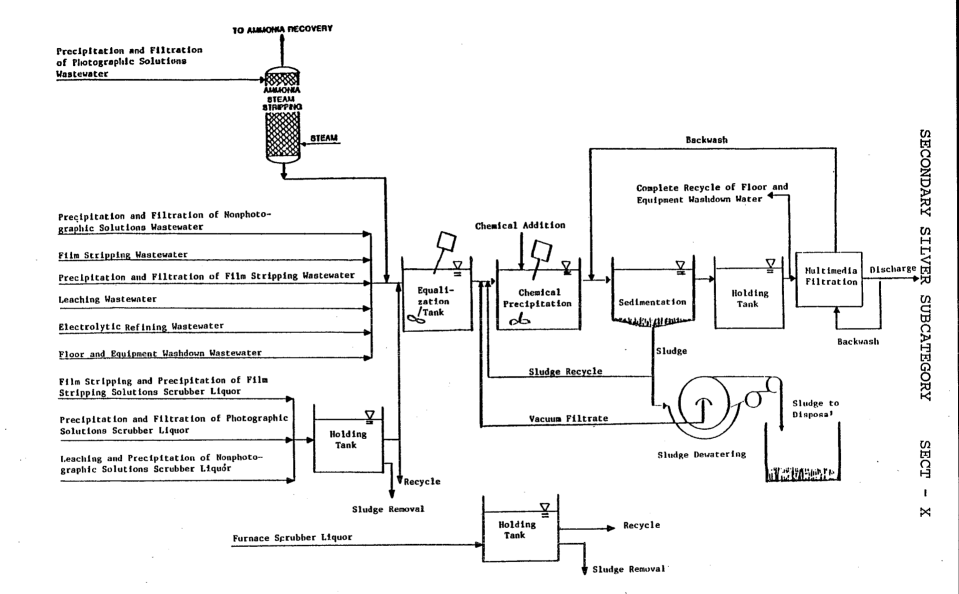
Pollutant or Pollutant Pr	coperty	Maximum for Any One Day	Maximum for Monthly Average
,	mg/troy ounce	of silver production	
Antimony		0.000	0.000
Arsenic		0.000	0.000
Cadmium		0.000	0.000
Chromium		0.000	0.000
*Copper		0.000	0.000
Lead		0.000	0.000
Nickel		0.000	0.000
Selenium		0.000	0.000
Silver		0.000	0.000
Thallium		0.000	0.000
*Zinc	· ,	0.000	0.000
*Ammonia (as	N)	0.000	0.000



BAT TREATMENT SCHEME OPTION A SECONDARY SILVER SUBCATEGORY



BAT TREATMENT SCHEME OPTION B SECONDARY SILVER SUBCATEGORY



BAT TREATMENT SCHEME OPTION C SECONDARY SILVER SUBCATEGORY

SECONDARY SILVER SUBCATEGORY SECT - XI

SECTION XI

NEW SOURCE PERFORMANCE STANDARDS

This section describes the control technology for treatment of wastewater from new sources and presents mass discharge limitations of regulatory pollutants for NSPS in the secondary silver subcategory based on the described control technology. New plants have the opportunity to design the best and most efficient production processes and wastewater treatment technologies, without facing the added costs and restrictions encountered in retrofitting an existing plant.

TECHNICAL APPROACH TO BDT

All of the treatment technology options applicable to a new source were previously considered for the BAT options. For this reason, three options were considered for BDT after proposal, all identical to the BAT options discussed in Section X.

Treatment and control technologies used for the BDT options are:

OPTION A

- Preliminary treatment for precipitation and filtration of photographic solutions wastewater with ammonia steam stripping concentrations
- o Chemical precipitation and sedimentation
- Complete recycle of floor and equipment washdown wastewater after treatment

OPTION B

- In-process flow reduction of wet air pollution control water
- Preliminary treatment for precipitation and filtration
 of photographic solutions wastewater with ammonia steam
 stripping
- o Chemical precipitation and sedimentation
- o Complete recycle of floor and equipment washdown wastewater after treatment

OPTION C

- In-process flow reduction of wet air pollution control water
- Preliminary treatment for precipitation and filtration of photographic solutions wastewater with ammonia steam stripping
- o Chemical precipitation and sedimentation
- o Complete recycle of floor and equipment washdown wastewater after treatment
- o Multimedia filtration

Partial or complete recycle and reuse of wastewater is an essential part of all three options. Recycle and reuse can precede or follow end-of-pipe treatment. A more detailed discussion of the treatment options is presented in Section X.

BDT OPTION SELECTION

EPA is promulgating a best available demonstrated technology for the secondary silver subcategory equal to Option C (in-process reduction, preliminary treatment with ammonia steam ng, lime precipitation, sedimentation, and multimedia flow steam stripping, filtration end-of-pipe treatment with complete recycle of treated This technology is floor and equipment washdown wastewater). equivalent to the proposed NSPS technology (although flow allowances for casting and casting wet air pollution have been eliminated, as explained earlier). The Agency recognizes that new sources have the opportunity to implement more advanced levels of treatment without incurring the costs of retrofitting and the costs of partial or complete shutdown necessary for installation of the new equipment that existing plants should of the subcategory indicates that no Review have. new demonstrated technologies that improve on BAT exist.

Activated carbon adsorption technology (Option E) was eliminated; it is not necessary since toxic organic pollutants are not selected for limitation in this subcategory. (Refer to the discussion of exclusion of toxic organic pollutants in Sections VI and X.)

Dry scrubbing is not demonstrated for controlling emissions from film stripping, precipitation and filtration of film stripping solutions, precipitation and filtration of photographic solutions, reduction furnaces, leaching and precipitation and filtration. The nature of these emissions (acidic fumes, hot particulate matter) technically precludes the use of dry scrubbers.

Therefore, EPA is including an allowance for these sources at NSPS equivalent to that promulgated for BAT Option C. The Agency also does not believe that new plants could achieve any additional flow reduction beyond that proposed for BAT.

REGULATED POLLUTANT PARAMETERS

The Agency has no reason to believe that the pollutants that will be found in treatable concentrations in processes within new sources will be any different than with existing sources. Accordingly, pollutants and pollutant parameters selected for limitation under NSPS, in accordance with the rationale of Suction VI and X, are identical to those selected for BAT. The conventional pollutant parameters TSS and pH are also selected for limitation.

NEW SOURCE PERFORMANCE STANDARDS

The NSPS discharge flows for each wastewater source are the same as the discharge rates for BAT and are listed in Table XI-1 (page 2824). The mass of pollutant allowed to be discharged per mass of product is calculated by multiplying the appropriate effluent concentration by the production normalized wastewater discharge flows (1/troy ounce). New source performance standards are presented in Table XI-2 (page 2826).

Table XI-1

NSPS WASTEWATER DISCHARGE RATES FOR THE SECONDARY SILVER SUBCATEGORY

	NSPS Normalized Discharge Rate		Production' Normalizing
Wastewater Stream	<pre>1/troy ounce</pre>		Parameter
Film stripping	50.35		troy ounces of silver produced from film stripping
Film stripping wet air pollution control and precipitation and fileration of film stripping solutions wet air pollution control	0.97	2 ⁴	troy ounces of silver produced from precipitation and filtration of film stripping solutions
Precipitation and filtration of film stripping solutions	57.57		troy ounces of silver precipitated
Precipitation and filtration of photo- graphic solutions	26.6		troy ounces of silver precipitated
Precipitation and filtration of photo- graphic solutions wet air pollution control	12.14		troy ounces of silver precipitated
Electrolytic refining	0.76		troy ounces of silver refined
Famace wet air pollution control	0		troy ounces of silver smelted, roasted, or dried

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2.826

XI

Table XI-1 (Continued)

NSPS WASTEWATER DISCHARGE RATES FOR THE SECONDARY SILVER SUBCATEGORY

NSPS Normalized Discharge Rate

1/troy ounce

0.086

4.43

3.07

A

Wastewater Stream

Leaching

Leaching wet air pollution control and precipitation of nonphotographic solutions wet air pollution control

Precipitation and filtration of nonphotographic solutions

Floor and equipment washdown wastewater

Production Normalizing Parameter

troy ounces of silver produced from leaching

troy ounces of silver produced from leaching or precipitation

troy ounces of silver precipitated

troy ounces of silver produced

SECT

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TABLE XI-2

NSPS EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(a) Film Stripping NSPS

2

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
mg/troy ounce of si	lver from film st	ripping
Antimony	97.180	43.300
Arsenic	69.990	31.220
Cadmium	10.070	4.028
Chromium	18.630	7.553
*Copper	64.450	30.710
Lead	14.100	6.546
Nickel	27.690	18.630
Selenium	41.290	18.630
Silver	14.600	6.042
Thallium	70.490	30.710
*Zinc	51.360	21.150
*Ammonia (as N)	6712.000	2951.000
*TSS	755.300	604.200
*pH Within the ra	ange of 7.5 to 10	.0 at all times
(b) <u>Film Stripping Wet Air Po</u> and <u>Filtration</u> <u>of Film St</u> <u>Control</u> NSPS	llution Control a ripping Solutions	nd <u>Precipitation</u> Wet <u>Air</u> Pollution
Pollutant or Pollutant Property	Maximum for	Maximum for
	Any One Day	Monthly Average

mg/troy ounce of silver from precipitation and filtration of film stripping solutions

Antimony Arsenic		1.872 1.348	0.834 0.601
Cadmium		0.194	0.078
Chromium		0.359	0.146
*Copper		1.242	0.592
Lead		0.272	0.126
Nickel		0.534	0.359
Selenium		0.795	0.359
Silver		0.281	0.116
Thallium		1.358	0.592
*Zinc		0.989	0.407
*Ammonia (as N)		129.300	56.840
*TSS		14.550	11.640
*pH	Within the range	of 7.5 to 1	10.0 at all times

NSPS EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(c) <u>Precipitation</u> and <u>Filtration</u> of <u>Film</u> <u>Stripping</u> <u>Solutions</u>

· · · · · · · · · · · · · · · · · · ·	•	1		
Pollutant or Pollutant Property		Maximum for Any One Day	Maximum Monthly	
mg/troy	ounce of	silver precipit	tated	
Antimony		111.100	49.510	
Arsenic		80.020	35.690	· · ·
Cadmium		11.510	4.606	
Chromium	й. 	21.300	8.636	
*Copper		73.690	35.120	
Lead	·	16.120	7.484	and in the second second
Nickel		31.660	21.300	· · ·
Selenium	· · ·	47.210	21.300	an an Arana an Arana. An an Arana an Arana
Silver		16.700	6.908	
Thallium		80.600	35.120	and the second sec
*Zinc	2	58.720	24.180	
*Ammonia (as N)		7674.000	3374.000	
*TSS *pH Wit		863.600	690.800	er i le composition de la comp
(d) Precipitation and		ange of 7.5 to 1 on <u>of</u> Photograph		•
(d) <u>Precipitation</u> and Pollutant or		<u>on of Photograph</u> Maximum for	<u>lic</u> Solutions Maximum	NSP S for
(d) <u>Precipitation</u> and Pollutant or Pollutant Property	<u>Filtrati</u>	on of Photograph	<u>lic</u> Solutions Maximum Monthly	NSP S for
(d) <u>Precipitation</u> and Pollutant or Pollutant Property mg/troy Antimony	<u>Filtrati</u>	on of Photograph Maximum for Any One Day silver precipit 51.340	<u>lic</u> Solutions Maximum Monthly	NSP S for
(d) <u>Precipitation</u> and Pollutant or Pollutant Property mg/troy Antimony Arsenic	<u>Filtrati</u>	on of Photograph Maximum for Any One Day silver precipit 51.340 36.970	Maximum Monthly ated 22.880 16.490	NSP S for
(d) <u>Precipitation and</u> Pollutant or Pollutant Property mg/troy Antimony Arsenic Cadmium	<u>Filtrati</u>	on of Photograph Maximum for Any One Day silver precipit 51.340 36.970 5.320	Maximum Monthly ated 22.880 16.490 2.128	NSP S for
(d) <u>Precipitation and</u> Pollutant or Pollutant Property mg/troy Antimony Arsenic Cadmium Chromium	<u>Filtrati</u>	on of Photograph Maximum for Any One Day silver precipit 51.340 36.970 5.320 9.842	nic Solutions Maximum Monthly ated 22.880 16.490 2.128 3.990	NSP S for
(d) <u>Precipitation and</u> Pollutant or Pollutant Property mg/troy Antimony Arsenic Cadmium Chromium Copper	<u>Filtrati</u>	on of Photograph Maximum for Any One Day silver precipit 51.340 36.970 5.320 9.842 34.050	<u>nic</u> <u>Solutions</u> Maximum Monthly ated 22.880 16.490 2.128 3.990 16.230	NSP S for
(d) <u>Precipitation and</u> Pollutant or Pollutant Property mg/troy Antimony Arsenic Cadmium Chromium *Copper Lead	<u>Filtrati</u>	on of Photograph Maximum for Any One Day silver precipit 51.340 36.970 5.320 9.842 34.050 7.448	<u>nic</u> <u>Solutions</u> Maximum Monthly ated 22.880 16.490 2.128 3.990 16.230 3.458	NSP S for
(d) <u>Precipitation and</u> Pollutant or Pollutant Property mg/troy Antimony Arsenic Cadmium Chromium Copper Lead Nickel	<u>Filtrati</u>	on of Photograph Maximum for Any One Day silver precipit 51.340 36.970 5.320 9.842 34.050 7.448 14.630	<u>Maximum</u> Monthly ated 22.880 16.490 2.128 3.990 16.230 3.458 9.842	NSP S for
(d) <u>Precipitation and</u> Pollutant or Pollutant Property mg/troy Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium	<u>Filtrati</u>	<u>on of Photograph</u> <u>Maximum for</u> Any One Day silver precipit 51.340 36.970 5.320 9.842 34.050 7.448 14.630 21.810	<u>Maximum</u> Monthly ated 22.880 16.490 2.128 3.990 16.230 3.458 9.842 9.842	NSP S for
<pre>(d) Precipitation and Pollutant or Pollutant Property mg/troy Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver</pre>	<u>Filtrati</u>	<u>on of Photograph</u> <u>Maximum for</u> Any One Day silver precipit 51.340 36.970 5.320 9.842 34.050 7.448 14.630 21.810 7.714	<u>Maximum</u> Monthly ated 22.880 16.490 2.128 3.990 16.230 3.458 9.842 9.842 3.192	NSP S for
<pre>(d) Precipitation and Pollutant or Pollutant Property mg/troy Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium</pre>	<u>Filtrati</u>	<u>on of Photograph</u> <u>Maximum for</u> Any One Day silver precipit 51.340 36.970 5.320 9.842 34.050 7.448 14.630 21.810 7.714 37.240	<u>nic</u> Solutions Maximum Monthly ated 22.880 16.490 2.128 3.990 16.230 3.458 9.842 9.842 3.192 16.230	NSP S for
<pre>(d) Precipitation and Pollutant or Pollutant Property mg/troy Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc</pre>	<u>Filtrati</u>	<u>on of Photograph</u> <u>Maximum for</u> Any One Day silver precipit 51.340 36.970 5.320 9.842 34.050 7.448 14.630 21.810 7.714 37.240 27.130	<u>Maximum</u> Monthly ated 22.880 16.490 2.128 3.990 16.230 3.458 9.842 9.842 3.192 16.230 11.170	NSP S for
<pre>(d) Precipitation and Pollutant or Pollutant Property mg/troy Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc *Ammonia (as N)</pre>	<u>Filtrati</u>	<u>on of Photograph</u> <u>Maximum for</u> Any One Day silver precipit 51.340 36.970 5.320 9.842 34.050 7.448 14.630 21.810 7.714 37.240 27.130 3546.000	<u>Maximum</u> Monthly ated 22.880 16.490 2.128 3.990 16.230 3.458 9.842 9.842 3.192 16.230 11.170 1559.000	NSP S for
<pre>(d) Precipitation and Pollutant or Pollutant Property mg/troy Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc *Ammonia (as N) *TSS</pre>	Filtratio	<u>on of Photograph</u> <u>Maximum for</u> Any One Day silver precipit 51.340 36.970 5.320 9.842 34.050 7.448 14.630 21.810 7.714 37.240 27.130	<u>Maximum</u> Monthly ated 22.880 16.490 2.128 3.990 16.230 3.458 9.842 9.842 3.192 16.230 11.170 1559.000 319.200	NSPS for Average

*Regulated Pollutant

NSPS EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(e) <u>Precipitation and Filtration of Photographic Solutions</u> <u>Wet Air Pollution Control</u> NSPS

Pollutant Pollutant	or Proper	ty		Maximum Any One		Maximum Monthly	
mg/troy	ounce o	f silver phote	from p ographi	recipita c solutio	tion and ons	filtratio	on of
Antimony				23.4	30	10.440	
Arsenic				16.8		7.527	
Cadmium				2.4	28	0.971	
Chromium				4.4	92	1.821	
*Copper	,			15.5	40	7.405	
Lead				3.3	99	1.578	
Nickel				6.6	77	4.492	
Selenium				9.9	55	4.492	
Silver				3.5	21	1.457	
Thallium				17.0	00	7.405	
*Zinc				12.3	80	5.099	
*Ammonia	(as N)	¢		1618.00	00	711.400	
*TSS	· ·			182.1		145.700	
*pH		Within	the ran	ge of 7.	5 to 10.0	at all H	times

(f) Electrolytic Refining NSPS

Pollutant or Pollutant Prope	erty	Maximum for Any One Day	Maximum Monthly	-
mg/troy o	ounce of silver	from electrolytic	refining	
Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc *Ammonia (as N) *TSS *pH	Within the ra	1.467 1.056 0.152 0.281 0.973 0.213 0.418 0.623 0.220 1.064 0.775 101.300 11.400 nge of 7.5 to 10.0	0.654 0.471 0.061 0.114 0.464 0.099 0.281 0.281 0.281 0.091 0.464 0.319 44.540 9.120 at all ti	Lmes

SECONDARY SILVER SUBCATEGORY SECT - XI

TABLE XI-2 (Continued)

NSPS EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(g) Furnace Wet Air Pollution Control NSPS

Pollutant or Pollutant Prope	rtv	• •	Maximu Any On			ximum nthlv	for Average
· · · · · ·	-		_				-
mg/troy ounce o	f silver	from s:	ilver ro	asted	, smelte	d, or	dried
Antimony			0.	000	(0.000	÷
Arsenic			0.	000		0.000	
Cadmium	6		0.	000	I	0.000	ъ
Chromium	*		0.	000	1	0.000	*
*Copper	÷.,		Ο.	000		0.000	•
Lead			0.	000	· ·	0.000	
Nickel			0.	000		0.000	
Selenium			0.	000		0.000	1
Silver	.		0.	000		0.000	•
Thallium			Ο.	000		0.000	
*Zinc			0.	000		0.000	
*Ammonia (as N)			0.	000	I	0.000	• 1 1 1
*TSS			0.	000		0.000	
*pH	Within	the ran	nge of 7	.5 to	10.0 at	all	times
-			-				۰.

Leaching NSPS (h)

Pollutant Pollutant	or Property	· .	-	Maximum for Any One Day	Maximum Monthly	
mg/troy	ounce of	silver	from	silver produced	from leach	ning
Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc *Ammonia (*TSS *pH	as N)	Wit	hin t	0.166 0.120 0.017 0.032 0.110 0.024 0.047 0.071 0.025 0.120 0.088 11.460 1.290 the range of 7.5 at all times	0.074 0.053 0.007 0.013 0.052 0.011 0.032 0.032 0.032 0.036 5.040 1.032 to 10.0	

NSPS EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(i) Leaching Wet Air Pollution Control and Precipitation of Nonphotographic Solutions Wet Air Pollution Control NSPS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of silver from of s	ilver produced	from leaching
-	8.550 6.158 0.886 1.639 5.670 1.240 2.437 3.633 1.285 6.202 4.519 590.500 66.450 ge of 7.5 to 10.0	
(j) <u>Precipitation</u> and <u>Filtratic</u> NSPS	on of Nonphotogra	phic Solutions
Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of s	silver precipitat	ed
Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc *Ammonia (as N) *TSS *pH Within the rand	5.925 4.267 0.614 1.136 3.930 0.860 1.689 2.517 0.890 4.298 3.131 409.200 46.050 ge of 7.5 to 10.0	2.640 1.903 0.246 0.461 1.873 0.399 1.136 1.136 0.399 1.873 1.289 179.900 36.840) at all times

SECONDARY SILVER SUBCATEGORY SECT - XI

TABLE XI-2 (Continued)

NSPS EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(k) Floor and Equipment Washdown Water NSPS

Pollutant or Pollutant Pr	operty		num for Dne Day	Maximum Monthly	
	mg/troy ounc	e of silve	product	ion	
Antimony			0.000	0.000	
Arsenic	χ.		0.000	0.000	
Cadmium		· · · ·	0.000	0.000	
Chromium			0.000	0.000	
*Copper			0.000	0.000	
Lead			0.000	0.000	
Nickel	· · · · · · · · · · · · · · · · · · ·		0.000	0.000	
Selenium			000.000	0.000	
Silver			0.000	0.000	
Thallium		· · · ·	000	0.000	1
*Zinc		and the second sec	000	0.000	
*Ammonia (às	N)		0.000	0.000	
*TSS	-	(0.000	0.000	
*pH	Within	the range of	of 7.5 to	10.0 at all	times

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SECTION XII

PRETREATMENT STANDARDS

This section describes the control and treatment technologies for pretreatment of process wastewaters from existing sources and new sources in the secondary silver subcategory. Pretreatment standards for regulated pollutants are presented based on the selected control and treatment technologies.

PSES are designed to prevent the discharge of pollutants which pass through, interfere with, or are otherwise incompatible with the operation of publicly owned treatment works (POTW). The Clean Water Act of 1977 requires pretreatment for pollutants, as toxic metals, that limit POTW sludge such management alternatives. New indirect discharge facilities, like new direct discharge facilities, have the opportunity to incorporate the available demonstrated technologies, including best process and end-of-pipe in-plant controls, treatment changes, technologies, and to use plant site selection to ensure adequate treatment system installation. Pretreatment standards are to be technology based, analogous to the best available technology for removal of toxic pollutants.

TECHNICAL APPROACH TO PRETREATMENT

Before proposing pretreatment standards, the Agency examines whether the pollutants discharged by the industry pass through the POTW or interfere with the POTW operation or its chosen sludge disposal practices. In determining whether pollutants pass through a well-operated POTW, achieving secondary treatment, the Agency compares the percentage of a pollutant removed by POTW with the percentage removed by direct dischargers applying the best available technology economically achievable. A pollutant is deemed to pass through the POTW when the average percentage removed nationwide by well-operated POTW meeting secondary treatment requirements, is less than the percentage removed by direct dischargers complying with BAT effluent limitations guidelines for that pollutant. (See generally, 46 FR at 9415-16 (January 28, 1981).)

definition of pass through satisfies two competing This objectives set by Congress: (1) that standards for indirect dischargers be equivalent to standards for direct dischargers, while at the same time, (2) that the treatment capability and performance of the POTW be recognized and taken into account in regulating the discharge of pollutants from indirect dischargers. The Agency compares percentage removal rather than the mass or concentration of pollutants discharged because the latter would take into account the mass of pollutants discharged to the not POTW from non-industrial sources nor the dilution of the pollutants in the POTW effluent to lower concentrations due to the addition of large amounts of non-industrial wastewater.

PRETREATMENT STANDARDS FOR EXISTING AND NEW SOURCES

Options for pretreatment of wastewaters are based on increasing the effectiveness of end-of-pipe treatment technologies. All inplant changes and applicable end-of-pipe treatment processes have been discussed previously in Sections X and XI. The options for PSES and PSNS, therefore, are the same as the BAT options discussed in Section X.

A description of each option is presented in Section X, while a more detailed discussion, including pollutants controlled by each treatment process and achievable treatment concentration for each option, is presented in Section VII of Vol. I.

Treatment technology used for the PSES and PSNS options are:

OPTION A

- Preliminary treatment for precipitation and filtration of photographic solutions wastewater with ammonia steam stripping
- o Chemical precipitation and sedimentation
- o Complete recycle of floor and equipment washdown wastewater after treatment

OPTION B

- o In-process flow reduction of wet air pollution control water
- Preliminary treatment for precipitation and filtration of photographic solutions wastewater with ammonia steam stripping
- o Chemical precipitation and sedimentation
- o Complete recycle of floor and equipment washdown wastewater after treatment

OPTION C

- o In-process flow reduction of wet air pollution control water
- Preliminary treatment for precipitation and filtration of photographic solutions wastewater with ammonia steam stripping
- o Chemical precipitation and sedimentation
- o Complete recycle of floor and equipment washdown wastewater after treatment
- o Multimedia riltration

INDUSTRY COST AND POLLUTANT REMOVAL ESTIMATES

The industry cost and pollutant removal estimates of each treatment option were used to determine the most cost-effective option. The methodology applied in calculating pollutant removal estimates and plant compliance costs is discussed in Section X.

Table XII-1 (page 2837) shows the estimated pollutant removals for indirect dischargers, while compliance costs for indirect discharges are presented in Table VIII-2 (page 2838).

PSES OPTION SELECTION - PROPOSAL

At proposal, EPA selected in-process flow reduction, ammonia steam stripping preliminary treatment, lime precipitation, and sedimentation (Option B) and in-process flow reduction, ammonia steam stripping preliminary treatment, chemical precipitation, sedimentation, and multimedia filtration (Option C) as alternative pretreatment standards for existing sources for this subcategory. This selection follows from the rationale used in selecting alternative options as the basis for BAT. (Refer to Section X.)

Activated carbon adsorption technology (Option E) was eliminated because it is not necessary since toxic organic pollutants are not selected for limitation in this subcategory. (Refer to the discussion of selection of pollutants for limitation in Section X.)

PSES OPTION SELECTION - PROMULGATION

EPA is promulgating PSES equal to promulgated BAT for this subcategory. The technology basis for PSES thus is lime precipitation and sedimentation, ammonia steam stripping, wastewater flow reduction, and filtration. Flow reduction for the selected technology represents a 23 percent reduction in flow over current discharge rates.

Based on revised calculations, EPA estimates that implementation of the promulgated PSES limitations would remove annually an estimated 4,259 kg of toxic pollutants and 42,900 kg of ammonia The final PSES effluent mass over estimated raw discharge. limitations will remove 13 kg/yr of toxic metals over the intermediate PSES option considered, which lacks filtration. Both options are economically achievable. The Agency believes the incremental removal justifies selection of filtration as part of PSES model technology. Filtration is currently demonstrated by eight indirect discharging secondary silver plants. Capital cost for achieving proposed PSES is \$634,000 (March, 1982 dollars), and annual cost of \$422,000 (March, 1982 dollars).

PSNS OPTION SELECTION

EPA has selected in-process flow reduction, preliminary treatment with ammonia steam stripping, lime precipitation, sedimentation, recycle of treated floor and equipment complete washdown wastewater, and multimedia filtration (Option C) as the technology basis for PSNS. The Agency recognizes that new sources have the opportunity to implement more advanced levels of treatment without incurring the costs of retrofitting and the costs of partial or complete shutdown necessary for installation of the new equipment that existing plants should have.

EPA has not identified any demonstrated technology that provides more efficient pollutant removal than PSNS technology. No additional flow reduction for new sources is feasible because dry scrubbing is not demonstrated for controlling emissions from film stripping, precipitation and filtration of photographic solutions, reduction furnaces, leaching and precipitation and filtration. The nature of these emissions (acidic fumes, hot particulate matter) technically precludes the use of dry scrubbers. Since PSNS does not include any additional costs compared to NSPS, the Agency does not believe PSNS will be a barrier to entry for new facilities.

REGULATED POLLUTANT PARAMETERS

Pollutants and pollutant parameters selected for limitation for PSES and PSNS, in accordance with the rationale of Section VI and X, are identical to those selected for limitation for BAT. EPA is proposing PSNS for copper, zinc, and ammonia to prevent passthrough. The conventional pollutants, TSS and pH, are not limited under PSES and PSNS because they are effectively controlled by POTW.

PRETREATMENT STANDARDS

The PSES and PSNS discharge flows are identical to the BAT discharge flows for all processes. These discharge flows are listed in Table XII-2 (page 2838). The mass of pollutant allowed to be discharged per mass of product is calculated by multiplying the achievable treatment concentration (mg/l) by the normalized wastewater discharge flow (l/troy ounce). Pretreatment standards for existing and new sources, as determined from the above procedure, are shown in Tables XII-3 and XII-4 (pages 2840 and 2846) for each waste stream.

Mass-based standards are promulgated for the secondary silver subcategory to ensure that the standards are achieved by means of pollutant removal rather than by dilution. They are particularly important since the standards are based upon flow reduction. Pollutant limitations associated with flow reduction cannot be measured any way but as a reduction of mass discharged. Massbased PSES without alternative concentration-based standards are promulgated in this subcategory, although the flow reduction for the entire subcategory is not great. However, several plants grossly exceed the flow basis of PSES. Mass-based standards are needed to ensure that these plants reduce their water usage. Mass-based PSNS are promulgated in this subcategory because PSNS for secondary silver is based on 99 percent flow reduction of raw wastewater by recycle, and new plants would lack incentive to achieve these reductions without a mass-based standard.

Table XII-1

POLLUTANT REMOVAL ESTIMATES FOR SECONDARY SILVER INDIRECT DISCHARGERS

POLLUTANT	TOTAL Raw Waste (kg/yr)	OPTION A Discharged (kg/yr)	OPTION A Removed (kg/yr)	OPTION B DISCHARGED (kg/yr)	OPTION B Removed (kg/yr)	OPTION C DISCHARGED (kg/yr)	OPTION C REMOVED (kg/yr)
Arsenic Antimony Cadmium Chromium Lead Nickel Selenium Siiver Thallium	2.1 7.8 11.6 34.0 59.4 129.1 441.0 7.3 1.2	2.1 7.8 1.5 1.6 2.3 14.3 5.8 1.9 1.9	0.0 0.0 10.1 32.4 57.1 114.8 435.2 5.3 0.0	2.1 7.8 1.5 1.6 2.3 14.3 5.8 1.9 1.2	0.0 0.0 10.1 32.4 57.1 114.8 435.2 5.3 0.0	2.1 7.8 0.9 1.4 1.5 4.2 3.9 1.4 1.2	$\begin{array}{c} 0.0\\ 0.0\\ 10.7\\ 32.7\\ 57.9\\ 124.8\\ 437.2\\ 5.9\\ 0.0\\ \end{array}$
Copper Zinc	184.7 3,417.2	11.2	173.5 3,410.8	11.2 6.4	173.5 3,410.8	7.5 4.4	177.2 3,412.8
TOTAL TOXIC METALS	4,295.5 43,519.2	618.0	4,239.3 42,901.2	56.2 618.0	4,239.3 42,901.2	36.4 618.0	4,259.1 42,901.2
TOTAL NONCONVENTIONALS	43,519.2	618.0	42,901.2	618.0	42,901.2	618.0	42,901.2
TSS	13,602.8	231.8	13,371.0 13,371.0	231.8	13,371.0 13,371.0	50.2	13,552.6 13,552.6
TOTAL CONVENTIONALS TOTAL POLLUTANTS	13,602.8 61,417.5	906.0	60,511.6	906.0	60,511.6	704.6	60,712.9
FLOW (1/yr)		19,313,320		19,313,320	· · · · ·	19,313,320	

TOTAL TOXIC METALS = Arsenic + Antimony + Cadmium + Chromium + Lead + Nickel + Selenium + Silver + Thallium NOTE: + Copper + Zinc

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TOTAL NONCONVENTIONALS = Ammonia
TOTAL CONVENTIONALS = TSS
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TOTAL POLLUTANTS - Total Toxic Metals + Total Nonconventionals + Total Conventionals

OPTION A = Ammonia Steam Stripping, Lime Precipitation, and Sedimentation OPTION B = Option A, plus in-Process Flow Reduction OPTION C - Option B, plus Multimedia Filtration

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Table XII-2

PSES AND PSNS WASTEWATER DISCHARGE RATES FOR THE SECONDARY SILVER SUBCATEGORY

	PSES and PSNS Normalized Discharge Rate	Production
Wastewater Stream	<u>l/troy ounce</u>	Normalizing Parameter
Fi.m stripping	50.35	troy ounces of silver produced from film stripping
Film stripping wet air pollution control and precipitation and filtration of film stripping solutions wet air pollution control	0.97	troy ounces of silver produced from precipitation and filtration of film stripping solutions
Precipitation and filtration of film stripping solutions	57.57	troy ounces of silver precipitated
Precipitation and filtration of photo- graphic solutions	26.6	troy ounces of silver precipitated
Precipitation and filtration of photo- graphic solutions wet air pollution control	12.14	troy ounces of silver precipitated
Electrolytic refining	0.76	troy ounces of silver refined
Furnace wet air pollution control	0	troy ounces of silver smelted, roasted, or dried

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SECONDARY SILVER SUBCATEGORY SECT

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XII

Table XII-2 (Continued)

PSES AND PSNS WASTEWATER DISCHARGE RATES FOR THE SECONDARY SILVER SUBCATEGORY

PSES and PSNS Normalized Production Discharge Rate Normalizing Parameter 1/troy ounce Wastewater Stream troy ounces of 0.086 Leaching silver produced from leaching troy ounces of 4.43 Leaching wet air pollution control silver produced and precipitation of nonphotofrom leaching graphic solutions wet air polluor precipitation tion control troy ounces of 3.07 Precipitation and filtration of nonsilver precipitated photographic solutions troy ounces of Floor and equipment washdown wastewater 0 silver produced

SECT - XII

TABLE XII-3

PSES EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(a) Film Stripping PSES

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average

mg/troy ounce of silver from film stripping

Antimony Arsenic Cadmium	97.180 69.990 10.070	43.300 31.220 4.028
Chromium	18.630	7.553
*Copper	64.450	30.710
Lead	14.100	6.546
Nickel	27.690	18.630
Selenium	41.290	18.630
Silver	14.600	6.042
Thallium	70.490	30.710
*Zinc	51.360	21.150
*Ammonia (as N)	6712.000	2951.000

(b) <u>Film Stripping Wet Air Pollution</u> <u>Control and Precipitation</u> <u>and Filtration</u> <u>of Film Stripping</u> <u>Solutions Wet Air Pollution</u> Control PSES

	· · · · · · · · · · · · · · · · · · ·		
Pollutant or	Maximum	for Maximum f	or
Pollutant Property	Any One	Day Monthly A	verage
		·	

mg/troy ounce of silver from precipitation and filtration of film stripping solutions

Antimony	1.872	0.834	
Arsenic	1.348	0.601	
Cadmium	0.194	0.078	τ.
Chromium	0.359	0.146	
*Copper	1.242	0.592	
Lead	0.272	0.126	
Nickel	0.534	0.359	
Selenium	0.795	0.359	
Silver	0.281	0.116	
Thallium	1.358	0.592	
*Zinc	0.989	0.407	
*Ammonia (as N)	129.300	56.840	

PSES EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(c) <u>Precipitation</u> and <u>Filtration</u> of <u>Film</u> <u>Stripping</u> <u>Solutions</u> PSES

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce	of silver precipita	ated
Antimony	111.100	49.510
Arsenic	80.020	35.690
Cadmium	11.510	4.606
Chromium	21.300	8.636
*Copper	73.690	35.120
Lead	16.120	7.484
Nickel	31.660	21.300
Selenium	47.210	21.300
Silver	16.700	6.908
Thallium	80.600	35.120
*Zinc	58.720	24.180
*Ammonia (as N)	7674.000	3374.000

(d) <u>Precipitation</u> and <u>Filtration</u> of <u>Photographic</u> <u>Solutions</u> PSES

Pollutant or	coperty	Maximum for	Maximum for
Pollutant Pr		Any One Day	Monthly Average
· .	mg/troy ounce of	of silver precipita	ted
Antimony		51.340	22.880
Arsenic		36.970	16.490
Cadmium		5.320	2.128
Chromium		9.842	3.990
*Copper		34.050	16.230
Lead		7.448	3.458
Nickel		14.630	9.842
Selenium		21.810	9.842
Silver		7.714	3.192
Thallium		37.240	16.230
*Zinc		27.130	11.170
*Ammonia (as		3546.000	1559.000

PSES EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(e) <u>Precipitation and Filtration of Photographic Solutions</u> <u>Wet Air Pollution Control PSES</u>

Pollutant	or	Maximum for	Maximum for
Pollutant	Property	Any One Day	Monthly Average
mg/troy	ounce of silver fr photogr	com precipitation and caphic solutions	filtration of
Antimony	as N)	23.430	10.440
Arsenic		16.880	7.527
Cadmium		2.428	0.971
Chromium		4.492	1.821
*Copper		15.540	7.405
Lead		3.399	1.578
Nickel		6.677	4.492
Selenium		9.955	4.492
Silver		3.521	1.457
Thallium		17.000	7.405
*Zinc		12.380	5.099
*Ammonia (1618.000	711.400

(f) Electrolytic Refining PSES

Pollutant Pollutant		perty				ximum y One		Maximum Monthly	
mg/	troy	ounce	of	silver	from	elect	crolytic	refining	
Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc *Ammonia (a	as N))			1	1.40 1.05 0.15 0.28 0.97 0.21 0.41 0.62 0.22 1.06 0.77	56 52 31 73 13 13 18 23 20 54 75	0.654 0.471 0.061 0.114 0.464 0.099 0.281 0.281 0.281 0.991 0.464 0.319 44.540	· · ·

PSES EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(g) <u>Furnace Wet Air Pollution Control</u> PSES

Pollutant Pollutant		t y	•		lmum for One Day	Maximum Monthly	
mg/troy	ounce of	silver	from	silver	roasted,	smelted, or	dried
Antimony	,		· .		0.000	0.000	•
Arsenic					0.000	0.000	
Cadmium					0.000	0.000	÷
Chromium	, L				0.000	0.000	
*Copper	-	÷ .			0.000	0.000	· .
Lead	÷.,				0.000	0.000	
Nickel					0.000	0.000	
Selenium	1				0.000	0.000	
Silver	•				0.000	0.000	· .
Thallium	1				0.000	0.000	
*Zinc	•				0.000	0.000	
*Ammonia	(as N)		-		0.000	0.000	· , <i>'</i>

(h) Leaching PSES

Pollutant or Pollutant Property				Maximum for Any One Day	Maximum Monthly	
mg/troy oun	ce of	silver	from	silver produced	from leach	ing
Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc *Ammonia (as N				0.166 0.120 0.017 0.032 0.110 0.024 0.047 0.071 0.025 0.120 0.088 11.460	0.074 0.053 0.007 0.013 0.052 0.011 0.032 0.032 0.010 0.052 0.036 5.040	

*Regulated Pollutant

PSES EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(i) Leaching Wet Air Pollution Control and Precipitation of Nonphotographic Solutions Wet Air Pollution Control PSES

Pollutant or	coperty	Maximum for	Maximum for
Pollutant Pr		Any One Day	Monthly Average
mg/troy ou		from silver produced or silver	from leaching
Antimony	N)	8.550	3.810
Arsenic		6.158	2.747
Cadmium		0.886	0.354
Chromium		1.639	0.665
*Copper		5.670	2.702
Lead		1.240	0.576
Nickel		2.437	1.639
Selenium		3.633	1.639
Silver		1.285	0.532
Thallium		6.202	2.702
*Zinc		4.519	1.861
*Ammonia (as		590.500	259.600

(j) <u>Precipitation</u> and <u>Filtration</u> of <u>Nonphotographic</u> <u>Solutions</u> PSES

Pollutant of Pollutant	or	Maximum for	Maximum for
	Property	Any One Day	Monthly Average
	mg/troy ounce	of silver precipitat	ed
Antimony		5.925	2.640
Arsenic		4.267	1.903
Cadmium		0.614	0.246
Chromium		1.136	0.461
*Copper		3.930	1.873
Lead		0.860	0.399
Nickel		1.689	1.136
Selenium		2.517	1.136
Silver	as N)	0.890	0.399
Thallium		4.298	1.873
*Zinc		3.131	1.289
*Ammonia (a		409.200	179.900

PSES EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(k) Floor and Equipment Washdown Water PSES

Pollutant Pollutant	or Property		Maximum for Any One Day	Maximum Monthly	for Average
· ·	mg/troy o	ounce of	silver production		· .
Antimony			0.000	0.000	
Arsenic			0.000	0.000	
Cadmium			0.000	0.000	
Chromium			0.000	0.000	
-			0.000	0.000	
*Copper			0.000	0.000	
Lead Nickel	•		0.000	0.000	1. A
	4 - 4 		0.000	0.000	
Selenium			0.000	0.000	
Silver			0.000	0.000	;
Thallium	·		0.000	0.000	
*Zinc *Ammonia (as N)		0.000	0.000	

TABLE XII-4

PSNS EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(a) Film Stripping PSNS

Pollutant or		, , , , , , , , , , , , , , , , , , ,	Maximum for		Maximum	
Pollutant	Property		Any One	e Day	Monthly	Average
	mg/troy	ounce of	silver from	film	stripping	
Antimony			97.1	.80	43.300	
Arsenic			69.9	90	31.220	
Cadmium			10.0	70	4.028	
Chromium		1 1	18.6	30	7.553	• .
*Copper			64.4	50	30.710	
Lead			14.1	.00	6.546	
Nickel			27.6	90	18.630	5 5
Selenium			· 41.2	90	18.630	
Silver			14.6	00	6.042	
Thallium			70.4	90	30.710	
*Zinc			51.3	60	21.150	
*Ammonia (a	IS N)	¢	6712.0	00	2951.000	

(b) Film Stripping Wet Air Pollution Control and Precipitation and Filtration of Film Stripping Solutions Wet Air Pollution Control PSNS

Pollutant or Pollutant Property	Maximum for Any One Day	Maximum Monthly	
mg/troy ounce of silver fr	om precipitation and ipping solutions	filtration	of film
Antimony Arsenic	1.872	0.834	
Cadmium	1.348 0.194	0.601 0.078	¢
Chromium	0.359	0.146	
*Copper Lead	1.242 0.272	0.592	
Nickel	0.534	0.126 0.359	
Selenium	0.795	0.359	
Silver Thallium	0.281	0.116	
*Zinc	0.989	0.592 0.407	
*Ammonia (as N)	129.300	56.840	

*Regulated Pollutant

PSNS EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(c) <u>Precipitation</u> and <u>Filtration</u> of <u>Film</u> <u>Stripping</u> <u>Solutions</u> PSNS

Pollutant o Pollutant	or Property		Maximum for Any One Day	Maximum Monthly	
	mg/troy	ounce of	silver precipitate	đ	
Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc *Ammonia (as N)		111.100 80.020 11.510 21.300 73.690 16.120 31.660 47.210 16.700 80.600 58.720 7674.000	49.510 35.690 4.606 8.636 35.120 7.484 21.300 21.300 6.908 35.120 24.180 3374.000	

(d) <u>Precipitation</u> and <u>Filtration</u> of <u>Photographic</u> <u>Solutions</u> PSNS

Pollutant or	operty	Maximum for	Maximum for
Pollutant Pr		Any One Day	Monthly Average
<u></u>	mg/troy ounce	e of silver precipit	ated
Antimony	N)	51.340	22.880
Arsenic		36.970	16.490
Cadmium		5.320	2.128
Chromium		9.842	3.990
*Copper		34.050	16.230
Lead		7.448	3.458
Nickel		14.630	9.842
Selenium		21.810	9.842
Silver		7.714	3.192
Thallium		37.240	16.230
*Zinc		27.130	11.170
*Ammonia (as		3546.000	1559.000

*Regulated Pollutant

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PSNS EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(e) <u>Precipitation and Filtration of Photographic Solutions</u> <u>Wet Air Pollution Control PSNS</u>

Pollutant	or	Maximum for	Maximum for
Pollutant	Property	Any One Day	Monthly Average
mg/troy	ounce of silver photo	from precipitation and graphic solutions	filtration of
Antimony	as N)	23.430	10.440
Arsenic		16.880	7.527
Cadmium		2.428	0.971
Chromium		4.492	1.821
*Copper		15.540	7.405
Lead		3.399	1.578
Nickel		6.677	4.492
Selenium		9.955	4.492
Silver		3.521	1.457
Thallium		17.000	7.405
*Zinc		12.380	5.099
*Ammonia (1618.000	711.400

(f) Electrolytic Refining PSNS

Pollutant or Pollutant Property		Maximum for Any One Day	Maximum Monthly	,
mg/troy ounce o	of silver	from electrolytic	refining	
Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc *Ammonia (as N)	· ·	1.467 1.056 0.152 0.281 0.973 0.213 0.418 0.623 0.220 1.064 0.775 101.300	0.654 0.471 0.061 0.114 0.464 0.099 0.281 0.281 0.281 0.281 0.464 0.319 44.540	

*Regulated Pollutant

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PSNS EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(g) Furnace Wet Air Pollution Control PSNS

Pollutant Pollutant	or Property			imum for One Day	Maximum Monthly	for Averag e
mg/troy o	unce of sil	ver fro	om silver	roasted,	smelted, or	dried
Antimony	•	4	· · · · ·	0.000	0.000	
Arsenic		* .		0.000	0.000	
Cadmium				0.000	0.000	
Chromium				0.000	0.000	
*Copper				0.000	0.000	
Lead		·	•	0.000	0.000	• [*] •
Nickel				0.000	0.000	
Selenium				0.000	0.000	
Silver	i -	۰. ۱		0.000	0.000	
Thallium			×	0.000	0.000	
*Zinc	1			0.000	0.000	
*Ammonia (as N)	• * * · · · ·		0.000	0.000	
(h) <u>Leach</u>	ing PSNS					
Pollutant	or			imum for One Day	Maximum Monthly	
Pollutant Pollutant mg/troy	or Property	ailver f	Any	One Day er produc	Monthly ed from leach	Average
Pollutant Pollutant mg/troy Antimony	or Property	ilver f	Any	One Day er produc 0.166	Monthly ed from leac 0.074	Average
Pollutant Pollutant mg/troy Antimony Arsenic	or Property	ilver f	Any	One Day er produc 0.166 0.120	Monthly ed from leac 0.074 0.053	Average
Pollutant Pollutant mg/troy Antimony Arsenic Cadmium	or Property	ilver f	Any	One Day er produc 0.166 0.120 0.017	Monthly ed from leac 0.074 0.053 0.007	Average
Pollutant Pollutant mg/troy Antimony Arsenic Cadmium Chromium	or Property	ilver f	Any	One Day er produc 0.166 0.120 0.017 0.032	Monthly ed from leac 0.074 0.053 0.007 0.013	Average
Pollutant Pollutant mg/troy Antimony Arsenic Cadmium Chromium *Copper	or Property	ilver f	Any	One Day er produc 0.166 0.120 0.017 0.032 0.110	Monthly ed from leach 0.074 0.053 0.007 0.013 0.052	Average
Pollutant Pollutant mg/troy Antimony Arsenic Cadmium Chromium *Copper Lead	or Property	ilver f	Any	One Day er produc 0.166 0.120 0.017 0.032 0.110 0.024	Monthly ed from leac 0.074 0.053 0.007 0.013	Average
Pollutant Pollutant mg/troy Antimony Arsenic Cadmium Chromium *Copper	or Property	ilver f	Any	One Day er produc 0.166 0.120 0.017 0.032 0.110 0.024 0.047	Monthly ed from leach 0.074 0.053 0.007 0.013 0.052 0.011	Average
Pollutant Pollutant mg/troy Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium	or Property	ilver f	Any	One Day er produc 0.166 0.120 0.017 0.032 0.110 0.024 0.047 0.071	Monthly ed from leach 0.074 0.053 0.007 0.013 0.052 0.011 0.032	Average
Pollutant Pollutant mg/troy Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver	or Property	ilver f	Any	One Day er produc 0.166 0.120 0.017 0.032 0.110 0.024 0.047	Monthly ed from leach 0.074 0.053 0.007 0.013 0.052 0.011 0.032 0.032	Average
Pollutant Pollutant mg/troy Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium	or Property	ilver f	Any	One Day er produc 0.166 0.120 0.017 0.032 0.110 0.024 0.047 0.071 0.025	Monthly ed from leach 0.074 0.053 0.007 0.013 0.052 0.011 0.032 0.032 0.010	Average

*Regulated Pollutant

PSNS EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(i) <u>Leaching Wet Air</u> <u>Pollution</u> <u>Control</u> <u>and</u> <u>Precipitation</u> <u>of</u> <u>Nonphotographic</u> <u>Solutions</u> <u>Wet Air</u> <u>Pollution</u> <u>Control</u> <u>PSNS</u>

Pollutant o Pollutant	r Property	•	Maximum for Any One Day	Maximum for Monthly Average
mg/troy	ounce of	silver	from silver produced or silver	from leaching
Antimony Arsenic Cadmium Chromium *Copper Lead Nickel Selenium Silver Thallium *Zinc *Ammonia (a	s N)	÷ •	8.550 6.158 0.886 1.639 5.670 1.240 2.437 3.633 1.285 6.202 4.519 590.500	3.810 2.747 0.354 0.665 2.702 0.576 1.639 1.639 0.532 2.702 1.861 259.600

(j) <u>Precipitation</u> and <u>Filtration</u> of <u>Nonphotographic</u> <u>Solutions</u> PSNS

Pollutant or	Maximum for	Maximum for
Pollutant Property	Any One Day	Monthly Average
mg/troy ounce of silver prec	ipitated	
Antimony	5.925	2.640
Arsenic	4.267	1.903
Cadmium	0.614	0.246
Chromium	1.136	0.461
*Copper	3.930	1.873
Lead	0.860	0.399
Nickel	1.689	1.136
Selenium	2.517	1.136
Silver	0.890	0.399
Thallium	4.298	1.873
*Zinc	3.131	1.289
*Ammonia (as N)	409.200	179.900

*Regulated Pollutant

SECONDARY SILVER SUBCATEGORY

SECT - XII

TABLE XII-4 (Continued)

PSNS EFFLUENT LIMITATIONS FOR THE SECONDARY SILVER SUBCATEGORY

(k) Floor and Equipment Washdown Water PSNS

Pollutant or Pollutant Property		Maximum for Any One Day	Maximum for Monthly Average
mg/troy ounce of s	lver product	ion	
Antimony		0.000	0.000
Arsenic	. · · · ·	0.000	0.000
Cadmium		0.000	0.000
Chromium	Sing a prime and a second s	0.000	0.000
*Copper	_ + *_	0.000	0.000
Lead		0.000	0.000
Nickel	i. P	0.000	0.000
Selenium		0.000	0.000
Silver		0.000	0.000
Thallium		0.000	0.000
*Zinc	en e	0.000	0.000
*Ammonia (as N)		0.000	0.000

*Regulated Pollutant

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SECONDARY SILVER SUBCATEGORY

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SECT - XII

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SECONDARY SILVER SUBCATEGORY

SECTION XIII

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BEST CONVENTIONAL POLLUTANT CONTROL TECHNOLOGY

EPA is not promulgating best conventional pollutant control technology (BCT) for the secondary silver subcategory at this time.

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NONFERROUS METALS MANUFACTURING FOINT SOURCE CATEGORY

DEVELOPMENT DOCUMENT SUPPLEMENT

for the

Secondary Mercury Subcategory

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May 1989

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SECTION I

SUMMARY

This document provides the technical basis for promulgating pretreatment standards for new indirect dischargers (PSNS) and standards of performance for new source direct dischargers (NSPS). The secondary mercury subcategory is comprised of four plants. Two plants achieve zero discharge of process wastewater, and two plants do not generate process wastewater.

EPA first studied the secondary mercury subcategory to determine differences in whether raw materials, final products, manufacturing processes, equipment, age and size of plants, or water usage, required the development of separate effluent limitations and standards for different segments of the subcategory. This involved a detailed analysis of wastewater discharge and treated effluent characteristics, including the sources and volume of water used, the processes used, the sources of pollutants and wastewaters in the plant, and the constituents of wastewaters including priority pollutants. As a result, three subdivisions have been identified for this subcategory that warrant separate effluent limitations. These include:

1. Spent battery electrolyte,

2. Acid wash and rinse water, and

3. Furnace wet air pollution control.

Several distinct control and treatment technologies (both inplant and end-of-pipe) applicable to the secondary mercury subcategory were identified. The Agency analyzed both historical and newly generated data on the performance of these technologies, including their nonwater quality environmental impacts and air quality, solid waste generation, and energy requirements. EPA also studied various flow reduction techniques reported in the data collection portfolios (dcp) and plant visits.

Engineering costs were prepared for each of the control and treatment options considered for the subcategory. These costs were then used by the Agency to estimate the implementing the various options on the subcategory. impact of For each control and treatment option that the Agency found to be effective and technically feasible in controlling the discharge pollutants, the number of potential closures, number of of employees affected, and impact on price were investigated. These results are reported in a separate document entitled "The Economic Impact Analysis of Effluent Limitations and Standards for the Nonferrous Metals Manufacturing Industry."

No plants in the secondary mercury subcategory discharge process wastewater. This is achieved by 100 percent recycle on-site or by contractor disposal of process wastewater, or is a result of a production process that generates no process wastewater. Therefore, BPT, BAT, BCT, and PSES are not being promulgated for this subcategory. The secondary mercury subcategory is regulated under New Source Performance Standards and Pretreatment Standards for New Sources.

After examining the various treatment technologies, the Agency has identified best demonstrated technology, which is the technical basis of NSPS, to represent the best existing technology in the nonferrous metals manufacturing category. Metals removal based on chemical precipitation, sedimentation, and multimedia filtration technology is the basis for the NSPS limitations. In selecting NSPS, EPA recognizes that new plants have the opportunity to implement the best and most efficient manufacturing processes and treatment technologies available.

PSES is not being promulgated for this subcategory because there are no existing indirect dischargers in the secondary mercury subcategory. For PSNS, the Agency selected end-of-pipe treatment techniques equivalent to NSPS.

BCT is not being promulgated because there are no direct dischargers.

The mass limitations for NSPS and PSNS are presented in Section II.

SECONDARY MERCURY SUBCATEGORY SECT - II

SECTION II

CONCLUSIONS

EPA has divided the secondary mercury subcategory into three subdivisions or building blocks for the purpose of effluent limitations and standards. These subdivisions are:

- (a) Spent battery electrolyte,
- (b) Acid wash and rinse water, and
- (c) Furnace wet air pollution control.

BPT is not being promulgated because there are no direct dischargers in the secondary mercury subcategory.

BAT is not being promulgated because there are no direct dischargers in the secondary mercury subcategory.

NSPS are promulgated based on the performance achievable by the application of chemical precipitation, sedimentation, and multimedia filtration technology. The following effluent standards are promulgated for new sources:

(a) Spent Battery Electrolyte NSPS

Pollutant	or	Maximum	for	Maximum for
Pollutant	Property	Any One	Day	Monthly Average

mg/kg (lb/million lbs) of mercury produced from batteries

Lead		0.030		0.014	
Mercury		0.016		0.006	
TSS		1.590		1.272	
pH	Within the	range of	7.5 to	10.0 at all	times

(b) Acid Wash and Rinse Water NSPS

Pollutant	or	Maximum fo	or Maximu	n for
Pollutant	Property	Any One Da	ay Monthly A	Average

mg/kg (lb/million lbs) of mercury washed and rinsed

Lead			0.00	0056	5			0.0	0002	5
Mercury			0.00	0030) .		104	0.0	0001:	2
TSS	•		0.03	300(D			0.0	0240	0
pH	Within	the	range	of	7.5	to	10.0	at	a 11	times

SECT - II

(c) <u>Furnace Wet Air Pollution Control</u> NSPS

Pollutant Pollutant		Maximum for Any One Day	Maximum for Monthly Average	
mg/kg	(lb/million	lbs) of mercury	processed through furnad	ce
Lead Mercury TSS pH	Within (0.000 0.000 0.000 the range of 7.5	0.000 0.000 0.000 to 10.0 at all times	

PSES is not being promulgated because there are no indirect dischargers in the secondary mercury subcategory.

PSNS are promulgated based on the performance achievable by the application of chemical precipitation. sedimentation, and multimedia filtration technology. The following pretreatment standards are promulgated for new sources:

(a) Spent Battery Electrolyte PSNS

Pollutant or	r	Maximum	for	Maximum	for
Pollutant Pr	roperty	Any One	Day	Monthly Av	verage

mg/kg (lb/million lbs) of mercury produced from batteries

Lead	0.030	0.014
Mercury	0.016	0.006

(b) Acid Wash and Rinse Water PSNS

Pollut Pollut	ant or ant Pro		Maximum Any One			ximum hly Av		
	mg/kg	(lb/millio	on lbs)	of r	mercury	washed	and r	insed
Lead Mercur	У			0056 0030			00026 00012	

SECONDARY MERCURY SUBCATEGORY SECT - II

(c) <u>Furn</u>	ace <u>Wet</u> Air	<u>Pollu</u>	ution Contr	ol PSNS	۰ ۲۰۰۰ (۲۰۰۰)	
Pollutant Pollutant	or Property		mum for One Day	Maximum Monthly Av		
mg/kg	(lb/million	lbs)	of mercury	processed	through	furnace
Lead Mercury			0.000		000	

BCT is not being promulgated for the secondary mercury subcategory at this time.

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SECTION III

SUBCATEGORY PROFILE

This section of the secondary mercury supplement describes the raw materials and processes used in producing secondary mercury and presents a profile of the secondary mercury plants identified in this study. For a discussion of the purpose, authority, and methodology for this study, and a general description of the nonferrous metals manufacturing category, refer to Section III of Vol. I.

Mercury is used in numerous agricultural, chemical and electrical applications. Mercury is used extensively in the chemical industry, particularly in the production of chlorine and caustic soda. Mercury compounds are also used extensively in paints and as catalysts. Agricultural uses of mercury include germicides for seed protection and weed control, and fungicidal fruit sprays. Electrical applications include low-pressure and high pressure mercury vapor lamps, power control switches, and drycell batteries. Other uses are in barometers, thermometers, as a vibration damper, and as a coolant. Mercury produced from secondary sources is used in many applications, such as those described above.

DESCRIPTION OF SECONDARY MERCURY PRODUCTION

The production of secondary mercury can be divided into three distinct stages: separation of gross impurities, distillation, and acid washing. The actual processes used in each stage vary with the type and purity of the raw material used. The secondary mercury production process is presented schematically in Figure III-1 (page 2876) and is described below.

RAW MATERIALS

Mercury can be reclaimed from a variety of raw materials. including thermometers, switches, filters. controls, zinc and silver amalgams, mercuric oxide battery cells, and other types of scrap. Secondary mercury annually supplies the United States with approximately 20 percent of domestic requirements. Several plants refining secondary mercury also refine prime virgin mercury. Although prime virgin mercury can be considered to be a primary raw material, its refining is included with secondary mercury, because it is refined on-site with secondary mercury using the same equipment and production processes.

SEPARATION OF GROSS IMPURITIES

Depending on the type of raw material being processed, gross impurities, such as glass from mercury thermometers, or spent electrolyte from mercuric oxide battery cells, may have to be

SECONDARY MERCURY SUBCATEGORY SECT - III

separated from the mercury. The separation of gross impurities must occur prior to distilling the mercury. Raw materials such as thermometers, switches, filters, controls, and zinc and silver amalgams may be separated from their gross impurities by roasting in a furnace. The mercury is separated from impurities by vaporizing it, and then recovering mercury by condensation The nonvolatilized solids are removed from the furnace after all the mercury has been removed. A water scrubber may be used to control air emissions from the mercury furnace and condenser, and the scrubber may have a discharge from it.

Before mercury can be recovered from mercuric oxide battery cells, the battery electrolyte must be removed. On a small scale, this is most likely accomplished by manually draining the spent electrolyte from each cell. Spent electrolyte removed in this step is a waste stream.

DISTILLATION

Mercury distillation columns, also known as retorts stills, or kettles, are used to produce high-purity mercury. No wastewater is generated by this process. A typical distillation process consists of charging raw, impure mercury into the bottom of a still and heating the charge to a prescribed temperature, some. what less than the boiling point of mercury, 356.9°C. While heating the charge, air may be bubbled through the still in order to oxidize metallic impurities, such as lead, zinc, cadmium, copper or tin. When the charge reaches the critical temperature. the mercury begins to vaporize, and the mercury is recovered in water cooled condensing an overhead, system. Mercury distillation may be run batchwise or continuously, and in both cases it can be considered a dry process. None of the water used in the condensing coils contacts the mercury.

Multiple distillation units may be operated in series to produce high purity (approximately 99.999999 percent) mercury. Like the single distillation process, no wastewater is generated by multiple distillation units.

ACID WASHING

Another method for further purifying mercury is acid washing and rinsing. In this method, a small amount of dilute nitric acid is used to wash the distilled mercury product, and then a small amount of distilled water is used to wash the residual acid from the mercury product. Mercury of 99.9 percent purity can be produced in this manner. The acid wash and rinse water may be discharged from this process as a waste stream.

PROCESS WASTEWATER SOURCES

Although a variety of processes are involved in secondary mercury production the process wastewater sources can be subdivided as follows:

- 1. Spent battery electrolyte,
- 2. Acid wash and rinse water, and
- 3. Furnace wet air pollution control.

OTHER WASTEWATER SOURCES

There are other wastewater streams associated with the secondary mercury subcategory. These wastewater streams may include stormwater runoff, maintenance and cleanup water, and noncontact cooling water. These waste streams are not considered as a part of this rulemaking. EPA believes that the flows and pollutant loadings associated with these waste streams are insignificant relative to the waste streams selected, or are best handled by the appropriate permit authority on a case-by-case basis under authority of Section 402 of the Clean Water Act.

AGE, PRODUCTION, AND PROCESS PROFILE

Figure III-2 (page 2877) shows the locations of the four secondary mercury plants operating in the United States. Two of the four plants are located near the industrial centers of the Northeast, one is in Illinois, and one in California.

Table III-1 (page 2874) shows the relative age and discharge status of the mercury plants and illustrates that all the plants were built after World War II. The average plant age is 30 years old. From Table III-2 (page 2875), it can be seen that two plants produce between 50 and 100 tons per year of metal, while one plant produces less than 25 tons per year. Mean production is about 55 tons per year.

Table III-3 (page 2875) provides a summary of the number of plants generating wastewater for the waste streams associated with various processes and the number of plants with the process.

Table III-1

INITIAL OPERATING YEAR (RANGE) SUMMARY OF PLANTS IN THE SECONDARY MERCURY SUBCATEGORY BY DISCHARGE TYPE

	Initial (P			
Type of Plant	1982- 1968 (0-15)	lant Age in Yea 1967- 1958 (16-25)	1957- 1948 (26-35)	Total
Direct	0	0	0	0
Indirect	0	0	0	0
Zero	0	0	2	2
Dry	0	1,	0	2*
	—	·	_	
TOTAL	0	0	1	4

*One plant did not report initial operating year

SECONDARY MERCURY SUBCATEGORY SECT - III

Table III-2

PRODUCTION RANGES FOR THE SECONDARY MERCURY SUBCATEGORY

Type of Plant	Mercury Pro 0-25 (tons/yr)	oduction Rang 25-50 (tons/yr)	ge for 1982 50-100 (tons/yr)	Total Number <u>of</u> <u>Plants</u>
Direct	0	0	0	0
Indirect	0	0	0	0
Zero	· · 1	0	l	2
Dry	0	0	1	2*
÷				-

* One plant did not report mercury production.

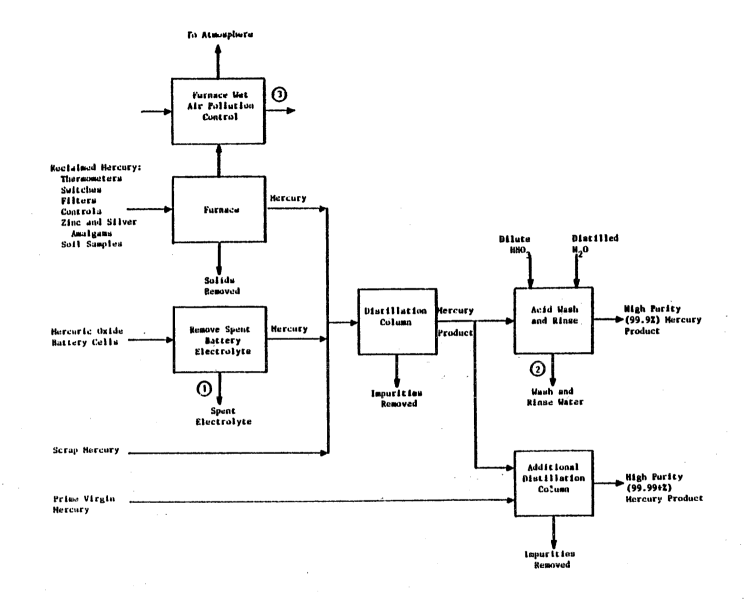
Table III-3

SUMMARY OF SUBCATEGORY PROCESSES AND ASSOCIATED

WASTE STREAMS

Process or Waste Stream	Number of Plants With Process or Waste Stream	Number of Plants Reporting Generation of Wastewater*
Spent battery electrolyte	1	1
Furnace wet air pollution control	1	0
Distillation	4	0
Acid wash and rinse water	1	1

*Through reuse or evaporation practices, a plant may "generate" a wastewater from a particular process but not discharge it.



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SECONDARY MERCURY SUBCATEGORY SECT

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III

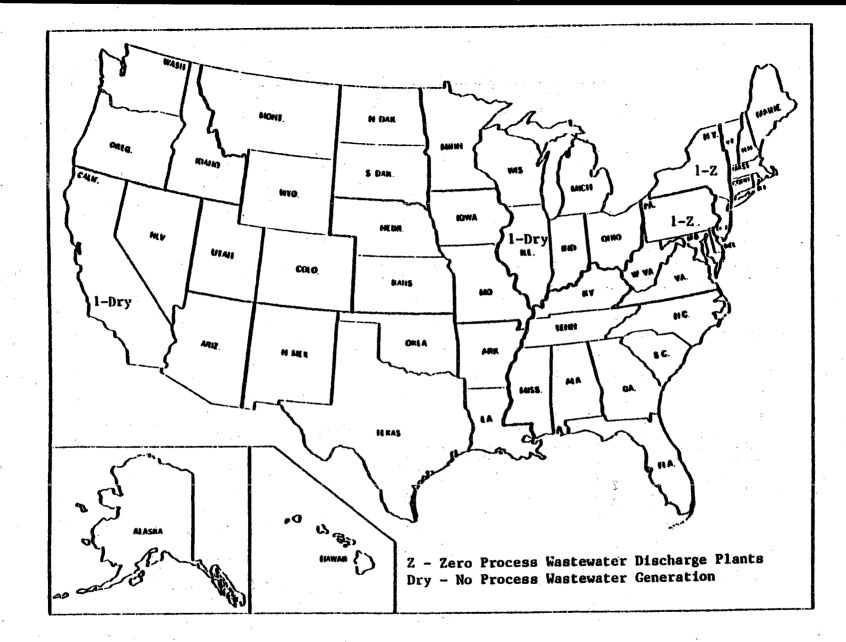


Figure III-2

GEOGRAPHIC LOCATIONS OF THE SECONDARY MERCURY SUBCATEGORY PLANTS

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SECT - III

SECONDARY MERCURY SUBCATEGORY

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SECONDARY MERCURY SUBCATEGORY SECT - IV

SECTION IV

SUBCATEGORIZATION

This section summarizes the factors considered during the designation of the secondary mercury subcategory and its related subdivisions. Production normalizing parameters for each subdivision will also be discussed.

FACTORS CONSIDERED IN SUBDIVIDING THE SECONDARY MERCURY SUBCATEGORY

The general factors for subcategorization listed previously were each evaluated when considering subdivision of the secondary mercury subcategory. In the discussion that follows, the factors will be described as they pertain to this subcategory.

The rationale for considering subdivision of the secondary mercury subcategory is based primarily on differences in the production processes and raw materials used. Within this subcategory, a number of different operations are performed, which may or may not have a water use or discharge, and which may require the establishment of separate effluent limitations. While secondary mercury is considered a single subcategory, a more thorough examination of the production processes has illustrated the need for limitations and standards based on specific flow allowances for the following subdivisions:

- 1. Spent battery electrolyte,
- 2. Acid wash and rinse water, and
- 3. Furnace wet air pollution control.

These subdivisions follow directly from differences within the three distinct production states of secondary mercury: separation of gross impurities, distillation, and additional purification. A secondary mercury plant may have one, two, or all three of these production states.

Separation of gross impurities such as spent battery electrolyte or glass from thermometers gives rise to the first and third subdivisions: spent battery electrolyte and furnace wet air pollution control. A plant which recovers mercury from mercuric oxide battery cells must first drain the spent electrolyte from the cells. This wastewater may be discharged. A plant which recovers mercury from recycled thermometers, switches, filters, and amalgams may remove the mercury from the unwanted solids by vaporizing mercury in a furnace. After condensing the product mercury, the air emissions may be controlled with a scrubber. The furnace scrubber may have a discharge, and this creates the need for the third subdivision.

Additional purification of the mercury product gives rise to the second subdivision: acid wash and rinse water. After distilling

the mercury, it may be washed with acid and rinsed with water to increase its purity. The acid wash and rinse water may be discharged as a waste stream.

OTHER FACTORS

The other factors considered in this evaluation were shown to be inappropriate bases for subdivision. Air pollution control methods, treatment costs, and total energy requirements are functions of the selected subcategorization factors -- metal product, raw materials, and production processes. Therefore, they are not independent factors and do not affect the subcategorization which has been applied. Certain other factors, such as plant age, plant size, and the number of employees were also evaluated and determined to be inappropriate for use as bases for subdivision of secondary mercury plants.

PRODUCTION NORMALIZING PARAMETERS

As discussed previously, the effluent limitations and standards developed in this document establish mass limitations for the discharge of specific pollutant parameters. To allow these regulations to be applied to plants with various production capacities, the mass of pollutant discharged must be related to a unit of production. This factor is known as the production normalizing parameter (PNP).

In general, for each production process which has a wastewater associated with it, the actual mass of mercury product or intermediate produced will be used as the PNP Thus, the PNPs for the three subdivisions are as follows:

Subdivision

PNP

1. Spent battery electrolyte

mercury produced from batteries

2. Acid wash and rinse water

mercury washed and rinsed

3. Furnace wet air pollution control

mercury processed through furnace

Other PNPs were considered. The use of production capacity instead of actual production was eliminated from consideration because the mass of the pollutant produced is more a function of true production than of installed capacity.

SECTION V

WATER USE AND WASTEWATER CHARACTERISTICS

This section describes the characteristics of the wastewaters associated with the secondary mercury subcategory. Water use and discharge rates are explained and then summarized in tables at the end of this section. Data used to characterize the wastewaters are presented. Finally, the specific source, water use and discharge flows, and wastewater characteristics for each separate wastewater source are discussed.

The two principal data sources were used in the development of effluent limitations and standards for this subcategory are data collection portfolios (dcp) and field sampling results. Data collection portfolios contain information regarding wastewater flows and production levels.

In order to quantify the pollutant discharge from secondary mercury plants, the levels of priority pollutants in the wastewaters must be known. Since field sampling was not performed at any plants in the secondary mercury subcategory, analytical data, presented in Section V of the supplement for the primary precious metals and mercury subcategory, were transferred from a primary mercury plant to characterize wastewater in the secondary mercury subcategory. In general, the samples were analyzed for two classes of pollutants (including 13 of the 126 priority pollutants): priority metal pollutants and criteria pollutants (which includes both conventional and nonconventional pollutants). Because the analytical standard for TCDD was judged to be too hazardous to be made generally available, samples were never analyzed for this pollutant. Samples were also never analyzed for asbestos or cyanide. There is no reason to expect that TCDD, asbestos, or cyanide would be present in secondary mercury wastewater.

As described in Section IV of this supplement, the secondary mercury subcategory has been divided into three wastewater source oriented subdivisions or building blocks, so that the promulgated regulation contains mass discharge limitations and standards for three manufacturing processes that discharge process wastewater. Differences in the wastewater characteristics associated with these subdivisions are to be expected. For this reason, wastewater streams corresponding to each subdivision are addressed separately in the discussions that follow. These wastewater sources are:

- 1. Spent battery electrolyte,
- 2. Acid wash and rinse water, and
- 3. Furnace wet air pollution control.

No additional sampling data for this subcategory were obtained from EPA sampling efforts or industry comments between proposal and promulgation. Characterization of secondary mercury

SECONDARY MERCURY SUBCATEGORY SECT - V

subcategory wastewaters (Section V), and selection of pollutant parameters for limitation (Section VI) are based upon the same data used for proposal.

WASTEWATER FLOW RATES

Data supplied by dcp responses were evaluated, and two flow-toproduction ratios, water use and wastewater discharge flow, were calculated for each stream. The two ratios are differentiated by the flow value used in calculation. Water use is defined as the volume of water or other fluid required for a given process per mass of mercury product and is therefore based on the sum of recycle and make-up flows to a given process. Wastewater flow discharged after pretreatment or recycle (if these are present) is used in calculating the production normalized flow -- the volume of wastewater discharged from a given process to further treatment, disposal, or discharge per mass of mercury produced. Differences between the water use and wastewater flows associated with a given stream result from recycle, evaporation, and carryover on the product. The production values used in calculation correspond to the production normalizing parameter, PNP, assigned to each stream, as outlined in Section IV. As an example, acid wash and rinse water flow is related to the amount of mercury washed and rinsed. As such, the discharge rate is expressed in liters of acid wash and rinse water per metric ton of mercury washed and rinsed (gallons of acid wash and rinse water per ton of mercury washed and rinsed).

The production normalized discharge flows were compiled and statistically analyzed by stream type. These production normalized water use and discharge flows are presented by subdivision in Tables V-1 through V-3 (pages 2885 - 2886). Where appropriate, an attempt was made to identify factors that could account for variations in water use and discharge rates. These variations are discussed later in this section by subdivision. A similar analysis of factors affecting the wastewater flows is presented in Sections XI and XII where representative NSPS and pretreatment flows are selected for use in calculating the effluent limitations.

WASTEWATER CHARACTERISTICS DATA

Data used to characterize the various wastewaters associated with secondary mercury production come from two sources -- data collection portfolios and analytical data from field sampling trips.

DATA COLLECTION PORTFOLIOS

In the data collection portfolios, the mercury plants that generate wastewater were asked to specify the presence of priority pollutants in their wastewater. No plants indicated that any priority organic pollutants were present. However, one of the two plants stated that they either knew priority metals to be present or they believed the metals to be present. The

responses for the metals and cyanide are summarized below:

Pollutant	Known Present	Believed Present
Antimony	0	0
Arsenic	Ô ·	0
Beryllium	0	0
Cadmium	0	· • 0
Chromium	0	0
Copper	• • • O	0
Cyanide	0	· O
Lead	0	0
Mercury	1	1
Nickel	0	0
Selenium	0	0
Silver	0	. · · · O
Thallium	0 1	• 0
Zinc	0	0

FIELD SAMPLING DATA

In order to quantify the concentrations of pollutants present in wastewater from secondary mercury plants, wastewater samples were collected at one primary mercury plant, which roasts mercury ore to produce mercury metal. Analytical data from the primary mercury plant are presented in the supplement for the primary precious metals and mercury subcategory. Primary mercury and secondary mercury field sampling data are expected to show similar characteristics because of similarities in raw materials and production processes. Both plants roast or distill а mercury-containing raw material and use wet scrubbers to control emissions, and also wash their product to increase its purity.

WASTEWATER CHARACTERISTICS AND FLOWS BY SUBDIVISION

Since secondary mercury production involves three principal sources of wastewater and each has potentially different characteristics and flows, the wastewater characteristics and discharge rates corresponding to each subdivision will be described separately. A brief description of why the associated production processes generate a wastewater and explanations for variations of water use within each subdivision will also be discussed.

SPENT BATTERY ELECTROLYTE

One plant recovers mercury from mercuric oxide battery cells. The first step in this recovery is to drain the spent electrolyte from the cells. Spent battery electrolyte may be discharged as a wastewater stream. Production normalized water use and discharge rates for this waste stream are shown in Table V-1 (page 2885), in liters per metric ton of mercury produced from batteries. This subdivision is similar to spent battery electrolyte from lead batteries (see the battery cracking subdivision of the

secondary lead subcategory), however, secondary mercury spent electrolyte is not expected to have similar pollutant characteristics nor similar production normalized flows.

Although spent battery electrolyte was not sampled, wastewater from the primary mercury industry should have similar characteristics to this waste stream. Spent battery electrolyte should contain treatable concentrations of priority metals, total suspended solids, and exhibit a low pH.

ACID WASH AND RINSE WATER

After recovering mercury in a distillation system, the product may be washed with dilute nitric acid and rinsed with distilled water in order to further purify it. Acid washing and water rinsing produces a high-purity (99.9 percent) mercury product, and also generates a wastewater stream which may be discharged. The production normalized water use and discharge rates for acid wash and rinse water are given in Table V-2 (page 2885) in liters per metric ton of mercury washed and rinsed.

Although acid wash and rinse water was not sampled, data from the primary mercury industry should be similar to this waste stream. Acid wash and rinse water should contain treatable concentrations of priority metals, total suspended solids, and exhibit a low pH.

FURNACE WET AIR POLLUTION CONTROL

One plant recovers mercury from sources such as thermometers, switches, contacts, and amalgams by heating the raw materials in a furnace in order to vaporize the mercury. After condensing the mercury product, air emissions from the furnace may be controlled with a wet scrubber. The furnace scrubber may have a discharge associated with it. Water use and discharge rates for furnace wet air pollution control are presented in Table V-3 (page 2886). Only one plant has this process and operates its scrubber at 100 percent recycle.

Table V-1

WATER USE AND DISCHARGE RATES FOR SPENT BATTERY ELECTROLYTE

(1/kkg of mercury produced from batteries)

Percent Plant Code Recycle		Production Normalized Water Use	Production Normalized Discharge Flow
1161	0	106	106

Table V-2

WATER USE AND DISCHARGE RATES FOR ACID WASH AND RINSE WATER

(1/kkg of mercury washed and rinsed)

Plant Code	Percent Recycle	Production Normalized Water Use	Production Normalized Discharge Flow
· · · · · · · · · · · · · · · · · · ·			
1161	0	2.0	2.0

Table V-3

WATER USE AND DISCHARGE RATES FOR FURNACE WET AIR POLLUTION CONTROL

(1/kkg of mercury processed through furnace)

Plant Code	Percent Recycle	Production Normalized Water Use	Production Normalized Discharge Flow	
1011	100	Unknown	0	
	•			

SECTION VI

SELECTION OF POLLUTANT PARAMETERS

Although wastewater from secondary mercury facilities was not sampled, it should have similar characteristics to wastewater from a primary mercury facility. Analytical data from a primary mercury plant are presented in Section V of the supplement for primary precious metals and mercury. This section examines that data and discusses the selection or exclusion of pollutants for potential limitation.

The basis for the regulation of toxic and other pollutants along with a discussion of each pollutant selected for potential limitation is discussed in Section VI of the General Development Document. That discussion provides information concerning the nature of the pollutant (i.e., whether it is a naturally occurring substance, processed metal, or a manufactured compound); general physical properties and the form of the pollutant; toxic effects of the pollutant in humans and other and behavior of the pollutant in POTW animals; at the concentrations expected in industrial discharges.

The discussion that follows presents and briefly discusses the selection of conventional pollutants for effluent limitations. Also described is the analysis that was performed to select or exclude priority pollutants for further consideration for limitations and standards. Pollutants will be considered for limitation if they are present in concentrations treatable by the technologies considered in this analysis. The treatable concentrations used for the priority metals were the long-term performance values achievable by chemical precipitation, sedimentation, and filtration.

CONVENTIONAL POLLUTANT PARAMETERS

This study examined samples for the secondary mercury subcategory for three conventional pollutant parameters (oil and grease, total suspended solids, and pH).

CONVENTIONAL POLLUTANT PARAMETERS SELECTED

The conventional pollutants or pollutant parameters selected for limitation in this subcategory are:

total suspended solids (TSS) pH

-

No nonconventional pollutants or pollutant parameters are selected for limitation in this subcategory.

TSS are expected to be present in secondary mercury wastewaters in concentrations exceeding that achievable by identified

treatment technologies (2.6 mg/l). In the primary mercury plant's wastewater, TSS concentrations ranged from 4 mg/l to 3,700 mg/l. Furthermore, most of the specific methods used to remove toxic metals do so by converting these metals to precipitates, and these toxic-metal-containing precipitates should not be discharged. Meeting a limitation on total suspended solids helps ensure that removal of these precipitated toxic metals has been effective. For these reasons, total suspended solids are selected for limitation in this subcategory.

Spent battery electrolyte and acid wash and rinse water are expected to have pH values less than pH 7.5, which is outside the pH 7.5 to 10 range considered desirable for discharge to receiving waters. Four of the six primary mercury wastewater samples had pH values between 2.3 and 2.6. Many deleterious effects are caused by extreme pH values or rapid changes in pH. Also, effective removal of toxic metals by precipitation requires careful control of pH. Since pH control within the desirable limits is readily attainable by available treatment, pH is selected for limitation in this subcategory.

TOXIC PRIORITY POLLUTANTS

Raw wastewater from secondary mercury plants was not sampled, however, raw wastewater samples from the primary mercury industry should be representative of the wastewater from secondary mercury plants. These data provide the basis for the categorization of specific pollutants, as discussed below. Treatment plant samples were not considered in the frequency count.

TOXIC POLLUTANTS NEVER DETECTED

The priority pollutants listed in Table VI-1 (page 2891) were not detected or not analyzed for in any raw wastewater samples; therefore, they are not selected for consideration in establishing limitations.

TOXIC POLLUTANTS NEVER FOUND ABOVE THEIR ANALYTICAL QUANTIFICATION CONCENTRATION

The priority pollutants listed below were never found above their analytical quantification concentration in any raw wastewater samples; therefore, they are not selected for consideration in establishing limitations.

- 114. antimony
- 117. beryllium
- 119. chromium (Total)
- 120. copper
- 124. nickel
- 125. selenium
- 126. silver

TOXIC POLLUTANTS PRESENT BELOW CONCENTRATIONS ACHIEVABLE BY TREATMENT

The pollutants listed below are not selected for consideration in establishing limitations because they were not found in any raw wastewater samples above concentrations considered achievable by existing or available treatment technologies. These pollutants are discussed individually following the list.

115. arsenic 118. cadmium

Arsenic was detected above the quantification concentration but below the treatable concentration in one sample analyzed. The sample contained 0.32 mg/l arsenic which is below the 0.34 mg/l treatable concentration. Therefore, arsenic is not selected for limitation.

Cadmium was detected above the quantification concentration in one sample analyzed. The sample indicated a cadmium concentration of 0.04 mg/l. This is below the 0.049 mg/l treatable concentration, thus cadmium is not selected for limitation.

TOXIC POLLUTANTS SELECTED FOR FURTHER CONSIDERATION IN ESTABLISHING LIMITATIONS AND STANDARDS

The priority pollutants listed below are selected for further consideration in establishing limitations and standards for this subcategory. The toxic pollutants selected for further consideration for limitation are each discussed following the list.

122. lead 123. mercury 127. thallium 128. zinc

Lead was detected above its treatable concentration of 0.08 mg/l in one sample. This sample indicated a lead concentration of 22 mg/l. Lead is also expected to be present in wastewaters from this industry because it is a contaminant of the raw materials used for mercury recovery. Thus, lead is selected for further consideration for limitation.

Mercury was present above treatable concentrations in the wastewater from this industry. One sample showed a concentration of 360 mg/l of mercury. In the recovery of secondary mercury, mercury contacts various aqueous streams in which it is partially soluble. For these reasons, mercury is selected for further consideration for limitation.

Thallium was detected above its treatable concentration of 0.34 mg/l in one sample. This sample indicated 0.61 mg/l of thallium. Thus, thallium is selected for consideration for limitation.

VI

Zinc was detected above treatable concentrations in one sample indicating 0.73 mg/l. Treatable concentration for zinc is 0.23 mg/l. Zinc is also expected to be present in wastewaters from this industry because it is present in batteries which are used as raw materials for secondary mercury recovery. Therefore, zinc is selected for further consideration for limitation.

SECONDARY MERCURY SUBCATEGORY

SECT - VI

TABLE VI-1

TOXIC POLLUTANTS NEVER DETECTED

.

1.	acenaphthene*
2.	acrolein*
3.	acrylonitrile*
4.	benzene*
5.	benzidine*
6.	carbon tetrachloride (tetrachloromethane)*
7.	chlorobenzene*
8.	1,2,4-trichlorobenzene*
9.	hexachlorobenzene*
10.	1,2-dichloroethane*
11.	1,1,1-trichloroethane*
12.	hexachloroethane*
13.	1,1-dichloroethane*
14.	1,1,2-trichloroethane*
15.	1,1,2,2-tetrachloroethane*
16.	chloroethane*
17.	bis (chloromethyl) ether (Deleted)*
18.	bis (2-chloroethyl) ether*
19.	2-chloroethyl vinyl ether (mixed)*
20.	2-chloronaphthalene*
21.	2,4,6-trichlorophenol*
22.	parachlorometa cresol*
23.	chloroform (trichloromethane)*
24.	2-chlorophenol*
25.	1,2-dichlorooenzene*
26.	1,3-dichlorobenzene*
27.	1,4'-dichlorobenzene*
28.	3.3 -dichlorobenzidine*
29.	1.1-dichloroethylene*
30.	1,2-trans-dichloroethylene*
31.	2,4-dichlorophenol*
32.	1,2-dichloropropane*
33.	1.2-dichloropropylene (1.3-dichloropropene)*
34.	2,4-dimethylphenol*
35.	2,4-dinitrotoluene*
36.	2,6-dinitrotoluene*
37.	1,2-diphenylhydrazine*
38.	ethylbenzene*
39.	fluoranthene*
40.	4-chlorophenyl phenyl ether*
41.	4-bromopĥenyl pĥenyl ether*
42.	bis(2-chloroisopropyl) ether*
43.	bis(2-choroerhoxy) methane*
44.	methylene chloride (dichloromethane)*
45.	methyl chloride (chloromethane)*
46.	methyl bromide (bromomethane)*
	hunnetten (tuthunnethene)*

47. bromoform (tribromomethane)*

TABLE VI-1 (Continued)

TOXIC POLLUTANTS NEVER DETECTED

trichlorofluoromethane (Deleted)* 49. dichlorodifluoromethane (Delered)* 50. chlorodibromomethane* 51. 52. hexachlorooutadiene* hexachlorocyclopentadiene* 53. 54. isophorone* 55. naphthalene* 56. nitrobenzene* 57. 2-nitrophenol* 58. 4-nitrophenol* 2,4-dinitrophenol* 59. 4,6-dinitro-o-cresol* 60. N-nitrosodimethylamine* 61. 62. N.nitrosodiphenylamine* 63. N-nitrosodi-n-propylamine* 64. pentachlorophenol* 65. phenol* bis(2-ethylhexyl) phthalate* 66. butyl benzyl phthalate* 67. di.n-butyl phthalate* 68. 69. di-n-octyl phthalate* diethyl phthalate* 70. dimethyl phthalate* 71. benzo (a)anthracene (1,2-benzanthracene)* 72. 73. benzo (a)pyrene (3,4-benzopyrene)* 3,4-benzofluoranthene* 74. benzo(k)fluoranthane (ll,l2-benzofluoranthene)* 75. 76. chrysene* 77. acenaphthylene* 78. anthracene* 79. benzo(ghi)perylene (1,11-benzoperylene)* 80. fluorene* 81. phenanthrene* 82. dibenzo (a,h)anthracene (1,2.5,6-dibenzanthracene)* 83. indeno (1.2,3-cd)pyrene (w,e,-o-phenylenepyrene)* 84. pyrene* tetrachloroethylene* 85. toluene* 86. 87. trichloroethylene* vinyl chloride (chloroethylene)* 88. 89. aldrin* 90. dieldrin* 91. chlordane (technical mixture and metabolites)* 92. $4, 4' - DDT^*$ 4,4'-DDE(p,p'DDX)*93.

94. 4,4'-DDD(p,p'TDE)*

48.

dichlorobromomethane*

TABLE VI-1 (Continued)

TOXIC POLLUTANTS NEVER DETECTED

Alpha-endosulfan*
Beta-endosulfan*
endosulfan sulfate*
endrin*
endrin aldehyde*
heptachlor*
heptachlor epoxide*
Alpha-BHC*
Beta-BHC-*
Gamma-BHC (lindane)*
Delta-BHC*
PCB-1242 (ArochIor 1242)*
PCB-1254 (Arochlor 1254)*
PCB-1221 (Arochlor 1221)*
PCB-1Z?Z (Arochlor 1232)*
PCB-12?8 (Arochlor 1248)*
PCB-1260 (Arochlor 1260)*
PCB-1016 (Arochlor 1016)*
toxaphene*
asbestos (Fibrous)
cyanide (Total)*
2,3,7,8-tetra chlorodibenzo-p-dioxin (TCDD)

*We did not analyze for these pollutants in samples of raw wastewater from this subcategory. These pollutants are not believed to be present based on the Agency's best engineering judgment which includes consideration of raw materials and process operations.

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SECTION VII

CONTROL AND TREATMENT TECHNOLOGIES

The preceding sections of this supplement discussed the sources, flows, and characteristics of the wastewaters from secondary mercury plants. This section summarizes the description of these wastewaters and indicates the treatment technologies which are currently practiced in the secondary mercury subcategory for each waste stream. Secondly, this section presents the control and treatment technology options which were examined by the Agency for possible application to the secondary mercury subcategory.

CURRENT CONTROL AND TREATMENT PRACTICES

This section presents a summary of the control and treatment technologies that are currently being applied to each of the sources generating wastewater in this subcategory. As discussed in Section V, wastewater associated with the secondary mercury subcategory is characterized by the presence of the toxic metal pollutants and suspended solids. This analysis is supported by the raw (untreated) wastewater data presented for primary mercury sources as well as raw materials and production processes as shown in Section VI. Generally, these pollutants are present in each of the waste streams at concentrations above treatability, and these waste streams are commonly combined for treatment. Construction of one wastewater treatment system for combined. treatment allows plants to take advantage of economic scale and in some instances to combine streams of different alkalinity to reduce treatment chemical requirements. No plants in this subcategory currently treat their wastewater. One plant emplovs contractor disposal of their wastewater, and one plant employs 100 percent recycle of scrubber liquor. The options selected for consideration for NSPS and pretreatment based on combined treatment of these compatible waste streams will be summarized toward the end of this section.

SPENT BATTERY ELECTROLYTE

Mercury may be reclaimed from recycled mercuric oxide battery cells. Before distilling the mercury contained in the battery, the spent electrolyte must be drained. One plant processes recycled batteries, and has their spent battery electrolyte hauled away by an approved contractor.

ACID WASH AND RINSE WATER

After recovering mercury from recycled batteries by distillation, the mercury product may be further purified. Purification is effected by washing the mercury with dilute nitric acid, and then rinsing it with water. One plant generates an acid wash and rinse wastewater stream in this manner, and disposes of it by having a contractor haul it away.

FURNACE WET AIR POLLUTION CONTROL

Mercury may be reclaimed from scrap such as thermometers, switches, filters, controls, amalgams, and soil samples by vaporizing the mercury in a furnace. After recovering the vaporized mercury by condensation, the air emissions from the furnace may be controlled with a wet scrubber. One plant practices furnace wet air pollution control, and recycles 100 percent of the scrubber liquor. There is no liquid effluent from this process.

CONTROL AND TREATMENT OPTIONS

The Agency examined two control and treatment technology options that are applicable to the secondary mercury subcategory. The options selected for evaluation represent a combination of endof-pipe treatment technologies. The effectiveness of these technologies is detailed in Section VII of Vol. I.

OPTION A

Option A for the secondary mercury subcategory requires control and treatment technologies to reduce the discharge of wastewater pollutant mass.

The Option A treatment scheme consists of chemical precipitation and sedimentation technology. Specifically, lime or some other alkaline compound is used to precipitate metal ions as metal hydroxides. The metal hydroxides and suspended solids settle out and the sludge is collected. Vacuum filtration is used to dewater sludge.

OPTION C

Option C for the secondary mercury subcategory consists of all control and treatment requirements of Option A (chemical precipitation and sedimentation) plus multimedia filtration technology added at the end of the Option A treatment scheme. Multimedia filtration is used to remove suspended solids, including precipitates of metals, beyond the concentration attainable by gravity sedimentation. The filter suggested is of the gravity, mixed media type, although other forms of filters, such as rapid sand filters or pressure filters would perform satisfactorily. The addition of filters also provides consistent removal during periods in which there are rapid increases in flows or loadings of pollutants to the treatment system.

SECTION VIII

COSTS, ENERGY, AND NONWATER QUALITY ASPECTS

This section presents a summary of compliance costs for the secondary mercury subcategory and a description of the treatment options and subcategory-specific assumptions used to develop these estimates. Together with the estimated pollutant removal performance presented in Section XI of this supplement, these cost estimates provide a basis for evaluating each regulatory option. These cost estimates are also used in determining the probable economic impact of regulation on the subcategory at different pollutant discharge levels.

there are no existing direct or indirect dischargers in this As subcategory, plant-by-plant compliance cost estimation was not Rather, based on analysis of the appropriate. production data from plants presently in sampling ርhe 👘 subcategory, compliance costs for new source model plants were estimated for of the considered treatment options. each Since no new information or data were received from industry comments between proposal and promulgation, new source model plant compliance costs for promulgation are the same as for proposal.

In addition, this section addresses nonwater quality environmental impacts of wastewater treatment and control alternatives, including air pollution, solid wastes, and energy requirements, which are specific to the secondary mercury subcategory.

TREATMENT OPTIONS FOR NEW SOURCES

As discussed in Section VII, two treatment options have been developed and considered in proposing standards for the secondary mercury subcategory. These options are summarized below and schematically presented in Figures XI-1 and XI-2 Pages 2914 - 2915).

OPTION A

The Option A treatment scheme consists of chemical precipitation and sedimentation technology.

OPTION C

Option C for the secondary mercury subcategory consists of all control and treatment requirements of Option A (chemical precipitation and sedimentation) plus multimedia filtration technology added at the end of the Option A treatment scheme.

COST METHODOLOGY

A detailed discussion of the methodology used to develop the

compliance costs is presented in Section VIII of Vol. I. Projected compliance costs for new source model plants in the secondary mercury subcategory have been determined and are presented in the administrative record supporting this regulation. The costs developed for the promulgated regulation are presented in Table VIII-1 (page 2901) for model new sources in the secondary mercury subcategory.

Each of the general assumptions used to develop compliance costs is presented in Section VIII of Vol. I. Each subcategory contains a unique set of waste streams requiring certain subcategoryspecific assumptions to develop compliance costs. Three major assumptions relevant to the cost estimation of new source model plants in the secondary mercury subcategory are discussed briefly below.

- (1) Operating hours are assumed to be 2,000 hours per year (8 hrs/day, 250 days/yr).
- (2) Treatment of the furnace wet air pollution control wastewater stream is not included in the cost estimate because it is considered a process step in the recovery of mercury from furnace scrubber liquor.
- (3) Pollutant concentration data for the two wastewater streams included in the treatment scheme were transferred from the calciner venturi scrubber in the primary mercury subcategory.

NONWATER QUALITY ASPECTS

Nonwater quality impacts specific to the secondary mercury subcategory, including energy requirements, solid waste, and air pollution are discussed below.

ENERGY REQUIREMENTS

The methodology used for determining the energy requirements for the various options is discussed in Section VIII of the General Development Document. Energy requirements for new source model plants are estimated at 2,300 kwh/yr for Option A and 3,500 kwh/yr for Option C. Option C energy requirements increase over those for Option A because filtration is being added as an endof-pipe treatment technology. Both options represent less than one percent of a typical existing plant's energy usage. It is therefore expected that the energy requirements of the treatment options considered will have no significant impact on total plant energy consumption for new sources.

SOLID WASTE

Sludge generated in the secondary mercury subcategory is due to the precipitation of metal hydroxides and carbonates using lime. Sludges associated with the secondary mercury subcategory will necessarily contain quantities of toxic metal pollutants. Wastes

generated by secondary metal industries can be regulated as hazardous. However, the Agency examined the solid wastes that would be generated at secondary nonferrous metals manufacturing plants by the suggested treatment technologies and believes they not hazardous wastes under the Agency's are regulations implementing Section 3001 of the Resource Conservation and Recovery Act. None of the secondary mercury wastes are listed specifically as hazardous, nor are they likely to exhibit a characteristic of hazardous waste. This judgment is made based the recommended technology of lime precipitation on and filtration. By the addition of a small (5-10%) excess of lime during treatment, similar sludges, specifically toxic metalbearing sludges, generated by other industries such as the iron and steel industry passed the Extraction Procedure (EP) toxicity 40 CFR \$261.24. Thus, the Agency believes that the test. See wastewater sludges will similarly not be EP toxic if the recommended technology is applied.

Although it is the Agency's view that solid wastes generated as a result of these guidelines are not expected to be hazardous generators of these wastes must test the waste to determine if the wastes meet any of the characteristics of hazardous waste (see 40 CFR 262.11).

If these wastes should be identified or are listed as hazardous, they will come within the scope of RCRA's "cradle to grave" hazardous waste management program, requiring regulation from the point of generation to point of final disposition. EPA's standards would require generators of generator hazardous nonferrous metals manufacturing wastes to meet containerization, labeling, recordkeeping, and reporting requirements; if plants dispose of hazardous wastes off-site, they would have to prepare a manifest which would track the movement of the wastes from the generator's premises to a permitted off-site treatment, storage, or disposal facility. See 40 CFR 262.20 45 FR 33142 (May 19, 1980), as amended at 45 FR 86973 (December 31, 1980). The transporter regulations require transporters of hazardous wastes to comply with the manifest system to assure that the wastes are delivered to a permitted facility. See 40 CFR 263.20 45 FR 33151 (May 19, 1980), as amended at 45 FR 86973 (December 31, 1980). Finally, RCRA regulations establish standards for hazardous waste treatment, storage, and disposal facilities allowed to receive such wastes. See 40 CFR Part 464 46 FR 2802 (January 12, 1981), 47 FR 32274 (July 26, 1982).

Even if these wastes are not identified as hazardous, they still must be disposed of in compliance with the Subtitle D open dumping standards, implementing 4004 of RCRA. See 40 FR 53438 (September 13, 1979). It is estimated that a new source model plant in the secondary mercury subcategory would generate an estimated 12 kg/yr of sludge when implementing the promulgated NSPS treatment technology, based on a production level of 50 metric tons of mercury per year. The Agency has calculated as part of the costs for wastewater treatment the cost of hauling and disposing of solid wastes.

AIR POLLUTION

There is no reason to believe that any substantial air pollution problems will result from implementation of chemical precipitation, sedimentation, and multimedia filtration. These technologies transfer pollutants to solid waste and are not likely to transfer pollutants to air.

TABLE VIII-1

COST OF COMPLIANCE FOR NEW SOURCE MODEL PLANTS IN THE SECONDARY MERCURY SUBCATEGORY*

(March, 1982 Dollars)

Option	Total Required Capital Cost	Total Annual Cost	
A	1,237	3,070	
С	3,162	4,530	

*Based on production of 50 metric tons of mercury per year.

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SECTION IX

BEST PRACTICABLE CONTROL TECHNOLOGY CURRENTLY AVAILABLE

The wastewater handling practices of the plants within the secondary mercury subcategory were studied. BPT was found to be not applicable to this industrial subcategory. Existing performance of plants in the secondary mercury subcategory is such that no discharge of process wastewater is presently practiced. This is achieved by 100 percent recycle on-site or by contractor disposal of process wastewater, or is a result of a production process that generates no process wastewater. Since there are no discharge from secondary mercury producers, BPT and mass limitations, with their corresponding treatment BAT technologies, need not be promulgated for this subcategory. Rather, the secondary mercury subcategory will be regulated under New Source Performance Standards in Section XI, and Pretreatment Standards for New Sources in Section XII.

SECONDARY MERCURY SUBCATEGORY SECT - X

SECTION X

BEST AVAILABLE TECHNOLOGY ECONOMICALLY ACHIEVABLE

As described in Section IX, BAT is not applicable to the secondary mercury subcategory because no plants in the data base discharge process wastewater. Regulation of the secondary mercury subcategory is covered in Section XI under New Source Performance Standards, and Section XII under Pretreatment Standards for New Sources.

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SECTION XI

NEW SOURCE PERFORMANCE STANDARDS

This section describes the technologies for treatment of wastewater from new sources and presents mass discharge standards for regulated pollutants for NSPS in the secondary mercury subcategory, based on the selected treatment technology.

TECHNICAL APPROACH TO NSPS

New source performance standards are based on the most effective and beneficial technologies currently available. The Agency reviewed and evaluated a wide range of technology options, and elected to examine two technology options, applied to combined wastewater streams, which could be applied to the secondary mercury subcategory as alternatives for the basis of NSPS.

Treatment technologies considered for the NSPS options are summarized below:

OPTION A (Figure XI-1, page 2914) is based on:

o Chemical precipitation and sedimentation

OPTION C (Figure XI-2 page 2915) is based on:

- o Chemical precipitation and sedimentation
- o Multimedia filtration

As explained in Section IV, the secondary mercury subcategory has been subdivided into three potential wastewater sources. Since the water use, discharge rates, and pollutant characteristics of each of these wastewaters is potentially unique, effluent limitations will be developed for each of the three subdivisions.

For each of the building blocks, a specific approach was followed for the development of NSPS. The first requirement to calculate limitations is to account for production and flow these variability from plant to plant. Therefore, a unit of production or production normalizing parameter (PNP) was determined for each wastewater stream which could then be related to the flow from the process to determine a production normalized flow. Selection of the PNP for each process element is discussed in Section IV. Each process within the subcategory was then analyzed to determine (1) which subdivisions were present, (2) the specific flow rates generated for each subdivision, and (3) the specific production normalized flows for each subdivision. This analysis is discussed in detail in Section V. Nonprocess wastewaters such as rainfall runoff and noncontact cooling water are not considered in the analysis.

Production normalized flows for each subdivision were analyzed to determine which flow was to be used as part of the basis for

NSPS. The selected flow (sometimes referred to as the NSPS regulatory flow or NSPS discharge flow) reflects the water use controls which are practiced within the subcategory. The NSPS normalized flow is based on the average of all applicable data. Nothing was found to indicate that the wastewater flows and characteristics of new plants would not be similar to those from existing plants, since the processes used by new sources are not expected to differ from those used at existing sources.

For the development of NSPS, mass loadings were calculated for each wastewater source or building block. This calculation was made on a stream-by-stream basis, primarily because plants in this subcategory may perform one or more of the operations in various combinations. The mass loadings (milligrams of pollutant per metric ton of production unit - mg/kkg) were calculated by multiplying the NSPS normalized flow (l/kkg) by the treatment effectiveness concentration using the NSPS treatment system (mg/l) for each pollutant parameter to be limited under NSPS.

The mass loadings which are allowed under NSPS for each plant will be the sum of the individual mass loadings for the various building blocks which are found at particular plants. Accordingly, all the wastewater generated within a plant may be combined for treatment in a single or common treatment system, but the effluent limitations for these combined wastewaters are based on the various wastewater sources which actually contribute to the combined flow. This method accounts for the variety of combinations of wastewater sources and production processes which may be found at secondary mercury plants.

The Agency usually establishes wastewater limitations in terms of mass rather than concentration. This approach prevents the use of dilution as a treatment method (except for controlling pH). The production normalized wastewater flow (1/kkg) is a link between the production operations and the effluent limitations. The pollutant discharge attributable to each operation can be calculated from the normalized flow and effluent concentration achievable by the treatment technology and summed to derive an appropriate limitation for each plant.

INDUSTRY COST AND POLLUTANT REMOVAL ESTIMATES

As one means of evaluating each technology option, EPA developed estimates of the pollutant removals and the compliance costs associated with each option. The methodologies are described below. For both pollutant removals and compliance costs, estimates reported at promulgation are the same as those reported at proposal. This is because new information or data were not received from industry during the comment period between proposal and promulgation.

POLLUTANT REMOVAL ESTIMATES

Since there are no existing discharging plants in the secondary

mercury subcategory, the pollutant removal analysis was carried out for new source model plants.

A complete description of the methodology used to calculate the estimated pollutant removal achieved by the application of the various treatment options is presented in Section X of Vol. I. short, sampling data used to characterize the major In waste streams considered for regulation was production normalized for each unit operation (i.e., mass of pollutant generated per mass product manufactured). This value, referred to as the raw of was used to estimate the mass of toxic pollutants waste within the secondary mercury subcategory. The generated . removal estimates were calculated for each plant bv pollutant estimating the total mass of each pollutant in the first untreated wastewater. This was calculated by multiplying the raw waste values by the corresponding new source model plant production value for that stream and then summing these values for each pollutant for every stream generated by the plant.

the volume of wastewater discharged after the application Next, of each treatment option was estimated for each operation at each plant by comparing the actual discharge to the regulatory flow. smaller of the two values was selected and summed with the The The mass of pollutant discharged was then other plant flows. estimated by multiplying the achievable concentration values attainable with the option (mg/l) by the estimated volume of of process wastewater discharged by the subcategory. The mass pollutant removed is the difference between the estimated mass of pollutant generated within the subcategory and the mass of pollutant discharged after application of the treatment option. The pollutant removal estimates for the new source model plant in the secondary mercury subcategory are presented in Table XI-1 (page 2911).

COMPLIANCE COSTS

In estimating subcategory-wide compliance costs, the first step was to develop a cost estimation model, relating the total costs with installation and operation of wastewater associated treatment technologies to plant process wastewater discharge. EPA applied the model to each plant. The plant's investment and operating costs are determined by what treatment it has in place and by its individual process wastewater discharge flow. discussed above, this flow is either the actual or the As BDT regulatory flow, whichever is lesser. The final step was to annualize the capital costs, and to sum the annualized capital and the operating and maintenance costs for each plant, costs, yielding the cost of compliance for the subcategory. The compliance costs associated with each option are presented in (page 2912) for new source model plants mercury subcategory. These costs were us in the Table XI-2 These costs were used in secondary assessing economic achievability.

NSPS OPTION SELECTION - PROPOSAL

EPA selected Option C for the proposed NSPS, which includes chemical precipitation, sedimentation, and filtration.

The pollutants proposed for limitation under NSPS were lead, mercury, total suspended solids, and pH. The estimated capital cost of proposed NSPS was \$3,162 and the estimated annual cost was \$4,530 (1982 dollars) for new facilities.

NSPS OPTION SELECTION - PROMULGATION

The Agency received no industry comments on the proposed regulation for the secondary mercury subcategory. Therefore, the promulgated regulation is equivalent to the proposed regulation for the secondary mercury subcategory.

EPA is promulgating NSPS based on the best available demonstrated technology for the secondary mercury subcategory which is equivalent to Option C (chemical precipitation, sedimentation, and multimedia filtration). This selection is based on an economic analysis of the two NSPS options and their impact on the cost of building new production plants within the scope of this subcategory. ERA believes the promulgated NSPS are economically achievable, and that they are not a barrier to entry of new plants into this subcategory. The estimated capital cost of promulgated NSPS for new source model plants is \$3,162, and the estimated annual cost is \$4,530 (1982 dollars), based on production of 50 metric tons of mercury per year. The end-ofpipe treatment configuration for Option C is presented in Figure XI-2 (page 2915).

WASTEWATER DISCHARGE RATES

A NSPS discharge rate is calculated for each subdivision based on the average of the flows of the existing plants, as determined from analysis of dcp. The discharge rate is used with the achievable treatment concentrations to determine NSPS. Since the discharge rate may be different for each wastewater source, separate production normalized discharge rates for each of the three wastewater sources are discussed below and summarized in Table XI-3 (page 2912). The discharge rates are normalized on a production basis by relating the amount of wastewater generated to the mass of the product which is produced by the process associated with the waste stream in question. These production normalizing parameters, or PNPs, are also listed in Table XI-3.

Section V of this document further describes the discharge flow rates and presents production normalized water use and discharge rates for each plant by subdivision in Tables V-1 through V-3 (pages 2885 - 2886). For all subdivisions, the proposed and promulgated NSPS discharge rates are equivalent. These rates are discussed below.

SPENT BATTERY ELECTROLYTE

The proposed and promulgated NSPS wastewater discharge rate for spent battery electrolyte is 106 liters per kkg of mercury produced from batteries. This rate is allocated only for those plants which drain electrolyte from mercuric oxide batteries prior to recovering mercury. Water use and wastewater discharge rates are presented in Table V-1 (page 2885). One plant drains spent battery electrolyte, and generates 106 1/kkg.

ACID WASH AND RINSE WATER

The proposed and promulgated NSPS wastewater discharge rate for acid wash and rinse water is 2.0 liters per kkg of mercury washed and rinsed. This rate is allocated only for those plants which further purify their mercury product by washing with acid and then rinsing with water. Water use and wastewater discharge rates are presented in Table V-2 (page 2885). One plant further purifies their mercury product in this manner, and generates 2.0 1/kkg.

FURNACE WET AIR POLLUTION CONTROL

No NSPS wastewater discharge rate for furnace wet air pollution control is provided based on 100 percent recycle of furnace scrubber water, as demonstrated at the one plant operating this process. This is shown in Table V-3 (page 2886).

REGULATED POLLUTANT PARAMETERS

The raw wastewater concentrations from individual operations and the subcategory as a whole were examined to select certain pollutant parameters for limitation. This examination and evaluation was presented in Section VI. A total of four pollutants or pollutant parameters are selected for limitation under NSPS and are listed below:

122.	lead
123.	mercury
	TSS
	рН

The Agency has chosen not to regulate all four priority pollutants selected in Section VI for further consideration.

The high cost associated with analysis for priority metal pollutants has prompted EPA to develop an alternative method for regulating and monitoring priority pollutant discharges from the nonferrous metals manufacturing category. Rather than developing specific effluent mass limitations and standards for each of the priority metals found in treatable concentrations in the raw wastewater from a given subcategory, the Agency is promulgating effluent mass limitations only for those pollutants generated in the greatest quantities as shown by the pollutant removal analysis. By establishing limitations and standards for certain toxic metal pollutants, dischargers will attain the same degree of control over toxic metal pollutants as they would have been required to achieve had all the priority metal pollutants been directly limited.

This approach is technically justified since the treatable concentrations used for chemical precipitation and sedimentation technology are based on optimized treatment for concomitant multiple metals removal. Thus, even though metals have somewhat different theoretical solubilities, they will be removed at very nearly the same rate in a chemical precipitation and sedimentation treatment system operated for multiple metals removal. The mass limits established for lead and mercury will ensure that thallium and zinc, the other two priority metals selected for further consideration, will be adequately removed by a lime and settle unit.

NEW SOURCE PERFORMANCE STANDARDS

The treatable concentrations achievable by application of the promulgated NSPS are discussed in Section VII of Vol. I and summarized there in Table VII-21 (page 248). These treatable concentrations (both one day maximum and monthly aver-age values) are multiplied by the NSPS normalized discharge flows summarized in Table XI-3 (page 2912) to calculate the mass of pollutants allowed to be discharged per mass of product. The results of these calculations in milligrams of pollutant per kilogram of product represent the NSPS effluent standards and are presented in Table XI-4 (page 2913) for each individual waste stream.

Table XI-1

POLLUTANT REMOVAL ESTIMATES FOR NEW SOURCE MODEL PLANTS*

Pollutant	Raw Waste (kg/yr)	Option A Discharge (kg/yr)	Option A Removed (kg/yr)	Option C Discharge (kg/yr)	Option C Removed (kg/yr)
Antimony	0	0	0	0	0
Arsenic	0.0017 0.0002	0.0017	U U	0.0017 0.0002	0
Cadmium Chromium (Total)	0	0	Ŏ	0	Ő
Copper	0 0.0119	0.0006	0 0.0112	0.0004	0.0114
Lead Mercury	1.9440	0.0003	1.9437	0.0002	1.9438
Nickel	0	0	0	0	0
Selenium Silver	0	Ŭ Ŭ	0	Ŏ	Ŏ
Thallium	0.0033 0.0039	0.0027 0.0018	0.0006	0.0018	0.0015 0.0027
Zinc	0.003%	0.0018	· ·		
TOTAL PRIORITY POLLUTANTS	1.9651	0.0074	1.9577	0.0056	1.9594
TSS	0.4320	0.0646	0.3674	0.0140	0.4180
TOTAL CONVENTIONALS	0.4320	0.0646	0.3674	0.0140	0.4180
TOTAL POLLUTANTS	2.3971	0.0720	2.3250	0.0196	2.3774

*Based on production of 50 metric tons mercury per year.

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SECONDARY MERCURY SUBCATEGORY SECT 1 XI

TABLE XI-2

COST OF COMPLIANCE FOR NEW SOURCE MODEL PLANTS IN THE SECONDARY MERCURY SUBCATEGORY*

(March, 1982 Dollars)

Option	Total Required Capital Cost	Total Annual Cost
A	1,237	3,070
С	3,162	4,530

*Based on production of 50 metric tons of mercury per year.

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TABLE XI-3

NSPS WASTEWATER DISCHARGE RATES FOR THE SECONDARY MERCURY SUBCATEGORY

Wastewater Stream	NSPS Normalized Discharge Rate l/kkg gal/ton		Production Normalized Parameter
Spent battery electrolyte	106	25.5	mercury produced from batteries
Acid wash and rinse water	2.0	0.5	mercury washed and rinsed
Furnace wet air pollution	0	0	mercury control processed through furnace

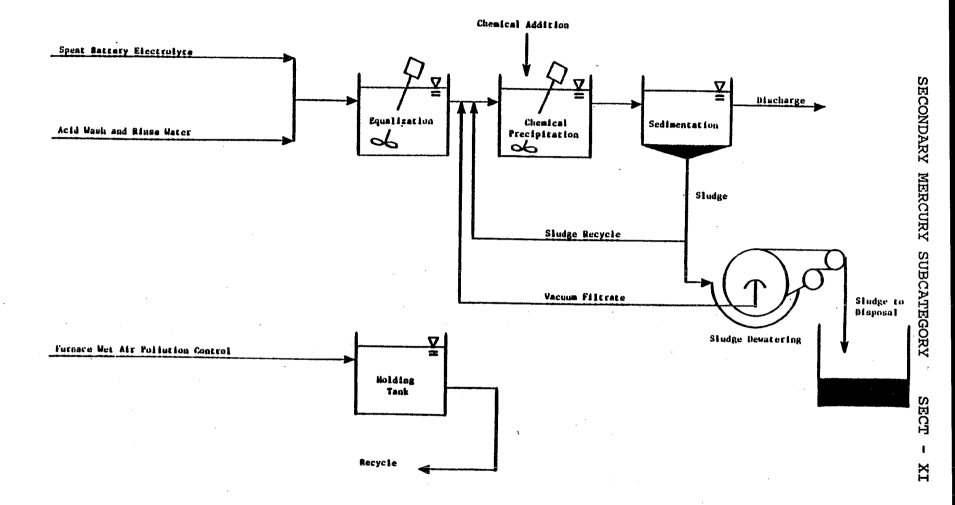
TABLE XI-4

NSPS FOR THE SECONDARY MERCURY SUBCATEGORY

(a) <u>Spent</u> <u>Battery</u> <u>Electrolyte</u> NSPS

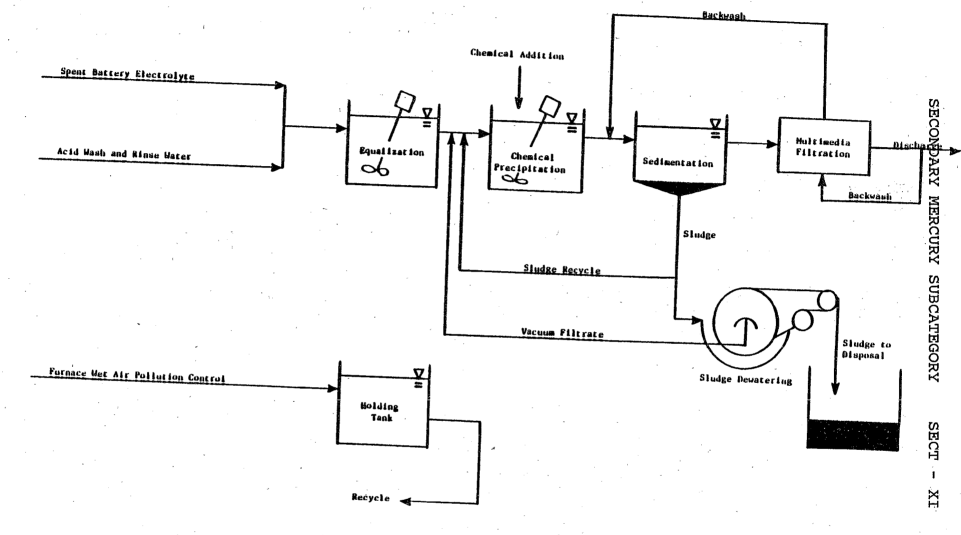
Pollutant	or	Maxi	mum for	Maximum	for
	property		one day	monthly	average
Ollucanc	propercy	any	one day		
mg/kg	(lb/millio	n lbs)	of mercur	y produced	from batterie
Lead			0.030		0.014
			0.016	· · ·	0.006
Mercury			0.148		0.065
Thallium	1				0.045
Zinc			0.108		
TSS	8.9 ± 1.		1.590	7 5 +0 10 0	1.272) at all times
Hq*	WICH	in the	range or	7.5 10 10.0	
(b) Acid 1	Wash and Ri	nse Wat	er NSPS		•
D) Actu			· · ·		
Pollutant			mum for	Maximum	
ollutant	property	any	one day	monthly	average
				· · · · · · · · · · · · · · · · · · ·	}
mg	/kg (lb/mil	lion lb	s) of mer	cury washe	d and rinsed
- hT a a d	-	۰ ۶	0.00056		0.00026
*Lead			0.00030	٠	0.00012
*Mercury	· .		0.00280		0.00122
Thallium					
Zinc			0.00204		0.00084
Zinc *TSS			0.00204 0.03000) () at all	0.00084 0.02400
Zinc *TSS	thin the ra	nge of	0.00204 0.03000).0 at all	0.00084 0.02400
Zinc *TSS *pH Wi	thin the ra	<u>,</u>	0.00204 0.03000 7.5 to 10		0.000 84 0.02400
Zinc *TSS *pH Wi (c) <u>Furna</u>	thin the ra ce Wet <u>Air</u>	Polluti	0.00204 0.03000 7.5 to 10	<u>51</u>	0.00084 0.02400 times
Zinc *TSS *pH Wi (c) <u>Furna</u> Pollutant	thin the ra <u>ce Wet Air</u> or	Polluti Maxi	0.00204 0.03000 7.5 to 10 on <u>Contro</u> mum for	<u>ol</u> Maximum	0.00084 0.02400 times for
Zinc *TSS *pH Wi (c) <u>Furna</u> Pollutant	thin the ra ce Wet <u>Air</u>	Polluti Maxi	0.00204 0.03000 7.5 to 10	<u>ol</u> Maximum	0.00084 0.02400 times
Zinc *TSS *pH Wi (c) <u>Furna</u> Pollutant	thin the ra <u>ce Wet Air</u> or property	Polluti Maxi any	0.00204 0.03000 7.5 to 10 on <u>Contro</u> mum for one day	<u>ol</u> Maximum monthly	0.00084 0.02400 times for average
Zinc TSS PH Wi (c) <u>Furna</u> Pollutant	thin the ra <u>ce Wet Air</u> or property	Polluti Maxi any on lbs	0.00204 0.03000 7.5 to 10 on <u>Contro</u> mum for	<u>Dl</u> Maximum monthly rcury cont	0.00084 0.02400 times for
Zinc TSS PH Wi (c) <u>Furna</u> Collutant collutant mg/kg	thin the ra <u>ce Wet Air</u> or property	Polluti Maxi any on lbs	0.00204 0.03000 7.5 to 10 on Contro mum for one day	<u>Maximum</u> monthly cury cont	0.00084 0.02400 times for average
Zinc TSS PH Wi (c) <u>Furna</u> Collutant collutant mg/kg	thin the ra <u>ce Wet Air</u> or property	Polluti Maxi any on lbs	0.00204 0.03000 7.5 to 10 on Contro mum for one day a) of men brough fun 0.000	<u>Maximum</u> monthly cury cont	0.00084 0.02400 times for average rol processe 0.000
Zinc TSS PH Wi (c) <u>Furna</u> Collutant Collutant mg/kg *Lead *Mercury	thin the ra <u>ce Wet Air</u> or property (lb/milli	Polluti Maxi any on lbs	0.00204 0.03000 7.5 to 10 0.001 0.000 0.000 0.000	<u>Maximum</u> monthly cury cont	0.00084 0.02400 times for average rol processe 0.000 0.000
Zinc *TSS *pH Wi (c) <u>Furna</u> Pollutant pollutant mg/kg *Lead *Mercury Thallium	thin the ra <u>ce Wet Air</u> or property (lb/milli	Polluti Maxi any on lbs	0.00204 0.03000 7.5 to 10 0.00 <u>Contro</u> mum for one day 0.000 0.000 0.000 0.000	<u>Maximum</u> monthly cury cont	0.00084 0.02400 times for average rol processe 0.000 0.000 0.000
Zinc *TSS *pH Wi (c) <u>Furna</u> Pollutant collutant mg/kg *Lead *Mercury Thallium Zinc	thin the ra <u>ce Wet Air</u> or property (lb/milli	Polluti Maxi any on lbs	0.00204 0.03000 7.5 to 10 0.00 <u>Contro</u> mum for one day 0.000 0.000 0.000 0.000 0.000	<u>Maximum</u> monthly cury cont	0.00084 0.02400 times for average rol processe 0.000 0.000 0.000 0.000 0.000
Zinc *TSS *pH Wi (c) <u>Furna</u> Pollutant pollutant mg/kg *Lead *Mercury Thallium	thin the ra <u>ce Wet Air</u> or property (lb/milli	Polluti Maxi any on lbs th	0.00204 0.03000 7.5 to 10 0.00 <u>Contro</u> mum for one day 0.000 0.000 0.000 0.000 0.000 0.000	<u>Dl</u> Maximum monthly rcury cont rnace	0.00084 0.02400 times for average rol processe 0.000 0.000 0.000

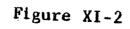
*Regulated Pollutant



NSPS TREATMENT SCHEME FOR OPTION A

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NSPS TREATMENT SCHEME FOR OPTION C

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SECTION XII

PRETREATMENT STANDARDS

This section describes the control and treatment technologies for pretreatment of process wastewaters from sources in the secondary subcategory. Pretreatment standards mercury for regulated pollutants are presented based on the selected control and treatment technology. Pretreatment standards for existing sources will not be promulgated for the secondary (PSES) mercury subcategory because there are no existing indirect dischargers in this subcategory. However, pretreatment standards for new sources (PSNS) for this subcategory will be promulgated.

TECHNICAL APPROACH TO PRETREATMENT

Before proposing and promulgating pretreatment standards, the Agency examines whether the pollutants discharged by the industry pass through the POTW or interfere with the POTW operation or its sludge disposal practices. In determining chosen whether pollutants pass through a well-operated POTW achieving secondary treatment, the Agency compares the percentage of a pollutant removed by POTW with the percentage removed by direct dischargers applying the best available technology economically achievable. A pollutant is deemed to pass through the POTW when the average percentage removed nationwide by well-operated POTW meeting secondary treatment requirements is less than the percentage removed by direct dischargers complying with BAT effluent limitations guidelines for that pollutant. (See generally, 46 FR at 9415-16 (January 28, 1981).)

This definition of pass-through satisfies two competing objectives set by Congress: (1) that standards for indirect discharg-ers be equivalent to standards for direct dischargers while at the same time, (2) that the treatment capability and performance of the POTW be recognized and taken into account in regulating the discharge of pollutants from indirect dischargers.

The Agency compares percentage removal rather than the mass or concentration of pollutants discharged because the latter would not take into account the mass of pollutants discharged to the POTW from non-industrial sources or the dilution of the pollutants in the POTW effluent to lower concentrations due to the addition of large amounts of non-industrial wastewater.

PRETREATMENT STANDARDS FOR NEW SOURCES

Options for pretreatment of wastewaters from new sources are based on increasing the effectiveness of end-of-pipe treatment technologies. All in-plant changes and applicable end-of-pipe treatment processes have been discussed previously in Section XI. The options for PSNS, therefore, are the same as the NSPS options discussed in Section XI.

A description of each option is presented in Section XI, while a more detailed discussion, including pollutants controlled by each treatment process, is presented in Section VII of Vol. I.

Treatment technologies considered for the PSNS options are:

OPTION A

o Chemical precipitation and sedimentation

OPTION C

- o Chemical precipitation and sedimentation
- o Multimedia filtration

PSNS OPTION SELECTION

Option C (chemical precipitation, sedimentation, and multimedia filtration) has been selected as the treatment technology for the proposed and promulgated pretreatment standards for new sources (PSNS) on the basis that it achieves effective removal of toxic pollutants at a reasonable cost. In addition, filtration is demonstrated in the nonferrous metals manufacturing category at 25 plants, and will not result in adverse economic impacts.

The wastewater discharge rates for promulgated PSNS are identical to the promulgated NSPS discharge rates for each waste stream. The PSNS discharge rates are shown in Table XII-1 (page 2920). No additional flow reduction measures for PSNS are feasible beyond the flow allowances given for NSPS.

REGULATED POLLUTANT PARAMETERS

The toxic pollutants selected for limitation, in accordance with the rationale of Sections VI and XI, are identical to those selected for limitation for NSPS. It is necessary to promulgate PSNS to prevent the pass-through of lead and mercury, which are the limited pollutants. The toxic pollutants are removed by a well operated POTW achieving secondary treatment at an average of 59 percent, while PSNS level technology removes approximately 99 percent.

PRETREATMENT STANDARDS FOR NEW SOURCES

Pretreatment standards for new sources are based on the treatable concentrations from the selected treatment technology (Option C), and the discharge rates determined in Section XI for NSPS. A mass of pollutant per mass of product (mg/kg) allocation is given for each subdivision within the subcategory. This pollutant allocation is based on the product of the treatable concentration from the promulgated treatment (mg/1) and the production normalized wastewater discharge rate (l/kkg). The achievable treatment concentrations for NSPS are identical to those for PSNS. These concentrations are listed in Table VII-21 (page 248) of Vol. I. PSNS are presented in Table XII-2 (page 2921).

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TABLE XII-1

PSNS WASTEWATER DISCHARGE RATES FOR THE SECONDARY MERCURY SUBCATEGORY

Wastewater Stream	PSNS Norm Discharge 1/kkg g		Production Normalized Parameter
Spent battery electrolyte	106	25.5	mercury produced from batteries
Acid wash and rinse water	2.0	0.5	mercury washed and rinsed
Furnace wet air pollution control	0:	0	mercury control processed through furnace

Table XII-2

PSNS FOR THE SECONDARY MERCURY SUBCATEGORY

(a) Spent Battery Electrolyte PSNS

Pollutant or	Maximum for	Maximum	for
pollutant property	any one day		average
mg/kg (lb/million	n lbs) of mercury	produced	from batterie
*Lead	0.030		0.014
*Mercury	0.016		0.006
Thallium	0.148		0.065
Zinc	0.108		0.045
(b) <u>Acid Wash and Ri</u>	nse Water PSNS	10 <u>1</u>	
Pollutant or	Maximum for	Maximum	for
pollutant property	any one day	monthly	average
*Lead	lion lbs) of merce 0.00056	ury washed	0.00026
*Lead		ury washed	•
*Lead *Mercury Thallium Zinc	0.00056 0.00030 0.00280 0.00204		0.00026 0.00012 0.00122
*Lead *Mercury Thallium Zinc (C) <u>Fugnace Wet Air</u> <u>F</u>	0.00056 0.00030 0.00280 0.00204	PSNS	0.00026 0.00012 0.00122 0.00084
*Lead *Mercury Thallium Zinc (c) <u>Furnace Wet Air F</u> Pollutant or	0.00056 0.00030 0.00280 0.00204 Pollution Control	PSNS Maximum monthly	0.00026 0.00012 0.00122 0.00084
*Lead *Mercury Thallium Zinc (c) <u>Furnace Wet Air E</u> Pollutant or pollutant property	0.00056 0.00030 0.00280 0.00204 Pollution Control Maximum for any one day	PSNS Maximum monthly iry contro	0.00026 0.00012 0.00122 0.00084
*Lead *Mercury Thallium Zinc (c) <u>Fugnace Wet Air E</u> Pollutant or pollutant property mg/kg (lb/mill	0.00056 0.00030 0.00280 0.00204 Pollution Control Maximum for any one day	PSNS Maximum monthly iry contro	0.00026 0.00012 0.00122 0.00084
*Lead *Mercury Thallium Zinc (c) <u>Furnace Wet Air E</u> Pollutant or pollutant property mg/kg (lb/mill *Lead *Mercury	0.00056 0.00030 0.00280 0.00204 Pollution Control Maximum for any one day fuion 1bs) of mercu through furna 0.000 0.000	PSNS Maximum monthly iry contro	0.00026 0.0012 0.00122 0.00084 for average
*Lead *Mercury Thallium Zinc (c) <u>Furnace Wet Air E</u> Pollutant or pollutant property mg/kg (lb/mill	0.00056 0.00030 0.00280 0.00204 Pollution Control Maximum for any one day fuion lbs) of mercu through furna 0.000	PSNS Maximum monthly iry contro	0.00026 0.00012 0.00122 0.00084 for average ol processed 0.000

*Regulated Pollutant

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SECTION XIII

BEST CONVENTIONAL POLLUTANT CONTROL TECHNOLOGY

EPA is not promulgating best conventional pollutant control technology (BCT) for the secondary mercury subcategory at this time.

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