DEVELOPMENT DOCUMENT FOR PROPOSED EFFLUENT LIMITATIONS GUIDELINES AND NEW SOURCE PERFORMANCE STANDARDS FOR THE

PETROLEUM REFINING

POINT SOURCE CATEGORY



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY
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Publication Notice

This is a development document for proposed effluent limitations guidelines and new source performance standards. As such, this report is subject to changes resulting from comments received during the period of public comments of the proposed regulations. This document in its final form will be published at the time the regulations for this industry are promulgated.

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and

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ABSTRACT

This development document presents the findings of an extensive study of the Petroleum Refining Industry for the purposes of developing effluent limitation guidelines, standards of performance, and pretreatment standards for the industry to implement Sections 304, 306 and 307 of the Federal Water Pollution Control Act of 1972, (PL 92-500). Guidelines and standards were developed for the overall petroleum refining industry, which was divided into six subcategories.

Effluent limitation guidelines contained herein set forth the degree of reduction of pollutants in effluents that is attainable through the application of best practicable control technology currently available (BPCTCA), and the degree of reduction attainable through the application of best available technology economically achievable (BATEA) by existing point sources for July 1, 1977, and July 1, 1983, respectively. Standards of performance for new sources are based on the application of best available demonstrated technology (BADT).

Annual costs for the petroleum refining industry for achieving BPCTCA Control by 1977 are estimated at \$244,000,000, and the additional annual costs for attaining BATEA Control by 1980 are estimated at \$250,000,000. The estimated annual costs for BADT for new sources is \$26,000,000.

Supporting data and rationale for the development of proposed effluent limitation guidelines and standards of performance are contained in this velopment document.

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SECTION I

CONCLUSIONS

This study covered the products included in the Petroleum Refining Industry (SIC 2911). The 247 U.S. petroleum refineries currently process 2.2 million cubic meters (14 million barrels) of crude oil per stream day. U.S. refineries vary in complexity from the very small, with simple atmospheric fractionation, or topping, to the very large integrated refineries manufacturing a multitude of petroleum and petrochemical products from a variety of feedstocks. The raw waste water load is dependent upon the types of processes employed by the refinery, justifying the utilization of production process groupings, as delineated by their effects on raw waste water as the basis for the subcategorization. The subcategories developed for the petroleum refining industry for the purpose of establishing effluent limitations are as follows:

Subcategory Basic Refinery Operations Included

Topping Topping and catalytic reforming

Low-Cracking Topping and cracking, with fresh feed (non-recycle)

to the cracking and hydroprocessing of less than 50

percent of the feedstock throughput.

High-Cracking Topping cracking, with a fresh feed (nonrecycle) to

the cracking and hydroprocessing of greater than 50

percent of the feedstock throughput.

Petrochemical Topping, cracking and petrochemicals operations.*

Lube Topping, cracking and lubes.**

Integrated Topping, cracking, lubes and petrochemicals

operations.*

- * Petrochemical operations Production of greater than 15 percent of the feedstock throughput in first generation petrochemicals and isomerization products (benzene, toluene, xylene, olefins, cyclohexane, etc.) and/or production of second generation petrochemicals (cumene, alcohols, ketones, etc.).
- ** Lubes the production of less than 12 percent of the feedstock throughput as lubes. Refineries with greater than 12 percent lubes are being considered speciality refineries and the guidelines for these specialty refineries will be set at a later date.

All six subcategories generate waste waters which contain similar constituents. However, the concentration and loading of the constitutents, termed "raw waste load," vary between the subcategories. Existing control and treatment technology, as practiced by the industry, includes both end-of-pipe treatment and in-plant reductions. Many of the individual wastewater streams, such as sour waters, have a deleterious effect on biological treatment facilities and/or receiving waters. Consequently, these individual streams are pretreated in-plant, prior to discharge to waste water facilities. Current technology for end-of-pipe treatment involves biological treatment and granular media filtration. Biological treatment systems employed include activitated sludge plants and aerated lagoons and stabilization pond systems.

SECTION II

RECOMMENDATIONS

The significant waste water constituents are BOD5, COD, TOC, total suspended solids, oil and grease, phenolic compounds, ammonia (N), sulfides, total and hexavalent chromium and zinc. These waste water constituents were selected to be the subject of the effluent limitations.

Effluent limitations commensurate with the best practical control technology currently available are proposed for each subcategory. These limitations, listed in Table 1, are explicit numerical values for the allowable discharges within each subcategory. Implicit in BPCTCA in-process technology is segregation of non-contact waste waters from process waste water. BPCTCA end-of-pipe technology is based on the application of the existing waste water treatment processes currently used in the Petroleum Refining Industry. These consist of equilization and storm diversion; initial oil and solids removal (API separators or baffle plate separators); further oil and solids removal (clarifiers, dissolved air flotation, or filters); carbonaceous waste removal (activated sludge, aerated lagoons, oxidation ponds, trickling filter, activated carbon, or combinations of these); and filters (sand, dual media; or multi-media) following biological treatment methods. variability of performance of biological waste water treatment systems been recognized in the development of the BPCTCA effluent mitations.

Effluent limitations commensurate with the best available technology economically achievable are proposed for each subcategory. These effluent limitations are listed in Table 2. The limitations are explicit numerical values for the allowable discharges within each subcategory. The primary end-of-pipe treatment proposed for BATEA effluent limitations is activated carbon adsorption, as further treatment in addition to BPCTCA control technology. Also implicit in BATEA technology are achievable reductions in waste water flow.

New source performance standards commensurate with the best available demonstrated technology are based on the flows achievable with BATEA technology, and the end-of-pipe control technology achievable with BPCTCA technology. These BADT effluent limitations are listed in Table 3. Activated carbon adsorption has not been included as BADT technology, since the use of this technology has not been sufficiently demonstrated, at this time, on petroleum refining waste water to insure its applicability and reliability on secondary effluent waste waters from refineries.

Table 1
BPCTCA
Petroleum Refining Industry Effluent Limitations
Kilograms of Pollutants/1000 Cubic Meters of Feedstock (1) Per Stream Day
(Pounds of Pollutants/1000 BBL of Feedstock Per Stream Day)

BOD5		<u>DD5</u>	<u>C</u>	OD	TOC		
Refinery	Monthly	Daily	Monthly -	Daily	Monthly -	Daily	
Subcategory	Average	Maximum	Average	Maximum	Average	Maximum	
Topping	7.3 (2.6)	9.0 (3.2)	28.3 (10.0)	31.7 (11.2)	6.3 (2.2)	7.7 (2.7)	
Low-Cracking	10.2 (3.6)	12.6 (4.4)	62.5 (22.1)	78.1 (27.6)	14.0 (4.9)	17.2 (6.1)	
High-Cracking	13.5 (4.4)	16.7 (5.9)	108.7 (38.4)	135.8 (48.0)	18.4 (6.5)	22.6 (8.0)	
Petroleum	15.5 (5.4)	19.1 (6.7)	95.1 (33.6)	118.9 (42.0)	21.0 (7.4)	25.8 (9.1)	
Lube	18.4 (6.5)	22.7 (8.0)	152.6 (53.9)	190.7 (67.4)	25.0 (8.8)	30.8 (10.9)	
Integrated	27.5 (9.7)	34.0 (12.0)	198.8 (70.2)	248.5 (87.8)	37.9 (13.4)	46.6 (16.5)	
Runoff (2)	0.025 (0.21	0.031(0.26)	0.24 (1.6)	0.30 (2.0)	0.035 (0.29	93)0.043(0.360)	
Ballast (3)	0.025 (0.21	0.031(0.26)	0.19 (2.0)	0.24 (2.5)	0.035 (0.29	93)0.043(0.360)	

- (1) Feedstock crude oil and/or natural gas liquids.
- (2) The additional allocation being allowed for contaminated storm runoff flow (kg/1000 liters (1b/1000 gallons) shall be based solely on that storm flow which passes through the treatment system. All additional storm runoff, that has been segregated from the main waste stream, shall not show a visible sheen or exceed a TOC concentration of 15 mg/l when discharged.
- (3) This is an additional allocation, based on ballast water intake (daily average)-per 1000 liters (per 1000 gallons)

Table 1 (continued) BPCTCA

Total Suspended Solids		ded Solids	Oil & Grease		Phenolic Compounds		
Refinery	Monthly	Daily	Monthly	Daily	Monthly	Daily	
Subcategory	Average	Maximum	Average	Maximum	<u>Average</u> -	Maximum	
Topping	4.6 (1.6)	5.8(2.0	2.2 (0.8)	2.8 (1.0)	0.048 (0.017)	0.070 (0.025)	
Low-Cracking	6.4 (2.2)	8.0 (2.8)	3.2 (1.1)	4.0 (1.4)	0.068 (0.024)	0.096 (0.034	
High-Cracking	8.2 (2.9)	10.2 (3.6)	4.0 (1.4)	5.0 (1.8)	0.088 (0.031)	0.125 (0.044)	
Petrochemical		12.0.(4.2)	5.0 (1.8)	6.2 (2.2)	0.110 (0.039)	0.156 (0.055)	
Lube	14.1(5.0)	17.6 (6.2)	6.9 (2.4)	8.6 (3.0)	0.150 (0.053)	0.211 (0.074)	
∪ Integrated	17.3(6.1)	21.6 (7.6)	8.6 (3.0)	10.8 (3.8)	0.188 (0.066)	0.266 (0.094)	
Runoff (2)	0.016(0.13)	0.020(0.17)	0.0080(0.067)	0.010(0.084)			
Ballast (3)	0.016(0.13)	0.020(0.17)	0.0080(0.067)	0.010(0.084)			

Table 1 (continued) BPCTCA

	Hexavalent Chromium				Zinc			
Refinery	Mo	onthly	Dai	Īy	Mont	hly		ily
Subcategory	Av	verage	Maxim	num	Aver	age	Max	<u>cimum</u>
Topping	0.0023	(0.00080)	0.0028	(0.0010)	0.23	(.080)	0.28	(0.10)
Low-Cracking	0.0032	(0.0011)	0.0040	(0.0014)	0.32	(0.11)	0.40	(0.14)
High-Cracking			0.0051	(0.0018)	0.41	(0.14)	0.51	(0.18)
Petrochemica			0.0057	(0.0020)	0.45	(0.16)	0.57	(0.20)
Lube	0.0068	(0.0024)	0.0085	(0.0030)	0.68	(0.24)	0.85	(0.30)
Integrated	0.0091	(0.0032)	0.011	(0.0040)		(0.32)	1.1	(0.40)
Runoff (2)	-			_		_		_
Ballast (3)				_		-		_

Table 1 (continued) BPCTCA

		Ammo	nia (N)	Sulfi	de	Total Chro	mium
	Refinery Subcategory	Monthly Average	Daily Maximum	Monthly Average	Daily <u>Maximum</u>	Monthly Average	Daily Maximum
7	Topping Low-Cracking High-Cracking Petroleum Lube Integrated	1.5 (0.53) 3.0 (1.1) 6.9 (2.4) 10.2 (3.6) 6.9 (2.4) 10.6 (3.8)	2.0 (0.70) 4.0 (1.4) 9.2 (3.2) 13.6 (4.8) 9.2 (3.2) 14.2 (5.0)	0.07 (0.025) 0.085 (0.029) 0.13 (0.042)	0.07 (0.022) 0.09 (0.031) 0.11 (0.040) 0.13 (0.046) 0.20 (0.066) 0.24 (0.086)	0.115 (0.040) 0.16 (0.056) 0.20 (0.070) 0.235 (0.083) 0.35 (0.123) 0.445 (0.157)	0.14 (0.050) 0.20 (0.070) 0.25 (0.088) 0.295 (0.104) 0.435 (0.154) 0.555 (0.196)
	Runoff (2) Ballast (1					 	

Month1v

Refinery

 $\boldsymbol{\omega}$

Petroleum Refining Industry Effluent Limitations Kilograms of Pollutants/1000 Cubic Meters of Feedstock (1) Per Stream Day

Monthly

(Pounds of Pollutants/1000 BBL of Feedstock Per Stream Day)

Daily

Subcategory	Average	Maximum	Average	Maximum	Average	Maximum
Topping	1.4(0.50)	1.7(0.61)	3.7(1.3)	4.5(1.6)	3.1(1.1)	4.0(1.4)
Low-Cracking	2.2(0.78)	2.7(0.97)	12.7(4.5)	15.8(5.6)	5.1(1.8)	6.2(2.2)
High-Cracking	2.8(0.99)	3.5(1.22)	20.4(7.2)	25.5(9.0)	6.5(2.3)	8.2(2.9)
Petroleum	3.0(1.07)	3.7(1.32)	11.3(4.0)	14.2(5.0)	7.1(2.5)	8.5(3.0)
Lube	6.1(2.16)	7.6(2.67)	37.6(13.3)	47.0(16.6)	13.9(4.9)	17.3(6.1)
Integrated	6.3(2.23)	7.8(2.75)	34.0(12.0)	42.5(15.0)	14.4(5.1)	17.5(6.2)
			/			
Runoff(2)	0.0085(0.071)	0.0(0.088)	0.023(0.19)	0.028(0.23)	0.019(0.16)	0.024(0.20)
) 0.0085(0.071		0.028(0.23)	0.035(0.29)	0.019(0.16)	0.024(0.20)

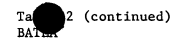
- (1) Feedstock crude oil and/or natural gas liquids.
- (2) The additional allocation being allowed for contaminated storm runoff flow (kg/1000 liters (1bs/1000 gallons) shall be based solely on that storm flow which passes through the treatment system. All additional storm runoff, that has been segregated from the main waste stream, shall not show a visible sheen or exceed a TOC concentration of 15 mg/l when discharged.

Daily

Daily

Monthly

(3) This is an additional allocation, based on ballast water intake (daily average) per 1000 liters (per 1000 gallons)



Total Suspended Solids		nded Solids	Oil & Grease		Phenolic Compounds	
Refinery	Monthly	Daily	Monthly	Daily	Monthly	Daily
Subcategory	Average	Maximum	Average	Maximum	Average-	Maximum
Topping	1.3(0.46)	1.6(0.58)	0.28(0.10)	0.34(0.12	0.0050(0.0018)	0.0073(0.0026)
Low-Cracking	2.1(0.74)	2.6(0.92)	0.40(0.14)	0.51(0.18)	0.0084(0.0030)	0.012(0.0043)
High-Cracking	2.6(0.93)	3.3(1.2)	0.54(0.19)	0.68(0.24)	0.011 (0.0039)	0.0155(0.055)
Petrochemical	2.8(1.0)	3.6(1.3)	0.59(0.21)	0.74(0.26)	0.012 (0.0042)	0.017 (0.0060)
Lube	5.6(2.0)	7.1(2.5)	1.1(0.40)	1.4 (0.50)	0.025 (0.0087)	0.034 (0.012)
Integrated	5.9(2.1)	7.4(2.6)	1.2(0.42)	1.5(0.52)	0.0255(0.0090)	0.037 (0.013)
Runoff (2)	0.0079(0.06	6)0.010(0.083)	0.0016(0.04)	0.0020(0.017)		
Ballast (3)	0.0079(0.06	66)0.010(0.083)	0.0016(0.014)	0.0020(0.017)		

Table 2 (continued)
BATEA

	<u>Ammonia (N)</u>		Sulf	ide	Total Chromium	
Refinery	Monthly	Daily	Monthly	Daily	Monthly	Daily
Subcategory	Average	Maximum	Average	Maximum	Average	Maximum
Topping	0.34(0.12)	0.45(0.16)	0.024(0.0084)	0.037(0.013)	0.065(0.023)	0.085(0.030)
Low-Cracking	0.76(0.27)	1.0(0.36)	0.036(0.013)	0.056(0.020)	0.105(0.037)	0.13(0.046)
High-Cracking	1.3(0.46)	1.6(0.58)	0.048(0.017)	0.075(0.026)	0.13(0.046)	0.16(0.058)
Petroleum	2.5(0.87)	3.3(1.2)	0.075(0.027)	0.12(0.042)	0.14(0.050)	0.175(0.062)
Lube	2.3(0.80)	3.0(1.1)	0.10(0.036)	0.16(0.057)	0.29(0.102)	0.36(0.128)
Integrated	2.8(1.0)	3.8(1.4)	0.10(0.036)	0.16(0.057)	0.30(0.106)	0.37(0.132)
Runoff(2)				. -		
Ballast(3)						

Table 2 (continued) BATEA

	Hexavalent	Chromium	Zinc		
Refinery	Monthly	Daily	Monthly	Daily	
Subcategory	Average	Maximum	Average	Maximum	
Topping	0.0013(0.00046)	0.0016(0.00058)	0.13(0.046)	0.16(0.050)	
Low-Cracking	0.0020(0.00072)	0.0025(0.00090)	0.20(0.072)	0.25(0.090)	
High-Cracking	0.0026(0.00092)	0.0034(0.00120)	0.26(0.092)	0.34(0.120)	
Petrochemical	0.0028(0.00099)	0.0035(0.00124)	0.28(0.099)	0.35(0.124)	
Lube	0.0059(0.0020)	0.0072(0.0025)	0.44(0.15)	0.57(0.20)	
Integrated	0.0059(0.0021)	0.0074(0.0026)	0.46(0.16)	0.59(0.21)	
Runoff (2)					
Ballast (3)					

Table 3
BADT
Petroleum Refining Industry Effluent Limitations
Kilograms of Pollutants/1000 Cubic Meters of Feedstock (1) Per Stream Day
(Pounds of Pollutants/1000 BBL of Feedstock Per Stream Day)

		BOD5		COD	Ţ	OC
Refinery	Month:	ly Daily	Monthly	Daily	Monthly	Daily
Subcategory	Averag	ge <u>Maximum</u>	Average	Maximum	Average	Maximum
Topping	4.3(1.5)	5.2(1.85)	15.0(5.3)	18.7(6.6)	3.7(1.3)	4.5(1.6)
Low-Cracking	5.8(2.4)	8.3(2.9)	40.2(14.2)	50.4(17.8)	9.3(3.3)	11.3(4.0)
High-Cracking	8.8(3.1)	10.8(3.8)	75.3(25.6)	90.6(32.0)	11.9(4.2)	14.4(5.1)
Petrochemical	9.1(3.2)	11.3(4.0)	57.2(20.2)	71.3(25.2)	12.5(4.4)	15.3(5.4)
Lube	14.9(5.3)	18.4(6.5)	125(44.3)	157(55.4)	20.6(7.3)	25.5(9.0)
Integrated	18.8(6.6)	23.2(8.2)	136(48.2)	170(60.2)	26.1(9.2)	32.3(11.4)
·		0.001(0.00)	0.01(0.7()	2 - 2 (2 2)		
		0.031(0.26)	0.91(0.76)	0.11(0.94)	0.023(0.19)	0.028(0.23)
Ballast(3)(0.025(0.21)	0.031(0.26	0.11(0.95)	0.14(1.2)	0.023(0.19)	0.028(0.23)

- (1) Feedstock crude oil and/or natural gas liquids.
- (2) The additional allocation being allowed for contaminated storm runoff flow (kg/1000 liters (1bs/1000 gallons shall be based solely on that storm flow which passes through the treatment system. All additional storm runoff, that has been segregated from the main waste stream, shall not show a visible sheen or exceed a TOC concentration of 15 mg/1 when discharged.
- (3) This is an additional allocation, based on ballast water intake (daily average) per 1000 liters (per 1000 gallons)

Table 3 (continued) BADT

<u>Total Suspende</u>		nded Solids Oil & Grease		Grease	Phenolic Compounds	
Refinery	Monthly	Daily	Monthly	Daily	Monthly	Daily
Subcategory	Average	Maximum	Average	Maximum	Average-	Maximum
Topping	2.6(0.93)	3.3(1.2)	1.3(0.46)	1.6(0.58)	0.02(0.0099)	0.040(0.014)
Low-Cracking	4.2(1.5)	5.2(1.8)	2.1(0.74)	2.6(0.92)	0.045(0.016)	0.062(0.022)
High-Cracking	5.4(1.9)	5.8(2.4)	2.6(0.93)	3.3(1.2)	0.057(0.020)	0.082(0.029)
Petrochemical	5.9(2.1)	7.4(2.6)	2.8(1.0)	3.6(1.3)	0.059(0.021)	0.085(0.030)
Lube	11.9(4.2) 14	4.7(5.2)	5.7(2.0)	7.1(2.5)	0.125(0.044)	0.177(0.062)
Integrated	11.9(4.2) 14	4.7(5.2)	5.9(2.1)	7.4(2.6)	0.130(0.046)	0.183(0.065)
Runoff (2)	0.016(0.13)	0.020(0.17)	0.0080(0.066)	0.010(0.083)		
Ballast (3)	0.016(0.13)	•	0.0080(0.066)	0.010(0.083)		

Table 3 (continued) BADT

		Ammonia (N)		Sulfic	de	Total Chromium	
	Refinery	Monthly	Daily	Monthly Monthly	Daily	Monthly	Daily
	Subcategory	Average	Maximum	Average	Maximum	Average	Maximum
	Topping	0.85(0.30)	1.1(0.40)	0.023(0.0081)	0.037(0.013)	0.065(0.023)	0.085(0.030)
	Low-Cracking	1.9(0.68)	2.5(0.90)	0.037(0.013)	0.057(0.020)	0.105(0.037)	0.13(0.046)
	High-Cracking	4.7(1.7)	6.2(2.2)	0.048(0.017)	0.074(0.026)	0.13(0.046)	0.16(0.058)
	Petrochemical	5.9(2.1)	7.9(2.8)	0.051(0.018)	0.079(0.028)	0.14(0.050)	0.175(0.062)
	Lube	5.7(2.0)	7.4(2.6)	0.103(0.026)	0.162(0.057)	0.29(0.102)	0.36(0.128)
	Integrated	7.2(2.6)	9.6(3.4)	0.107(0.038)	0.168(0.059)	0.30(0.106)	0.37(0.132)
	Runoff(2)						
	Ballast(3)						

Table 3 (continued) BADT

	Hexavalent	Chromium	Zinc		
Refinery	Monthly	Daily	Monthly	Daily	
Subcategory	Average	Maximum	Average	Maximum	
Topping	0.0013(0.00046)	0.0016(0.00058	0.13(0.046)	0.16(0.050)	
Low-Cracking	0.0020(0.00072)	0.0025(0.00090)	0.20(0.072)	0.25(0.090)	
High-Cracking	0.0026(0.00092)	0.0034(0.00120)	0.26(0.092)	0.34(0.120)	
Petrochemical	0.0028(0.00099)	0.0035(0.00124)	0.28(0.099)	0.35(0.124)	
Lube	0.0057(0.0020)	0.0072(0.0025)	0.44(0.15)	0.57(0.20)	
Integrated	0.0059(0.0021)	0.0074(0.0026)	0.46(0.16)	0.59(0.21)	
Runoff (2)					
Ballast (3)					

SECTION III

INTRODUCTION

Purpose and Authority

Section 301(b) of the Act requires the achievement by no later than July 1, 1977, of effluent limitations for point sources, other than publicly owned treatment works, which are based on the application of the best practicable control technology currently available as defined by the Administrator, pursuant to section 304(b) of the Act. Section 301(b) also requires the achievement by not later than July 1, 1983, of effluent limitations for point sources, other than publicly owned treatment works, which are based on the application of the best available technology economically achievable, which will result in reasonable further progress toward the national goal of eliminating discharge of all pollutants, as determined in accordance with regulations issued by the Administrator, pursuant to section 304(b) of the Act. Section 306 of the Act requires the achievement by new sources of a Federal standard of performance providing for the control of the discharge of pollutants which reflects the greatest degree of effluent reduction which the Administrator determines to be achievable through application of the best available demonstrated technology, processes, operative methods or other alternatives, including, where practicable, a standard permitting no discharge of pollutants.

section 304(b) of the Act requires the Administrator to publish within one year of enactment of the Act, regulations providing guidelines for effluent limitations setting forth the degree of effluent reduction attainable through the application of the best practicable control technology currently available and the degree of effluent reduction attainable through the application of the best control measures and practices achievable including treatment techniques, process and procedure innovations, operation methods and other alternatives. The regulations proposed herein set forth effluent limitations guidelines pursuant to section 304(b) of the Act for the petroleum refining industry source category.

Section 306 of the Act requires the Administrator, within one year after a category of sources is included in a list published pursuant to section 306(b) (1)(A) of the Act, to propose regulations establishing Federal standards of performance for new sources within such categories. The Administrator published in the Federal Register of January 16, 1973 (38 F.R. 1624), a list of 27 source categories. Publication of the list constituted announcement of the Administrator's intention of establishing, under section 306, standards of performance applicable to new sources within the petroleum refining industry source category which was included in the list published January 16, 1973.

Methods Used for Development of the Effluent Limitations Guidelines Standards of Performance

and

The Office of Air and Water Programs of the Environmental Protection Agency has been given the responsibility for the development of effluent limitation guidelines and new source standards as required by the Act. In order to promulgate the required guidelines and standards, the following procedure was adopted.

The point source category was first categorized for the purpose of determining whether separate limitations and standards are appropriate for different segments within a point source category. Such subcategorization was based upon raw materials used, products produced, manufacturing processes employed, raw waste loads, and other factors. This included an analysis of (1) the source and volume of water used in the plant and the sources of waste and waste waters in the plant; and (2) the constituents (including thermal) of all waste waters (including toxic constituents and other constituents) which result in taste, odor, and color in water or aquatic organisms. The constituents of waste waters which should be subject to effluent limitations guidelines and standards of performance were identified.

full range of control and treatment technologies existing within The each subcategory was identified. This included an identification of each distinct control and treatment technology, including both inplant and end-of-pipe technologies, which are existent or capable of being designed for each subcategory. It also included an identification, terms of the amount of constituents (including thermal) and the chemical, physical, and biological characteristics of pollutants, of the effluent level resulting from the application of each of the treatment and control technologies. The problems, limitations, and reliability of each treatment and control technology, and the required implementation In addition, the nonwater quality identified. time was also environmental impact (such as the effects of the applicubation of technologies upon other pollution problems, including air, solid waste, noise, and radiation) was also identified. The energy requirement of each of the control and treatment technologies was identified, as well as the cost of the application of such technologies.

The information, as outlined above, was then evaluated in order to determine methods or other alternatives. In identifying such technologies, various factors were considered. These included the total cost of application of technology in relation to the effluent reduction benefits to be achieved from such application, the age of equipment and facilities involved, the processes employed, the engineering aspects of the application of various types of control techniques, process changes, nonwater quality environmental impact (including energy requirements) and other factors.

During the initial phases of the study, an assessment was made of the ilability, adequacy, and usefulness of all existing data sources. Data on the identity and performance of waste water treatment systems within the petroleum refining industry were known to be included in:

- National Petroleum Refining Waste Water Characterization Studies and the Petroleum Industry Raw Waste Load Survey of 1972.
- 2. Environmental Protection Agency (Refuse Act) Permit Application.
- 3. Self-reporting discharge data from various states.
- 4. Monitoring data on individual refineries, collected by state agencies and/or regional EPA offices.

A preliminary analysis of these data indicated an obvious need for additional information. Although approximately 135 refineries were surveyed during the 1972 Raw Waste Load Survey, five activated sludge treatment plants were subjected to intensive sampling for identification of waste water treatment plant effluent performance. Identification of the types of treatment facilities used by the other individual refineries included no performance data.

Refuse Act Permit Application data are limited to identification of the atment systems used and reporting of final concentrations (which were luted with cooling waters in many cases); consequently, operating performance could not be established.

Self-reporting data was available from Texas, Illinois, and Washington. These reports show only the final effluent concentrations and identify the systems in use; rarely is there production information available which would permit the establishment of unit waste loads.

Monitoring data from the individual states and/or regional EPA offices again show only the final effluent concentrations and identify the systems in use. Rarely is production information available to permit the establishment of unit waste loads.

Additional data in the following areas were therefore required: 1) currently practiced or potential in-process waste control techniques; 2) identity and effectiveness of end-of-pipe waste control techniques; and 3) long-term data to establish the variability of performance of the end-of-pipe waste control techniques. The best source of information was the petroleum refineries themselves. New information was obtained from direct interviews and inspection visits to petroleum refinery facilities. Verification of data relative to long-term performance of waste control techniques was obtained by the use of standard EPA reference samples to determine the reliability of data submitted by the

petroleum refineries, and by comparison of the refinery data wit monitoring data from the state agencies and/or regional EPA offices.

The selection of petroleum refineries as candidates to be visited was guided by the trial categorization, which was based on the 1972 Raw Waste Load Survey. The final selection was developed from identifying information available in the 1972 Raw Waste Load Survey, EPA Permit Applications, state self-reporting discharge data, and contacts within regional EPA offices and the industry. Every effort was made to choose facilities where meaningful information on both treatment facilities and manufacturing processes could be obtained.

Survey teams composed of project engineers conducted plant visits. Information on the identity and performance of waste water treatment systems were obtained through:

- 1. Interviews with plant water pollution control personnel.
- 2. Examinations of treatment plant design and historical data (flow rates and analyses of influent and effluent).
- 3. Inspection of operations and analytical procedures, including verification of reported analyses by the use of EPA standard reference samples and by comparison of the refinery data with monitoring data from state agencies and/or regional EPA offices.

Information on process plant operations and associated waste water characteristics were obtained through:

- Interviews with plant operating personnel.
- Examination of plant design and operating data.
- Inspection of in-plant waste water controls.

The data base obtained in this manner was then utilized to develop recommended effluent limitations and standards of performance for the petroleum refining industry. All of the references utilized are included in Section XIII of this report. The data obtained during the field data collection program are included in Supplement B.

General Description of the Industry

The industrial waste water profile covers the petroleum refining industry in the United States, as defined by Standard Industrial Classification (SIC) Code 2911 of the U.S. Department of Commerce. Intermediates and finished products in this industry are numerous and varied. Table 4 is a partial listing of these products. The production of crude oil or natural gas from wells, or the production of natural

TABLE 4

Intermediates and Finished Products Frequently Found in the Petroleum Refining Industry

SIC 2911

Acid Oil
Alkylates
Aromatic Chemicals
Asphalt and Asphaltic Materials:
Semi-Solid and Solid
Benzene
Benzol
Butadiene
Coke (Petroleum)
Fuel Oils
Gas, Refinery or Still Oil
Gases, (LPG)
Gasoline, except natural gasoline
Greases: Petroleum, mineral jelly,
lubricative, etc.

Jet Fuels
Kerosene
Mineral Oils, natural
Mineral Waxes, natural
Naphtha
Napthenic Acids
Oils, partly refined
Paraffin Wax
Petrolatums, nonmedicinal
Road Oils
Solvents
Tar or Residuum

gasoline and other operations associated with such production, as covered under SIC Code 1311, are not within the scope of this study. This study also does not include distribution activities, such as gasoline service stations. Transportation of petroleum products is covered only to the extent that it is part of refinery pollution control, such as the treatment of ballast water. Other activities outside the scope of the SIC Code 2911 were included in the development of raw waste load data, and are listed as auxiliary processes which are inherent to an integrated refinery operation. Some of these include soap manufacture for the production of greases, steam generation, and hydrogen production.

A petroleum refinery is a complex combination of interdependent operations engaged in the separation of crude molecular constituents, molecular cracking, molecular rebuilding and solvent finishing to produce the products listed under SIC Code 2911. The refining operations may be divided among 12 general categories, where each category defines a group of refinery operations. The categories are listed in Table 5.

The characteristics of the waste water differ considerably for different processes. Considerable knowledge is available that can be used to make meaningful qualitative interpretations of pollutant loadings refinery processes. Such information is presented in Table 6, a semigraphic outline of the major sources of pollutants within a refinery. In order to set forth the character of the waste derived from each of the industry categories established in Section IV, it is essential study the sources and contaminants within the individual production processes and auxiliary activities. Each process is itself a series of unit operations which causes chemical and/or physical changes in the feedstock or products. In the commercial synthesis of a single product from a single feedstock, there generally are sections of the process associated with: the preparation of the feedstock, the chemical reaction, the separation of reaction products, and the final purification of the desired product. Each unit operation may have drastically different water usages associated with it. The type and quantity of contact waste water are therefore directly related to the nature of the various processes. This in turn implies that the types and quantities of waste water generated by each plant's total production mix are unique. The processes and activities along with brief process descriptions, trends in applications, and a delineation of waste water sources, are as follows:

TABLE 5

Major Refinery Process Categories

- 1. Storage and Transportation
- 2. Crude Desalting
- 3. Fractionation
- 4. Cracking
- 5. Hydrocarbon Rebuilding
- 6. Hydrocarbon Rearrangement
- 7. Solvent Refining or Extraction
- 8. Hydrotreating
- 9. Grease Manufacturing
- 10. Asphalt Production
- 11. Product Finishing
- 12. Auxiliary Activities (Not listed under SIC Code 2911)

TABLE 6

Qualitative Evaluation of Wastewater Flow and Characteristics
by Fundamental Refinery Processes

Production Processes	Flow	80D	COD	Pheno1	Sulfide	<u>011</u>	Emulsified 011	ρΗ	Temp.	Am- monia	<u>Chloride</u>	Acidity	Alkalinity	Susp. Solid
Crude Oil and Product Storage	xx	·x	xxx	x		ххх	xx	0	, 0	0		O		xx
Crude Desalting X	хх	XX	ХХ	x	xxx	×	xxx	; x	XXX	ХХ	xxx	0	X	XXX
Crude Distill- X	xxx	x	x	xx	XXX	xx	XXX	X	xx	xxx	X	0	x	X
Thermal Cracking	X.	x	x	×	x	x .		xx	xx	x	x	0	XX	x
Catalytic Cracking	xxx	ХХ	, XX	xxx	xxx	x .	x	xxx	xx	xxx	X	0	XXX	x ,
Hydrocracking	x			ХХ	XX .				XX	xx		•		•
Polymerization	x	x	X	0	x	. X	0	X	X	· x	x	x	0	x
Alkylation	xx	X	.x	0	xx	X	0	xx	1	×	xx	xx	. 0	xx
isomerization	x								•		•			
Reforming	x	0	. 0	X	x	X	. 0	0	x .	x	0	0	0	o
Solvent Refining	x		x	X	0		X	x	0		•	0	X	
Asphalt Blowing	xxx	xxx	xxx	X		XXX								
Dewaxing	X	xxx	xxx	x	0	x	0							
Hydrotreating	x	x	x		XX	•	0	XX		XX	0	0	x	0
Drying and Sweete ing	xxx	xxx	x	xx	0	0	x	XX	0	x	0	x	X	хx

STORAGE AND TRANSPORTATION

CRUDE OIL AND PRODUCT STORAGE

Process Description

Crude oil, intermediate, and finished products are stored in tanks of varying size to provide adequate supplies of crude oils for primary fractionation runs of economical duration, to equalize process flows and provide feedstocks for intermediate processing units, and to store final products prior to shipment in adjustment to market demands. Generally, operating schedules permit sufficient detention time for settling of water and suspended solids.

Wastes

Waste waters associated with storage of crude oil and products are mainly in the form of free and emulsified oil and suspended solids. During storage, water and suspended solids in the crude oil separate. The water layer accumulates below the oil, forming a bottom sludge. When the water layer is drawn off, emulsified oil present at the oilwater interface is often lost to the sewers. This waste is high in COD and contains a lesser amount of BOD5. Bottom sludge is removed at infrequent intervals. Additional quantities of waste result from leaks, spills, salt "filters" (for product drying), and tank cleaning.

stermediate storage is frequently the source of polysulfide bearing ste waters and iron sulfide suspended solids. Finished product storage can produce high BOD5, alkaline waste waters, as well as tetraethyl lead. Tank cleaning can contribute large amounts of oil, COD, and suspended solids, and a minor amount of BOD5. Leaks, spills and open or poorly ventilated tanks can also be a source of air pollution, through evaporation of hydrocarbons into the atmosphere.

Trends

Many refineries now have storage tanks equipped to minimize the release of hydrocarbons to the atmosphere. This trend is expected to continue probably accelerate. Equipment to minimize the release of hydrocarbon vapors includes tanks with floating-roof covers, pressurized tanks, and/or connections to vapor recovery systems. Floating-roof covers add to the waste water flow from storage tanks. refineries impose strict Bottom Sediment and Water (BS&W) specifications on crude oil supplies, and frequently have mixed-crude storage tanks; little or no waste water should originate from modern consequently, crude storage. Another significant trend is toward increased use of dehydration or drying processes preceding product finishing. processes significantly reduce the water content of finished product, thereby minimizing the quantity of waste water from finished product storage.

B. BALLAST WATER

Process Description

Tankers which are used to ship intermediate and final products generally arrive at the refinery in ballast (approximately 30 percent of the cargo capacity is generally required to maintain vessel stability).

Wastes

The ballast waters discharged by product tankers are contaminated with product materials which are the crude feedstock in use at the refinery, ranging from water soluble alcohol to residual fuels. In addition to the oil products contamination, brackish water and sediments are present, contributing high COD, and dissolved solids to the refinery waste water. These waste waters are generally discharged to either a ballast water tank or holding ponds at the refinery. In many cases, the ballast water is discharged directly to the waste water treatment system, and constitutes a shock load on the system.

Trends

As the size of tankers and refineries increases, the amount of ballast waters discharged to the refinery waste water system will also increase. The discharge of ballast water to the sea or estuary without treatment, as had been the previous practice by many tankers, is no longer a practical alternative for disposal of ballast water. Consequently, ballast water will require treatment for the removal of pollutants proto discharge. The use of larger ballast water storage tanks or ponds, for control of flow into the waste water treatment system, should increase as ballast water flow increases.

2. CRUDE DESALTING

Process Description

Common to all types of desalting are an emulsifier and settling tank. Salts can be separated from oil by either of two methods. In the first method, water wash desalting in the presence of chemicals (specific to the type of salts present and the nature of the crude oil) is followed by heating and gravity separation. In the second method, water wash desalting is followed by water/oil separation under the influence of a high voltage electrostatic field acting to agglomerate dispersed droplets. In either case, waste water containing various removed impurities is discharged to the waste stream, while clean desalted crude oil flows from the upper portion of the holding tank. A process flow schematic of electrostatic desalting is shown in Figure 1.

Wastes

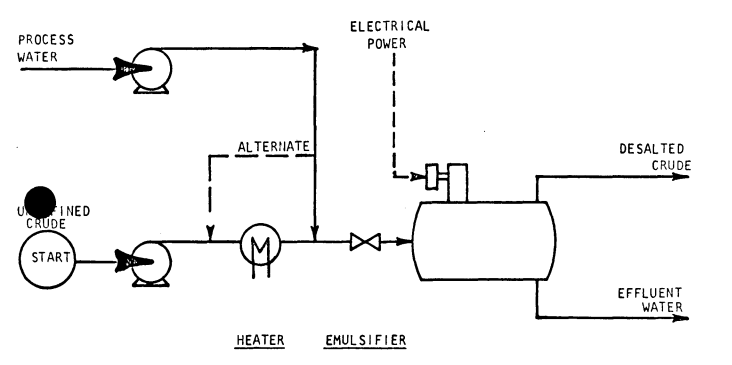


Figure 1
Crude Desalting
(Electrostatic Desalting)

Much of the BS&W content in crude oil is caused by the "Load-on-Top" procedure used on many tankers. This procedure can result in one demore cargo tanks containing mixtures of sea waters and crude oil, which cannot be separated by decantation while at sea, and are consequently retained in the crude oil storage at the refinery. While much of the water and sediment are removed from the crude oil by settling during storage, a significant quantity remains to be removed by desalting prior to processing of the crude in the refinery.

The continuous waste water stream from a desalter contains emulsified, and occasionally free oil, ammonia, phenol, sulfides, and suspended solids. These pollutants produce a relatively high BOD5 and COD. This waste water also contains enough chlorides and other dissolved materials to contribute to the dissolved solids problem in the areas where the waste water is discharged to fresh water bodies. There are also potential thermal pollution problems because the temperature of the desalting waste water often exceeds 95°C (200°F).

Trends

Electrical desalting is currently used much more than chemical desalting. In the future, chemical methods are expected to be used only as a supplement where the crude has a very high salt content. Two stage electrical desalting will become a more prevelant process, as dirtier crude feedstocks are processed in refineries. The growth in capacity of desalting units will parallel the growth of crude oil capacity.

3. CRUDE OIL FRACTIONATION

Fractionation serves as the basic refining process for the separation of crude petroleum into intermediate fractions of specified boiling point ranges. The several alternative subprocesses included are prefractionation and atmospheric fractionation, vacuum fractionation, vacuum flashing, and three-stage crude distillation.

Process Description

Prefractionation and Atmospheric Distillation (Topping or Skimming)

Prefractionation is an optional distillation process to separate economical quantities of very light distillates from the crude oil. Lower temperature and higher pressure conditions are used than would be required in atmospheric distillation. Some process water can be carried over to the prefractionation tower from the desalting process.

Atmospheric Distillation breaks the heated crude oil as follows:

1. Light overhead products (C5 and lighter) as in the case of prefractionation.

- 2. Sidestream distillate cuts of kerosene, heating and gas oil can be separated in a single tower or in a series of topping towers, each tower yielding a successively heavier product stream.
- 3. Residual or reduced crude oil.

Vacuum Fractionation

The asphaltic residuum from the atmospheric distillation amounts to 37 percent (U.S. average) of the crude charged. This material is sent to vacuum stills, which recover additional heavy gas oil and deasphalting feedstock from the bottoms residue.

Three Stage Crude Distillation

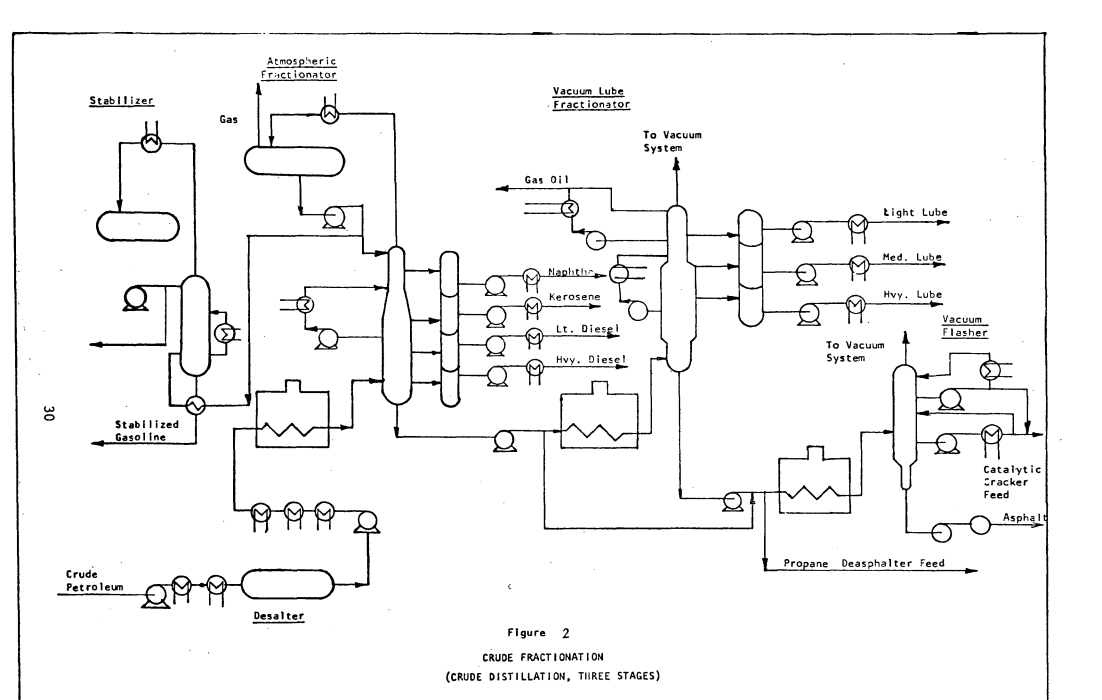
Three stage crude distillation, representing only one of many possible combinations of equipment, is shown schematically in Figure 2. The process consists of:

- 1. An atmospheric fractioning stage which produces lighter oils;
- 2. An initial vacuum stage which produces well-fractioned, lube oil base stocks plus residue for subsequent propane deasphalting;
- 3. A second vacuum stage which fractionates surplus atmospheric bottoms not applicable for lube production, plus surplus initial vacuum stage residuum not required for deasphalting. This stage adds the capability of removing catalytic cracking stock from surplus bottoms to the distillation unit.

Crude oil is first heated in a simple heat exchanger, then in a direct-fired crude charge heater. Combined liquid and vapor effluent flow from the heater to the atmospheric fractionating tower, where the vaporized distillate is fractionated into gasoline overhead product and as many as four liquid sidestreams products:naphtha, kerosene, light and heavy diesel oil. Part of the reduced crude from the bottom of the atmospheric tower is pumped through a direct-fired heater to the vacuum lube fractionator. Bottoms are combined and charged to a third direct-fired heater. In the tower, the distillate is subsequently condensed and withdrawn as two sidestreams. The two sidestreams are combined to form catalytic cracking feedstocks, with an asphalt base stock withdrawn from the tower bottom.

Wastes

The waste water from crude oil fractionation generally comes from three sources. The first source is the water drawn off from overhead accumulators prior to recirculation or transfer of hydrocarbons to other



fractionators. This waste is a major source of sulfides and ammonia, pecially when sour crudes are being processed. It also contains significant amounts of oil, chlorides, mercaptans and phenols.

A second waste source is discharged from oil sampling lines. This should be separable but may form emulsions in the sewer.

A third possible waste source is the very stable oil emulsions formed in the barometric condensers used to create the reduced pressures in the vacuum distillation units. However, when barometric condensers are replaced with surface condensers, oil vapors do not come in contact with water; consequently, emulsions do not develop.

Trends

The general industry trend to larger and more complete refineries has been reflected also in larger and more complete crude fractionation units. Thus, simple atmospheric "topping" units are being replaced by the atmospheric- vacuum combinations with an increasing number of sidestream products. Installed vacuum fractionation capacity now totals, 0.8 million cu m/day (5 million bbl/day). (3) Modern refineries are installing surface condensers to significantly reduce waste water loads from vacuum operations.

4. CRACKING

THERMAL CRACKING

Process Description

This fundamental process is defined in this study to include visbreaking and coking, as well as regular thermal cracking. In each of these operations, heavy gas oil fractions (from vacuum stills) are broken down into lower molecular weight fractions such as domestic heating oils, catalytic cracking stock, and other fractions by heating, but without the use of a catalyst. Typical thermal cracking conditions are 480° -603°C, (900° - 1100°F) and 41.6 - 69.1 atm (600-1000 psig). The high pressures result from the formation of light hydrocarbons in the cracking reaction (olefins, or unsaturated compounds, are always formed in this chemical conversion). There is also always a certain amount of heavy fuel oil and coke formed by polymerization and condensation reactions.

Wastes

The major source of waste water in thermal cracking is the overhead accumulator on the fractionator, where water is separated from the hydrocarbon vapor and sent to the sewer system. This water usually contains various oil and fractions and may be high in BOD5, COD, ammonia, phenol, and sulfides, and may have a high alkalinity.

Trends

Regular thermal cracking, which was an important process before the development of catalytic cracking, is being phased out. Visbreaking and coking units are still installed but, because of product sulfur restrictions, to a lesser extent than before. With the trends toward dirtier crudes containing more sulfur, hydrocracking and propane deasphalting are receiving more attention to recover salable products with low sulfur content from the residuum.

B. CATALYTIC CRACKING

Process Description

Catalytic cracking, like thermal cracking, breaks heavy fractions, principally gas oils, into lower molecular weight fractions. This is probably the key process in the production of large volumes of high-octane gasoline stocks; furnace oils and other useful middle molecular weight distillates are also produced. The use of a catalyst permits operations at lower temperatures and pressures than with thermal cracking, and inhibits the formation of undesirable polymerized products. Fluidized catalytic processes, in which the finely powdered catalyst is handled as a fluid, have largely replaced the fixed bed and moving bed processes, which use a beaded or pelleted catalyst. A schematic flow diagram of fluid catalytic cracking is shown in Figure 3.

The process involves at least four types of reactions: 1) thern decomposition; 2) primary catalytic reactions at the catalyst surface, 3) secondary catalytic reactions between the primary products, and 4) removal of polymerizable products from further reactions by absorption onto the surface of the catalyst as coke. This last reaction is the key to catalytic cracking because it permits decomposition reactions to move closer to completion than is possible in simple thermal cracking. Cracking catalysts include synthetic and/or natural silica-alumina, treated bentonite clay, Fuller's earth, aluminum hydrosilicates and bauxite. These catalysts are in the form of beads, pellets, and powder, and are used in either a fixed, moving or fluidized bed. The catalyst is usually heated, lifted into the reactor area by the incoming oil feed which, in turn, is immediately vaporized upon contact. Vapors from the reactors pass upward through a cyclone separator, which removes most of the entrained catalyst. These vapors then enter the fractionator, where the desired products are removed and heavier fractions recycled to the reactor.

Wastes

Catalytic cracking units are one of the largest sources of sour and phenolic wastewaters in a refinery. Pollutants from catalytic cracking generally come from the steam strippers and overhead accumulators on

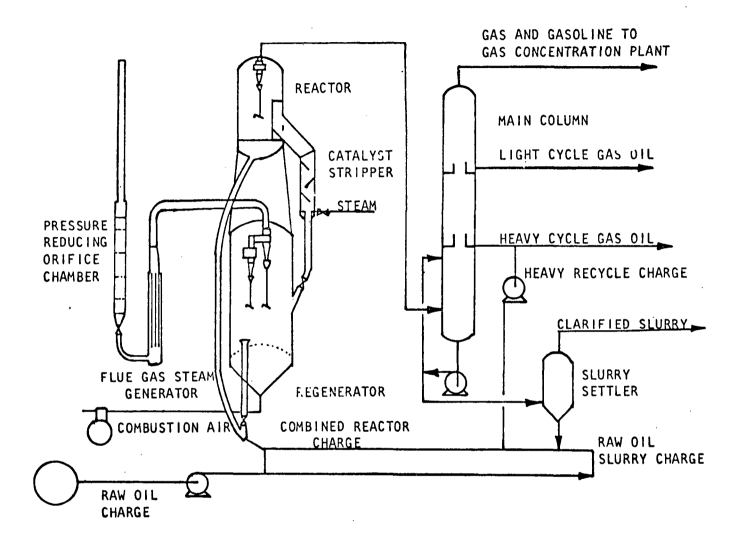


Figure 3

CATALYTIC CRACKING

(FLUID CATALYTIC CRACKING)

fractionators, used to recover and separate the various hydrocarbonations produced in the catalytic reactors.

The major pollutants resulting from catalytic cracking operations are oil, sulfides, phenols, cyanides, and ammonia. These pollutants produce an alkaline waste water with high BOD5 and COD concentrations. Sulfide and phenol concentrations in the waste water vary with the type of crude oil being processed, but at times are significant. Regeneration of spent catalyst may produce enough carbon monoxide and catalyst fines to constitute an air pollution problem.

Trends

Recycle rates have been declining since 1968, and the trend is expected to continue due to the development of higher activity catalysts (molecular sieve catalysts, as opposed to high surface area silica-alumina catalysts). The trend in subprocesses is toward greater use of large fluid catalytic cracking in preference to moving or fixed-bed cracking. Catalytic cracking units are also being supplanted by hydrocracking and hydrotreating processes. During 1972, a decline of 1.4 percent in fresh feed catalytic cracking capacity was experienced in the United States. (3)

C. HYDROCRACKING

Process Description

This process is basically catalytic cracking in the presence of hydrogen, with lower temperatures and higher pressures than fluid catalytic cracking. Hydrocracking temperatures range from 203° - 425°C (400° - 800°F), while pressures range from 7.8 - 137.0 atm (100 to 2000 psig). Actual conditions and hydrogen consumption depend upon the feedstock, and the degree of hydrogenation required. The molecular weight distribution of the products is similar to catalytic cracking, but with the reduced formation of olefins.

Wastes

At least one waste water stream from the process should be high in sulfides, since hydrocracking reduces the sulfur content of the material being cracked. Most of the sulfides are in the gas products which are sent to a treating unit for removal and/or recovery of sulfur and ammonia. However, in product separation and fractionation units following the hydrocracking reactor, some of the HS will dissolve in the waste water being collected. This water from the separator and fractionator will probably be high in sulfides, and possibly contain significant quantities of phenols and ammonia.

Trends

Hydrocracking has greater flexibility than catalytic cracking in justing operations to meet changing product demands. For the last few years, it has been one of the most rapidly growing refining processes. This trend is expected to continue.

HYDROCARBON REBUILDING

A. POLYMERIZATION

Process Description

Polymerization units are used to convert olefin feedstocks (primarily propylene) into higher octane polymer units. These units generally consist of a feed treatment unit (remove H2S, mercaptans, nitrogen compounds), a catalytic reactor, an acid removal section, and a gas stabilizer. The catalyst is usually phosphoric acid, although sulfuric acid is used in some older methods. The catalytic reaction occurs at 147° - 224°C (300° - 435°F), and a pressure of 11.2 - 137.0 atm (150 - 2000 psig). The temperature and pressure vary with the individual subprocess used.

Wastes

Polymerization is a rather dirty process in terms of pounds of pollutants per barrel of charge, but because of the small polymerization capacity in most refineries, the total waste production from the process small. Even though the process makes use of acid catalysts, the ste stream is alkaline, because the acid catalyst in most subprocesses is recycled, and any remaining acid is removed by caustic washing. Most of the waste material comes from the pretreatment of feedstock to the reactor. The waste water is high in sulfides, mercaptans, and ammonia. These materials are removed from the feedstock in caustic acid.

Trends

Polymerization is a marginal process, since the product octane is not significantly higher than that of the basic gasoline blending stocks, and does not provide much help in upgrading the overall motor fuel pool. In addition, alkylation yields per unit of olefin feed are much better than polymerization yields. Consequently, the current polymerization downtrend is expected to continue.

B. ALKYLATION

Process Description

Alkylation is the reaction of an isoparaffin (usually isobutane) and an olefin (propylene, butylene, amylenes) in the presence of a catalyst at carefully controlled temperatures and pressures to produce a high octane alkylate for use as a gasoline blending component. Propane and butane

are also produced. Sulfuric acid is the most widely used catalyst although hydrofluoric acid is also used. The reactor products a separated in a catalyst recovery unit, from which the catalyst recycled. The hydrocarbon stream is passed through a caustic and water wash before going to the fractionation section.

Wastes

The major discharge from sulfuric acid alkylation are the spent caustics from the neutralization of hydrocarbon streams leaving the sulfuric acid alkylation reactor. These waste waters contain dissolved and suspended solids, sulfides, oils, and other contaminants. Water drawn off from the overhead accumulators contains varying amounts of oil, sulfides, and other contaminants, but is not a major source of waste in this subprocess. Most refineries process the waste sulfuric acid stream from the reactor to recover clean acids, use it as if for neutralization of other waste streams, or sell it.

Hydrofluoric acid alkylation units have small acid rerun units to purify the acid for reuse. HF units do not have a spent acid or spent caustic waste stream. Any leaks or spills that involve loss of fluorides constitute a serious and difficult pollution problem. Formation of fluosilicates has caused line plugging and similar problems. The major sources of waste material are the overhead accumulators on the fractionator.

Trends

Alkylation process capacity is currently declining slowly, but this trend may be reversed, as the demand for low lead, high octane gasoline increases.

6. HYDROCARBON REARRANGEMENTS

A. ISOMERIZATION

Process Description

Isomerization is a process technique for obtaining higher octane motor fuel by converting light gasoline stocks into their higher octane isomers. The greatest application has been, indirectly, in the conversion of isobutane from normal butane, for uses as feedstock for the alkylation process. In a typical subprocess, the desulfurized feedstock is first fractionated to separate isoparaffins from normal paraffins. The normal paraffins are then heated, compressed, and passed through the catalytic hydrogenation reactor which isomerizes the n-paraffin to its respective high octane isomer. After separation of hydrogen, the liquids are sent to a stabilizer, where motor fuel blending stock or synthetic isomers are removed as products.

Wastes

Isomerization waste waters present no major pollutant discharge problems. Sulfides and ammonia are not likely to be present in the effluent. Isomerization waste waters should also be low in phenolics and oxygen demand.

Trends

The requirements for units to isomerize n-butane to isobutane will not be as great in refineries where hydrocracking is being installed, as the hydrocracking process yields an off-gas rich in isobutane. However, the isomerization capacity of U.S. refiners is not expected to decrease, but to continue to grow as the demand for motor fuel grows.

B. REFORMING

Process Description

Reforming converts low octane naphtha, heavy gasoline, and napthene-rich stocks, to high octane gasoline blending stock, aromatics for petrochemical use, and isobutane. Hydrogen is a significant by-product of the process. Reforming is a mild decomposing process, since some reduction occurs in molecular size and boiling range of the feedstock. Feedstocks are usually hydrotreated for the removal of sulfur and nitrogen compounds prior to charging to the reformer, since the platinum talysts widely used are readily poisoned.

The predominant reaction during reforming is the dehydrogenation of naphthenes. Important secondary reactions are the isomerization and dehydrocyclization of paraffins. All three reactions result in higher octane products.

One subprocess may be divided into three parts: the reactor heater section, in which the charge plus recycle gas is heated and passed over the catalyst in a series of reactions; the separator drum, in which the reactor effluent is separated into gas and liquid streams, the gas being compressed for recycle; and the stabilizer section, in which the separated liquid is stabilized to the desired vapor pressure. There are many variations in subprocesses, but the essential, and frequently the only, difference is the composition of the catalyst involved.

Wastes

Reforming is a relatively clean process. The volume of waste water flow is small, and none of the waste water streams has high concentration of significant pollutants. The waste water is alkaline, and the major pollutant is sulfide from the overhead accumulator on the stripping tower used to remove light hydrocarbon fractions from the reactor effluent. The overhead accumulator catches any water that may be

contained in the hydrocarbon vapors. In addition to sulfides, the waster contains small amounts of ammonia, mercaptans and oil.

Trends

Reforming capacity in the U.S. is currently growing at about the same rate as total crude capacity. This growth rate may increase, however, as the demand for motor fuel grows.

7. SOLVENT REFINING

Refineries employ a wide spectrum of contact solvent processes, which are dependent upon the differential solubilities of the desirable and undesirable feedstock components. The principal steps are: countercurrent extraction, separation of solvent and product by heating and fractionation, and solvent recovery. Napthenics, aromatics unsaturated hydrocarbons, sulfur and other inorganics are separated, with the solvent extract yielding high purity products. Many of the solvent processes may produce process waste waters which contain small amounts of the solvents employed. However, these are ususally minimized, because of the economic incentives for reuse of the solvents.

Process Description

The major processes include:

Solvent Deasphalting - The primary purpose of solvent deasphalting is recover lube or catalytic cracking feedstocks from asphaltic residuals, with asphalt as a by-product. Propane deasphalting is the predominant technique. The vacuum fractionation residual is mixed in a fixed proportion with a solvent in which asphalt is not soluble. The solvent is recovered from the oil via steam stripping and fractionation, and is reused. The asphalt produced by this method is normally blended into fuel oil or other asphaltic residuals.

Solvent Dewaxing - Solvent dewaxing removes wax from lubricating oil stocks by promoting crystallization of the wax. Solvents which are used include: furfural, phenol, cresylic acid - propane (Duo-Sol), liquid sulfur dioxide (Eleleanu process), B-B - dichloroethyl ether, methyl ethyl ketone, nitrobenzene, and sulfur-benzene. The process yields deoiled waxes, wax-free lubricating oils, aromatics, and recovered solvents.

Lube Oil Solvent Refining - This process includes a collection of subprocesses for improving the quality of lubricating oil stock. The raffinate or refined lube oils obtain improved temperature, viscosity, color and oxidation resistance characteristics. A particular solvent is selected to obtain the desired quality raffinate. The solvents include: furfural, phenol, sulfur dioxide, and propane.

Aromatic Extraction - Benzene, toluene, and xylene (BTX) are formed as -products in the reforming process. The reformed products are ractionated to give a BTX concentrate cut, which in turn is extracted from the napthalene and the paraffinics with a glycol base solvent.

Butadiene Extraction - Approximately 15 percent of the U.S. supply of butadiene is extracted from the C4 cuts from the high temperature petroleum cracking processes. Furfural or cuprous ammonia acetate (CAA) are commonly used for the solvent extraction.

Wastes

The major potential pollutants from the various solvent refining subprocesses are the solvents themselves. Many of the solvents, such as phenol, glycol, and amines, can produce a high BOD5. Under ideal conditions the solvents are continually recirculated with no losses to the sewer. Unfortunately, some solvent is always lost through pump seals, flange leaks, and other sources. The main source of waste water is from the bottom of fractionation towers. Oil and solvent are the major waste water constituents.

Trends

Solvent extraction capacities can be expected to slowly increase as quality requirements for all refinery products become more stringent, as the demand for lube oils grows, and as the petrochemicals industry nation to require increasing quantities of aromatics.

8. HYDROTREATING

Process Description

Hydrotreating processes are used to saturate olefins, and to remove sulfur and nitrogen compounds, odor, color and gum-forming materials, and others by catalytic action in the presence of hydrogen, from either straight-run or cracked petroleum fractions. In most subprocesses, the feedstock is mixed with hydrogen, heated, and charged to the catalytic reactor. The reactor products are cooled, and the hydrogen, impurities and high grade product separated. The principal difference between the many subprocesses is the catalyst; the process flow is similar for essentially all subprocesses.

Hydrotreating processes are used to reduce the sulfur content of product streams from sour crudes by approximately 90 percent or more. Nitrogen removal requires more severe operating conditions, but generally 80 - 90 percent, or better, reductions are accomplished.

The primary variables influencing hydrotreating are hydrogen partial pressure, process temperature, and contact time. An increase in hydrogen pressure gives a better removal of undesirable materials and a

better rate of hydrogenation. Make-up hydrogen requirements are generally high enough to require a hydrogen production unit. Excessive temperatures increase the formation of coke, and the contact time is set to give adequate treatment without excessive hydrogen usage and/or undue coke formation. For the various hydrotreating processes the pressures range from 7.8 - 205.1 atm (100 to 3000 psig). Temperatures range from less than 177°C (350°F) to as high as 450°C (850°F), with most processing done in the range of 314°C (600°F) to 427°C (800°F). Hydrogen consumption is usually less than 5.67 NM3 (200 scf) per barrel of charge.

Principal hydrotreating subprocesses are used as follows:

- 1. Pretreatment of catalytic reformer feedstock.
- 2. Naphtha desulfurization.
- 3. Lube oil polishing.
- 4. Pretreatment of catalytic cracking feedstock.
- 5. Heavy gas-oil and residual desulfurization.
- 6. Naphtha saturation.

Wastes

The strength and quantity of waste waters generated by hydrotreating depends upon the subprocess used and feedstock. Ammonia and sulfides are the primary contaminants, but phenols may also be present, if the feedstock boiling range is sufficiently high.

Trends

The use of hydrotreating is increasing and should continue to increase at a greater rate than crude capacity since the process can be applied to almost any sour feedstock, is flexible, and eliminates contaminants of concern to the refining industry from an operating standpoint, and to the general public from an aesthetic standpoint.

9. GREASE MANUFACTURING

Process Description

Grease manufacturing processes require accurate weight or volumetric measurements of feed components, intimate mixing, rapid heating and cooling, together with milling, dehydration and polishing in batch reactions. The feed components include soap and petroleum oils, with inorganic clays and other additives.

Grease is primarily a soap and lube oil mixture. The properties of grease are determined in large part by the properties of the soap component. For example, sodium metal base soaps are water soluble and would then not be suitable for water contact service. A calcium soap

erease can be used in water service. The soap may be purchased as a raw erial or may be manufactured on site as an auxiliary process.

Wastes

Only very small volumes of waste water are discharged from a grease manufacturing process. A small amount of oil is lost to the waste water system through leaks in pumps. The largest waste loading occurs when the batch units are washed, resulting in soap and oil discharges to the sewer system.

Trends

Because of an increase in sealed grease fittings in automobiles and longer lasting greases, a slight decline in grease production is expected through 1975.

10. ASPHALT PRODUCTION

Process Description

Asphaltic feedstock (flux) is contacted with hot air at 203°C (400°F) to 280°C (550°F) to obtain desirable asphalt product. Both batch and continuous processes are in operation at present, but the batch process is more prevalent because of its versatility. Nonrecoverable catalytic compounds include: Copper sulfate, zinc chloride, ferric chloride, minum chloride, phosphorous pentoxide, and others. The catalyst will normally contaminate the process water effluent.

Wastes

Waste waters from asphalt blowing contain high concentrations of oils, and have high oxygen demand. Small quantities of phenols may also be present.

11. PRODUCT FINISHING

A. DRYING AND SWEETENING

Process Description

Drying and sweetening is a relatively broad process category primarily used to remove sulfur compounds, water and other impurities from gasoline, kerosene, jet fuels, domestic heating oils, and other middle distillate products. "Sweetening" pertains to the removal of hydrogen sulfide, mercaptans and thiophenes, which impart a foul odor and decrease the tetra-ethyl lead susceptibility of gasoline. The major sweetening operations are oxidation of mercaptans or disulfides, removal of mercaptans, and destruction and removal of all sulfur compounds. Drying is accomplished by salt filters or absorptive clay beds.

Electric fields are sometimes used to facilitate separation of product.

the

Wastes

The most common waste stream from drying and sweetening operations is spent caustic. The spent caustic is characterized as phenolic or sulfidic, depending on which is present in the largest concentration. Whether the spent caustic is actually phenolic or sulfidic is mainly determined by the product stream being treated. Phenolic spent caustics contain phenol, cresols, xylenols, sulfur compounds, and some neutral oils. Sulfidic spent caustics are rich in sulfides, but do not contain any phenols. These spent caustics have very high BOD5 and COD. The phenolic caustic streams are usually sold for the recovery of phenolic materials.

Other waste streams from the process result from water washing of the treated product and regeneration of the treating solution such as sodium plumbite (No2 PbO2) in doctor sweetening. These waste streams will contain small amounts of oil and the treating material, such as sodium plumbite (or copper from copper chloride sweetening).

The treating of sour gases produces a purified gas stream, and an acid gas stream rich in hydrogen sulfide. The H2S rich stream can be flared, burned as fuel, or processed for recovery of elemental sulfur.

Trends

As air pollution agencies increase their efforts to control sulfur emissions to the atmosphere, the restrictions on sulfur content in fuels can be expected to tighten. This will generate a strong trend to replacement of the sweetening processes by more hydrotreating (desulfurization), because hydrotreating removes almost all sulfur compounds and not just hydrogen sulfide, mercaptans, and elemental sulfur. Nevertheless, on certain feedstocks sweetening will continue to be used because it will be as effective as, and more economical than, hydrotreating. Those processes producing high waste loads (Doctor Sweetening, etc.) are being replaced by lower waste-producing processes.

B. LUBE OIL FINISHING

Process Description

Solvent refined and dewaxed lube oil stocks can be further refined by clay or acid treatment to remove color-forming and other undesirable materials. Continuous contact filtration, in which an oil-clay slurry is heated and the oil removed by vacuum filtration, is the most widely used subprocess.

Wastes

Acid treatment of lubricating oils produces acid bearing wastes occuring as rinse waters, sludges, and discharges from sampling, leaks and utdowns. The waste streams are also high in dissolved and suspended solids, sulfates, sulfonates, and stable oil emulsions.

Handling of acid sludge can create additional problems. Some refineries burn the acid sludge as fuel. Burning the sludge produces large volumes of sulfur dioxide that can cause air pollution problems. Other refineries neutralize the sludge with alkaline wastes and discharge it to the sewer, resulting in both organic and inorganic pollution. The best method of disposal is probably processing to recover the sulfuric acid, but this also produces a waste water stream containing acid, sulfur compounds and emulsified oil.

Clay treatment results in only small quantities of waste water being discharged to the sewer. Clay, free oil, and emulsified oil are the major waste constituents. However, the operation of clay recovery kilns involves potential air pollution problems of hydrocarbon and particulate emissions. Spent clays usually are disposed of by landfill.

Trends

Acid and clay treatment of lube oils is gradually being replaced by hydrotreating methods. Acid treatment in particular is being phased out rather rapidly.

C. BLENDING AND PACKAGING

ocess Description

Blending is the final step in producing finished petroleum products to meet quality specifications and market demands. The largest volume operation is the blending of various gasoline stocks (including alkylates and other high-octane components) and anti-knock (tetra-ethyl lead), anti-rust, anti-icing, and other additives. Diesel fuels, lube oils, and waxes involve blending of various components and/or additives. Packaging at refineries is generally highly-automated and restricted to high volume, consumer-oriented products such as motor oils.

Wastes

These are relatively clean processes because care is taken to avoid loss of product through spillage. The primary source of waste material is from the washing of railroad tank cars or tankers prior to loading finished products. These wash waters are high in emulsified oil.

Tetra-ethyl lead is the major additive blended into gasoline and it must be carefully handled because of its high toxicity. Sludges from finished gasoline storage tanks can contain large amounts of lead and should not be washed into the wastewater system.

Trends

There will be an increased use of automatic proportioning facilities for product blending with a trend toward contracting out of packaging of lower-volume products that are less suitable to highly-automated operation.

12. AUXILIARY ACTIVITIES

A. HYDROGEN MANUFACTURE

Process Description

The rapid growth of hydrotreating and hydrocracking has increased the demand for hydrogen beyond the level of by-product hydrogen available from reforming and other refinery processes. The most widely used process for the manufacture of hydrogen in the refinery is steam reforming, which utilizes refinery gases as a charge stock. The charge is purified to remove sulfur compounds that would temporarily deactivate the catalysts.

The desulfurized feedstock is mixed with superheated steam and charged to the hydrogen furnace. On the catalyst the hydrocarbons are converted to hydrogen, carbon monoxide, and carbon dioxide. The furnace supplies the heat needed to maintain the reaction temperature.

The gases from the furnace are cooled by the addition of condensate and steam, and then passed through a converter containing a high- or lotemperature shift catalyst depending on the degree of carbon monoxide conversion desired. Carbon dioxide and hydrogen are produced by the reaction of the monoxide with steam.

The gas mixture from the converter is cooled and passes to a hydrogen purifying system where carbon dioxide is absorbed into amine solutions and later driven off to the atmosphere by heating the rich amine solution into the reactivator.

Since some refining processes require a minimum of carbon oxides in the product gas, the oxides are reacted with hydrogen in a methanation step. This reaction takes place in the methanator over a nickel catalyst at elevated temperatures.

Hydrocarbon impurities in the product hydrogen usually are not detrimental to the processes where this hydrogen will be used. Thus, a small amount of hydrocarbon is tolerable in the effluent gas.

Wastes

Information concerning wastes from this process are not available. However, the process appears to be a relatively clean one. In the steam

reforming subprocess a potential waste source is the desulfurization unit, which is required for feedstock that has not already been sulfurized. This waste stream would contain oil, sulfur compounds, and phenol. In the partial oxidation subprocess free carbon is removed by a water wash. Carbon dioxide is discharged to the atmosphere at several points in the subprocess.

Trends

Hydrogen requirements of the rapidly growing hydrocracking and hydrotreating processes in many instances exceed the by-product hydrogen available from catalytic reforming units. Since hydrocracking and hydrotreating are expected to grow more rapidly than other refinery processes, the demand for hydrogen manufacturing units should continue to be strong.

B. UTILITIES FUNCTION

Utility functions such as the supply of steam and cooling water generally are set up to service several processes. Boiler feed water is prepared and steam is generated in a single boiler house. Non-contact steam used for surface heating is circulated through a closed loop whereby varying quantities are made available for the specific requirements of the different processes. The condensate is nearly always recycled to the boiler house, where a certain portion is discharged as blowdown.

e three major uses of steam generated within a refinery plant are:

- 1. For non-contact process heating. In this application, the steam is normally generated at pressures of 9.5 to 45.2 atm (125 to 650 psig).
- 2. For power generation such as in steam-driven turbines, compressors, and pumps associated with the process. In this application, the steam is normally generated at pressures of 45.2 to 103 atm (650 to 1500 psig) and requires superheating.
- 3. For use as a diluent, stripping medium, or source of vacuum through the use of steam jet ejectors. This steam actually contacts the hydrocarbons in the manufacturing processes and is a source of contact process waste water when condensed. It is used at a substantially lower pressure than the foregoing and frequently is exhaust steam from one of the other uses.

Steam is supplied to the different users throughout the plant either by natural-circulation, vapor-phase systems, or by forced-circulation

liquid heat-transfer systems. Both types of systems discharge some condensate as blowdown and require the addition of boiler make-up water. The main areas of consideration in boiler operation are normally boil efficiency, internal deposits, corrosion, and the required steam quality.

Boiler efficiency is dependent on many factors. One is the elimination of boiler-tube deposition that impedes heat transfer. The main contributors to boiler deposits are calcium, magnesium, silicon, iron, copper, and aluminum. Any of these can occur in natural waters, and some can result from condensate return-line corrosion or even from make-up water pretreatment. Modern industrial boilers are designed with efficiencies on the order of 80 percent. A deposit of 0.32 cm (1/8 inch) in depth will cause a 2-3 percent drop in this efficiency, depending on the type of deposit.

Internal boiler water treatment methods have advanced to such a stage that corrosion in the steam generation equipment can be virtually eliminated. The control of caustic embrittlement in boiler tubes and drums is accomplished through the addition of sodium nitrate in the correct ratio to boiler water alkalinity. Caustic corrosion in high heat transfer boilers can also be controlled by the addition of chelating agents. This type of solubilizing internal boiler water treatment has been shown to be more effective than previous precipitation treatment using phosphate.

Other factors influencing boiler efficiency include reduction of the amount of boiler blowdown by increasing cycles of concentration of the boiler feedwater, efficiency of the blowdown heat-recovery equipment and the type of feed used.

Steam purity is of prime importance if:

- 1. The boilers are equipped with superheaters.
- The boilers supply power-generation equipment.
- 3. The steam is used directly in a process where contamination could affect product quality or destroy some material (such as a catalyst) essential to the manufacture of the product.

The minimum purity required for contact steam (or contact process water) varies from process to process. Limits for suspended solids, total solids, and alkalinity vary inversely with the steam pressure. The following tabulation summarizes boiler water concentration limits for a system providing a steam purity of 0.5 - 1.0 ppm total solids, which is required for most non-contact steam uses. It should be noted that the boiler operation must incorporate the use of antifoam agents and steam separation equipment for the concentrations shown to be valid.

Boiler Water Concentration Required to Maintain Steam Purity at 0.5 - 1.0 ppm Total Solids

Boiler Pressure, atm.

Parameters	21.4	21.5 - 31.6	31.7 -41.8	41.9 - 52.0
Total Solids (mg/l)	6,000	5,000	4,000	2,500
Suspended Solids (mg/l)	1,000	200	100	50
Total Alkalinity (mg/l)	1,000	900	800	750

Water conditioning or pretreatment systems are normally part of the utilities section of most plants. From the previous discussions, it is obvious that the required treatment may be quite extensive. Ion-exchange demineralization systems are very widely employed, not only for conditioning water for high-pressure boilers, but also for conditioning various process waters. Clarification is also widely practiced and usually precedes the ion-exchange operation.

Non-contact cooling water also is normally supplied to several processes from the utilities area. The system is either a loop which utilizes one or more evaporative cooling towers, or a once-through system with direct discharge.

Cooling towers accomplish the cooling of water circulated over the tower by moving a predetermined flow of ambient air through the tower with large fans. The air water contact causes a small amount of the water to be evaporated by the air. Thus, through latent heat transfer, the mainder of the circulated water is cooled.

Approximately 252 kg cal (1,000 BTU) are removed from the total water circulation by the evaporation of 0.454 kg (1 lb) of water. Therefore, if 45.4 kg (100 lbs) of water are introduced at the tower inlet and 0.454 kg (1 lb) is evaporated to the moving air, the remaining 44.9 kg (99 lbs) of water are reduced in total heat content by 252 kg cal (1,000 BTU), of water leaving the tower have been cooled 3.24°C/kg/kg cal (1°F/lb/BTU removed, and the exit temperature is reduced by about 5,500 (10°F). This leads to the common rule of thumb: 1 percent evaporation loss for each 5.5°C (10°F) cooling.

Since cooling is primarily by transfer of latent heat, cooling tower selection is based on the total heat content or enthalpy of the entering air. At any one enthalpy condition, the wet bulb temperature is constant. Therefore, cooling towers are selected and guaranteed to cool a specific volume of water from a hot-water temperature to a cold-water temperature while operating at a design wet-bulb temperature. Design wet-bulb temperatures vary from 15.6°C (60°F) to 35°C (85°F) depending on the geographic area, and are usually equaled or exceeded only 2.5 percent to 5 percent of the total summer operating time.

Hot water temperature minus cold water temperature is termed cooling range, and the difference between cold water and wet-bulb temperature is called approach.

A closed system is normally used when converting from once-through river cooling of plant processes. In the closed system, a cooling tower is used for cooling all of the hot water from the processes. With the closed system, make-up water from the river is required to replace evaporation loss at the tower.

Two other water losses also occur. The first is drift, which is droplet carryover in the air as contrasted to evaporative loss. The cooling tower industry has a standarized guarantee that drift loss will not exceed 0.2 percent of the water circulated. The second loss in the closed system is blowdown to sewer or river. Although blowdown is usually taken off the hot water line, it may be removed from the cold water stream in order to comply with regulations that limit the temperature of water returned to the stream. Blowdown from a tower system will vary depending on the solids concentration in the make-up water, and on the occurrence of solids that may be harmful to equipment. Generally, blowdown will be about 0.3 percent per 5.5°C (10°F) of cooling, in order to maintain a solids concentration in the recirculated water of three to four times that of the make-up water.

The quantity and quality of the blowdown form boilers and cooling towers depend on the design of the particular plant utility system. The heat content of these streams is purely a function of the heat recovery equipment associated with the utility system. The amounts of waste brine and sludge produced by ion exchange and water treatment systement and on both the plant water use function and the intake source. None of these utility waste streams can be related directly to specific process units.

Quantitative limitations on parameters such as dissolved solids, hardness, alkalinity, and temperature, therefore, cannot be allocated on a production basis. The limitations on such parameters associated with non-contact utility effluents should be established on the basis of the water quality criteria of the specific receiving water body or an EPA study of all industries, to define specific utility effluent limitations.

Refinery Distribution

There are a total of 247 operating petroleum refineries in the United States, as of January 1, 1973, with a combined capacity of 2.24 million cu m/day (14 million barrels/day of) crude oil processing (see Figure 4 and Table 7). The capacity of these plants range from 32 cu m/day (200 bbl/day) to 69,000 cu m/day (434,000 bbl/day) of crude oil.

Figure 4
Geographical Distribution of Petroleum
Refineries in the
United States

TABLE 7

Crude Capacity of Petroleum Refineries by States as of Jan. 1, 1973 (3)

Rated Crude Capacity

State	Number of Plants	Cubic Meters/day	Barrels/day
Alabama	5	6,460	40,650
Alask a	· 4	8,950	56,300
Arkans as	.4	7,600	47,800
Californi: a	34	285,675	1,796,700
Colorad <mark>o</mark>	3	8,580	53,950
Delawa re	Ī	23,850	150,000
Florida	i	875	5,500
Georgia	2	2,050	12,900
Hawa ii	2	11,290	•
Illinois	າ້າ	173,070	71,000 1,088,500
Índiana	· 7	88,250	
Kansas	τί	64,475	555,000 405,500
Kentucky	3	26,300	405,500
Louisiana	18	255,350	165,400
Maryland	2	4,040	1,606,200
Michigan	2	21,340	25,400
Minnesota	6 3 5	28,380	134,190
Mississippi) r	52,260	178,500
Missouri	5	16,880	328,700
Montana	 		106,200
Nebrask a	8	23,780	149,575
New Jersey	 	875	5,500
New Mexico	5 6 2 2	98,700	620,800
New York	0	7,740	48,700
North Dakota	2	16,850	106,000
Ohio		8,750	55,000
Oklahoma		98,850	584,000
	12	75,370	474,000
Oregon	.!	2,860	18,000
Pennsyl vania Rhode Island	1]	108,870	684,715
Tennessee		1,590	10,000
	1	4,770	30,000
Texas	40	579,640	3, 645,550
Utah Vimalala	5	19,875	125,000
Virginia	1	-7.7950	50,000
Washington	7	55,520	349,200
West Virg inia	3	3,260	20,500
Wisconsin	1	5,800	36,500
Wyoming	9	23,810	149,750
·	247	2,230,535	13,991,580

Within the United States, refineries are concentrated in areas of major crude production (California, Texas, Louisiana, Oklahoma, Kansas), and in major population areas (Illinois, Indiana, Ohio, Pennsylvania, Texas, d California).

There are an almost unlimited number of process combinations possible within the process area, or "Battery Limits", of the typical refinery. Selection of the processing route for the manufacture of a particular product mix at a particular location or time is a decision based on the particular refiner's unique situation. In order to illustrate the diversity of operations which may be included within a refinery, Figure 5 shows the schematic flow diagram for a hypothetical 15,900 cu m/day (100,000 bbl/day) integrated refinery. This hypothetical refinery includes essentially all production processes previously outlined; hence the hypothetical refinery shown in Figure 5 is completely integrated for current U.S. refinery capacity. Inspection of Table 8 demonstrates the general distribution of refining processes.

The trend in the petroleum refining industry is toward fewer and larger refineries, which are integrated with satellite or companion industries. This consolidation trend for a six-year period (1967-1973) is shown Refineries with capacities over 15,900 cu m/day (100,000 bbl/day) (11.5 percent of the total) represented 48 percent of the domestic refinery crude capacity in 1967; in 1972, 16.6 percent of the refineries had capacities of 15,900 cu m/day (100,000 bbl/day) or more, and represented 58 percent of the domestic refinery crude capacity. Growth of the large refinery was a result of the annual need for increased fuel capacity, and the imposed load due to the phasing out of aller refineries. Refineries are increasing capacities for reforming, drotreating, cracking, and isomerization processes to obtain higher octane gasoline in lieu of adding lead. Desulfurization of heavy fuels, longer process catalyst life requirements, and high quality, low sulfur, light fuels and lubes are factors in the rapid growth of the hydrogen treating process. The complexity and size of the typical refinery be expected to increase at a rate comparable to the period 1967 through 1972 for the near future, and no major technological breakthroughs expected that would drastically alter petroleum processes.

Anticipated Industry Growth

The petroleum refining industry is presently facing a shortage of crude oil. There have been scattered shortages of gasoline and fuel oil. Since demand continues to grow and very little refinery expansion work is under way, shortages will become more severe over the next few years. Consumption of petroleum products will keep growing, and supplies must be generated to satisfy these growing demands. (1972 consumption of petroleum products, shown in Table 10, was approximately 2.56 million cu m/day (16.1 million bbl/day). The growth rate in consumption has been 5.2 percent per year; the projected growth in consumption over the next

FIGURE 5
HYPOTHETICAL 100,000 BARREL/STREAM DAY INTEGRATED REFINERY

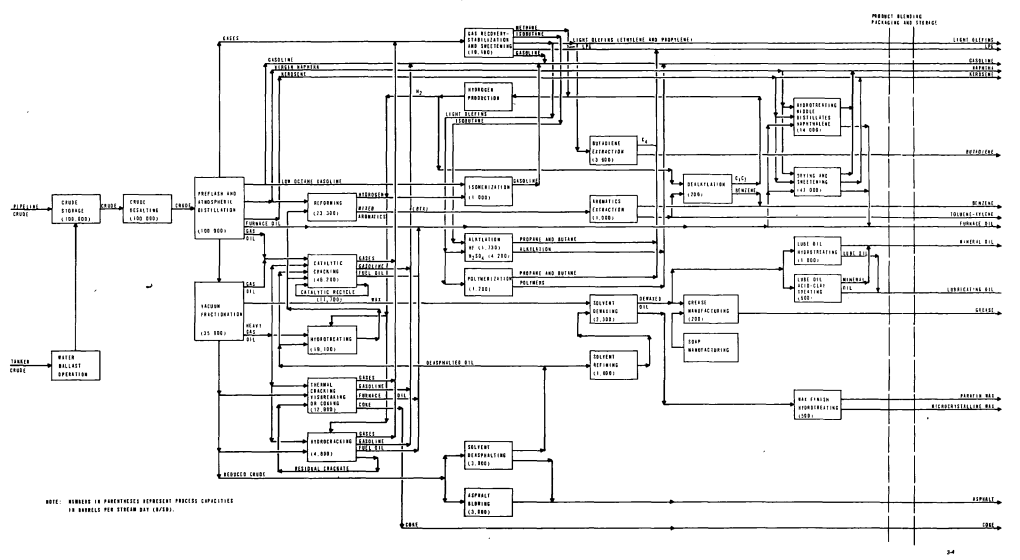


TABLE 8

Process Employment Profile of Refining Processes as of January 1, 1973 (3)

Production Processes	Number of R Employing a by Crude Ca	Producti			Percent of Refineries Employing a Production Process by Crude Capacity Classification			
	All Refineries	<35 MB/SD	35 to 100 MB/SD	>100 MB/SD	All Refineries	<35 MB/SD	35 to 100 MB/SD	>100 MB/SD
Storage: Crude & Product	247	141	65	41	100	100	100	100
Crude Desalting	247	141	65	41	100	100	100	100
Atmospheric Distillation	247	141	65	41	100	100	100	100
Vacuum Distillation	175	79	55	41	70	56	85	100
л Thermal Cracking	87	27	32	28	35	19	49	68
" Catalytic Cracking	141	41	59	41	57	29	91	100
Hydrocracking	45	11	11	23	18	8	17	56
Hydrotreating: Cat Reformer and Cat Crack Feed	-	42	52	35	52	30	80	85
Middle Distillates & Napth	a 54	11	21	22	22	8	32	54
Lubes	13	2	2	9	5	1	3	22
Heavy Oils and Residuals	5	2	3	•	2	1	5	-
Other Feedstocks	57	14	17	26	23	10	26	63
Alkylation	125	30	57	38	51	21	88	93
Isomerization	30	4	19	7	12	3	29	17
Reforming	166	65	60	41	67	46	92	100
Aromatics	35	3	16	16	14	2	25	39
Lubes	44	16	10	18	18	11	15	44
Asphalt	111	58	30	23	. 45	41	46	. 56

TABLE 9

<u>Trend in Domestic Petroleum Refining from 1967 to 1973</u> (3,3a)

	January 1, 1967	January 1, 1973	Percent Change
Crude Capacity, M ³ /SD (bb1/SD) 1,853,618	(11,657,975)	2,224,661(13,991,580)	+ 20
Total Companies	146	132	(- 10)
Total Refineries	269	247	(- 8)
Refineries with Capacity ≥ 100 Mbbl/SD	31	41	+
Refineries with Capacity <35 Mbb1/SD	159	141	(- 11)
Total Capacity of All ≥100 Mbb1/SD Refineries	5,597,300	8,167,200	+ 46
Average Refinery Capacity, M ³ /SD (bbl/S	D) 6890 (43,338)	9006 (56,646)	+ 31

TABLE 10

1972 Consumption of Petroleum Products (63)

Products	1972 Consumption, Million Cubic Meters/Day
	(Million Barrels/Day)
Motor Gasoline	1.02 (6.4)
Aviation Fuel	0.17 (1.1)
W1111 B4 . 111 .	0 (0 (0 1)
Middle Distillates	0.49 (3.1)
Residual Fuels	0.40 (2.5)
Residual rueis	0.40 (2.3)
All Other Products	0.49 (3.1)
	0.17 (3.2)

eight years is 43 percent, or a compounded growth rate of 4.6 percent per year.

Supplies of refinery feedstocks and products will show a rapid increase in imports. Table 11 indicates current and projected 1980 sources of feedstocks and products. In 1972, imports accounted for 29 percent of the total supply; in 1980, imports are projected at 55 percent of the total supply.

Refinery runs of crude oil are projected to increase from 1.86 million cu m/day (11.7 million bbl/day) in 1972 to 2.73 million cu m/day (17.2 million bbl/day) in 1980. Refinery capacity in 1972 was about 2.23 million cu m/day (14.0 million bbl/day). By 1980 the national refinery capacity must increase to 3.18 million cu m/day (20.0 million bbl/day) to satisfy the projected requirements. The need for 0.95 million cu m/day (6.0 million bbl/day) of new refinery construction for real growth, plus 0.64 million cu m/day (4.0 million bbl/day) of new construction for replacement, indicates a total of 1.59 million cu m/day (10.0 million bbl/day) of new refinery construction is required by 1980.

Because of crude supply limitations, most new refinery capacity will be designed to process higher sulfur crudes. (A partial list of analyses of crude oils from major oil fields around the world is given in Table 12.) The use of sour crude feedstock from outside the United States will require not only a change in processing equipment, but changes in in-plant waste water control and treatment operations. Some refineries currently consuming sweet crude stocks are not employing strippers to remove minimal amounts of ammonia and hydrogen sulfide from their was waters. When processing sour crude within these refineries, sour water strippers will be required prior to discharge of the waste waters to biological waste water treatment facilities. Two stage desalting will become more prevalent. Other changes will be required within the refinery to minimize corrosion, treat more sour heavy bottoms, and reduce emissions of sour gases.

TABLE 11
Sources of Supply for U.S. Petroleum Feedstocks

Source	Supply, Million Cubic Meter 1972	s/Day(Million Barrels/Day 1980 (Projected)
		•
Domestic Crude Oil Production	1.51 (9.5)	1.35 (8.5)
Domestic Natural Gas Liquids	0.27 (1.7)	0.24 (1.5)
Crude Oil Imports	0.35 (2.2)	1.38 (8.7)
Residual Fuel Imports	0.27 (1.7)	0.40 (2.5)
Other Imports	0.13 (0.8)	0.24 (1.5)
Miscellaneous Sources	0.06 (0.4)	0.08 (0.5)

TABLE 12

Characteristics of Crude Oils from Major Fields Around the World (40, 43)

Country	Gravity, API	Sulfur, Percent	Nitrogen, Percent
Abu Dhabi	39.3		
Algeria	46 - 48	0.15	
Brunei	21 - 37	0.1	
Canada			
Alberta			
Bonnie Glen	41 - 42	0.25	
Golden Spike	36 - 39	0.23	
Judy Creek	42 - 43	•	
Pembina	35 - 42	0.42	
Swan Hills	41	0.80	
Saskatchew an			
Midale	28 - 32	1.89	
Weyburn	24 - 33	2.12	
Indonesia	35	0.10	
<u> Iran</u>	31 - 38	1.12 - 1.66	
Iraq	35 - 36	1.97	
<u>Libya</u>	37 - 41	0.23 - 0.52	
Mexico			
Ebano Panu c o	12	5.38	
Naranjos-Cerro-Azul	20	3.80	
Poza Řica	35	1.77	
Peru	33.5 - 35.5	0.12	
Saudi Arabia	27 - 38	1.30 - 3.03	
United States			
Alaska			
Cook Inlet	36	0.0	
Prudhoe Bay	30.5		
Swanson River	29.7	0.16	0.203
Arkansas Smackover	22.2	2.10	·0.080
Jing GROV OF			0.000

TABLE 12 (Continued)

Country	Gravity, API	Sulfur, Percent	Nitrogen, Percent
California			
Elk Hills	22.5	0.68	0.472
Huntington Beach	22.6	1.57	0.048
Kern River	12.6	1.19	0.604
Midway-Sunset	22.6	0.94	3.00
San Ardo	11.1	2.25	0.913
Wilmington	22.1	1.44	0.515
Colorado			
Rangely	34.8	0.56	0.073
Kansas	J.10	0.30	0.075
Bemis Shutts	34.6	0.57	0.162
Louisiana	J	0.37	0.102
Bayou Sale	36.2	0.16	•
Caillou Isl.	35.4	0.23	0.040
Golden Meadow	37.6	0.18	0.040
Grand Bay	35	0.31	
Lake Barre	40.4	0.14	0.02
Lake Washington	28.2	0.37	0.146
West Bay	32.1	0.27	
Bay Marchand Blk. 2	20.2	0.46	0.071
Main Pass Blk. 69	30.6		0.009
		0.25	0.098
South Pass Blk, 24	32.3	0.26	0.068
South Pass Blk. 27	35.6 34.4	0.18	0.069
Timbalier Bay		0.33	0.081
West Delta Blk. 30	27	0.33	0.09
Mississippi	17 1	0.71	0.111
Baxterville	17.1	2.71	0.111
New Mexico	25	0.05	0.075
Vacuum	35	0.95	0.075
Oklahoma	h2 1	0.11	
Golden Trend Texas	42.1	0.11	
	22 2	0.22	0.01.1
Anahuac Conroe	33.2 37.6	0.23	0.041
Diamond M	45.4	0.15	
·	39.4	0.20	,
East Texas	31.0	0.32	0.00
Hastings Hawkins	26.8	0.15	0.02
Headlee	51.1	2.19	0.076
		<0.10	0.083
Kelly Snyder	38.6	0.29	0.066
Levelland	31.1	2.12	0.136
Midland Farms	39.6	0.13	0.080
Panhandle	40.4	0.55	0.067
Seeliason	41.3	<0.10	0.014

TABLE 12 (Continued)

Country	Gravity, API	Sulfur, Percent	Nitrogen, Percent
Tom O'Connor	31.1	0.16	0.03
Wasson	31.9	1.40	0.07
Webster	29.3	0.21	0.046
Yates	30.2	1.54	0.150
Utah			
Aneth	40.4	0.20	0.059
Venezuela			
Bachaquero	21.3	2.62	
Boscan	10.5	5.53	
Lagunillas	24.8	2.18	
Mene Grande	18.4	2.65	
Tia Juana	20.2	1.49	
Oficina	21.4	0.59	
Los Claros	10.5		

SECTION IV

INDUSTRY SUBCATEGORIZATION

Discussion of the Rationale of Subcategorization

The goal of this study is the development of effluent limitations commensurate with different levels of pollution control technology. These effluent limitations will specify the quantity of pollutants which will ultimately be discharged from a specific manufacturing facility and will be related to the quantity of raw materials consumed and the production methodology.

The diverse range of products and manufacturing processes to be covered suggests that separate effluent limitations be designated for different segments with the industry. To this end, a subcategorization of the Petroleum Refining Industry has been developed. The subcategorization is process oriented, with a delineation between subcategories based upon raw waste load characteristics in relation to the complexity of refinery operations.

Today's petroleum refinery is a very complex combination of interdependent operations and systems. In the development of a pollution profile for this industry, twelve major process categories were listed as fundamental to the production of principal oil products (see listing in Table 5). As was indicated in the qualitative pluation of refinery process flows and pollutant characteristics in the processes - crude desalting, distillation, and cracking contribute most heavily to a refinery's pollution load. It is felt that any new method of classification must recognize at least one of these process technologies.

The American Petroleum Institute (API) has developed a classification system which utilizes this technology breakdown. They have tentatively divided U.S. refineries into 5 classifications, which primarily recognize varying degrees of processing complexity and resultant distribution of products. The present API classification system is as follows:

Class	Process Complexity
A	Crude Topping
В	Topping and Cracking
С	Topping, cracking, and petrochemicals
D	"B" Category, and lube oils processing

Development of Industry Subcategorization

Age, size, and waste water treatability of refineries were considered during the subcategorization of the refining industry. However, subcategorization by age is not necessarily useful, as additions to and modifications of refineries are the industry's principal form of expansion. Since most of the technology employed within the industry is of an evoluationary nature, refinery age was not a major factor in refinery subcategorization.

While the size of a refinery is important in terms of economical waste water treatment, the control technology employed in smaller refineries need not be as sophisticated a technology to achieve parity with larger refineries within the same subcategory. Thus, size of refinery was not used as a criterion for refinery classification.

Treatability characteristics of refinery waste waters indicate that these waste waters are generally amenable to excellent degrees of removal of pollutants. Since this is an industry-wide characteristic, the proper place to evaluate the subcategorization of the industry is with the raw waste load delivered to the refinery waste water treatment plant. The 1972 National Petroleum Refining Waste Water Characterization Studies of 135 refinery API separator effluents, provides a major tool for this evaluation. Attempts to explain and justify the differences based solely on type and method of cooling, inplant pretreatment, and housekeeping practices were also fruitles However, generally speaking those refineries with good practices in a these areas did have the lower waste loadings.

In an attempt to determine the effects of process technology, a further analysis was made of the API individual or combined categories to evaluate the raw waste load as a function of the degree of cracking employed within the refinery. The operations included in degree of cracking were: thermal operations, catalytic cracking and hydroprocessing. The degree of cracking was expressed as percentage capacity of the total feedstock processing capacity within the refinery. The data for evaluating the net raw waste loads by this criteria was obtained by analyzing the raw waste load surveys supplied by refineries, literature sources, and analysis of the 1972 National Petroleum Refining Waste Water Characterization Studies.

1. The net waste load (total raw waste load minus quality of influent water) from the API categories, with corrections supplied by EPA, were statistically analyzed; determining the 50 percent probability-of-occurrence loading for the key waste water parameters (BOD5, Oil/Grease, Phenol, and Ammonia). These parameters are representative of the major contaminants discharged by refineries and therefore, could serve as a valid

basis for screening correlations of variations in oil separator performance, housekeeping, severity of cracking, and other factors. The 50 percent probability-of-occurrence numbers were chosen, as they reflect the median net raw waste load performance of the entire subcategory.

2. A subcategory-to-subcategory comparison was then made to determine if, based upon these median levels of performance, significant differences in waste water loadings between subcategories of refineries existed. By comparing the median values on a subcategory-to-subcategory basis, internal differences in separator performance, housekeeping, and other factors are minimized. Those subcategories which exhibited a high degree of similarity in median net raw waste loads for the key parameters were then combined and re-analyzed to develop new median values for the combinations.

In an additional attempt to determine the effects of process technology, an analysis was made of the lube oil manufacturing refineries. It was found that these refineries split into two separate groups; (1) large, complex refineries with a low overall percentage lube production, and (2) small, specialty lube refineries. The first group is made up of 27 refineries, with capacities of 36,000 bbl/day and higher and lube productions from 0.6 - 9.4 percent of the feedstock throughput. The second group includes 18 refineries ranging in capacity from 1,000 - 10,400 bbl/day and lube productions from 14 - 100 per cent of feedstock throughput.

bcategorization Results

Using the procedures outlined above, many trials were performed in order to obtain a subcategorization of the petroleum refining industry which is reflective of the net raw waste load with respect to type of refinery (function), process technology employed, and severity of operations. The final subcategorization obtained from this analysis is indicated below in Table 13. Detailed probability plots for the development of the subcategorization are contained in supplement B.

For each of these new subcategories the parameters for the selected median values are indicated in Table 14. A further enumeration of overall net raw waste load characteristics is given in Section V.

In addition to the subcategorization made by raw waste load, the splitting out of the speciality lube plants (as outlined above) is being made because of the special nature of these plants.

Analysis of the Subcategorization

Topping subcategory

Table 13

Subcategorization of the Petroleum Refining Industry Reflecting Significant Differences in Wastewater Characteristics

Subcategory Basic Refinery Operations Included

Topping and Catalytic reforming

Low-Cracking Topping and cracking, with fresh feed

(non-recycle) to the cracking and hydroprocessing of less than 50% of the feed-

stock throughout.

High-Cracking Topping cracking, with a fresh feed

(non-recycle to the cracking and hydroprocessing of greater than 50% of the

feedstock throughout.

Petrochemical Topping, cracking and petrochemicals

operations.*

Lube Topping, cracking and lubes.**

Integrated Topping, cracking, lubes and petrochemicals

operations.*

* Petrochemical operations - Production of greater than 15 % of the feedstock throughout in first generation petrochemicals and feedstock isomerization products (BTX, olefins, cyclohexane, etc.) and/or production of second generation petrochemicals (cumene alcohols, ketones, etc.)

** Lubes - the production less than 12% of the feedstock throughout as lubes. Refineries with greater than 12% lubes are being considered speciality refineries and are to be handled on an individual basis.

TABLE 14

Median Net Raw Waste Loads from Petroleum Refining *

Industry Categories

	<u>Median Net</u>	Raw Waste Lo	ad, kg/1000 m ³	(1b/1000 BBL)
Subcategory	BOD ₅	Oil/Grease	<u>Phenol</u>	<u>Ammonia</u>
Topping	7.1 (2.5)	5.1 (1.8)	0.029 (0.01)	1.43 (0.5)
Low-Cracking	71.3 (25)	27.4 (9.6)	2.85 (1.0)	10.0 (3.5)
High-Cracking	82.7 (29)	31.4 (11)	5.1 (1.8)	32.8 (11.5)
Petrochemical	148.4 (52)	45.6 (16)	10.3 (3.6)	34.2 (12.0)
Lube	184.3 (65)	136.1 (48)	6.2 (2.2)	22.1 (7.8)
Integrated	215.5 (76)	131.8 (46.5)	5.1 (1.8)	35.4 (12.5)

^{*} Supplement B contains probability plots containing distributions of individual parameters for each category.

The topping subcategory is similar to the previous API category A. Refineries in this subcategory are relatively simple in operation, operating only crude oil distillation or topping units and catalytime reforming. These processes are common to all other subcategories.

High & low cracking subcategories

API Category B includes refineries which contain topping, reforming, and cracking operations. Also included are all first generation conventional refinery-associated products or intermediates, such as benzene-toluene-xylene (BTX), alkanes, alkenes, alkynes, and other miscellaneous items such as sulfur, hydrogen, coke, and ammonia.

Category B has been subdivided on the basis of degree of cracking. The primary differentiation between the high and low cracking subcategories is in the degree of cracking operations performed on the feedstock. An analysis of all refineries for which cracking data was available was made in order to determine if the amount of cracking employed within the refinery had a demonstrable effect upon the net raw waste load. No direct correlation, relating percent cracking of crude to the resultant raw waste load, was obtained. However, a breakpoint appeared at approximately 50 percent cracking, based on feedstock charge.

The division of API category B refineries into two subcategories, high and low cracking, using 50 percent cracking as the breakpoint, was made in order to more accurately reflect actual raw waste load conditions within this portion of the petroleum refining industry. While this division of the refineries is not the ultimate answer to subcategorization, it nevertheless recognizes the existence of an inhered difference in raw waste loads which result from the increasing complexity of refinery operations in API category B.

Another change from the API classification, is that the inclusion of first generation petrochemicals shall only be for those whose production amounts to less than 15 percent of the refinery throughput.

Petrochemical subcategory

The petrochemical subcategory is similar to the API category C. Operations included within this subcategory are topping, cracking, and petrochemical operations. Petrochemical operations include first generation conventional refinery-associated production, as described under high and low cracking and category B, but only when it amounts to greater than 15 percent of the refinery throughput. This takes into consideration the additional cooling tower blowdown from this operation. Intermediate chemical production, including such typical products as cumene, phthalic anhydride, alcohols, ketones, trimer, and styrene, shall also be considered petrochemical operations and classify a refinery in this subcategory.

Attempts to correlate degree of cracking data for the petrochemical subcategory refineries were unsuccessful. Only a small data base was vailable for analysis, and no conclusions could be drawn from this udy as to the effect of increased cracking on the petrochemical subcategory refinery raw waste loads. The analysis is also probably overshadowed by the presence of petrochemicals operations within these refineries.

Lube subcategory

The new lube subcategory is the same as the API category D, except that those refineries with greater than 12 percent of their throughput going to lube production have been split out for separate consideration, with limits to be set at a later date.

In the lube subcategory, the operations included under the high and low cracking subcategories are expanded to include lube operations. Lube operations in this subcategory require the production of lube oil blending stocks via operations such as dewaxing, lube hydrotreating, or clay treatment. Lube oil processing excludes formulating blended oils and additives.

Again, for the new lube subcategory, no correlation or breakpoint for degree of cracking was observable. This may be due in part to: the small data base; the presence of lube operations; or the size and complexity of refineries within this subcategory.

Integrated subcategory

The new integrated subcategory is the same as API category E, except for the new definition of petrochemical operatons specified in the petrochemical subcategory.

Conclusion

The subcategorization of the petroleum refinery industry presented above allows for the definition of logical segments of the industry in terms of factors which effect generated API separator effluent waste water quality. It allows for rapid identification of the expected median net raw waste loads as a basis for developing effluent guidelines for the discharge from the individual refinery. The subcategorization determined above is used throughout this report as the basis for development of effluent limitations and guidelines.

SECTION V

WASTE CHARACTERIZATION

General

an understanding of the fundamental production developing processes and their inter-relationships in refinery operations. the best method of characterizing determination of refinerv discharges will enhance the interpretation of the industry pollution profile. If unit raw waste loads could be developed for each production process, then the current effluent waste water profile could obtained by simply adding the components, and future profiles by projecting the types and sizes of refineries. However, the information required for such an approach is not available. Essentially all of the available data on refinery waste waters apply to total API separator effluent, rather than to effluents from specific processes.

Another factor detracting from the application of a summation of direct subprocess unit raw waste loads, is the frequent practice of combining specific waste water streams discharging from several units for treatment and/or reuse. Thus, such streams as sour waters, caustic washes, etc., in actual practice are generally not traceable to a specific unit, but only to a stripping tower or treatment unit handling wastes from several units. The size, sequence, and combination of contributing processes are so involved that a breakdown by units would be extremely difficult to achieve.

In view of the limitations imposed by the summation of waste water data from specific production process, the evaluation of refinery waste loads was based on total refinery effluents discharged through the API (Oil) separator, which is considered an integral part of refinery process operations for product/raw material recovery prior to final waste water treatment.

Raw Waste Loads

The information on raw waste loading was compiled from the 1972 National Petroleum Refining Waste Water Characterization Studies and plant visits. The data are considered primary source data, i.e., they are derived from field sampling and operating records. The raw waste data for each subcategory of the petroleum refining industry, as subcategorized in Section IV, have been analyzed to determine the probability of occurrence of mass loadings for each considered parameter in the subcategory. These frequency distributions are summarized in Tables 15 through 20 for each subcategory.

Waste water Flows

As shown in Table 15 through 20, the waste water flows associated with raw waste loads can vary significantly. However, the loadings of pollutants tend to vary within fairly narrow limits, independent flow.

Since the inter-refinery data suggest that the pollutant loading to be expected from a refinery is relatively constant in concentration, an examination of water use practices was made. The waste water flow frequencies reported in Tables 15 through 20 are dry-weather flows, and in many cases include large amounts of once-through cooling water. Refineries with more exemplary waste water treatment systems are probably making a greater effort to control waste loads and flows. Conversely, refineries with very high water usages and/or raw waste loads either do not have identifiable waste water treatment plants, or have them under construction.

The primary methods for reduction of the waste water flows to the API separator are either segregation of once-through cooling waters, or by installation of recycle cooling towers and/or air coolers. In order to estimate the flows that should be attainable in refineries with good water practices, a statistical analysis was made of flows from refineries in which 3 percent or less of the total heat removal load is accomplished by once-through cooling water. Data for this analysis were obtained from the tabulation of refinery cooling practices contained in the 1972 National Petroleum Refining Waste Water Characterization Studies. These frequency distributions are summarized in Table 21.

Basis for Effluent Limitations

The median (50 percent probability-of-occurrence) raw waste loads outlined in Tables 15 through 20 are reflective of the performance of median refineries within each subcategory. At the same time, attainable process waste water flows, as reflected by the median water usage for refineries in which 3 percent or less of the total heat removal load is accomplished by once-through cooling water, are indicative of equitable process waste water loadings which require waste water treatment.

Consequently, these median (50 percent probability-of-occurrence) waste water loadings and estimated process waste water flows were selected as one basis for developing effluent limitations, and are used in subsequent sections to define these effluent limitations.

TABLE 15 Topping Subcategory Raw Waste Load** Effluent from Refinery API Separator Net Kilograms/1000 m³ (LB/1000 bbls) of Feedstock Throughput

Parameter			•	Occurrence,	
Tar and Co	-			n or equal to	
BOD ₅	0.26	(0.09)	7.1	(Median) (2.5)	90% 286 (100)
COD	0.72	(0.25)	24.0	(8.4)	858 (300)
TOC	0.086	(0.03)	4.9	(1.7)	269 (98)
TSS	0.30	(0.105)	6.6	(2.3)	123 (43)
	0.97	(0.34)	5.1	(1.8)	45.8 (16)
Phenols	0.001	(0.00035)	0.029	(0.01)	0.92(0.32)
Ammonia	0.18	(0.064)	1.43	(0.5)	11.4 (4.0)
Sulfides	0.0037	(0.0013)	1.57	(0.55)	7.2 (2.5)
Flow *	0.024	(1.0)	0.53	(22)	12.0 (500)

^{* 1000} cubic meters/1000 m³ Feedstock Throughput (gallons/bbl)
** Probability plots are contained in Supplement B.

TABLE 16 Low-Cracking Subcategory Raw Waste Load** Effluent from Refinery API Separator Net Kilograms/1000 m³ (LB/1000 bbls) of Feedstock Throughput

Probability of Occurrence,

Parameter	Percen	it less than or equal to	
	10%	<u>50%</u> (Median)	90%
BOD ₅	15.7 (5.5)	71.5 (25)	343 (120)
COD	54 (19)	200 (70)	744 (260)
тос	7.4 (2.6)	45.7 (16)	286 (100)
TSS	3.7 (1.3)	27 (9.6)	200 (70)
Oily	5.4 (1.9)	27 (9.6)	137 (48)
Pheno1 s	0.36 (0.125)	2.86 (1.0)	22.9(8.0)
Ammonia	1.9 (0.67)	10.0 (3.5)	51.5(18)
Sulfides	0.00049(0.00017)	1.0 (0.35)	22,308(7800)
			(10(070)
Flow	0.10 (4.3)	0.79 (33)	6.48(270)

^{* 1000} Cubic Meters/1000 m³ Feedstock Throughout (gallons/bbl)
** Probability plots are contained in Supplement B.

TABLE 17
High Cracking Subcategory Raw Waste Load

Effluent from Refinery API Separator

Net Kilograms/1000 m³ (LB/1000 bbls) of Feedstock Throughput

Probability of Occurrence, Percent less than or equal to

rarameter	Perc	cent ress than or equal to	
	10%	50% (Median)	90%
BOD ₅ .	28.0 (9.8)	82.9 (29)	235 (82)
COD	45.8 (16)	260 (91)	1487 (520)
	7.4 (2.6)	52.9 (18.5)	372 (130)
TSS	0.054 (0.019)	32.3 (11.3)	212 (74)
Oil	6.7 (2.35)	31.4 (11.0)	120 (42)
Phenols	0.57 (0.20)	5.1 (1.8)	48.6(17)
Ammonia	6.6 (2.3)	32.8 (11.5)	166 (58)
Sulfides	0.0049(0.0017)	1.28(0.45)	315 (110 ₎
Flow	0.12 (4.8)	0.62(26)	48.0(2000)

^{*1000} Cubic Meters/1000 m³ Feedstock Throughput (gallons/bbl)
** Probability plots are contained in Supplement B.

TABLE 18

Petrochemical Subcategory Raw Waste Load **

Effluent from Refinery API Separator

Net Kilograms/1000 m³ (LB/1000 bbls) of Feedstock Throughput

Probability of Occurrence,

Parameter	Per	cent less than or equal t	0
	10%	<u>50%</u> (median)	90%
BOD ₅	34.3 (12)	149 (52)	629 (220)
COD	137 (48)	372 (130)	2888 (800)
TOC	31.5 (11)	117 (41)	443 (155)
TSS	4.0 (1.4)	44.3(15.5)	887 (310)
Oily	7.4 (2.6)	45.8(16)	315 (110)
Phenols	2.2 (0.78)	10.3(3.6)	48.6(17)
Ammonia	6.3 (2.2)	34.3(12)	189 (66)
Sulfides	0.011(0.004)	1.69(0.59)	229 (80)
Flow*	0.11 (4.6)	0.96(40)	12.0(500)

^{* 1000} Cubic Meters/1000 m³ Feedstock Throughput (gallons/bbl)

^{**} Probability polots are contained in Supplement B.

TABLE 19

Lube Subcategory Raw Waste Load**

Effluent from Refinery API Separator

Net Kilograms/1000 m³ (LB/1000 bbls) of Feedstock Throughput

Probability of Occurrence,

Parameters	Percen	t less than or equal to	
10.000013	10%	<u>50%</u> (median)	90%
BOD ₅	113 (40)	187 (66)	311 (110)
COD	200 (71)	382 (134)	1750 (618)
TOC	-	- -	-
TSS	3.4 (1.2)	79 (28)	1769 (625)
	48 (17)	136 (48)	396 (140)
Phenolics	0.2 (0.07)	6.2 (2.2)	20 (7.0)
Ammonia (N)	6.2 (2.2)	22 (7.8)	79 (28)
Sulfides	0.00017 (0.00006)	1.1 (0.4)	35.0 (12.4)
Flow	0.56(24.2)	0.91(39)	13.0 (560)

^{* 1000} Cubic Meters/1000 m³ Feedstock Throughput (gallons/bbl)

^{**} Probability plots are contained in Supplement B.

Integrated Subcategory Raw Waste Load**

Effluent from REfinery API Separator

Net Kilograms/1000 M³ (LB/1000 BBLs) of Feedstock Throughput

Probability of Occurrence, Percent less than or equal to Parameters 10% 50%(median) 45 (16) (84) 413 (146)BOD₅ 238 (406)COD 120 (42.4)-590 (208)1150 TOC 0.6(0.2)(10.2)340 (120)29 TSS 011 23.8 (8.4) 133 (47) 750 (26.5)(0.35)6.5 (2.3) 41 (14.5)Phenolics 100 Ammonia (N) 7.4(2.6)35.4 (12.5) (60) 170 Sulfides 0.00028 (0.00010) 1.7 (0.6) 45 (15.9)0.23(10.0) 1.8 (79) 25.6 (1100) Flow*

^{* 1000} Cubic Meters/1000 m³ Feedstock Throughput (gallons/bbl)

^{**} Probability plots are contained in Supplement B.



Wastewater Flow from Petroleum Refineries* Using 3 Percent or Less Once-Through Cooling Water for Heat Removal 1000 M³ / 1000 M³ (gallons/bbl) of Feedstock Throughput

Probability of Occurrence, Percent Less than or Equal to

Subcategory	10%	50% (median)	90%
Topping	0.046 (1.9)	0.29 (12)	2.16 (90)
Low-Cracking	0.18 (7.6)	0.41 (17)	1.46 (61)
High-Cracking	0.21 (8.8)	0.50 (21)	5.52 (230)
Petrochemical	0.19 (8.0)	0.60 (25)	1.58 (66)
Lube	0.53 (22)	0.89 (37)	1.25 (52)
Integrated	0.65 (27)	1.11 (46)	8.80 (365)

^{*} Probability plots are contained in Supplement B.

SECTION VI

SELECTION OF POLLUTANT PARAMETERS

Selected Parameters

The selection of the complete list of pollutant parameters which are discharged in significant quantities was based on a review of: the Environmental Protection Agency permits for discharge of waste waters from a number of refineries; reviews with personnel in regional EPA 1972 offices: the National Petroleum Refining Waste Characterization Studies; discussions with industry representatives and consultants; and literature survey data. The results of the above indicated the parameters shown in Table 22 are significant in describing the physical, chemical and biological characteristics of waste waters discharged by the petroleum refining industry, as defined in the Act.

The rationale and justification for inclusion of these parameters are discussed below. This discussion will provide the basis for selection of parameters upon which the actual effluent limitations were postulated and prepared. In addition, particular parameters were selected for discussion in the light of current knowledge as to their limitations from an analytical as well as from an environmental standpoint.

Oxygen Demand Parameters

Three oxygen demand parameters are discussed below: BOD5, COD, and TOC. should be noted that since separate limitations are specified for BOD5, COD, and TOC in sections IX, X, and XI for each subcategory.

Almost without exception, waste waters from petroleum refineries exert a significant and sometimes major oxygen demand. The primary sources are soluble biodegradable hydrocarbons and inorganic sulfur compounds. Crude distillation, cat cracking, and the product finishing operations, are the major contributors of BOD5. In addition, the combination of small leaks and inadvertent losses that occur almost continuously throughout a complex refinery can become principal BOD_ pollution sources.

Biochemical oxygen demand (BOD) refers to the amount of oxygen required to stabilize biodegradable matter under aerobic conditions. The BOD5 test has been used to gauge the pollutional strength of a waste water in terms of the oxygen it would demand if discharged into a watercourse. Historically, the BOD5 test has also been used to evaluate the performance of biological waste water treatment plants and to establish effluent limitation values. However, objections to the use of the BOD5 test have been raised.

TABLE 22

Significant Pollutant Parameters for the Petroleum Refining Industry

Biochemical Oxygen Demand (BOD5)

Chemical Oxygen Demand (COD)

Total Organic Carbon (TOC)

Oil and Grease (O&G)

Ammonia as Nitrogen (NH3-N)

Phenolic Compounds

Sulfides

Chromium

Zinc

The major objections are as follows:

- 1. The standard BOD5 test takes five days before the results are available, thereby negating its use as a day-to-day treatment plant operational indicator.
- 2. At the start of the BOD5 test, seed culture (microorganisms) is added to the BOD5 bottle. If the seed culture was not acclimated, i.e., exposed to a similar waste water in the past, it may not readily be able to biologically degrade the waste. This results in the reporting of a low BOD5 value. This situation is very likely to occur when dealing with complex industrial wastes, for which acclimation is required in most cases. The necessity of using "acclimated bacteria" makes it very important to take a seed from the biological plant treating the waste or downstream of the discharge in the receiving waterbody.
- 3. The BOD5 test is sensitive to toxic materials, as are all biological processes. Therefore, if toxic materials are present in a particular waste water, the reported BOD5 value may very well be erroneous. This situation can be remedied by running a toxicity test, i.e., subsequently diluting the sample until the BOD5 value reaches a plateau indicating that the material is at a concentration which no longer inhibits biological oxidation.

There has been much controversy concerning the use of BOD5 as a measure pollution, and there have been recommendations to substitute some ther parameter, e.g., COD or TOC. EPA has recently pointed out that some or all of the previously cited reasons make the BOD5 test a non-standard test, and ASTM's Subcommittee D-19 has also recommended withdrawal of the BOD5 test as a standard test.

However, some of the previously cited weaknesses of the BOD5 test also make it uniquely applicable. It is the only parameter now available which measures the amount of oxygen used by selected microorganisms in metabolizing a waste water. The use of COD or TOC to monitor the efficiency of BOD5 removal in biological treatment is possible only if there is a good correlation between COD or TOC and BOD. After consideration of the advantages, disadvantages and constraints, BOD5 will continue to be used as a pollutional indicator for the petroleum refining industry.

Typical raw waste load concentrations for each subcategory are listed below:

Subcategory	BOD5 RWL Range, mg/l
Topping Low Cracking	10 - 50 30 - 300
High Cracking	100 - 600
Petrochemical	50 - 800
Lube	100 - 700
Integrated	100 - 800

As a matter of reference, typical BOD5 values for raw municipal waste waters range between 100 and 300 mg/L.

COD

Chemical oxygen demand (COD) provides a measure of the equivalent oxygen required to oxidize the materials present in a waste water sample, under acid conditions with the aid of a strong chemical oxidant, such as potassium dischromate, and a catalyst (silver sulfate). One major advantage of the COD test is that the results are available normally in less than three hours. Thus, the COD test is a faster test by which to estimate the maximum oxygen exertion demand a waste can make on a stream. However, one major disadvantage is that the COD test does not differentiate between biodegradable and non-biodegradable organic material. In addition, the presence of inorganic reducing chemicals (sulfides, reducible metallic ions, etc.) and chlorides may interfere with the COD test.

The slow accumulation of refractory (resistant to biological decomposition) compounds in watercourses has caused concern among various environmentalists and regulatory agencies. However, until these compounds are identified, analytical procedures developed to quantify them, and their effects on aquatic plants and animals are documented, it may be premature (as well as economically questionable) to require their removal from waste water sources.

Typical raw waste load concentrations for each subcategory are listed below:

Subcategory	COD RWL	Range,	mg/l
Topping	50 -	150	
Low Cracking	150 -	300	
High Cracking	150 -	400	
Petrochemical	300 -	600	
Lube	400 -	700	
Integrated	300 -	600	

Typical COD values for raw municipal waste waters are between 200 mg/l and 400 mg/l.

* TOC

tal organic carbon (TOC) is a measure of the amount of carbon in the organic material in a waste water sample. The TOC analyzer withdraws a small volume of sample and thermally oxidizes it at 150°C. The water vapor and carbon dioxide from the combustion chamber (where the water vapor is removed) is condensed and sent to an infrared analyzer, where the carbon dioxide is monitored. This carbon dioxide value corresponds to the total inorganic value. Another portion of the same sample is thermally oxidized at 950°C, which converts all the carbonaceous material to carbon dioxide; this carbon dioxide value corresponds to the total carbon value. TOC is determined by subtracting the inorganic carbon (carbonates and water vapor) from the total carbon value.

The recently developed automated carbon analyzer has provided a rapid and simple means of determining organic carbon levels in waste water samples, enhancing the popularity of TOC as a fundamental measure of pollution. The organic carbon determination is free of many of the variables which plague the COD and BOD analyses, yielding more reliable and reproducible data. However, meaningful correlations between the three are sometimes hard to develop.

Typical raw waste concentrations for each subcategory are presented below:

Subcategory	TOC RWL Range, mg/1
Topping	10 - 50
Low Cracking	50 - 100
High Cracking	50 - 500
Petrochemical	100 - 250
Lube	100 - 400
Integrated	50 - 500

Typical values for raw municipal waste waters range between 50 $\,$ and 250 $\,$ mg/L.

TSS

In refineries, major sources of suspended matter are contributed by crude storage, alkylation, crude desalting and finishing operations. Quenching and removal operations in the production of coke can contribute significant amounts of suspended fines to the refinery effluent.

Total suspended solids, when discharged to a watercourse, settle to the bottom and can blanket spawning grounds and interfere with fish propagation. In addition, the solids which are organic will be metabolized and exert an oxygen demand. Total suspended solids, in large concentrations, can impede light transmittance and interfere with

aquatic photosynthesis, thereby affecting the oxygen content of a body of water.

Typical total suspended solids raw waste concentrations for subcategory are listed below:



Subcategory	TSS RWL Range, mg/l
Topping	10 - 40
Low Cracking	10 - 70
High Cracking	20 - 100
Petrochemical	50 - 200
Lube	80 - 300
Integrated	20 - 200

Total suspended solids concentrations for typical raw municipal waste waters range from 100 to 300 mg/l.

Freon Extractables - Oil and Grease

No solvent is known which will directly dissolve only oil or grease, thus the manual "Methods for the Chemical Analysis of Water and Wastes 1971" distributed by the Environmental Protection Agency states that their method for oil and grease determinations includes the freon extractable matter from waters.

In the petroleum refining industry, oils, greases, various other hydrocarbons and some inorganic compounds will be included in the extraction procedure. The majority of material removed by the procedure in a refinery waste water will, in most instances, be of a hydrocard These hydrocarbons, predominately oil and grease nature. compounds, will make their presence felt in the COD, TOC, TOD, and usually the BOD tests where high test values will result. The oxygen demand potential of these freon extractables is only one of the detrimental effects exerted on water bodies by this class of compounds. The water insoluble hydrocarbons and free floating emulsified oils in a waste water will affect stream ecology by interfering with oxygen transfer, by damaging the plumage and coats of water animals and fowls, and by contributing taste and toxicity problems. The effect of oil spills upon boats and shorelines and their production of oil slicks and iridescence upon the surface of waters is well known. The average freon extractable material recorded by a refinery survey for effluent waters from the refineries ranged from a maximum of 37 mg/l to a minimum of 4. mq/1.

Grease is defined in "Webster's Third New International Dictionary" as a thick lubricant. The class of refinery products known as greases are usually included in the freon extractable portions of a water analysis. Some thick heavy petroleum products coat the silt and sediment of a stream bottom samples which have been contaminated by oily products over a long period. An infrared scan of such an extract done on bottom

w automative grease. Such bottom contamination can, of course, exert influence upon the aquatic life of a stream, estuary, bay or other water dy. Typical oil and grease concentrations for each subcategory are listed below:

Subcategory	Freon Extractables as Oil and Grease RWL Range, mg/l
Topping Low Cracking High Cracking Petrochemical Lube Integrated	10 - 50 15 - 150 30 - 300 20 - 250 40 - 400 20 - 500

Ammonia as Nitrogen

Ammonia is commonly found in overhead condensates from distillation and cracking and from desalting. It is usually found combined with sulfide as an ammonium sulfide salt. The presence of even small amounts of ammonia in surface waters contributes to eutrophication - the growth of algae. Large growths of algae are unsightly, often interfere with swimming and boating, impart tastes and odors to water, and when they die in the early fall add a substantial organic load to the stream. Ammonia may exert a toxic effect on aquatic life which is usually more pronounced at a high pH value.

monia nitrogen is also the nitrogen and energy source for autotrophic rganisms (nitrifiers). The oxidation of ammonia to nitrite and then nitrate has a stoichiometric oxygen requirement of approximately 4.6 times the concentration of NH3-N. The nitrification reaction is much slower than the carbonaceous reaction and therefore, the dissolved oxygen utilization is observed over a much longer period.

Typcial ammonia as nitrogen raw waste concentrations for each subcategory are listed below:

Subcategory	NH3 - N RWL Range, 1	ng/l
Topping	0.05 - 20	
Low Cracking	0.5 - 200	
High Cracking	2 - 200	
Petrochemical	4 - 300	
Lube	1 - 120	
Integrated	1 - 250	

Phenolic Compounds

Catalytic cracking, crude distillation, and product finishing and treating, are the major sources of phenolic compounds. Catalytic cracking produces phenols by the decomposition of multi-cyclicaromatics such as anthracene and phenanthrene. Some solvent refining process use phenol as a solvent and although it is salvaged by recovery processes, losses are inevitable.

Phenols in waste water present two major problems: 1) at high concentrations, phenols act as bactericides, and 2) at very low concentrations, when disinfected with chlorine, chlorophenols are formed producing taste and odor. Past experience has indicated that biological treatment systems may be acclimated to phenol concentrations of 300 mg/l or more. However, protection of a biological treatment system against slug loads of phenol should be given careful consideration.

Typical phenolic raw waste concentrations for each subcategory are listed below:

Subcategory	Phenolics, RWL Range, mg/l
Topping	0-200
Low Cracking	0-20
High Cracking	0-100
Petroleum	0.5-50
Lube	0.1-25
Integrated	0.5-50

Sulfides

In the petroleum refining industry, major sources of sulfide wastes are crude desalting, crude distillation and cracking processes. Sulfides cause corrosion, impair product quality, and shorten the useful catlyst life. They are removed by caustic, diethanciamine, water or steam, or appear as sour condensate waters in these initial processing operations. Hydrotreating processes can be used to remove sulfides in the feedstock. Most removed and recovered sulfide is burned to produce sulfuric acid or elemental sulfur.

When present in water, soluble sulfide salts can reduce pH; react with iron and other metals to cause black precipitates; cause odor problems; and can be toxic to aquatic life. The toxicity of solutions of sulfides to fish increases as the pH value is lowered. Sulfides also chemically react with dissolved oxygen present in water, thereby lowering dissolved oxygen levels.

Typical sulfide raw waste concentrations for each subcategory are listed below:

Subcategory

Sulfide, RWL Range, mg/l

Topping	0~5
Low Cracking	0-400
High Cracking	0-20
Petroleum	0-200
Lube	0-40
Integrated	0-60

Total Chromium

Chromium may exist in water supplies in both the hexavalent and trivalent state. Chromium salts are used extensively in industrial processes and chromate compounds are frequently added to cooling water for corrosion control. The toxicity of chromium salts toward aquatic life varies widely with the species, temperature, pH, concentration, and synergistic or antagonisite effects of other water constituents, especially hardness.

It appears that about 1 mg/l of chromium is toxic to fish life in any water. A limit of .05 mg/l for hexavalent chromium is set by the USPHS Drinking Water Standards of 1962. A survey of refinery effluents sampled across the U.S. produced chromium values ranging from .02 to 1.45 mg/l. The median figure found for chromium in the effluents was .26 mg/l.

Hexavalent Chromium

hexavalent chromium content of potable water supplies within the U.S. has been reported to vary between 3 to 40 micrograms per liter. In the +6 oxidation state; chromium is usually combined with oxygen in the form of the oxide, chromium trioxide Cr03 or the Oxyanions chromate Cr04= and dichromate Cr207. Chromates will generally be present in a refinery waste stream when they are used as corrosion inhibitors in cooling water.

Zinc

Zinc is an essential and beneficial element in human metabolism when its intake to an organism is limited. At higher amounts zinc can lead to gastrointestinal irritation and large amounts of the metal have been reported to upset trickling filter and activated sludge waste treatment processes.

Zinc may also affect toxicity of an effluent water through its synergistic effects on other ions present, although research on such effects have been limited. Zinc's reported toxicity to fish varies from about .1 to 1.0 mg/l. Calcium content of the water is said to directly affect this toxicity.

Zinc compounds can be used as corrosion inhibitors for cooling water. In addition, zinc is produced in the combustion of fossil fuels and may find its way into refining waters by leaching processes.

A survey of effluents from petroleum refineries across the U.S. reports zinc concentrations of .04 to 1.84 mg/l in the effluent waters. The median concentration of zinc found in the effluents was .16 mg/l.

Other Pollutants

Other pollutants which were examined in this study of refining waste water practices included: total dissolved solids, cyanide, pH (acidity and alkalinity), temperature, various metallic ions, chloride, fluoride and phosphates.

It was determined that these parameters are generally found in refineries in small enough amounts as not to warrant accross the board treatment. Restrictions on these parameters may be required as a result of water quality requirements.

TDS

Dissolved solids in refinery waste waters consist mainly of carbonates, chlorides, and sulfates. U.S. Public Health Service Drinking Water Standards for total dissolved solids are set at 500 mg/L on the basis of taste thresholds. Many communities in the United States use water containing from 2,000 to 4,000 mg/l of dissolved solids. Such waters are not palatable and may have a laxative effect on certain people However, the geographic location and availability of potable water will dictate acceptable standards. The following is a summary of a literature survey indicating the levels of dissolved solids which should not interfere with the indicated beneficial use:

Domestic Water Supply	1,000 mg/l
Irrigation	700 mg/l
Livestock Watering	2,500 mg/l
Freshwater Fish and Aquatic Life	2.000 mg/1

Median total dissolved solids concentrations for refinery effluents are 400-700 mg/L. The extensive amount of process water recycle and reuse is primarily responsible for these high concentrations.

Because dissolved solids concentrations are intimately tied to process recycle and the quality of the process raw water source; it is recommended that this parameter be dictated by local water quality requirements.

Cyanides

U.S. Public Health Service Drinking Water Standards set a cyanide value of 0.01 mg/L of CN-. Cyanides, although toxic at high concentrations, letteriorate by bacterial action at lower concentrations.

Cyanide raw waste load data for the refining industry show median values of 0.0 - 0.18 mg/L for the six subcategories. Only occasionally are any values found above 1.0 mg/l. At these concentration ranges, no inhibition is expected in biological waste facilities. Consequently, the values are such that specific limitations are not required. Cyanides are on the EPA toxic materials list and limitations based on health effects will be made available at a later date.

pH (Acidity and Alkalinity)

The acidity of a waste is a measure of the quantity of compounds contained therein which will dissociate in an aqueous solution to produce hydrogen ions. Acidity in petroleum refining waste waters can be contributed by both organic and inorganic compound dissociation. Most mineral acids found in waste waters (sulfuric acid, hydrochloric acid, nitric acid, phosphoric acid) are typically strong acids. The most common weaker acids found include the organic acids such as carboxyl and carbonic.

Compounds which contribute to alkalinity in waste waters are those which dissociate in aqueous solutions to produce hydroxyl ions. Alkalinity is often defined as the acid-consuming ability of the waste water and is measured by titrating a given volume of waste with standard acid until of the alkaline material has reacted to form salts. In effect, talinity is the exact opposite of acidity; high alkalinities lower the hydrogen ion concentration of a solution and raise its pH.

Most refinery waste waters are alkaline due to the presence of ammonia and the use of caustic for sulfur removal. Cracking (both thermal and catalytic) and crude distillation are the principal sources of alkaline discharges. Alkylation and polymerization utilize acids as catalysts and produce severe acidity problems.

Extreme pH values are to be avoided because of effects on emulsification of oil, corrosion, precipitation, volatilization of sulfides and other gases, etc. In streams and water courses, extreme pH levels accentuates the adverse effects of other pollutants as well as causing toxicity itself.

The hydrogen ion concentration in an aqueous solution is represented by the pH of that solution. The pH is defined as the negative logarithm of the hydrogen ion concentration in a solution. The pH scale ranges from zero to fourteen, with a pH of seven, representing neutral conditions, i.e., equal concentrations of hydrogen and hydroxyl ions. Values of pH less than seven indicate increasing hydrogen ion concentration or acidity; pH values greater than seven indicate increasing alkaline

conditions. The pH value is an effective parameter for predicting chemical and biological properties of aqueous solutions. It should be emphasized that pH cannot be used to predict the quantities of alkali or acidic materials in a water sample. However, most effluent and stream standards are based on maximum and minimum allowable pH values rather than on alkalinity and acidity.

since pH RWL values are not additive, it is not always possible to predict the final pH of a process waste water made up of multiple discharges. In addition, the individual refinery's discharge characteristics will dictate final pH ranges, which may be kept within the acceptable range merely by equalization, or which may require more sophisticated neutralization facilities. However, it is recommended that a pH range of 6.0 to 9.0 be established as the effluent limitation.

Temperature

Crude desalting, distillation, and cracking contribute substantial thermal wasteloads.

Effluent heat loads can have adverse effects on the receiving waters. Water temperature is important in terms of its effect on aquatic life, the use of water for cooling purposes, and its influence on the self-purification processes in a stream. Increased temperature reduces the solubility of oxygen in water and speeds biological degradation processes, thus accelerating the demand on oxygen resources of the stream. Both of these phenomenon reduce the streams assimilative capacity for waste loads. High temperatures have also been reported intensify the effect of toxic substances.

Other Metallic Ions

Several metallic ions in addition to chormium and zinc may be found in refinery effluents. The major sources for their presence in waste water are from the crude itself and corrosion products. The concentration of metallic ions varies considerably dependent upon the effectiveness of catalyst recovery in production process. Table 23 lists those metals which may be commonly found in petroleum refinery effluents. Dissolved metallic ions create turbidity and discoloration, can precipitate to form bottom sludges, and can impart taste to water.

Metallic ions such as copper, and cadmium are toxic to microorganisms because of their ability to tie up the proteins in the key enzyme systems of the microogranisms.

Chlorides:

Chloride ion is one of the major anions found in water and produces a salty taste at a concentration of about 250 mg/l. Concentrations of

1000 mg/l may be undetectable in waters which contain appreciable amounts of calcium and magnesium ions.

water is invariably associated with naturally occurring hydrocarbons underground and much of this water contains high amounts of sodium chloride. The saltiest oil field waters are located in the mid-continent region of the country where the average dissolved solids content is 174,000 ppm; therefore, waters containing high levels of salt may be expected.

Copper chloride may be used in a sweetening process and aluminum chloride in catalytic isomerization. These products may also find their ways to waste streams.

The toxicity of chloride salts will depend upon the metal with which they are combined. Because of the rather high concentration of the anion necessary to initiate detrimental biological effects, the limit set upon the concentration of the metallic ion with which it may be tied, will automatically govern its concentration in effluents, in practically all forms except potassium, calcium, mganesium, and sodium.

Since sodium is by far the most common (sodium 75 percent, magnesium 15 percent, and calcium 10 percent) the concentration of this salt will probably govern the amount of chlorides in waste streams from petroleum refineries.

It is extremely difficult to pinpoint the exact amount of sodium loride salt necessary to result in toxicity in waters. Large accentrations have been proven toxic to sheep, swine, cattle or poultry.

In swine fed diets of swill containing 1.5 to 2.0% salt by weight, poisoning symptoms can be induced if water intake is limited and other factors are met. The time interval necessary to accomplish this is still about one full day of feeding at this level.

Since problems of corrosion, taste and quality of water necessary for industrial or agricultural purposes occur at sodium chloride concentration levels below those at which toxic effects are experienced, these factors will undoubtedly determine the amount of chlorides allowed to escape in waste streams from refining operations. The study of refinery effluents previously mentioned, placed net chloride levels at values ranging from 57 to 712 mg/l. The median value was 176 mg/l.

Fluoride: HF

Alkalation units (when hydrofluoric acid is used) can contribute fluoride ion to the plant's waste effluent. Since calcium and barium fluoride are insoluble in water the fluorides will by necessity be associated with other cations.

TABLE 23

Metallic Ions Commonly Found in Effluents from Petroleum Refineries

Aluminum
Arsenic
Cadmium
Chromium
Cobalt
Copper
Iron
Lead
Mercury
Nickel
Vanadium
Zinc

In concentrations of approximately 1 mg/l in potable water supplies fluorides have been found to be an effective preventor of dental characters. In concentrations greater than this amount, fluorides can cause molting of tooth enamel and may be incorporated into the bones.

Natural waters can contain levels of fluorides up to 10 ppm. If these waters are to be used for potable supplies or for certain industrial or agricultural purposes the fluoride levels must be reduced. Since many municipal waters are artificially fluoridated as a dental health aid, the U.S. Public Health Service has placed limits on the total amounts of fluorides a water supply may contain. Their recommended control levels depend upon temperature and are expressed as lower, optimum, and upper limits. Optimum limits range from .7 to 1.2 mg/l. If values exceed two times the optimum value, the supply must be rejected or the fluoride content lowered. Because refinery effluents may empty into water ways which may eventually become public supplies, the maximum permissible limits of fluorides present in an effluent will probably be derived from the USPHS control limits for drinking water.

Phosphate - Total

Various forms of phosphates find their way into refinery effluents. They range through several organic and inorganic species and are usually contributed by corrosion control chemicals. Plant cooling systems may contain 20 to 50 mg/l of phosphate ion.

Phosphorus is an element which is essential to growth of an organism. ay at times become a growth limiting nutrient in the biological system of a water body. In these instances an over abundance of the element contributed from an outside source may stimulate the growth of photosynthetic aquatic macro and micro-organisms resulting in nuisance problems. Since the forms of phosphorus in waters or industrial wastes are so varied, the term total phosphate has been used to indicate all the phosphate present in an analyzed sample regardless of the chemical Also, many phosphorus compounds tend to degrade rather readily, and in these less complex forms phosphate may be readily utilized in the aquatic life cycle. It is therefore reasonable to direct concern toward the total amount of phosphorus present rather than chemical structure it may assume, for in only very unusual cases may the form or concentration of the element present in a waste stream be toxic. Total phosphate values noted on a nationwide refinery survey were 9.49 mg/l maximum and .096 mg/l minimum for effluents. The median value was .68 mg/l expressed as phosphorus.

SECTION VII

CONTROL AND TREATMENT TECHNOLOGY

Petroleum refinery waste waters vary in quantity and quality from refinery to refinery. However, the wastes are readily treatable. The results of the industry survey indicate, as would be expected, that techniques for in-process control are general across the industry and the specific application of these techniques at individual plants determines their success. Local factors such as climate, discharge criteria, availability of land, or other considerations may dictate the use of different waste water treatment processes to reach an acceptable effluent. The survey has shown that although the end-of-pipe waste water treatment technologies used throughout the petroleum refining industry have a marked similarity in operational steps, a considerable variation in treatment results exist. The processes used for treating refinery waste water, however, are similar in purpose; namely--maximizing oil recovery and minimizing the discharge of other pollutants. The wastewater treatment technology described below is generally applicable across all industry subcategories.

In-Plant Control/Treatment Techniques

In-plant practices are the sole determinant of the amount of waste water to be treated. There are two types of in-plant practices that reduce flow to the treatment plant. First, reuse practices involving the use water from one process in another process. Examples of this are: using stripper bottoms for makeup to crude desalters; using blowdown from high pressure boilers as feed to low pressure boilers; and using treated effluent as makeup water whereever possible. Second, recycle systems that use water more than once for the same purpose. Examples of recycle systems are: the use of steam condensate as boiler feedwater; and cooling towers. The reduction or elimination of a waste stream allows the end-of-pipe processes to be smaller, provide better treatment, and be less expensive. Since no treatment process can achieve 100 percent pollutant removal from the individual stream, reduction in flow allows for a smaller pollutant discharge.

Housekeeping

In addition to reuse/recycle of water streams and reduction in flows by other in-plant techniques, another effective in-plant control is good housekeeping. Examples of good housekeeping practices are: minimizing waste when sampling product lines; using vacuum trucks or dry cleaning methods to clean up any oil spills; using a good maintenance program to keep the refinery as leakproof as possible; and individually treating waste streams with special characteristics, such as spent cleaning solutions.

The use of dry cleaning, without chemicals, aids in reducing water discharges to the sewer. Using vacuum trucks to clean up spills and charging of this recovered material to slop oil tanks, reduces the discharge of both oil and water to the waste water system. The oil on also be recovered for reprocessing. Process units should be curbed to prevent the contamination of clean areas with oily storm runoff and to prevent spills from spreading widely. Prompt cleanup of spills will also aid in reducing discharges to the sewer systems. Additionally, sewers should be flushed regularly to prevent the buildup of material in the sewer, eliminating sudden surges of pollutants during heavy rains. Collection vessels should also be provided whenever maintenance is performed on liquid processing units, to prevent accidental discharges to the sewers.

Operations during turnaround present special problems. Wastes generated by cleaning tanks and equipment should be collected, rather than draining directly to the sewer. The wastes from these holding tanks should be gradually bled to the sewer, after first pretreating as necessary to eliminate deleterious effects on the waste water treatment system. An alternative method of disposal is through the use of contract carriers.

While these are not all the examples of good housekeeping practices which can be cited for refinery operations, it is evident that housekeeping practices within a refinery can have substantial impact on the loads discharge to the waste treatment facilities. The application of good housekeeping practices to reduce waste loads requires judicious planning, organization and operational philosophy.

Process Technology

Many of the newer petroleum refining processes are being designed or modified with reduction of water use and subsequent minimization of contamination as design criteria; although no major innovations in basic refining technology are anticipated. Improvements which can be expected to be implemented in existing refineries are: primarily dedicated to better control of refinery processes and other operations; elimination of marginal processing operations, and specific substitution of processes and/or cooling techniques to reduce discharge loads to waste treatment facilities. Examples of the possible changes which may be implemented include:

- 1. Substitution of improved catalysts which have higher activity and longer life, consequently requiring less regeneration and resulting in lower waste water loads.
- 2. Replacement of barometric condensers with surface condensers or air fan coolers, reducing a major oil-water emulsion source. As an alternative, several refineries are using oily water cooling tower systems, with the barometric condensers, equipped

with oil separation/emulsion breaking auxiliary equipment.

- 3. Substitution of air fan coolers to relieve water cooling duties simultaneously reduces blowdown discharges.
- 4. Installation of hydrocracking and hydrotreating processes will allow generation of lower waste loadings than the units they replace. The rapid pace at which such units are being installed is exerting and will continue to exert a strong influence on the reduction of waste loadings, particularly sulfides and spent caustics.
- 5. Installation of automatic monitoring instrumentation, such as TOC monitors, will allow early detection of process upsets which result in excessive discharges to sewers.
- 6. Increased use of improved drying, sweetening, and finishing procedures will minimize spent caustics and acids, water washes, and filter solids requiring disposal.

Cooling Towers

Cooling towers eliminate large volumes of once through cooling water by passing heated water through heat exchange equipment. By recycling the cooling water many times, the amount of water used is greatly reduced. The number of times cooling water can be reused is determined by the total dissolved solids (TDS) content of the water, and the effects high dissolved solids have on process equipment. When the TDS becomes too ligh, scaling occurs and heat transfer efficiency decreases. The TDS level in the circulating water is controlled by discharging a portion of the steam (blowdown) from the system. The higher the allowable TDS level, the greater number of cycles of concentration and the less make-up water is required (87). Installation of cooling towers will reduce the amount of water used within the refinery by at least 90 percent (87).

There are three types of cooling towers (106); wet or evaporative, dry, and combined "wet-dry."

Evaporative Cooling Systems

Evaporative cooling systems transport heat by transfer of the latent heat of vaporization. This results in a temperature decrease of circulating water and a temperature and humidity increase of cooling air.

Spray ponds are an evaporative cooling system using natural air currents and forced water movement. Because of their inefficiency, spray ponds are used less in industry than cooling towers. Cooling towers have a higher efficiency because they provide more intimate contact between the air and water. As the water falls over the packing, it exposes a large

contact surface area. As the water heats up the air, the air can absorb more water. The more water evaporated, the more heat is transferred (106). Because an evaporative cooling tower is dependent on ambie temperatures and humidity, its performance is variable throughout tyear. There are three types of evaporative cooling towers: mecahnical draft towers; atmospheric towers, which use wind or natural air currents; and natural draft towers, which use tall stacks to move air by stack effect. Most refineries use mechanical draft towers, which have baffles, called drift eliminators, to separate entrained water from the air stream, thus reducing the amount of water carried into the air. The evaporative system is the least costly of all cooling towers.

Dry Cooling Systems

There are two types of dry air cooling systems. Either system can be used with either mechanical or natural draft cooling towers. Most refineries use mechanical draft towers on indirect condensing systems. The tubes used in dry cooling equipment have circumferential fins to increase the heat transfer area. Most tube designs have an outside to inside surface area ratio of 20:1. (106) The advantage of the dry air system is that it requires no makeup water and there is no water entrainment. Dry air cooling systems are being increasingly used to reduce the amount of water discharged to the waste water treatment plant. A disadvantage of the dry cooling process is that it has low rates of heat transfer requiring large amounts of land and uses more power than other cooling systems. The dry cooling tower is also more expensive to install than evaporative systems.

Wet-Dry Systems

The wet-dry systems use an evaporative and non-evaporative cooling tower in either series or parallel, each of which can be operated with a mechanical or natural draft tower. The series design has the evaporative cooling process preceding the dry process with respect to the air flow. This lowers the temperature of the air entering the dry process which would mean a smaller unit could be used. The problem with this method is that solids are deposited in the dry tower due to drift from the wet section. The parallel process uses a dry cooling tower upstream of the wet section, each of which has its own air supply. The two air streams are mixed and discharged, reducing the vapor plume.

Recycle/Reuse Practices

Recycle/reuse can be accomplished either by return of the waste water to its original use, or by using it to satisfy a lower quality demand. The recycle/reuse practices within the refining industry are extremely varied and only a few examples are described briefly below:

1. Reduction of once-through cooling water results in tremendously decreased total effluents.

- 2. Sour water stripper bottoms are being used in several refineries as make-up water for crude desalter operations. These sour water bottoms are initially recovered from overhead accumulators on the catalytic cracking units.
- 3. Regeneration of contact process steam from contaminated condensate will reduce the contact process waste water to a small amount of blowdown. This scheme can be used to regenerate steam in distillation towers or dilution steam stripping in pyrolysis furnaces.
- 4. Reuse of waste water treatment plant effluent as cooling water, as scrubber water, or as plant make-up water, reduces total make-up requirements.
- 5. Cooling tower blowdowns are frequently reused as seal water on high temperature pump service, where mechanical seals are not practicable.
- 6. Storm water retention ponds are frequently used as a source of fire water or other low quality service waters.

Many other conservation methods can be implemented, such as the use of stripped sour water as low pressure (LP) boiler make-up, and LP boiler blowdown as make-up water for crude desalting. However, these, and the other possible recycle/reuse cases outlined above must be examined by the individual refinery in light of its possible vantages/disadvantages, insofar as product quality or refining process capabilities are affected. For example, one refinery has reported that reuse of sour water stripper bottoms for desalting resulted in a desalted crude which was difficult to process downstream.

At-Source Pretreatment

Major at-source pretreatment processes which are applicable to individual process effluents or groups of effluents within a refinery are stripping of sour waters, neutralization and oxidation of spent caustics, ballast water separation, and slop oil recovery. The particular areas of application of these processes are discussed below.

Sour Water Stripping

Sour or acid waters are produced in a refinery when steam is used as a stripping medium in the various cracking processes. The hydrogen sulfide, ammonia and phenols distribute themselves between the water and hydrocarbon phases in the condensate. The concentrations of these pollutatns in the water vary widely depending on crude sources and processing involved.

The purpose of the treatment of sour water is to remove sulfides (as hydrogen sulfide, ammonium sulfide, and polysulfides) before the waste enters the sewer. The sour water can be treated by: stripping the steam or flue gas; air oxidation to convert hydrogen sulfide thiosulfates; or vaporization and incineration.

Sour water strippers are designed primarily for the removal of sulfides and can be expected to achieve 85-99 percent removal. If acid is not required to enhance sulfide stripping, ammonia will also be stripped with the percentage varying widely with stripping temperature and pH. If acid is added to the waste water, essentially none of the ammonia will be removed. Thus, ammonia removals in sour water strippers vary from 0 to 99 percent. Depending upon such conditions as waste water pH, temperature, and contaminant partial pressure; phenols and cyanides can also be stripped with removal as high as 30 percent. The bottoms from the stripper usually go to the desalter where most of the phenols are extracted and the waste water can be sent to the regular process water treating plant. COD and BOD5 are reduced because of the stripping out of phenol and oxidizable sulfur compounds.

The heated sour water is stripped with steam or flue gas in a single stage packed or plate-type column. Two-stage units are also being installed to enhance the separate recovery of sulfide streams and ammonia streams. Hydrogen sulfide released from the waste water can be recovered as sulfuric acid or sulfur, or may be burned in a furnace. The bottoms have a low enough sulfide concentration to permit discharge into the general waste water system for biological treatment. If the waste contains ammonia, it is neutralized with acid before stam stripping. The waste liquid passes down the stripping column while estripping gas passes upward. Most refiners now incinerate th sour water stripper acid gases without refluxing the stripper. This converts the ammonia to nitrogen with possibly traces of nitrogen oxides. Due to the high concentrations of sulfur dioxide produced more complex processing will probably be required in the future.

Several stripping processes are available. These include: Chevron WWT; ammonium sulfate production; a dual burner Claus sulfur plant; and the Howe-Baker ammonex process. Deep well injection and oxidation to the thiosulfate are also being used, but in the future probably won't do a good enough job.

The Chevron WWT process (37) is basically two stage stripping with ammonia pruification, so that the hydrogen sulfide and ammonia are separated. The hydrogen sulfide would go to a conventional Claus sulfur plant and the ammonia can be used as fertilizer.

Ammonium sulfate can be produced by treating with sulfuric acid but a very dilute solution is produced and concentrating it for sale as fertilizer is expensive. Again the hydrogen sulfide goes to a conventional Claus sulfur plant.

A dual burner Claus sulfur process is generally the answer in new plants, but adding the second burner to an existing sulfur plant is difficult. The second burner is required to handle the ammonia. A refluxed stripper is required to reduce the water vapor in the hydrogen sulfide-ammonia mixture and the line between the stripper and the Claus Unit must be kept at about 150°F to prevent precipitation of ammonium sulfide complexes.

Howe-Baker Engineers Inc. of Tyler, Texas have developed to the pilot plant stage a process they call "Ammonex". It is a solvent extraction process that basically is intended to complete with the Chevron WWT process. No commercial units have been built.

Another way of treating sour water is to oxidize by aeration. Compressed air is injected into the waste followed by sufficient steam to raise the reaction temperature to at least 190°F. Reaction pressure of 50-100 psig is required. Oxidation proceeds rapidly and converts practically all the sulfides to thiosulfates and about 10 percent of the thiosulfates to sulfates. Air oxidation, however, is much less effective than stripping in regard to reduction of the oxygen demand of sour waters, since the remaining thiosulfates can later be oxidized to sulfates by aquatic microorganisms.

The stripping of sour water is normally carried out to remove sulfides and hence, the effluent may contain 50-100 ppm of ammonia, or even considerably higher, depending on the influent ammonia concentration. Values of ammonia have been reported as low as 1 ppm, but generally the effluent ammonia concentration is held to approximately 50 ppm to provide nutrient nitrogen for the refinery biological waste treatment system (2,14,33,58).

Spent Caustic Treatment

Caustic solutions are widely used in refining. Typical uses are to neutralize and extract:

- a. Acidic materials that may occur naturally in crude oil.
- b. Acidic reaction products that may be produced by various chemical treating processes.
- c. Acidic materials formed during thermal and catalytic cracking such as hydrogen sulfide, phenolics, and organic acids.

Spent caustic solutions may therefore contain sulfides, mercaptides sulfates, sulfonates, phenolates, naphthenates, and other similar organic and inorganic compounds.

At least four companies process these spent caustics to market the phenolics and the sodium hyposulfide. However, the market is limited

and most of the spent caustics are very dilute so the cost of shipping the water makes this operation uneconomical.

Some refiners neutralize the caustic with spent sulfuric from oth refining processes, and charge it to the sour water stripper. This removes the hydrogen sulfide. The bottoms from the sour water stripper go to the desalter where the phenolics are extracted by the crude oil.

Spent caustics usually originate as batch dumps, and the batches may be combined and equalized before being treated and/or discharged to the general refinery waste waters. Spent caustic solutions can also be treated by neutralization with flue gas. In the treatment of spent caustic solutions by flue gas, hydroxides are converted to carbonates. Sulfides, mercaptides, phenolates, and other basic salts are converted by the flue gas stripping. Phenols can be removed and used as a fuel or can be sold. Hydrogen sulfide and mercaptans are usually stripped and burned in a heater. Some sulfur is recovered from stripper gases. treated solution will contain mixtures of carbonates, sulfates, sulfites, thiosulfates and some phenolic compounds. Reaction time of 16-24 hours is required for the neutralization of caustic solution with flue gas.

The oxidation phase of spent caustic treatment is aimed at the sulfide content of these wastes and achieves 85-99 percent sulfide removal. this process, sulfides are oxidized primarily to thiosulfates although in some variations there is partial oxidation of the sulfur compounds to Oxidation processes are not applied to phenolic caustics, as phenols inhibit oxidation. It should be noted that those processes which oxidize the sulfide only to thiosulfate, satisfy half of t oxygen demand of the sulfur, as thiosulfate can be oxidized biological, to sulfate. Neutralization of spent caustics is applied to both phenolic and sulfidic caustic streams; the sulfidic caustics are also steam stripped, after neutralization, to remove the sulfides. phenolic spent caustics are neutralized, crude acid oils or "crude carbolates" are sprung and thus removed from the waste water. The major part of the phenols will appear in the oil fraction, but a significant part may remain in the waste water as phenolates.

Fluid bed incineration is also now being used. This process was developed under an EPA demonstration grant (26) and at least two large units are under construction. Once the incinerator is started up, the sludge should provide the necessary heating value to keep the system operating. Oxidizing fuels may be required when the sludge is burnt, as ash remains in the bed of the incinerator. A constant bed level is maintained, so the sand bed originally in the incinerator is gradually replaced by the inert sludge ash (5). The gasses pass through a scrubber, so the fines and particulate matter can be recovered. The ash and fines can be landfilled. This landfill is cleaner than a sludge landfill, because there are no organic materials present to contaminate ground water or run-off.

In the past ocean dumping, deep well injection, evaporative lagoons, and simple dilution have all been used. These methods will no longer be ceptable.

Sewer System Segregation

Waste water quantity is one of the major factors that affect the cost of waste treatment facilities most directly. Water usage in the petroleum refining industry varies from less than 5 gallons of water per barrel of crude charge in the newer refineries to higher than 1000 gallons of water per barrel of crude charge in the older refineries. In order to provide efficient treatment to the wastes originating within a refinery, it is very important that segregation of concentrated waste streams be Segregation of waste streams frequently simplifies waste treating problems as well as reduces treatment facility costs. treatment of highly polluted waste streams at the source can prevent gross pollution of large volumes of relatively clean waste water. treatment is often a more economical solution of a problem than would be possible if wastes are discharged directly to the refinery sewers. Treatment at the source is also helpful in recovering by-products from the wastes which otherwise could not be economically recovered when the wastes are combined.

In areas where water supply is limited, reduced water requirements have been incorporated into the design and operation, thereby reducing total water usage.

minimize the size of the waste water treatment processes it is meaning the various sewer systems. There should be a sewer carrying process and blowdown waters that are treated continuously. A polluted storm water sewer should go to a storage area from which it can be gradually discharged to the treatment facilities. A sewer system containing clean storm water can be discharged directly to the receiving water. The sanitary system should be treated separately from the process water because of the bacteria present in this stream. Once through cooling water should be kept separate because of the large volumes of water involved and the low waste loadings encountered. A connection to the treatment plant should be provided in case of oil leaks into the system.

Storm Water Runoff

An additional source of pollution from a petroleum refinery area is caused by rainfall runoff. Size and age of refinery site, housekeeping, drainage areas, and frequency and intensity of rainfall are several of the factors which compound the assignment of allowable pollutional values.

There are several measures that refiners can provide to minimize storm water loads to their treatment system after diverting all extraneous drainage around the refinery area. The major consideration is a separate storm water sewer and holding system.

By providing separate collection facilities for storm water runoff, protection is afforded the operation of the separator and ancillary treatment systems by controlling the hydraulic load to be treated. Comingling of inorganic particles with oily waste water often times produces an emulsion which is difficult to break in the oil-water separator.

Design of this facility should be based on the maximum ten-year, twenty-four-hour rainfall runoff of the refinery drainage area. Diversion of the collected storm water runoff to the oil-water separator facilities can be provided when hydraulic flows return to normal operations. In the event of excessive collection due to a high intensity storm, diversion facilities should be provided to allow for emergency bypass capability to divert the trailing edge of the runoff hydrograph (the leading edge normally contained the mass of pollutants in urban runoff investigations). An oil retention baffle and an API type overflow weir should be provided to prevent the discharge of free and floating oil.

An alternate to the separate sewer system would be the provision of a storm surge pond that would receive the polluted waters when the flow to the oil-water separator exceeded 15 percent of the normal hydraulic flow. During normal periods, the collected storm water-refinery water could then be diverted to the oil-water separator (provided process flow did not equal or exceed the units hydraulic capacity).

The major cause of pollution by storm water runoff is the lack of housekeeping within the refinery confine. Proper procedures should be encouraged to prevent the accumulation of materials which contribute to pollution due to rainfall runoff. Some of the more common preventive measures are: (1) Provide curbing around process unit pads; (2) Prevent product sample drainage to sewers; (3) Repair pumps and pipes to prevent oily losses to the surface areas; (4) Contain spilled oil from turnarounds; (5) Dike crude and product tank areas and valve precipitation to the storm water sewer.

In the event the collected water needs to be released from the storm water detention pond due to overflow, samples of the water should be monitored for; (1) Sheen, (2) Organic analysis such as COD, TOC, or TOD.

Ballast Water Separation

Ballast water normally is not discharged directly to the refinery sewer system because the intermittent high-volume discharges. The potentially high oil concentrations, would upset the refinery waste water treatment facilities. Ballast waters may also be treated separately, with

heating, settling, and at times filtration as the major steps. The settling tank can also be provided with a steam coil for heating the contents to help break emulsions, and an air coil to provide ation. The recovered oil, which may be considerable, is generally sent to the slop oil system.

Slop Oil Treatment

Separator skimmings, which are generally referred to as slop oil, require treatment before they can be reused, because they contain an excess amount of solids and water. Solids and water contents in excess of about 1 percent generally interfere with processing.

In most cases slop oils are easily treated by heating to 190°F for 12 to 14 hours. At the end of settling, three definite layers exist: a top layer of clean oil; a middle layer of secondary emulsion; and a bottom layer of water containing soluble components, suspended solids, and oil. In some cases, it is advantageous or even necessary to use acid or specific chemical demulsifiers to break slop oil emulsions. The water layer resulting from acid and heat treatment has high BOD and COD_{ℓ} but also low pH $_{\ell}$ and must be treated before it can be discharged.

Slop oil can also be successfully treated by centrifugation or by precoat filtration using diatomaceous earth as the precoat.

Gravity Separation of Oil

ity separators remove a majority of the free oil found in refinery waste waters. Because of the large amounts of reprocessable oils which can be recovered in the gravity separators, these units must be considered an integral part of the refinery processing operation and not a waste water treatment process. The functioning of gravity-type separators depends upon the difference in specific gravity of oil and water. The gravity-type separator will not separate substances in solution, nor will it break emulsions. The effectiveness of a separator depends upon the temperature of the water, the density and size of the oil globules, and the amounts of characteristics of the suspended matter present in the waste water. The "susceptibility to separation" (STS) test is normally used as a guide to determine what portion of the influent to a separator is amenable to gravity separation.

The API separator is the most widely used gravity separator. The basic design is a long rectangular basin, with enough detention time for most of the oil to float to the surface and be removed. Most API separators are divided into more than one bay to maintain laminar flow within the separator, making the separator more effective. API separators are usually equipped with scrapers to move the oil to the downstream end of the separator where the oil is collected in a slotted pipe or on a drum. On their return to the upstream end, the scrapers travel along the

bottom moving the solids to a collection trough. Any sludge which settles can be dewatered and either incinerated or disposed of as landfill.

The gravity separator usually consists of a pre-separator (grit chamber) and a main separator, usually rectangular in shape, provided with influent and effluent flow distribution and stilling devices and with oil skimming and sludge collection equipment. It is essential that the velocity distribution of the approach flow be as uniform as possible before reaching the inlet distribution baffle.

Another type of separator finding increasing employment in refineries is the parallel plate separator. The separator chamber is subdivided by parallel plates set at a 45° angle, less than 6 inches apart. This increases the collection area while decreasing the overall size of the unit. As the water flows through the separator the oil droplets coalesce on the underside of the plates and travel upwards where the oil is collected. The parallel plate separator can be used as the primary gravity separator, or following an API separator.

Further Removal of Oil and Solids

If the effluent from the gravity separators is not of sufficient quality to insure effective treatment before entering the biological or physical-chemical treatment system, it must undergo another process to remove oils and solids. Most refineries use either clarifiers, dissolved air flotation units or filters to reduce the oil and solids concentration. Each of these processes has also been used to treat teffluent from a biological system.

Clarifiers

Clarifiers use gravitational sedimentation to remove oil and solids from a waste water stream. Often it is necessary to use chemical coagulants such as alum or lime to aid the sedimentation process. These clarifiers are usually equipped with a skimmer to remove any floating oil. Clarifiers used after a biological system normally do not have skimmers as there should be no floating oils at that point. The sludge from the clarifiers is usually treated before final disposal.

End-of-Pipe Control Technology

End-of-pipe control technology in the petroleum refining industry relies heavily upon the use of biological treatment methods. These are supplemented by appropriate pretreatment to insure that proper conditions, especially sufficient oil removal and pH adjustment, are present in the feed to the biological system. When used, initial treatment most often consists of neutralization for control of pH or equalization basins to minimize shock loads on the biological systems.

The incorporation of solids removal ahead of biological treatment is not as important as it is in treating municipal waste waters.

of the initial criteria used to screen refineries for the field survey, was degree of treatment provided by their waste water treatment facilities. Therefore, the selection of plants was not based on a cross-section of the entire industry, but rather was biased in favor of those segments of the industry that had the more efficient waste water treatment facilities. Table 24 indicates the types of treatment technology and performance characteristics which were observed during the survey. In most of the plants analyzed, some type of biological treatment was utilized to remove dissolved organic material. Table 25 summarizes the expected effluents from waste water treatment processes throughout the petroleum refining industry. Typical efficiencies for these processes are shown in Table 26.

During the survey program, waste water treatment plant performance history was obtained when possible. This historical data were analyzed statistically and the individual plant's performance evaluated in comparison to the original design basis. After this evaluation, a group of plants was selected as being exemplary and these plants were presented in Table 24. The treatment data in Table 26 represent the annual daily average performance (50 percent probability-of-occurrence).

There were enough plants involving only one subcategory to make the interpretation meaningful. In preparing the economic data base, however, all the waste water treatment plant data were analyzed to elop a basis for subsequent capital and operating costs.

The treatment data from the exemplary plants referred to previoulsy were analyzed to formulate the basis for developing BPCTCA effluent criteria. The effluent limitations were based on both these treatment data, other data included in Supplement B, and other sources as discussed in Section IX. These effluent limitations were developed for each subcategory individually and thus no common treatment efficiency was selected as being typical of the petroleum refining industry for use in the BPCTCA effluent limitations. A brief description of the various elements of end-of-pipe treatment follows.

Equalization

The purpose of equalization is to dampen out surges in flows and loadings. This is especially necessary for a biological treatment plant, as high concentrations of certain materials will upset or completely kill the bacteria in the treatment plant. By evening out the loading on a treatment plant, the equalization step enables the treatment plant to operate more effectively and with fewer maintenance problems. Where equalization is not present, an accident or spill

Observed Refinery Treatment System and Effluent Loadings

TABLE 24

SUBCATEGOR	A Y	B-1	B-2	B-2	B-2	B-2	С	C	C	D	D	E
Type of Treatment	OP	AL-PP	AL-F	E-DAF-AS	OP	DAF,AL,PP	DAF, AS	DAF, AS	DAF,AL,PP	E,TF,AS	E,AS	DAF,AS,PP
Refinery Observed Ar Effluent L Net-kg/100 feedstock (1b/1000 b) feedstock	oadings 0 m3 of k b1 of	R18		R27	R26	R7	R20	R8	R23	R24	R28	R25
BOD5		8(2.8)	8.0(4.4)	5.9(2.1)	10(3.6)	3.7(1.3)	13(4.6)	2.7(0.95)	2.6(0.91)	7.4(2.6)	14(5.0)	17.5(6.2)
COD		39(13.8)	68(24)	96(34)	71(25.0)	39(13.8)	67(23.5)		54 (19)	57(20)	136 (48)	320(113)
TSS			25(8.7)	34(12)	8.5(3.0)	4.2(1.5)	13.6(4.8)	8.5(3.0)	7(2.5)	12(4.3)	38(13.5)	36(12.7)
0&G	2.0(0.7)	2.3(0.8)	9(3.2)	4.0(1.4)		2.8(1.0)	6.5(2.3)			4(1.4)	7.2(2.55)	22(7.7)
NН3-N		:			4.8(1.7)	0.14(0.05)	4.5(1.6)	****	2(0.7)	1.2(0.44)		2.3(0.8)
Phenolic Compounds	0.14(0.05)	0.003(0.001)	0.4(0.145)	0.37(0.13) 0.05(0.018)	0.0006 (0.002)	0.06 (0.023)			0.17(0.06)		0.017(0.003)
Sulfide	0.03(0.009)	•	0.2(0.07)	0(0)	0.03(0.010)	0.014 (0.005)	0.05 (0.018)					0.20(.07)
Footnotes:	AS-activate	ed sludge ed air flotatio	OP-oxi n PP-pol	tration dation pond ishing pond ckling filt	B-1-I B-2-H	Copping Low cracking High-Cracking Petrochemical		rated				

TABLE 25
Expected Effluents from Petroleum Treatment Processes

EFFLUENT CONCENTRATION, mg/L										
PROCESS	PROCESS INFLUENT	BOD ₅	COD	TOC	SS	OIL	PHENOL	AMMONIA	SULFIDE	REFERENCES
1. API Separator	Raw Waste	250-350	260-700	NA	50 –200	20-100	6-100	15-150	NA	7,13,30,41,49,59
2. Clarifier	ì	45-200	130-450	NA	25-60	5 - 35	10-40	NA	An	34,48a,49
3. Dissolved Air Plotation	1	45-200	130-450	NA	25-60	5-20	10-40	NA	NA	13,29,32,48a,49
4. Granular Media Filter	1	40-170	100-400	NA	, 5 - 25	6–20	3–35	NA	- NA	17,41,48a,48
5. Oxidation Pond	1	10 - 60	50-300	NA	20-100	1.6-50	0.01-12	3 – 50	0-20	18,22,23,31,42,48a, 49,55,75,R18
6. Aerated Lagoon	2,3,4	10-50	50-200	NA	10-80	5-20	0.1-25	4-25	0-0.2	31,39,42a,48a,49, 55,59,R7,R23,R26
7. Activated Sludge	2,3,4	5–50	30-200	20-80	5–50	1-15	0.01-2.0	1-100	0-0.2	13,24,27,30,34,35, 42,48a,49,60,69,72 R8,R20,R24,R25,R27 R28,R29
8. Trickling Filter	1	25-50	80-350	NA	20-70	10-80	0.5-10	25-100	0.5-2	18,30,42,48a,49,
9. Cooling Tower	2,3,4	25-50	47-350	70-150	4.5-100	20-75	.1-2.0	1-30	NA	33,41
0. Activated Carbon	2,3,4	5-100	30-200	. NA	10-20	2-20	< 1	10-140	NA	17,21,27,48,48a,49, 53,62a
1. Granular Media Filter	5-9	NA	NA	25-61	3-20	3-17	0.35-10	NA	NA	17,48,54
2. Activated Carbon	5-9 and 11	3-10	30-100	1-17	1-15	0.8-2.5	0-0.1	1-100	0-0.2	17,21,27,48,48a,49, 53,62a

A - Data Not Available

TABLE 26

Typical Removal Efficiencies for Oil Refinery Treatment Processes

		PROCESS						L EFFICIENCY,		•	·
PROC	ESS	INFLUENT	BOD ₅	COD	TOC	SS	OIL	PHENOL	AMMONIA	SULFIDE	REFERENCES
1.	API Separator	Raw Waste	5-40	5-30	NA	10-50	60-99	0-50	NA	NA	7,13,30,41,49;59
2.	Clarifier	1	30-60	20-50	NA	50-80	60 - 95	0-50	NA .	NA	34,482,49
3.	Dissolved Air Flotation	1 .	20-70	10-60	NA	50-85	7 0-85	10-75	NA	NA	13,29,32,48a,49
4 .	Filter	1	40-70	20-55	NA	75 - 95	65-90	5–20	NA	NA	17,41,48a,49
5.	Oxidation Pond	1	40-95	30-65	60	20-70	50-90	60-99	0-15	70-100	18,22,23,31,42,48 49,55,75,R18
6.	Aerated Lagoon	2,3,4	· 75-95	60-85	NA	40-65	70-90	90-99	10-45	95-100	31,39,42,48a,49, 55,59,R7,R23,R26
7.	Activated Sludge	2,3,4	80-99	50 - 95	40-90	60-85	80-99	95-99+	33-99	97-100	13,24,R7,30,34,35 42,48a,49,60,69,7 2 R8,R20,R24,R25,R2 R28,R29
8.	Trickling Filter	1	60-85	30-70	NA	60-85	50-80	70-98	15-90	70-100	18,30,42,48a,49
	Cooling Tower	2,3,4	50-90	40-90	10-70	50-85	60-75	75-99+	60-95	NA	33,41
10 10.	Activated Carbon	2,3,4	7 0–95	70-90	50-80	60-90	75-95	90-100	7-33	NA	17,21,27,48,48a,4 9 49,53,62a
11.	Filter Granular Media	5 - 9	NA	NA	50-65	75 - 95	65-95	5-20	NA.	NA	17,48,54
12.	Activated Carbon	5-9 plus 11	91-98	86-94	50-80	60-90	70-95	90–99	33-87	NA	17,21,27,48,48a, 49,53,62a

NA - Data Not Available

within the refinery can greatly affect the effluent quality or kill the biomass (R7, R20).

e equalization step usually consists of a large pond that may contain mixers to provide better mixing of the wastes. In some refineries the equalization is done in a tank (55, R29). The equalization step can be before or after the gravity separator but is more effective before as it increases the overall efficiency of the separator. However, care must be taken to prevent anaerobic decomposition in the equalization facilities.

Dissolved Air Flotation

Dissolved air flotation consists of saturating a portion of the waste water feed, or a portion of the feed or recycled effluent from the flotation unit with air at a pressure of 40 to 60 psig. The waste water or effluent recycle is held at this pressure for 1-5 minutes in a retention tank and then released at atmospheric pressure to the flotation chamber. The sudden reduction in pressure results in the release of microscopic air bubbles which attach themselves to oil and suspended particles in the waste water in the flotation chamber. results in agglomerates which, due to the entrained air, have greatlyincreased vertical rise rates of about 0.5 to 1.0 feet/minute. floated materials rise to the surface to form a froth layer. Specially designed flight scrapers or other skimming devices continuously remove The retention time in the flotation chambers is usually froth. about 10-30 minutes. The effectiveness of dissolved air flotation depends upon the attachment of bubbles to the suspended oil and other rticles which are to be removed from the waste stream. The attraction tween the air bubble and particle is a result of the particle surface and bubble-size distribution.

Chemical flocculating agents, such as salts of iron and aluminum, with or without organic polyelectrolytes, are often helpful in improving the effectiveness of the air flotation process and in obtaining a high degree of clarification.

Dissolved air flotation is used by a number of refineries to treat the effluent from the oil separator. Dissolved air flotation using flocculating agents is also used to treat oil emulsions. The froth skimmed from the flotation tank can be combined with other sludges (such as those from a gravity separator) for disposal. The clarified effluent from a flotation unit generally receives further treatment in a biological unit, prior to discharge. In two refineries, dissolved air flotation is used for clarification of biologically treated effluents (29).

Oxidation Ponds

The oxidation pond is practical where land is plentiful and cheap. An oxidation pond has a large surface area and a shallow depth, usually not exceeding 6 feet. These ponds have long detention periods from 11 110 days.

The shallow depth allows the oxidation pond to be operated aerobically without mechanical aerators. The algae in the pond produce oxygen through photosynthesis. This oxygen is then used by the bacteria to oxidize the wastes. Because of the low loadings, little biological sludge is produced and the pond is fairly resistant to uspsets due to shock loadings.

Oxidation ponds are usually used as the major treatment process. Some refineries use ponds as a polishing process after other treatment processes.

Aerated Lagoon

The aerated lagoon is a smaller, deeper oxidation pond equipped with mechanical aerators or diffused air units. The addition of oxygen enables the aerated lagoon to have a higher concentration of microbes than the oxidation pond. The retention time in aerated lagoons is usually shorter, between 3 and 10 days. Most aerated lagoons are operated without final clarification. As a result, biota is discharged in the effluent, causing the effluent to have high BOD5 and solids concentrations. As the effluent standards become more strict, final clarification will be increasing in use.

Trickling Filter

A trickling filter is an aerobic biological process. It differs from other processes in that the biomass is attached to the bed media, which may be rock, slag, or plastic. The filter works by: 1) adsorption of organics by the biological slime 2) diffusion of air into the biomass; and 3) oxidation of the dissolved organics. When the biomass reaches a certain thickness, part of it sloughs off. When the filter is used as the major treatment process, a clarifier is used to remove the sloughed biomass.

The trickling filter can be used either as the complete treatment system or as a roughing filter. Most applications in the petroleum industry use it as a roughing device to reduce the loading on an activated sludge system.

Bio-Oxidation Tower

The bio-oxidation tower uses a cooling tower to transfer oxygen to a waste water. API (112) has called the bio-oxidation towers a modified activated sludge process, as most of the biomass is suspended in the

wastewater. Results from refineries indicate it is a successful process to treat portions or all of a refinery waste water (80, 81, 92).

Activated Sludge

Activated sludge is an aerobic biological treatment process in which concentrations (1500-3000 mg/L) of newly-grown and recycled microorganisms are suspended uniformly throughout a holding tank to which raw waste waters are added. Oxygen is introduced by mechanical aerators, diffused air systems, or other means. The organic materials the waste are removed from the aqueous phase by the microbiological growths and stabilized by biochemical synthesis and oxidation reactions. The basic activated sludge process consists of an aeration tank followed by a sedimentation tank. The flocculant microbial growths removed in the sedimentation tank are recycled to the aeration tank to maintain a high concentration of active microorganisms. Although microorganisms remove almost all of the organic matter from the waste being treated, much of the converted organic matter remains in the system in the form of microbial cells. These cells have a relatively high rate of oxygen demand and must be removed from the treated waste water before discharge. Thus, final sedimentation and recirculation of biological solids are important elements in an activated sludge system.

Sludge is wasted on a continuous basis at a relatively low rate to prevent build-up of excess activated sludge in the aeration tank. Shock organic loads usually result in an overloaded system and poor sludge actilities requires pretreatment to remove or substantially reduce oil, sulfides (which causes toxicity to microorganisms), and phenol concentrations. The pretreatment units most frequently used are: gravity separators and air flotation units to remove oil; and sour water strippers to remove sulfides, mercaptans, and phenol. Equalization also appears necessary to prevent shock loadings from upsetting the aeration basin. Because of the high rate and degree of organic stabilization possible with activated sludge, application of this process to the treatment of refinery waste waters has been increasing rapidly in recent years.

Many variations of the activated sludge process are currently in use. Examples include: the tapered aeration process, which has greater air addition at the influent where the oxygen demand is the highest; step aeration, which introduces the influent waste water along the length of the aeration tank; and contact stabilization, in which the return sludge to the aeration tank is aerated for 1-5 hours. The contact stabilization process is useful where the oxygen demand is in the suspended or colloidal form. The completely mixed activated sludge plant uses large mechanical mixers to mix the influent with the contents of the aeration basin, decreasing the possibility of upsets due to shock

loadings. The Pasveer ditch is a variation of the completely mixed activated sludge process that is widely used in Europe. Here brushes are used to provide aeration and mixing in a narrow oval ditch. The advantage of this process is that the concentration of the biota higher than in the conventional activated sludge process, and the wasted sludge is easy to dewater. There is at least one refinery using the Pasveer ditch type system.

The activated sludge process has several disadvantages. Because of the amount of mechanical equipment involved, its operating and maintenance costs are higher than other biological systems. The small volume of the aeration basin makes the process more subject to upsets than either oxidation ponds or aerated lagoons.

As indicated in Table 25, the activated sludge process is capable of achieving very low concentrations of BOD5, COD, TSS, and oil, dependent upon the influent waste loading and the particular design basis. Reported efficiencies for BOD5 removal are in the range of 80 to 99 percent.

Physical-Chemical Treatment

Physical-chemical treatment refers to treatment processes that are non-biological in nature. There are two types of physical-chemical processes; those that reduce the volume of water to be treated (vapor compression evaporators, reverse osmosis, etc.), and those that reduce the concentration of the pollutants (activated carbon).

Physical-chemical (P-C) processes require less land than biological processes. P-C processes are not as susceptible to upset due to shock loading as are biological processes. Another advantage of P-C is that much smaller amounts of sludge are produced.

Flow Reduction Systems

Flow reduction systems produce two effluents, one of relatively pure water and one a concentrated brine. The pure water stream can be reused within the refinery resulting in a smaller effluent flow. The brine is easier to treat as it is highly concentrated. Both of the processes described herein have been demonstrated on small flows only and at present the costs involved are extremely high (45, 52, 93).

In the vapor compression evaporator the waste water flows over heat transfer surfaces. The steam generated enters a compressor where the temperature is raised to a few degrees above the boiling point of the waste water. The compressed steam is used to evaporate more waste water while being condensed. The condensed steam is low in dissolved solids. The major process costs are the costs of electrical power, which is approximately \$1.0/1000 gallons of clean water (93).

The reverse osmosis process uses high pressures (400-800 psig) to force water through a semi-permeable membrane. The membrane allows the water pass through, but contains the other constituents in the waste water. Trently available membranes tend to foul and blind, requiring frequent cleaning and replacement. Until this problem is corrected, reverse osmosis is not a practicable process. The operating cost for a reverse osmosis unit is approximately 20-30¢/1000 gallons (45, 95).

Granular Media Filters

There are several types of granular media filters: sand, dual media, and multimedia. These filters operate in basically the same way, the only difference being the filter media. The sand filter uses relatively uniform grade of sand resting on a coarser material. The dual media filter has a course layer of coal above a fine layer of sand. Both types of filters have the problem of keeping the fine particles on the bottom. This problem is solved by using a third very heavy, very fine material, (usually garnet) beneath the coal and sand.

As the water passes down through a filter, the suspended matter is caught in the pores. When the pressure drop through the filter becomes excessive, the flow through the filter is reversed for removal of the collected solids loading. The backwash cycle occurs approximately once a day, depending on the loading, and usually lasts for 5-8 minutes. Most uses of sand filters have been for removing oil and solids prior to an activated carbon unit. There is one refinery that uses a mixed media filter on the effluent from a biological system. Granular media filters e shown to be capable of consistently operated with extremely low TSS and oil effluent discharges, on the order of 5-10 mg/L.

Activated Carbon

The activated carbon (AC) process utilizes granular activated carbon to adsorb pollutants from waste water. The adsorption is a function of the molecular size and polarity of the adsorbed substance. Activated carbon preferentially adsorbs large organic molecules that are non-polar.

An AC unit follows a solids removal process, usually a sand filter which prevents plugging of the carbon pores. From the filter the water flows to a bank of carbon columns arranged in series or parallel. As the water flows through the columns the pollutants are adsorbed by the carbon, gradually filling the pores. At intervals, portions of the carbon are removed to a furnace where the adsorbed substances are burnt off. The regenerated carbon is reused in the columns, with some makeup added, because of handling and efficiency losses.

Activated carbon processes currently have only limited usage in the refining industry. However, there are new installations in the planning construction stages. The increasing use of activated carbon has occured

because activated carbon can remove organic materials on an economically competitive basis with biological treatment. Activated carbon regeneration furnaces have high energy requirements.

Sludge Handling and Disposal

Digestion

Digestion is usually used preceding the other sludge concentration and disposal methods. The purpose of digestion is to improve the dewatering of the sludge. Digestion can occur aerobically or anaerobically. During digestion, bacteria decompose the organic material in the sludge producing methane, carbon dioxide and water. At the end of the digestion process, the sludge is stable and non-decomposable.

Vacuum Filtration

The various vacuum filters, usually a revolving drum, use a vacuum to dewater the sludge. The revolving drum type has a vacuum applied against a cloth. The water passes through the cloth and returns to the influent of the treatment plant. The sludge remains on the drum until it is scraped off with a knife.

Centrifugation

Centrifugation uses high speed rotation to separate sludge and water. The heavier sludge moves to the outside and is conveyed to one end, where it is collected for final disposal. The water flows out the opposite end and is returned to the treatment plant.

Sludge Disposal

From any waste water treatment plant, the sludge must be disposed of. The methods used are landfilling, landfarming, barging to sea, and incineration.

Landfilling

A landfill operation requires a large amount of land. Before landfilling, the sludge should be digested to avoid odor problems. The sludge is disposed of in an excavation site. After each batch is disposed of, it is covered with a layer of earth. When the site is filled to capacity it is covered with a thick layer of earth.

The largest problem of industrial landfills is the pollution to ground and surface waters by leaching. Leaching occurs when water percolates through the landfill. As it drains through the landfill site, the water carries with it dissolved and suspended solids and organic matter. This water can then contaminate underground or surface streams it comes in contact with.

Incineration

Exposal. The principal process is fluid bed incineration. In this process, a bed of sand is preheated with hot air to 482-538°C (900 - 1000°F). Torch oil is then used to raise the bed temperature to 649 - 705°C (1200 - 1300°F). At this point waste water sludge and/or sludge is introduced and the torch oil is stopped. The solid products of combustion remain in the bed which is a gradually withdrawn to maintain a constant bed height. Eventually, the bed will be composed of only ash.

The sludge fed to the incinerator usually contains inorganic as well as organic material. However, the sludge must contain a minimum amount of organics to maintain the combustion process. One refinery (26) suggests a minimum of 1,930,000 cal/cu m (29,000 Btu/gal) of sludge heating value is necessary to maintain the combustion process.

SECTION VIII

COST, ENERGY, AND NON-WATER QUALITY ASPECTS

The first part of this section summarizes the costs (necessarilly generalized) and effectiveness of end-of-pipe control technology for BPCTCA and BATEA and BADT-NSPS effluent limitations. Treatment costs for small, medium, and large refineries in each subcategory have been estimated for the technologies considered. The expected annual costs for existing plants in the petroleum refining industry in 1977 consistent with BPCTCA effluent limitations are estimated at \$225 million (end-of-pipe treatment only). For 1983, consistent with BATEA effluent limitations, the estimated additional annual costs are estimated at \$250 million (end-of-pipe treatment only). For BADT-NSPS the annual cost is estimated at \$26 million. These costs are summarized by subcategory in Table 27.

The effect of plant size relative to annual costs can be seen in Table 28 where the annual costs are summarized for application of BPCTCA and BATEA to small, medium, and large refineries in each subcategory. The cost, energy, and nonwater quality aspects of in-plant controls are intimately related to the specific processes for which they are developed. Although there are general cost and energy requirements for equipment items (e.g. surface air coolers), these correlations are usually expressed in terms of specific design parameters, such as the quired heat transfer area. Such parameters are related to the duction rate and specific situations that exist at a particular production site.

There is a wide variation in refinery sizes. When these size ranges are superimposed on the large number of processes within each subcategory, it is apparent that many detailed designs would be required to develop a meaningful understanding of the economic impact of process modifications. The decision to attain the limitations through in-plant controls or by end-of-pipe treatment should be left up to individual manufacturers. Therefore, a series of possible designs for the end-of-pipe treatment models is provided.

Alternative Treatment Technologies

The range of components used or needed for either best practicable or best available technology have been combined into five alternative end-of-pipe treatment steps, which are as follows:

A. Initial treatment, consisting of dissolved air flotation, equalization, neutralization, and nutrient (phosphoric acid) feed facilities.

TABLE 27

Estimated Total Annual Costs for End-of-Pipe Treatment
Systems for the Petroleum Refining Industry (Existing Refineries)

Subcategory		nual Cost, \$ Million
	<u> 1977</u>	1983
Topping	\$14.2	\$16.5
Low-Cracking	35.8	44.4
High-Cracking	45.5	48.1
Petrochemical	53.9	50.0
Lube '	70.1	66.2
Integrated	35.5	24.8
	Industry Total \$255.0	\$250.0

TABLE 28

Summary of End-of Pipe Wastewater Treatment Costs for Representative Plants in the Petroleum Refining Industry

Subcategory	Representative Refinery Size		Annua Level 3 \$/1000 m		Annual Additional Level II Costs \$/1000 m ³ \$/1000 gal	
	1000 m ³ /day	1000 BBL/day				
Topping	0.318	2	0.066	17.31	0.070	18.41
	1.11	7	0.030	7.86	0.034	9.06
	2.4	15	0.018	4.87	0.023	5.97
Low-Cracking	2.4	15	0.014	3.78	0.019	4.90
	5.09	32	0.010	2.53	0.012	3.25
	11.9	75	0.007	1.84	0.008	2.20
High Cracking	4.0	25	0.009	2.34	0.013	3.40
	10.18	64	0.007	1.84	0.008	2.16
	23.8	150	0.006	1.62	0.006	1.47
Petrochemical	4.0	25	0.009	2.32	0.010	2.65
	15.9	100	0.007	1.78	0.006	1.63
	31.8	200	0.005	1.35	0.005	1.20
Lube	4.0	25	0.009	2.33	0.010	2.57
	17.5	110	0.006	1.50	0.006	1.51
	39.8	250	0.005	1.25	0.004	0.93
Integrated	9.8	65	0.006	1.67	0.006	1.53
	23.0	152	0.005	1.28	0.005	1.05
	49.0	326	0.005	1.13	0.003	0.65

- B. Biological treatment, consisting of acitvated sludge units, thickness, digestors, and dewatering facilities.
- C. Granular media filtration, consisting of filter systems an associated equipment.
- D. Physical-chemical treatment facilities consisting of activated carbon adsorption.
- E. Alternative Biological treatment, consisting of aerated lagoon facilities.

Tables 29 through 46 are summaries of the costs of major treatment steps required to achieve different levels of technology for small, medium, and large refineries in each subcategory; using median raw waste loads and median "good water use" flow rates, for the end-of-pipe treatment models.

BPCTCA Treatment Systems Used for the Economic Evaluation

A general flow schematic for the BPCTCA waste water treatment facilities is shown in Figure 6. A summary of the general design basis is presented in Table 47 and a summary of the treatment system effluent limitations for each subcategory is presented in Table 1.

BATEA treatment Systems Used for the Economic Evaluation

BATEA treatment facilities are basically added on to the discharge pine from BPCTCA facilities. It is expected that flows will be reducted slightly by the application of BATEA in-plant technology, so that the activated carbon treatment unit may treat a smaller hydraulic load. However, the activated carbon system was sized for the same flow basis as in BPCTCA technology in order to establish a conservative basis for economic evaluation of proposed effluent limitations.

A general flow schematic diagram for the BATEA waste water treatment facilities is shown in Figure 7. A summary of the general design basis is presented in Table 48. and a summary of the treatment system effluent limitations for each subcategory is presented in Table 2.

TABLE 29

WATER EFFLUENT TREATMENT COSTS

PETROLEUM REFINING INDUSTRY

TOPPING SUBCATEGORY

Refinery Capacity 1000 cubic meters/day (1000 BBL/DAY)	0.318	(2)
Wastewater Flow cubic meters/cublic metric crude oil (gal/bbl)	0.286	(12)
Treatment Plant Size 1000 cubic meters/day	0.091	(0.025)

Costs in \$1000		Altern	ative Treatm	ent Steps
Initial Investment	1 <u>A</u> 183	$\frac{B}{155}$	<u>C</u> 52	<u>D</u> 295
ANNUAL COSTS:				
Capital Costs (10%)	18.3	15.5	5.2	30
Depreciation (20%)	36.6	31.0	10.4	59
Operating Costs	14.6	12.4	4.2	72.5
Energy	1.0	7.8	1.0	6.5
Total Annual Costs	70.5	66.7	20.8	168.0

	Raw Waste Load	Resulting Effluent (Design Average Kg		31_	
	Kg/1000 m ³ (LB/1000 m	BBL)	<u>C</u>	<u>D</u>	
BOD ₅	7.1 (2.5)	4.3	· - .	0.82	
COD	24.0 (8.4)	16.0		2.34	
Oil/Grease	5.1 (1.8)	2.0	1.4	0.17	
Phenol	0.028 (0.01)	0.03	-	0.003	
Sulfide	0.157 (0.055)	0.03	-	0.017	
Ammonia	1.43 (0.5)	1.0		0.23	
Suspended Solids	6.6 (2.5)	5.8	2.9	0.82	

TABLE 30
WATER EFFLUENT TREATMENT COSTS

PETROLEUM REFINING INDUSTRY

TOPPING SUBCATEGORY

Refinery Capacity 1000 cubic meters/day (1000 BBL/DAY)	1.11	(7)	
Wastewater Flow cubic meters/cubic meter crude oil (gal/bbl)	0.286	(12)	
Treatment Plant Size 1000 cubic meters/day (MGD)	0.32	(0.085)
Costs in \$1000	Alternative	Treatment	Steps
Initial Investment $\frac{A}{282}$	2 <u>8</u> 238	<u>C</u> 80	<u>D</u> 612
ANNUAL COSTS:			
Capital Costs (10%) 28 Depreciation (20%) 56	24 48	8 16	61 122
Operating Costs 23 Energy 2	19 12	6 2	89 9

Total Annual Costs

Effluent Quality

	Raw Waste Load Kg/1000 m ³ (LB/1000	Resulting F		
•	ку/1000 ш- (LB/1000	<u>B</u>	<u>c</u>	<u>D</u>
BOD ₅	7.1 (2.5)	4.3	-	0.82
COD	24.0 (8.4)	16.0	· -	2.34
0i1/Grease	5.1 (1.8)	2.0	1.4	0.17
Phenol	0.028 (0.01)	0.03	-	0.003
Sulfide	0.157 (0.055)	0.03	-	0.017
Ammonia	1.43 (0.5)	1.0	-	0.23
Suspended Solids	6.6 (2.3)	5.8	2.9	0.82

109

103

32 281

TABLE 31

PETROLEUM REFINING INDUSTRY

TOPPING SUBCATEGORY

Refinery Capacity 1000 cubic meters/day (1000 BBL/DAY)	2.4	(15)
Wastewater Flow cubic meters/cubic meter crude oil (gal/bbl)	0.28	6 (12)
Treatment Plant Size 1000 cubic meters/day (MGD)	0.68	(0.18)
Costs in \$1000	Alternative	Treatment Steps
Initial Investment ANNUAL COSTS: AND ANNUAL COSTS:		$\frac{C}{114}$ $\frac{D}{943}$
Depreciation (20%) 6 Operating Costs 2	4 32 8 64 8 26 3 19	11 94 23 187 17 101 2 10
Total Annual Costs 13	3 141	46 392

	Raw Waste	**	Resulting Effluent Levels			
	Load Kg/1000 m ³ (LB/1000	(Design Aver	rage Kg/100	00 m ³)		
		<u>B</u>	<u>c</u>	$\underline{\mathbf{D}}$		
BOD 5	7.1 (2.5)	4.3	-	0.82		
COD	24.0 (8.4)	16.0	· -	2.34		
011/Grease	5.1 (1.8)	2.0	1.4	0.17		
Pheno1	0.028 (0.01)	0.03	-	0.003		
Sulfide	0.157 (0.055)	0.03	-	0.017		
Ammonia	1.43 (0.5)	1.0	-	0.23		
Suspended Solids	6.6 (2.3)	5.8	2.9	0.82		

TABLE 32

PETROLEUM REFINING INDUSTRY

LOW CRACKING SUBCATEGORY

Refinery Capacity 1000 cubic meters/day (1000)	O BBL/DAY)	2.4 (15)
Wastewater Flow cubic meters/cubic meter cry	ude oil (gal/bbl)	0.405 (17)
Treatment Plant Size 1000 cubic meters/day	(MGD)	0.97 (0.26)

Costs in \$1000	Alte	rnative Tre	atment St	eps
Initial Investment	<u>A</u>	<u>B</u>	<u>C</u> .	D
ANNUAL COSTS:	368	375	133	1.164
Capital Costs (10%)	37	38	13	116
Depreciation (20%)	. 74	75	26	233
Operating Costs	29	30	11	106
Energy	2	21	3	10
Total Annual Costs	142	164	53	465

	Raw Waste Load ₃ Kg/1000 m (LB/1000	Resulting E (Design Avera BBL)	ffluent Levage Kg/1000	vels m3)
		<u>B</u>	<u>c</u>	<u>D</u>
BOD ₅	71 (25)	6.0	-	1.31
COD	200 (70)	39.1	-	8.0
0il/Grease	27.4 (9.6)	2.8	2.0	0.26
Pheno1	2.85 (1.0)	0.04		0.006
Sulfide	1.0 (0.35)	0.03	_	0.026
Ammonia	10.0 (3.5)	2.0	-	0.51
Suspended Solids	27.4 (9.6)	8.0	4.0	1.31

TABLE 33

PETROLEUM REFINING INDUSTRY

LOW CRACKING SUBCATEGORY

1000 cubic meters/day (1000 BBL/DAY)	5.09 (32)	
Wastewater Flow cubic meters/cubic meter crude oil (g	al/bbl) 0.405 (17)	
Treatment Plant Size meters/day (MGD)	2.06 (0.54	4)

Costs in \$1000	Alternative Treatment Steps				
Initial Investment	<u>A</u>	<u>B</u>	<u>c</u>	D	
ANNUAL COSTS:	487	548	179	1,164	
Capital Costs (10%)	49.	54	18	177	
Depreciation (20%)	98	109	36	354	
Operating Costs	39	44	14	94	
Energy	3	. 31	4	15	
Total Annual Costs	189	238	64	. 640	

	Raw Waste	Resulting	Effluent Le	vels
	Load	(Design Ave	erage Kg/100	$5 \text{ m}^3)$
	$Kg/1000 \text{ m} ^{3}(LB/1000)$	BBL)		
		<u>B</u>	<u>c</u>	<u>D</u>
BOD ₅	71 (25)	6.0	- .	1.31
COD	200 (70)	39.1	-	8.0
Oil/Grease	27.4 (9.6)	2.8	2.0	0.26
Pheno1	2.85 (1.0)	0.04	-	0.006
Sulfide	1.0 (0.35)	0.03	-	0.026
Ammonia	10.0 (3.5)	2.0	_	0.51
Suspended Solids	27.4 (9.6)	8.0	4.0	1.31

TABLE 34

PETROLEUM REFINING INDUSTRY

LOW CRACKING SUBCATEGORY

Refinery Capacity 1000 cubic meters/day (1000 BBL/DAY)	11.9 (75)
Wastewater Flow cubic meters/cubic meter crude oil (gal/bbl)	0.405 (17)
Treatment Plant Size 1000 cubic meters/day (MGD)	4.8 (1.3 MGD)

Costs in \$1000	Alternative Treatment Steps				
Initial Investment	<u>A</u> 787	<u>B</u> 1062	C 251	D 2,895	
ANNUAL COSTS:	707	1002			
Capital Costs (10%)	79	106	25	290	
Depreciation (20%)	158	212	50	579	
Operating Costs	64	86	20	152	
Energy	. 8	59	7	25	

Total Annual Costs

	Kg/10	Raw Waste Load 000 m ³ (LB/1000	(D	Effluent Leve	_
			<u>B</u>	<u>c</u>	<u>D</u>
BOD 5	71 (25)		6.0	· -	1.31
COD	200 (70)		39.1	- .	8.0
Oil/Grease	27.4 (9.6)		2.8	, 2.0	0.26
Phenol	2.85 (1.0)		0.04	-	0.006
Sulfide	1.0 (0.35)	,	0.03	-	0.026
Ammonia	10.0 (3.5)	•	2.0	-	0.51
Suspended Solids	27.4 (9.6)		8.0	4.0	1.31

TABLE 35

PETROLEUM REFINING INDUSTRY

HIGH CRACKING SUBCATEGORY

4.0 (25)

Wastewater Flow cubic meters/cubic meter crude oil (g	al/bbl)	0.5 (2	1)	
Treatment Plant Size 1000 cubic meters/day (MGD)		2.0 (0	.525)	
Costs in \$1000	Alte	rnative Trea	tment Ste	ps
Initial Investment	A	<u>B</u>	· <u>C</u>	<u>D</u> .
ANNUAL COSTS:	443	470	170	1,720
Capital Costs (10%)	44	47	17	1.72
Depreciation (20%)	89	94	34	344
Operating Costs	36	38	13	121
Energy	4	29	4	15
Total Annual Costs	173	208	68	652

Effluent Quality

Refinery Capacity

nery Capacity
1000 cubic meters/day (1000 BBL/DAY)

	Raw Waste Load Kg/1000 m ³ (LB/10	(Degian Av	Effluent Le erage Kg/10	
•		<u>B</u>	<u>C</u>	D
BOD 5	83 (29)	8.9	-	1.65
COD	260 (91)	68.0	-	12.8
Oil/Grease	31.4 (11)	3.5	2.5	0.34
Pheno1	5.1 (1.8)	0.05	-	0.006
Sulfide	1.28 (0.45)	0.05	· -	0.034
Ammonia	32.8 (11.5)	4.5	-	1.65
Suspended Solids	32.2 (11.3)	10.2 9	5.1	1.65

TABLE 36

PETROLEUM REFINING INDUSTRY

HIGH CRACKING SUBCATEGORY

Refinery Capacity 1000 cubic meters/day (1000 BBL/DAY)	,	10.18	(64)	
Wastewater Flow cubic meters/cubic meter crude oil (ga	ı1/bb1)	0.500	(21)	
Treatment Plant Size 1000 cubic meters/day (MGD)	·	5.09	(1.35)	
Costs in \$1000	Alte	ernative Trea	tment S	teps
Initial Investment	<u>A</u> 817	$1,\frac{B}{105}$	<u>C</u> 261	<u>D</u> 2,950
ANNUAL COSTS:		·		•
Capital Costs (10%) Depreciation (20%)	82 164	110 221	26 52	295 590
Operating Costs Energy	67 9	90 59	21 8	155 25
Total Annual Costs	322	480	107	1,065

·	Raw Waste Load Kg/1000 m ³ (LB/1000	Resulting Effluent Levels (Design Average Kg/1000 m ³) BBL)		
	,	<u>B</u>	<u>C</u>	D
BOD ₅	83 (29)	8.0	. -	1.65
COD	260 (91)	68.0	-	12.8
Oil/Grease	31.4 (11.0)	3.5	2.5	0.34
Pheno1	5.1 (1.8)	0.51	-	0.006
Sulfide	1.28 (0.45)	0.05	-	0.034
Ammonia	32.8 (11.5)	4.5	-	0.82
Suspended Solids	32.2 (11.3)	10.2	5.1	1.65

TABLE 37

PETROLEUM REFINING INDUSTRY

HIGH CRACKING SUBCATEGORY

1000 cubic meters/day (1000 BBL/DA	Y)	23.8 (150) a1/bb1) 0.500 (21)		
Wastewater Flow cubic meters/cubic meter crude oil	(gal/bb1)			
Treatment Plant Size 1000 cubic meters/day (MGD)		11.	9 (3.2)	
Costs in \$1000	Alte	rnative Tre	atment St	ceps
Initial Investment	1.403	<u>B</u> 2,763	<u>C</u> 367	<u>D</u> 4,890
ANNUAL COSTS:		- , ,,,,		.,
Capital Costs (10%)	140	276	37	489
Depreciation (20%)	280	553	74	987
Operating Costs	119	236	31	211
Energy	17	113	15	44

Total Annual Costs 556 1,178 157 1,722

Effluent Quality

Refinery Capacity

	Raw Waste Load Kg/1000 m ³ (LB/1000 B	Resulting Effluent Levels (Design Average Kg/1000 m ³) BBL)		_
		<u>B</u>	<u>C</u>	$\underline{\mathbf{D}}$
BOD ₅	83 (29)	8.0	· _	1.65
COD	260 (91)	68.0	-	12.8
Oil/Grease	31.4 (11)	3.5	2.5	0.34
Pheno1	5.1 (1.8)	0.51	-	0.006
Sulfide	1.28 (0.45)	0.05	_	0.034
Ammonia	32.8 (11.5)	4.5	-	0.82
Suspended Solids	32.2 (11.3)	10.2	5.1	1.65

TABLE 38

PETROLEUM REFINING INDUSTRY

PETROCHEMICAL SUBCATEGORY

Refinery Capacity 1000 cubic meters/day (1000 BBI	/DAY)	4.0	(25)	
Wastewater Flow cubic meters/cubic meter crude of	oil (gal/bbl)	0.59	95 (25)	
Treatment Plant Size 1000 cubic meters/day (MGD))	2.4	(0.625)	
Costs in \$1000	Alte	ernative Tro	eatment St	ерв
Initial Investment	<u>A</u>	$\frac{\mathbf{B}}{\mathbf{B}}$	<u>C</u>	<u>D</u>
ANNUAL COSTS:	495	595	190	1,875
Capital Costs (10%)	49	60	19	188
Depreciation (20%)	99	119	- 38	375
Operating Costs	39	48	15	125
Energy	5	34	4	16
Total Annual Cost	rs 192	261	76	604

	Raw Waste Load Kg/1000 m (LB/1000 BB	Resulting Effluent Levels (Design Average Kg/1000 m ³) BBL)		_
		В	<u>c</u>	D
BOD ₅	148 (52)	9.1	. ,	1.8
COD	371 (130)	59.5	-	7.1
0il/Grease	45.6 (16)	4.3	3.1	0.37
Pheno1	10.3 (3.6)	. 0.065	-	0.009
Sulfide	1.7 (0.59)	0.06	-	0.054
Ammonia	34.2 (12)	6.8	-	1.65
Suspended Solids	44.2 (15.5)	12.0	6.0	1.8

TABLE 39

PETROLEUM REFINING INDUSTRY

PETROCHEMICAL SUBCATEGORY

Refinery Capacity

1000 cubic meters/day (1000 BBL/DAY)

Wastewater Flow

cubic meters/cubic meter crude oil (gal/bbl)

Treatment Plant Size

1000 cubic meters/day (MGD)

Costs in \$1000	Alternative Treatment Steps		eps	
Initial Investment	$\frac{A}{1,165}$	$\frac{B}{2,410}$	<u>C</u> 335	<u>D</u> 4,200
ANNUAL COSTS:	1,100	2,410	333	4,200
Capital Costs (10%)	117	241	33	420
Depreciation (20%)	233	482	67	840
Operating Costs	98	203	29	192
Energy	15	93	12	38
Total Annual Costs	463	1,019	141	1,490

÷	Raw Waste Load Kg/1000 m ³ (LB/1000 B	Resulting Effluent Levels (Design Average Kg/1000 m ³) BBL)		_
		<u>B</u>	<u>c</u>	D
BOD 5	148 (52)	9.1	-	1.8
COD	371 (130)	59.05	. -	7.1
Oil/Grease	45.6 (16)	4.3	3.1	0.37
Pheno1	10.3 (3.6)	0.065	-	0.009
Sulfide	1.7 (0.59)	0.06	. •••	0.054
Ammonia	34.2 (12)	6.8	-	1.65
Suspended Solids	44.2 (15.5)	12.0	6.0	1.8

TABLE 40

PETROLEUM REFINING INDUSTRY

PETROCHEMICAL SUBCATEGORY

Refinery Capacity 1000 cubic meters/day (1000 BBL/DA	AY)	31.8	(200)	
Wastewater Flow cubic meters/cubic meter crude oil	(ga1/bb1)	0.59	95 (25)	
Treatment Plant Size 1000 cubic meters/day (MGD)		18.9	(5.0)	
Costs in \$1000		ternative Trea	tment St	eps
Initial Investment	A 1,6 <u>6</u> 2	<u>B</u> 3,785	<u>C</u> 423	<u>D</u> 6,220
ANNUAL COSTS:	4, 004	3,.03	3	0,0
Ćapital Costs (10%)	166	379	42	622
Depreciation (20%)	332 -	757	85	1,244
Operating Costs	145	329	37	270
Energy	25	155	20	60
Total Annual Costs	668	1,620	184	2,196

,	Raw Waste	•	Resulting Effluent Levels		
	Load Kg/1000 m ³ (LB/1000	(Design Ave	erage Kg/100	0 m^3	
	Rg/ 1000 m (111/ 1000	<i>DD1</i>)			
		B	<u>C</u>	$\overline{\mathbf{D}}$	
BOD ₅	148 (52)	9.1,	-	1.8	
COD	371 (130)	59.5	-	7.1	
011/Grease	45.6 (16)	4.3	3.1	0.37	
Pheno1	10.3 (3.6)	0.065	_	0.009	
Sulfide	1.7 (0.59)	0.06	-	0.054	
Ammonia	34.2 (12)	6.8	-	1.65	
Suspended Solids	44.2 (15.5)	12.0	6.0	1.8	

TABLE 41

PETROLEUM REFINING INDUSTRY

LUBE SUBCATEGORY

Refinery Capacity 1000 cubic meters/day (1000 BBL/DAY)		4.0	(25)	
Wastewater Flow cubic meters/cubic meter crude oil (gal/bbl)		0.881	(37)	
Treatment Plant Size 1000 cubic meters/day (MGD)		3.5	(0.92	5)
Costs in \$1000	Alternative	Treatme	ent St	eps
Initial Investment	<u>A</u> <u>B</u> 895		2 <u>C</u> 247	$2,\frac{D}{3}60$
ANNUAL COSTS:				
Depreciation (20%) Operating Costs Energy	78 90 56 179 52 72 6 47 02 388		24 49 20 5	236 472 139 20 867

	Raw Waste Load Kg/1000 m ³ (LB/1000 Bi		(Degian	Resulting Effluent Levels (Design Average Kg/1000 m ³) BL)		
			<u>B</u>	<u>c</u>	D	
BOD 5	200	(66)	10.3	-	3.6	
COD	382	(134)	95.5	-	23.5	
0il/Grease	136	(38)	6.0	4.3	0.7	
Pheno1	6.2	(202)	0.088	-	0.015	
Sulfide	1.1	(0.4)	0.088	-	0.073	
Ammonia	227 79	(7.8)	4.5	-	1.5	
spended Solids	54	(28)	17.6	8.8	3.55	

TABLE 42

PETROLEUM REFINING INDUSTRY

LUBE SUBCATEGORY

Refinery Capacity 1000 cubic meters/day (1000 BBL/DAY)		17	.5 (11	0)
Wastewater Flow cubic meters/cubic meter crude oil (ga	1/bb1)	. 0	.881 (3	7)
Treatment Plant Size 1000 cubic meters/day (MGD)		15	.4 (4.	07) ·
Costs in \$1000	Alter	native Trea	tment S	teps
Initial Investment	1 <u>, A</u> 1, <u>5</u> 01	$\frac{B}{3,316}$	<u>C</u> 413	<u>D</u> 5,600
ANNUAL COSTS:	1,501	3,310	413	3,600
Capital Costs (10%) Depreciation (20%) Operating Costs Energy	150 300 129 20	332 663 285 135	41 83 35 17	,
Total Annual Costs	599	1,415	176	1,968,00

	Raw Waste Load Kg/1000 m ³ (LB/1000		Resulting Effluent Levels (Design Average Kg/1000 m ³		
			<u>B</u>	<u>c</u>	<u>D</u>
BOD ₅	187	(66)	10.3	-	3.6
COD	382	(134)	95.5	-	23.5
0i1/Grease	136	(48)	6.0	4.3	0.7
Phenol	6.2	(2.2)	0.088	-	0.015
Sulfide	1.1	(0.4)	0.088	-	.0.073
Ammonia	22	(7.8)	4.5	_	1.5
Suspended Solids	79	(28)		8.8	3.55

TABLE 43

PETROLEUM REFINING INDUSTRY

LUBE SUBCATEGORY

Refinery Capacity 1000 cubic meters/day (1000 BBL/DAY)		39.8	(250))
Wastewater Flow cubic meters/cubic meter crude oil (gal,	/bb1)	0.8	81 (37))
Treatment Plant Size 1000 cubic meters/day (MGD)		35	(9.2	25)
Costs in \$1000	<u>Alte</u>	rnative Trea	tment St	eps.
Initial Investment	$\frac{A}{846}$	$6,\frac{B}{6}10$	<u>C</u> 534	<u>D</u> 8,890
ANNUAL COSTS:	-	•		
Capital Costs (10%) Depreciation (20%) Operating Costs Energy	285 570 256 45	661 1,322 595 245	53 106 48 35	889 1778 370 95
Total Annual Costs	1,156	2,823	242	3,132

	Raw Waste Load Kg/1000 m ³ (LB/1000		Resulting Effluent Levels (Design Average Kg/1000 m ³)		
	Kg/1000 m ²	(LB/1000	<u>B</u>	<u>c</u>	D
BOD ₅	187	(70	10.3	-	3.6
COD	382	(134)	95.5	• <u>-</u>	23.5
0il/Grease	136	(48)	6.0	4.3	0.7
Pheno1	6.2	(2.2)	0.088	-	0.015
Sulfide	1.1	(0.4)	0.088	_	0.073
Ammonia	22	(7.8)	4.5	-	1.5
Suspended Solids	. 79	(28)	17.6	8.8	3.55

TABLE 44

PETROLEUM REFINING INDUSTRY

INTEGRATED SUBCATEGORY

	ry Capacity 000 cubic m		day (100	O BBL/DA	Υ)		9.8	(65)
	ater Flow abic meters	/cubic	meter cr	ude oil	(gal/bbl)		1.1	(46)
	ent Plant S 000 cubic m		day (MG	D)			10.8	(3.0)
Costs :	in \$1000		٠		Alternative	Treatment	Steps	
Initia	l Investmen	ıt		1,256	2,920	<u>C</u> 23 ¹ 4	<u>D</u> 4,75	0
ANNUAL	COSTS:							
De Oj	apital Cost epreciation perating Co nergy	(20%		126 252 103 20	292 584 243 106	23 46 21 15	47 95 20 4	0
	Total A	nnual	Costs	501	1,225	. 105	1,67	<u>'</u> 4
Effluer	nt Quality							
···	· .	R	aw Waste	Res	sulting Efflu	ėnt Levels		
	Kg/1000 m		oad 1000 BBL)	(De	esign Average	Kg/1000 m ³	_)	
	<u>. </u>			•	<u>B</u>	<u>C</u>	$\overline{\mathbf{D}}$	
BOD ₅		238	(84)		16.1		3.7	
COD		590	(208)	,	124.5		21 .§	i
Oil/Gre	ease	133	(47)		8.0	5.4	0.6	3
Phenol		6.5	(2.3)		0.111	<u>L</u> .	0.0	15
Sulfide	:	1.7	(0.6		0.111	L	0.0	7 §
Ammonia		35.4	(12.5)		7.1		1.9	
Suspend	led Solids	29	(10.2)		21.6	10.8	3.7	

TABLE 45

PETROLEUM REFINING INDUSTRY

INTEGRATED SUBCATEGORY

Refinery Capacity 1000 cubic meters/day (10	23 (152)					
Wastewater Flow cubic meters/cubic meter crude oil (gal/bbl)							
Treatment Plant Size 1000 cubic meters/day (MGD)							
Costs in \$1000		Alternative Tr	eatment S	teps			
Initial Investment	2,2 4 9	<u>B</u> 5 ,2 23	<u>C</u> 418	7,600			
ANNUAL COSTS:			•				
Capital Costs (10%) Depreciation (20%) Operating Costs Energy	225 550 203 36	522 1,044 470 {188	42 84 38 21	760 1,520 329 68			
Total Annual Costs	1,014	2,224	192	2,677			

Raw Waste Load Kg/1000 m ³ (lb/1000 BBL)			Resulting Effluent Levels (Design Average Kg/1000 m ³):			
Kg/1000 iii	, (10/1	OOO DDL)	<u>B</u>	<u>C</u>	D	
BOD ₅	238	(84)	16.1		3.7	
COD	590 ((2 08)	124.5		21.3	
Oil/Grease	133	(47)	8.0	5.4	0.63	
Phenol	6.5	(2.3)	0.111		0.015	
Sulfide	1.7	(0.6)	0.111	•	0.073	
Ammonia	35.4	(12.5)	7.1		1.9	
Suspended Solids	29	(10.2	21.6	10.8	3.7	

TABLE 46

PETROLEUM REFINING INDUSTRY

INTEGRATED SUBCATEGORY

Refinery Capacity 1000 cubic m		day (1000	BBL/DAY)		49	(326)
Wastewater Flow cubic meters	/cubic	meter crue	de oil (ga	1/bbl)	1.	.1 (46)
Treatment Plant S 1000 cubic m		day (MGD)		54	.0 (15.0)
Costs in \$1000			<u>A</u>	lternative Tr	reatment S	teps
Initial Investmen	.t		4, <u>2</u> 32	9,8 <u>3</u> 1	<u>c</u> 787	<u>D</u> 10,100
ANNUAL COSTS:						
Capital Cost Depreciation Operating Co Energy	(20%		423 846 381 69	983 1,966 885 354	79 158 71 52	1,010 2,020 439 107
Total A	nnual	Costs	1,719	4,188	360	3,576
Effluent Quality						
Kg/1000 m	I	aw Waste		ting Effluent gn Average Kg	_	
Kg/1000 III	(10)	(144 0001		<u>B</u>	<u>C</u>	<u>D</u>
.BOD ₅	238	(84)		16.1		3.7
COD	590	(208)		124.5		21.3
Oil/Grease	133	(47)		8.0	5.4	0.63
Phenol	6.5	(2.3)		0.111		0.015
Sulfide	1.7	(0.6)		0.111		0.073
Ammonia	35.4	(12.5)		7.1	,	1.9
Suspended Solids	29	(10.2)		21.6	10.8	3.7

MODEL SYSTEM USED FOR THE ECONOMIC EVALUATION

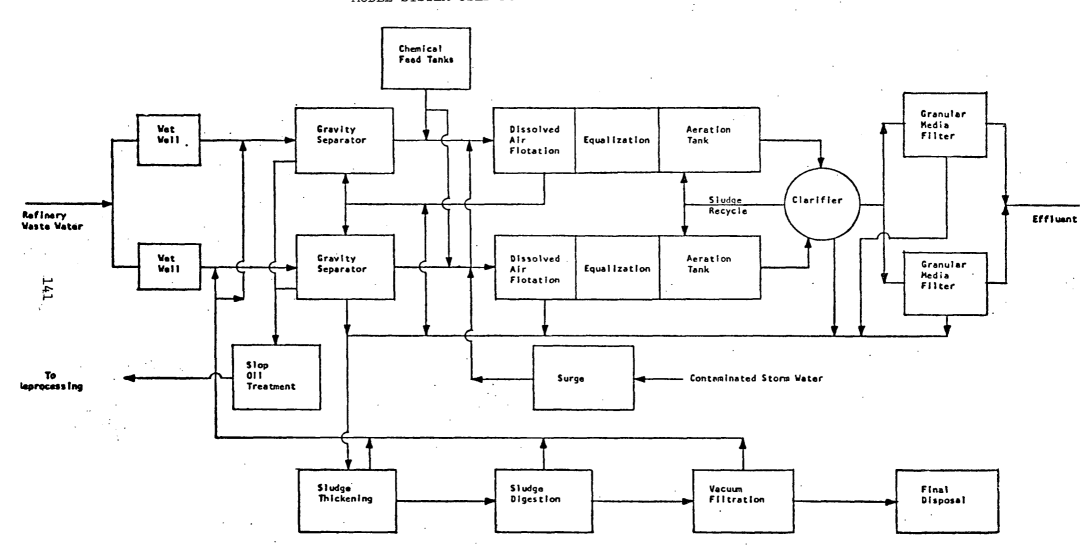


TABLE 47

BPCTCA - END OF PIPE TREATMENT SYSTEM MODEL USE FOR THE ECONOMIC EVALUATION DESIGN SUMMARY

Treatment System Hydraulic Loading

Treatment system hydraulic loadings are sized to represent the projected waste water flows from small, average, and large refineries in each subcategory. The flow range used in these estimates ranges from 95 to 38,000 m³/day (25,000 gpd to 10,000,000 gpd).

Dissolved Air Flotation

The flotation units are sized for an overflow rate of 570 m³/day/m² (1400 gpd/sq.ft)

Pump Station

Capacity to handle 200 percent of the average hydraulic flow.

Equalization

One day detention time is provided. Floating mixers are provided to keep the contents completely mixed.

<u>Neutralization</u>

The two-stage neutralization basin is sized on the basis of an average detention time of twenty minutes. The lime-handling facilities are sized to add 1,000 lbs. of hydrated lime per mgd of waste water, to adjust the pH. Bulk-storage facilities (based on 15 days usage) or bag storage is provided, depending on plant size. Lime addition is controlled by two pH probes, one in each basin. The lime slurry is added to the neutralization basin from a lime slurry recirculation loop. The lime-handling facilities are enclosed in a building.

Nutrient Addition

Facilities are provided for the addition of phosphoric acid to the biological system in order to maintain the ratio of BOD:P at 100:1.

<u>Aeration Basin</u>

Platform-mounted mechanical aerators are provided in the aeration basin. In addition, walkways are provided to all aerators for fan

access and maintenance. The following data were used in sizing the aerators.

1.5 kg O2/kg BOD Oxygen utilization (1.5 lbs 02/lb. BOD) removed L 0.8 В 0.9 20°C Waste water temperature 1.6 kg (3.5 lbs.) 02/hr./shaft HP at Oxygen transfer 20°C and zero D.O. in tap water Motor Efficiency 85 percent Minimum Basin D.O. 1 mg/1

Oxygen is monitored in the basins using D.O. probes.

Secondary Clarifiers

All secondary clarifiers are circular units. The side water depth is 3.0 meters (10 ft.) and the overflow rate is 500 gpd/sq. ft.). Sludge recycle pumps are sized to deliver 50 percent of the average flow.

Sludge Holding Tank-Thickener

For the smaller plants, a sludge-holding tank is provided, with sufficient capacity to hold 5 days flow from the aerobic digester. The thickener provided for the larger plants was designed on the basis of 29 kg/m²/day (6 lbs./sq. ft./day) and a side water depth of 3.0 meters (10 ft.)

Aerobic Digester

The aerobic digester is sized on the basis of a hydraulic detention time of 20 days. The sizing of the aerator-mixers was based on 0.044HP/m^3 (1.25/HP/1,000 cu.ft.) of digester volume.

Vacuum Filtration

The vacuum filters were sized on cake yield of 9.75 kg/m²/hr. (2 lbs./sq.ft./hr) and a maximum running time of 18 hrs./day. The polymer system was sized to deliver up to 0.005kg of polymer/kg of day solids (10 lbs. of polymer/ton dry solids).

Granular Media Filters

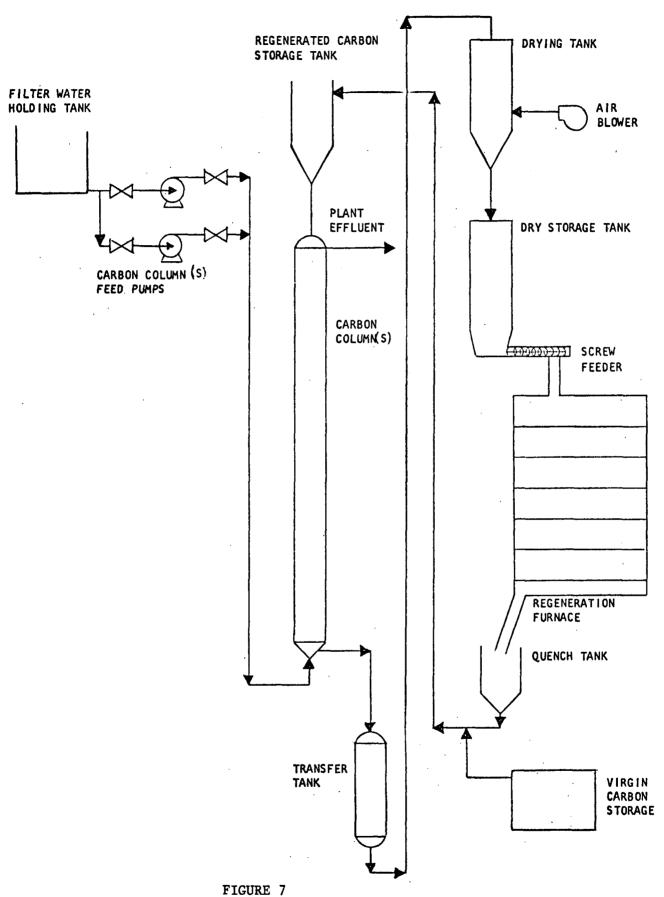
The filters are sized on the basis of an average hydraulic loading of 9.12m³/m²/min. (3 gpm/sq.ft.) Backwash facilities are sized to provide rate up to 0.82m³/m²/min. (20 gpm/sq.ft.) and a backwash cycle of up to 20 minutes duration.

Final Sludge Disposal

Sludge is disposed of at a sanitary landfill assumed to be 5 mile from the waste water treatment facility.

Design Philosophy

The plant's forward flow units are designed for parallel flow, i.e. either half of the plant can be operated independently. The sludge facilities are designed on the basis of series flow. All outside tankage is reinforced concrete. The tops of all outside tankage are assumed to be 12 ft above grade.



BATEA - PROPOSED TREATMENT 145

TABLE 48

BATEA - END OF PIPE TREATMENT SYSTEM DESIGN SUMMARY

Granular Carbon Columns

The carbon columns are sized on a hydraulic loading of 0.4-0.8 m³/m²/min. (10-20 gpm/sq. ft.) and a column detention time of 40 minutes. A backwash rate of (50 gpm/sq. ft.) was assumed for 40 percent bed expansion at $70^{\circ}F$.

Filter-Column Decant Sump

Tankage is provided to hold the backwash water and decant it back to the treatment plant over a 24-hour period. This will eliminate hydraulic surging of the treatment units.

Regeneration Furnace

An exhaustion rate of 1 kg of COD/kg carbon (1 lb COD/lb carbon) was used for sizing the regeneration facilities.

Regenerated Exhausted Carbon Storage

Tankage is provided to handle the regenerated and exhausted carbon both before and after regeneration.

Estimated Costs of Facilities

discussed previously, designs for the model treatment systems were sted out in order to evaluate the economic impact of the proposed effluent limitations. The design considerations resulted in the generation of cost data which would be conservative. However, relatively conservative cost numbers are preferred for this type of general, economic analysis.

Activated sludge followed by granular media filtration was used as the BPCTCA treatment system. The plant designs were varied to generate cost effectiveness data within each category. Activated carbon adsorption was used as the BATEA treatment.

Capital and annual cost data were prepared for each of the proposed treatment systems.

The capital costs were generated on a unit process basis, e.g. equalization, neutralization, etc. for all the proposed treatment systems. The following "percent add on" figures were applied to the total unit process costs in order to develop the total capital cost requirements:

Percent of Unit Process Capital Cost	-
12	
15	
8	
3	
uction	
15	
15	
	Process Capital Cost 12 15 8 3 uction 15

Land costs were computed independently and added directly to the total capital costs.

Annual costs were computed using the following cost basis:

Item	Cost Allocation
Amortization Depreciation Operations and Maintenance	10 percent of investment. 5 year-straight line with zero salvage value. Includes labor and supervision, chemicals sludge, hauling and disposal, insurance and taxes (computed at 2 percent of the capital cost), and maintenance (computed at 4 percent of the capital cost).
Power	Based on \$1.50/100 KWH for electrical power.

The short term capitalization and depreciation write-off period is .that which is presently acceptable under current Internal Revenue Service Regulations pertaining to industrial pollution control equipment.

All cost data were computed in terms of August, 1971 dollars, which corresponds to an Engineering News Records (ENR) value of 1580.

The following is a qualitative as well as a quantitative discussion of the possible effects that variations in treatment technology or design criteria could have on the total capital costs and annual costs.

Technology or Design Criteria

1. Use aerated lagoons and sludge dewatering lagoons in place of the proposed treatment system.

2. Use earthern basins with a plastic liner in place of reinforced concrete construction, and floating aerators versus platform-mounted aerators with permanent-access walkways.

Capital Cost Differential

- 1. The cost reduction could be to 70 percent of the proposed figures.
- 2. Cost reduction could be 10 to 15 percent of the total cost.
- 3. Place all treatment tankage above grade 3. Cost savings would depend to minimize excavation, especially if on the individual situation. a pumping station is required in any case. Use all-steel tankage to minimize capital cost.
- 4. Minimize flow and maximize concentrations through extensive in-plant recovery and water conservation, so that other treatment technologies (e.g. incineration) may be economically competitive.
- 4. Cost differential would depend on a number of items, e.g. age of plant, accessibility to process piping, local air pollution standards, etc.

The cost requirements for implementing BPCTCA effluent standards are presented in Tables 49 through 54. The additional cost requirements for implementing BATEA effluent standards are presented in Tables 55 and 60.

The following table summarizes the general ranges of sludge quantities generated by small, medium, and large refineries in each subcategory.

cu m/yr 1	cu yd/yr 1
2.3-15	3 - 20
76-380	100 - 500
380-2300	500 - 3000
460-3800	600 - 5000
610-6900	800 - 9000
760-9200	1000 - 12000
	2.3-15 76-380 380-2300 460-3800 610-6900

'Wet-weight basis

Particular plants within the petrochemical, lube, and integrated subcategories may be amenable to sludge incineration because of the large quantities of sludge involved. For example, sludge incineration would reduce the previous quantities by about 90 percent. Sludge cake is 80 percent water, which is evaporated during incineration, and more than half of the remaining (20 percent) solids are thermally oxidized during incineration. Sludge incineration costs were not evaluated for those specific cases, because the particular economics depend to a large degree on the accessibility of a sanitary landfill and the relative associated hauling costs.

The following discussion is presented to help visualize the complexities involved in evaluating cost effectiveness data. Every treatment system is composed of units whose design basis is primarily hydraulically dependent, organically dependent, or a combination of the two. The following is a list of the unit processes employed, and a breakdown of the design basis.

Hydraulically Dependent	Organically Dependent	Hydraulically and Organically Dependent
Pump station API separator Equalization Neutralization Nutrient addition Sludge recycle pump Clarifier	Thickener Aerobic Digestor Vacuum filter	Aeration basin Oxygen transfer equipment Air flotation Unit

The annual cost associated with the hydraulically dependent unit processes is not a function of effluent level. On the other hand, the sizing of the organically dependent units should theoretically vary in direct proportion to the effluent level: e.g. reducing the BOD5 removal from 95 to 85 percent should reduce the sizes of the sludge handling equipment by approximately 10 percent. However, there are two complicating factors: 1) only a relatively few sizes of commercially available equipment; and 2) broad capacity ranges. These two factors,

especially in regard to vacuum filters, tend to negate differentials. in capital cost with decreasing treatment levels.

The relationship between design varying contaminant levels and the design of aeration basins and oxygen transfer equipment is somewhat more complex. The levels are dependent on the hydraulic flow, organic concentration, sludge settleability, and the relationship between mixing and oxygen requirements. For example, to reach a particular effluent level, the waste water's organic removal kinetics will require a particular detention time at a given mixed-liquor concentration. The oxygen transfer capacity of the aerators may or may not be sufficient to keep the mixed liquor suspended solids in suspension within the aeration basin. Therefore, the required horsepower would be increased merely to fulfill a solids mixing requirement. Alternatively, the oxygen requirements may be such that the manufacturer's recommended minimum spacing and water depth requirements would require that the basin volume be increased to accommodate oxygen transfer requirements.

Non-Water Quality Aspects

The major nonwater quality consideration which may be associated with in-process control measures is the use of and alternative means of ultimate disposal of either liquid or solid wastes. As the process Raw Waste Load is reduced in volume, alternate disposal techniques such as incineration, ocean discharge, and deep-well injection are feasible. Recent regulations are tending to limit the applicability of ocean discharge and deep-well injection because of the potential long-term detrimental effects associated with these disposal procedures Incineration may be a viable alternative for highly concentrated wast However, associated air pollution and the need for auxiliary depending on the heating value of the waste, are considerations which must be evaluated on an individual basis for each use. nonwater quality aspects, such as noise levels, will not be perceptibly affected. Most refineries generate fairly high noise levels (85-95 within the battery limits because of equipment such as pumps, compressors, steam jets, flare stacks, etc. Equipment associated with in-process or end-of-pipe control systems would not add significantly to these levels. In some cases, substituting vacuum pumps for steam jets would in fact reduce plant noise levels. There are no radioactive nuclides used in the industry, other than in instrumentation. Thus no radiation problems will be expected. Compared to the odor emissions possible from other refinery sources, odors from the waste water treatment plants are not expected to create a significant problem. However, odors are possible from the waste water facilities, especially from the possible stripping of ammonia and sulfides in the air flotation units, and from accidental anaerobic conditions in biological facilities during upsets.

The extra power required for waste water treatment and control systems is negligible compared to the total power requirements of the petroleum fining equipment.

TABLE 49

BPCTCA ESTIMATED WASTEWATER TREATMENT COSTS FOR THE TOPPING SUBCATEGORY

(ENR 1580 - August, 1971 Costs)

	Flow, 1000 M3/Day (GPD)		
	0.091 (25,000)	0.32 (85,000)	0.68 (180,000)
Total Capital Cost	\$390,000	\$600,000	\$773,000
Annual Cost	•		
Capital Costs	\$ 39,000	\$ 60,000	\$ 77,000
Depreciation	\$ 78,000	\$120,000	\$155,000
Operating and Maintenance	\$ 31,200	\$ 48,000	\$64,000
Energy and Power	\$ 9,800	\$ 16,000	\$ 24,000
Total Annual Cost	\$158,000	\$244,000	\$320,000

TABLE 50

BPCTCA ESTIMATED WASTEWATER TREATMENT COSTS FOR THE LOW-CRACKING SUBCATEGORY

(ENR 1580 - August, 1971 Costs)

	Flow, 1000 M ³ /Day (GPD)		
	0.97 (260,000)	2.06 (540,000)	4.8 (1,300,000)
Total Capital Cost	\$876,000	\$1,214,000	\$2,100,000
Annual Cost	,		·
Capital Costs	\$ 88,000	\$ 121,000	\$ 210,000
Depreciation	\$175,000	\$ 243,000	\$ 420,000
Operating and Maintenance	\$ 70,000	\$ 97,000	\$ 170,000
Energy and `Power	\$ 26,000	\$ 38,000	\$ 74,000
Total Annual Cost	\$359,000	\$ 499,000	\$ 874,000

TABLE 51

BPCTCA ESTIMATED WASTEWATER TREATMENT COSTS FOR THE HIGH-CRACKING SUBCATEGORY

(ENR 1580 - August, 1971 Costs)

	Flow, 1000 M ³ /Day (GPD)		
	2.0 (525,000)	5.09 (<u>1,350,000</u>) 11.	9 (3,200,000)
Total Capital Cost	\$1,083,000	\$2,183,000	\$4,533,000
Annual Cost			٠.
Capital Costs	\$ 108,000	\$ 218,000	\$ 453,000
Depreciation	\$ 217,000	\$ 437,000	\$ 907,000
Operating and Maintenance	\$ 87,000	\$ 178,000	\$ 386,000
Energy and Power	\$ 37,000	\$ 76,000	\$ 145,000
Total Annual Cost	\$ 449,000	\$ 909,000	\$1,891,000

TABLE 30

BPCTCA ESTIMATED WASTEWATER TREATMENT COSTS FOR PETROCHEMICAL SUBCATEGORY

(ENR 1580 - August, 1971 Costs)

Flow, 1000 M^3/Day (GPD) 9.5 (2,500,000)2.4 (625,000) 18.9 (5,000,000) \$1,280,000 Total Capital Cost \$3,910,000 \$5,870,000 Annual Cost \$ 128,000 Capital Costs \$ 391,000 587,000 Depreciation \$ 256,000 782,000 \$1,174,000 102,000 330,000 Operating and 511,000 Maintenance Energy and 43,000 120,000 200,000 Power Total Annual Cost . \$1,623,000 529,000 \$2,472,000

TABLE 53

BPCTCA ESTIMATED WASTEWATER TREATMENT COSTS FOR THE LUBE SUBCATEGORY

(ENR 1580 - August, 1971 Costs)

	Flow, 1000 M ³ /Day (GPD)		
	3.5 (<u>925,000</u>)	15.4 (4,000,000)	35 <u>(9,250,000</u>)
Total Capital Cost	\$1,920,000	\$5,230,000	\$9,990,000
Annual Cost			
Capital Cost	\$ 192,000	\$ 523,000	\$ 999,000
Depreciation	\$ 384,000	\$1,046,000	\$1,998,000
Operating and Maintenance	\$ 154,000	\$ 449,000	\$ 899,000
Energy and Power	\$ 58,000	\$ 172,000	\$ 325,000
Total Annual Cost	\$ 788,000	\$2,190,000	\$4,221,000

TABLE 56

BPCTCA ESTIMATED WASTEWATER TREATMENT COSTS FOR THE INTEGRATED SUBCATEGORY

(ENR 1580 - August, 1971 Costs)

	Flow, 1000 M3/Day (GPD)		
	10.8 (3.000.000	25.5 (<u>7.000.000)</u>	54.0 <u>(15.000.00</u> 0)
Total Capital Cost	\$ 4,410,000	\$7,890,000	\$14,850,000
Annual Cost			
Capital Cost	\$ \h1 5000	\$ 789,000	\$ 1,485,000
Depreciation	\$ 882,000	\$1,578,000	\$ 2,970,000
Operating and Maintenance	\$ 367,000	\$ 657,000	\$ 1,237,000
Energy and Power	\$ 141,000	\$ 254 , 000	\$ 479,000
Total Annual Cost	1,831,600′	\$3,278,000	\$ 6,171,000

TABLE 55

ESTIMATED ADDITIONAL WASTEWATER TREATMENT COSTS FOR

BATEA TECHNOLOGY - TOPPING SUBCATEGORY

(ENR 1580 - August, 1971 Costs)

Flow, 1000 M^3/Day (GPD)

•	0.91 (25,000)	0.32 (<u>85,000)</u>	0.68 (180,000)
Total Capital Cost	\$295,000	\$612,000	\$943,000
Annual Cost			
Capital Cost	\$ 30,000	\$ 61,000	\$ 94,000
Depreciation	\$ 59,000	\$122,000	\$187,000
Operating and Maintenance	\$ 72,500	\$89,000	\$101,000
Energy and Power	\$ 6,500	\$ 9,000	\$ 10,000
Total Annual Cost	\$168,000	\$281,000	\$392,000

TABLE 56

ESTIMATED ADDITIONAL WASTEWATER TREATMENT COSTS FOR

BATEA TECHNOLOGY - LOW CRACKING SUBCATEGORY

(ENR 1580 - August, 1971 Costs)

	Flow, 1000 M3/Day (GPD)		
	0.97 (260,000)	2.06 (5 ¹ 40,000)	4.8 (1,300,000)
Total Capital Cost	\$1,164,00	\$1,769,000	\$2,895,000
Annual Cost			
Capital Cost	\$ 116,000	\$ 177,000	\$ 290,000
Depreciation	\$ 233,000	\$ 354,000	\$ 579,000
Operating and Maintenance	\$ 106,000	\$ 94,000	\$ 152,000
Energy and Power	\$ 10,000	\$ 15,000	\$ 25,000
Total Annual Cost	\$465,000	\$640,000	\$1,046,000

TABLE 57

ESTIMATED ADDITIONAL WASTEWATER TREATMENT COSTS FOR

BATEA TECHNOLOGY - HIGH CRACKING SUBCATEGORY

(ENR 1580 - August, 1971 Costs)

Flow, 1000 M^3/Day (GPD) 5.09 (1,350,000)11.9 (3,200,000) 2.0 (525,000) \$4,890,000 \$2,950,000 Total Capital Cost \$1,720,000 Annual Cost Capital Cost 172,000 \$ 295,000 489,000 Depreciation 344,000 978,000 590,000 211,000 Operating and 121,000 155,000 Maintenance 44,000 15,000 25,000 Energy and Power \$1,065,000 \$1,722,000 652,000 Total Annual Cost

TABLE 58

ESTIMATED ADDITIONAL WASTEWATER TREATMENT COSTS FOR

BATEA TECHNOLOGY - Petrochemical Subcategory

(ENR 1580 - August, 1971 Costs)

Flow,

	2.4 (625,000)	9.5 (<u>2,500,000</u>)	18.9(5,000,000)
Total Capital Cost	\$1,875,000	\$4,200,000	\$6,220,000
Annual Cost			
Capital Cost	\$ 188,000	\$ 420,000	\$ 622,000
Depreciation	\$ 375,000	\$ 840,000	\$1,244,000
Operating and Maintenance	\$ 125,000	\$ 192,000	\$ 270,000
Energy and Power	\$ 16,000	\$ 38,000	\$ 60,000
Total Annual Cost	\$ 604,000	\$1,490,000	\$2,196,000

TABLE **3**9

ESTIMATED ADDITIONAL WASTEWATER TREATMENT COSTS FOR
BATEA TECHNOLOGY - Lube Subcategory
(ENR 1580 - August, 1971 Costs)

Flow, 1000 M^3/Day (GPD) 15.4 (4,000,000) **3.**5 (925,000) 35 (9,250,000) \$8,890,000 \$2,360,000 Total Capital Cost \$5,600,000 Annual Cost \$ 236,000 \$ 560,000 \$ 889,000 Capital Cost \$ 472,000 \$1,120,000 \$1,778,000 Depreciation \$ 236,000 \$ 370,000 Operating and \$ 139,000 Maintenance 20,000 52,000 95,000 Energy and Power \$3,132,000 \$ 867,000 \$1,968,000 Total Annual Cost

TABLE 60

ESTIMATED ADDITIONAL WASTEWATER TREATMENT COSTS FOR

BATEA TECHNOLOGY - INTEGRATED SUBCATEGORY

(ENR - August, 1971 Costs)

	Flow, 1000 M ³ /Day (GPD)							
	10.8 (3,000,000)	25.5 (7,000,000)	54.0 (15,000,000)					
Total Capital Cost	\$4,750,000	\$7,600,000	\$10,100,000					
Annual Cost	÷							
Capital Cost	\$ 475,000	\$ 760,000	\$ 1,010,000					
Depreciation	\$ 950,000	\$1,520,000	\$ 2,020,000					
Operating and Maintenance	\$ 206,000	\$ 329,000	\$ 439,000					
Energy and Power	\$ 43,000	\$ 68,000	\$ 107,000					
Total Annual Cost	\$1,674,000	\$2,677,000	\$ 3,576,000					

SECTION IX

BEST PRACTICABLE CONTROL TECHNOLOGY CURRENTLY AVAILABLE--EFFLUENT LIMITATIONS

Based on the information contained in Sections III through VIII of this report, effluent limitations commensurate with the best practicable control technology currently available have been established for each petroleum refining subcategory. The limitations, which explicitly set numerical values for the allowable pollutant discharges within each subcategory, are presented in Table 1. The effluent limitations specify allowable discharges of BOD5, COD, TOC, total suspended solids, oil and grease, phenolic compounds, ammonia (N), sulfides, total and hexavalent chromium and zinc; based upon removals which are capable of being attained through the application of BPCTCA pollution control technology.

The best practicable control technology currently available is based on both in-plant and end-of-pipe technology. BPCTCA in-plant technology is based on control practices widely used within the petroleum refining industry, and include the following:

- 1. Installation of sour water strippers to reduce the sulfide and ammonia concentrations entering the treatment plant.
- 2. Elimination of once-through barometric condenser water by using surface condensers or recycle systems with oily water cooling towers.
- 3. Segregation of sewers, so that unpolluted storm runoff and oncethrough cooling waters are not treated normally with the process and other polluted waters.
- 4. Elimination of polluted once-through cooling water, by monitoring and repair of surface condensers or by use of wet and dry recycle systems.

BPCTCA end-of-pipe treatment technology is based on the existing waste water treatment processes currently used in the Petroleum Refining Industry. These consist of equalization and storm diversion; initial oil and solids removal (API separators or baffle plate separators); further oil and solids removal (clarifiers, dissolved air flotation, or filters); carbonaceous waste removal (activated sludge, aerated lagoons, oxidation ponds, trickling filter, activated carbon, or combinations of these); and filters (sand, dual media; or multi-media) following biological treatment methods. It must be recognized that specific treatability studies are required prior to the application of a specific treatment system to the individual refinery.

Granular media filtration or polishing ponds prior to final discharge are included so that the total suspended solids and oil concentrations in the final effluent can be generally maintained at approximately 10 mg/l and 5 mg/l, respectively. The final polishing step is consider BPCTCA for the petroleum refining industry since several refineries are now using polishing ponds, and granular media filters are becoming excepted technology with a few installations operating currently and several more now under construction.

In a petroleum refinery the waste water treatment plant should be used to treat only polluted waters. All once-through cooling water or storm runoff which is unpolluted should be segregated as it dilutes the polluted waters and requires treatment of a greater flow. Flows for BPCTCA were based on the 50 percent probability of occurance flows for plants practicing recycle with less than 3 percent heat removal by once-through cooling water (on a dry weather basis). Recognizing the additional flows and waste loads associated with rain runoff and ballast waters, allocations for these added flows must be given based on strict segregation of runoff and ballast waters treated.

PROCEDURE FOR DEVELOPMENT OF BPCTCA EFFLUENT LIMITATIONS

The effluent guideline limitations were determined using effluent data from refineries visited during this project or attainable effluent concentrations and the median flow from the refineries with 3 percent or less of the heat removed by once-through cooling water. In some cases the available data from the refineries visited was considered to be too stringent to be met by the industry in general. In these cases the flow and concentration procedure was used. The median flows are presented Table 21, Section V. The attainable concentrations for BPCTCA are presented in Table 61. Refinery data are presented in Tables 24+26, Section VII.

Several exceptions to this procedure were required to establish meaningful effluent limitations in specific cases. These are as follows:

Topping, Low Cracking, Petrochemical, Lube, and Integrated Subcategories - Ammonia as Nitrogen

The ammonia as nitrogen effluent limitations were calculated using an 80 percent reduction from the median raw waste loads in each subcategory.

Low Cracking, High Cracking, Petrochemial and Lube Subcategories -

Little data is available on the reduction of TOC. Available effluent data indicate an effluent TOC/BOD ratio of 1.8. Using this factor, effluent limitations for TOC, were based on BOD5 limitations. It is recognized that this ratio (TOC/BOD) is variable between the refineries,

and prior to use, an agreed upon correlation should be developed for the individual refinery.

Topping Subcategory - TOC

Application of the procedure outlined above yielded a TOC value of 2.7 lb/1000 bbl, which is higher than the median raw waste load for the topping subcategory. Therefore, the median raw waste load value was used for the topping subcategory TOC effluent limitations.

Topping Subcategory - COD

The COD effluent limitation was calculated using the COD/BOD ratio determined from published refinery data. This ratio was applied to the BOD5 removal for the topping subcategory A to determine the COD removal efficiency.

The long term (annual or design) average effluent limitations determined are contained in table 62.

Statistical Variability of a Properly Designed and Operated Waste Treatment Plant

The effluent from a properly designed and operated treatment plant changes continually due to a variety of factors. Changes in production mix, production rate and reaction chemistry influence the composition of raw wasteload and, therefore, its treatability. Changes in biological factors influence the efficiency of the treatment process. A common licator of the pollution characteristics of the discharge from a plant is the long-term average of the effluent load, however, the long-term (e.g., design or yearly) average is not a suitable parameter on which to base an enforcement standard. However, using data which show the variability in the effluent load, statistical analyses can be used to compute short-term limits (monthly or daily) which should never by exceeded, provided that the plant is designed and run in the proper way to achieve the desired long-term average load. It is these short-term limits on which make up the effluent guidelines.

In order to reflect the variabilities associated with properly designed and operated treatment plants for each of the parameters as discussed above, a statistical analysis was made of plants where sufficient data was available to determine these variances for day-to-day and month-to-month operations. The standard deviations for day-to-day and month-to-month operations were calculated. For the purpose of determing effluent limitation a variability factor was defined as follows:

Standard deviation = Q Long-term average (yearly or design) = x Variability factor $y = \frac{x+2Q}{y}$ The annual average is multiplied by the variability factor to determine the effluent limitation guideline for each parameter. The effluent limitation guideline as calculated by use of the variability fact based on two standard deviations is only exceeded 2-3 percent of the time for a plant that is attaining the long-term average. The data used for the variability analysis came from plants under voluntary operation. As a result of the application of mandatory requirements, the effluent limitation guidelines as discussed in this paragraph should never be exceeded by a properly designed and operated waste treatment facility.

The variability factors used are contained in Table 63. These factors for each parameter except total and hexavalent chromium and zinc were calculated from long-term refinery data. The factors for total chromium and zinc are the same as that used for suspended solids since both of these metallic ions are removed as insoluble salts. The variability factor for hexavalent chromium was based on the sulfide variability. The guidelines for BPCTCA presented in table 1 have taken into consideration the above variability factors.

TABLE 61

Attainable Concentrations from the Application of Best Practicable Control Technology Currently Available

Parameter		
BOD ₅	15	
COD	80	
TOC	*(1.8 x BOD ₅)	
SS	10	
O & G	5	
Pheno1	0.1	
NH ₃ -N	*(80% removal)	
Sulfide	0.1	
CrT	.25	
Cr6	.005	
Zħ	.5	

^{*}See Text

TABLE 62
BPCTCA

PETROLEUM REFINING INDUSTRY EFFLUENT LIMITATIONS Annual Daily Kilograms of Pollutants/1000 Cubic Meters Feedstock (1) Per Stream Day (Annual Average Daily Pounds of Pollutant/1000 BBL of Feedstock Per Stream Day)

Refinery Subcategory	BOD5	COD	TOC	Total Suspended Solids	Oil & Grease	Phenolic Compounds	Ammonia (N)	Sulfide	Total Chromium	Hexavalent Chromium	Zinc
Topping	4.3(1.5)	16.0(5.6)	4.9(1.7)	2.9(1.0)	1.4(0.5)	0.03(0.01)	1.0(0.35)	0.028(0.010)	0.071(0.025)	0.0014(0.0005)	0.142(0.050)
Low-Cracking	6.0(2.1)	39.1(13.8)	10.8(3.8)	4.0(1.4)	2.0(0.7)	0.040(0.014)	2.0(0.70)	0.040(0.014)	0.100(0.035)	0.0020(0.0007)	0.202(0.071)
High-Cracking	8.0(2.8)	68.0(24.0)	14.4(5.0)	5.1(1.8)	2.5(0.88)	0.051(0.018)	4.5(1.60)	0.050(0.018)	0.115(0.044)	0.0025(0.0009)	0.249(0.088)
Petrochemical	9.1(3.2)	59.5(21.0)	16.3(5.7)	6.0(2.1)	3.1(1.10)	0.065(0.023)	6.8(2.46)	0.060(0.021)	0.148(0.052)	0.0029(0 0010)	0.297(0.104)
Lube	10.7(3.8)	95.5(33.7)	19.4(6.0)	8.8(3.1)	4.3(1.5)	0.088(0.031)	4.5(1.60)	0.088(0.031)	0.219(0.077)	0.0044(0.0015)	0.436(0.154)
Intergrated	16.1(5.7)	124.5(43.9)	29.1(10.3)	10.8(3.8)	5.4(1.9)	0.111(0.039)	7.1(2.5)	0.111(0.039)	0.278(0.098)	0.0055(0.0020)	0.555(0.196)
Runoff(2)	0.015(0.125)	0.12(1.0)	0.027(0.225)	0.010(0.083)	0.0050(0.042)	·					
Ballast(3)	0.015(0.125)	0.15(1.250)) 0.027(0.225)	0.010(0.083)	0.0050(0.042)						

- (1) Feedstock -crude oil and/or natural gas liquids
- (2) The additional allocation being allowed for contaminated storm runoff flow (kg/1000 liters (lbs/1000 gallons) shall be based solely on that storm flow which passes through the treatment system. All additional storm runoff, that has been segregated from the main waste stream, shall not show a visible sheen or exceed a TOC concentration of 15 mg/l when discharged.
- (3) This is an additional allocation, based on ballast water intake (daily average) kilograms per 1000 liters (pounds per 1000 gallons)

TABLE 63

VARIABILITY FACTORS BASED ON PROPERLY DESIGNED AND OPERATED WASTE TREATMENT FACILITIES

	BOD ₅	COD	TOC	<u>TSS</u>	O & G	Phenol	Ammonia	Sulfide	$\underline{\mathtt{CrT}}$	<u>Cr6</u>	\underline{Zn}
Daily Variability	2.1	2.0	1.6	2.0	2.0	2.4	2.0	2.2	2.0	2.2	2.0
Monthly Variability	1.7	1.6	1.3	1.7	1.6	1.7	1.5	1.4	1.7	1.4	1.7

SECTION X

BEST AVAILABLE TECHNOLOGY ECONOMICALLY ACHIEVABLE -- EFFLUENT LIMITATIONS

The application of best available technology economically achievable is being defined as further reductions of water flows in-plant and the addition of a physical - chemical treatment step (activated carbon), end-of-pipe. The limitations, which set numerical values for the allowable pollutant discharges within each subcategory for BATEA are presented in Table 2. Although there are specific systems which can effectively reduce the water usage from a particular process to nearly zero, these "zero discharge" systems cannot be uniformly applied throughout the refinery to develop "zero discharge" criteria for the entire refinery.

BATEA in-plant technology is based on control practices now practiced by some plants in the petroleum refining industry, and include the following:

- (1) Use of air cooling equipment.
- (2) Reuse of sour water stripper bottoms in crude desalters.
- (3) Reuse of once-through cooling water as make-up to the water treatment plant.
- (4) Using waste water treatment plant effluent as cooling water, scrubber water, and influent to the water treatment plant.
- (5) Reuse of boiler condensate as boiler feedwater.
- (6) Recycle of water from coking operations.
- (7) Recycle of waste acids from alkylation units.
- (8) Recycle of overhead water in water washes.
- (9) Reuse overhead accumulator water in desalters.
- (10) Use of closed compressor and pump cooling water system.
- (11) Reuse of heated water from the vacuum overhead condensers to heat the crude. This reduces the amount of cooling water needed.
- (12) Use of rain runoff as cooling tower make-up or water treatment plant feed.
- (13) Other methods.

Flow

Flow reductions proposed for BATEA effluent limitations were derived from further analysis of the 1972 National Petroleum Waste Water Characterization Studies. The flows from refineries in each subcategory meeting the BPCTCA flow basis were averaged to determine the flow basis for establishment of BATEA effluent limitations. That these average flows are achievable within the petroleum refining industry is readily demonstrable, by determining the number and geographical distribution of

refineries in the United States currently at, or lower than, the proposed BATEA flows. There are 3 to 5 refineries in each of the six subcategories which have flows less than or equal to the proposed BATEA effluent limitations. These refineries range in size from 827,000 69,000,000 cubic meters per stream day (5,200 to 434,000 barrels per stream day), and range in cracking capacity from 0 to 106 percent. The geographical distribution of these refineries indicates that good water practices, and consequently low waste water flows, are not confined to water - short areas or cool climates, but are located throughout the United States. Within this group of refineries with low-water usage, there are refineries located in both high rainfall and dry areas (Washington and New Mexico) and areas of extreme temperatures (New Mexico and Texas to Alaska and Minnesota).

Consequently, these flows, shown in Table 64, were used as the basis for establishment of BATEA effluent limitations. The objective of this basis for flow is to provide inducement for in-plant reduction of both flow and contaminant loadings prior to end-of-pipe treatment. However, it is not the intent of these effluent limitations to specify either the unit waste water flow which must be achieved or the waste water treatment practices which must be employed at the individual petroleum refinery.

The end-of-pipe system proposed for BATEA technology is based on the addition of activated carbon adsorption in fixed bed columns, to the treatment system proposed as BPCTCA technology.

Procedure for Development of BATEA Effluent Limitations

The effluent limitations proposed for BATEA technology are based on refinery pilot plant data, which indicate the percentage reductions achievable or concentrations achievable for effluents from activated carbon adsorption systems. These data are presented in table 65.

These concentrations were then used in conjunction with the BATEA flows from Table 65 or the percentage reductions were applied to the BPCTCA effluent limit. The daily annual average effluent limitations determined are contained in Table 66.

Since these effluent limitations are based upon pilot plant data, which have not been fully demonstrated in full-scale installations as actual performance data becomes available, the effluent limitations presented in Table 2 may require revision.

Variability Allowance for Treatment Plant Performance

The effluent limitations presented in Table 2 have taken into consideration the variability factors, as in BPCTCA. Since there is not enough performance data from physical - chemical treatment systems

available at this time to determine variability, the ratios established for BPCTCA have been used.

TABLE 64

FLOW BASIS FOR DEVELOPING
BATEA EFFLUENT LIMITATIONS

Subcategory	Flow, per unit	throughout
	M3/M3	Gallons/BBL
Topping	0.17	. 7
Low-Cracking	0.26	11
High-Cracking	0.33	14
Petrochemical	0.36	15
Lube	0.73	30.5
Integrated	0.76	31.5



BATEA REDUCTIONS IN POLLUTANT LOADS ACHIEVABLE BY APPLICATION OF ACTIVATED CARBON TO MEDIA FILTRATION EFFLUENT BPCTCA

	<u>Parameter</u>	Type of Data	Achievable Refinery Eff mg/L	Fluent % Reduction	References
	BOD	Pilot Plant	5	-	21,27,31A,48,62A
	COD	Pilot Plant	-	75	21,27,31A,47,53,62A
•	тос	Pilot Plant	15	-	17,31A,48,62A
	TSS	Pilot Plant	5	· -	31A,48,53,62A
7	011	Pilot Plant	1-1.7	80	31A,48,62A
	Phenols	Pilot Plant	0.02	99	31A,48,62A
	Ammon i a	Pilot Plant		60	27 ,31A,62A
	Sulfides	No data	-	~	

TABLE 66
BATEA

Annual Average Daily Kilograms of Pollutants/1000 Cubic Meters of Feedstock (1) Per Stream Day (Annual Average Daily Pounds of Pollutants/1000 BBL of Feedstock Per Stream Day)

Refinery Subcategory	BCD5 COD	тос	Total Suspended Solids	Oil & Grease	Phenolic Compounds	Ammonia (N)	Sulfide	Total Chromium	Hexavalent Chromium	Zinc
Topping	0.82 (0.29) 2.3 (0.82)	2.5 (0.87)	0.82 (0.29)	0.17 (0.06)	0.0031 (0.0011)	0.23 (0.08)	0.017 (0.006	0.042 (0.015)	0.00084 (0.0003)	0.082 (0.029)
Low-Cracking	1.31 (0.46) 8.0 (2.8)	4.0 (1.4)	1.3 (0.46)	0.26 (0.09)	0.0050 (0.0018)	0.51 (0.18)	0.026 (0.009	0) 0.065 (0.023)	0.0013 (0.00045)	0.13 (0.046)
High-Cracking	1.65 (0.58) 12.8 (4.5)	5.1 (1.8)	1.7 (0.58)	0.34 (0.12)	0.0060 (0.0023)	0.82 (0.29)	0.034 (0.012	2) 0.082 (0.029)	0.0016 (0.00058)	0.16 (0.058)
Petrochemical	1.80 (0.63) 7.1 (2.5)	5.4 (1.9)	1.8 (0.63)	0.37 (0.13)	0.0090 (0.0025)	1.65 (0.58)	0.054 (0.019	0.088 (0.031)	0.0018 (0.00062)	0.18 (0.063)
Lube	3.59 (1.27) 23.5 (8.3)	10.8 (3.8)	3.6'(1.27)	0.71 (0.25)	0.014 (0.0051)	1.50 (0.53)	0.074 (0.026	5) 0.18 (0.064)	0.0037 (0.00127)	0.36 (0.127)
Integrated	3.70 (1.31) 21.2 (7.5)	11.0 (3.9)	3.7 (1.31)	0.74 (0.26)	0.015 (0.0053)	1.94 (0.68)	0.074 (0.026	0.066)	0.0037 (0.00131)	0.38 (0.131)
Runoff (2)	0.0050 (0.042) 0.014 (0.1	2) 0.016 (0.13)	0.0050 (0.042)	0.0010 (0.00	9)	,- .				
Ballast (3)	0.0050 (0.042) 0.019 (0.1	6) 0.016 (0.13)	0.0050 (0.042)	0.0010 (0.00	9)	~~				

⁽¹⁾ Feedstock - Crude oil and/or natural gas liquids.

⁽²⁾ The additional allocation being allowed for contaminated storm runoff flow (kg/1000 liters (1bs/1000 gallons) shall be based solely on that storm flow which passes through the treatment system. All additional storm runoff, that has been segregated from the main waste stream, shall not show a visible sheen or exceed a TOC concentration of 15 mg/l when discharged.

⁽³⁾ This is an additional allocation, based on ballast water intake (daily average) kilograms per 1000 liters (pounds per 1000 gallons).

SECTION XI

NEW SOURCE PERFORMANCE STANDARDS

Recommended effluent limitations for new source performance standards are based upon the application of BPCTCA control technology to the waste water flows used as the basis for BATEA effluent limitations. The proposed BADT effluent limitations are shown in Table 3.

The refining technology available today does not call for major innovations in refining processes. Basically, BADT refining technology consists of the same fundamental processes which are already in practice, with few modifications and additions. However, a major design criterion for new refinery capacity is reuse/ recycle of water streams to the greatest extent possible, in order to minimize discharges to waste water treatment facilities. Consequently, the water flow on which new source performance standards were based is identical to the best available technology economically achievable flow, which reflects the best water usage as demonstrated in the petroleum refining industry. These flows are shown in Table 65.

It should be clearly understood that no recommendations have been made, nor are any implied, regarding the substitution of processes which produce a lower raw waste load for others with higher raw waste load.

This is based on the consideration that the choice of a particular commercial route is governed largely by the availability of feedstocks on the conditions in the product markets. Companies produce a given a composition within the industry. The substitution of a cleaner process may be possible for new producers from a technical point of view, but completely impossible based on limited availability of the required alternative feedstocks or on the lack of viable markets for new coproducts.

The waste water treatment technology recommended for BADT effluent limitations is the same as called for by BPCTCA and does not include physical - chemical treatment, because that technology has not been sufficiently demonstrated by the petroleum refining industry.

Procedure for Development of BADT Effluent Limitations

The effluent limitations proposed for BADT technology are based on the concentrations considered achievable by BPCTCA and the flows from BATEA. The daily annual average effluent limitations thus determined are contained in Table 67.

Variability Allowance for Treatment Plant Performance

The guideline numbers presented in Table 3 have taken into consideration the variability factors, as in BPCTCA. Since the treatment technologiand process technology for BADT are the same as BPCTCA, the ratios established for BPCTCA have been used in BADT.

TABLE 67

NEW SOURCE PERFORMANCE STANDARDS FOR THE PETROLEUM REFINING INDUSTRY Annual Daily Kilograms of Pollutants/1000 Cubic Meters of Feedstock (1) Per Stream Day (Annual Average Daily Pounds of Pollutants/1000 BBL of Feedstock Per Streatm Day

Refinery Subcategory	BOD5	COD	<u> </u>	Total Suspended Solids	Oil & Grease	Phenolic Compounds	Armonia (N)	Sulfide	Total Chromium	Hexavalent Chromium	Zinc
Topping	2.5 (0.88)	9.4 (3.3)	2.9 (1.0)	1.65 (0.58)	0.83 (0.29)	0.017 (0.0058)	0.57 (0.20)	0.016 (0.0058)	0.042 (0.015)	0.00084 (0.0003)	0.082 (0.029)
Low-Cracking	4.0 (1.4)	25.2 (8.9)	7.1 (2.5)	2.6 (0.92)	1.32 (0.46)	0.026 (0.0092)	1.28 (0.45)	0.026 (0.0092)	0.065 (0.023)	0.0013 (0.00045)	0.13 (0.046)
High-Cracking	5.1 (1.8)	45.3 (16.0)	9.1 (3.2)	3.4 (1.2)	1.65 (0.58)	0.034 (0.012)	3.1 (1.1)	0.034 (0.012)	0.082 (0.029)	0.0016 (0.00058)	0.16 (0.058)
Petrochemical	5.4 (1.9)	35.7 (12.6)	9.7 (3.4)	3.7 (1.3)	1.8 (0.63)	0.036 (0.0125)	4.0 (1.4)	0.037 (0.013)	0.088 (0.031)	0.0018 (0.00062)	0.18 (0.063)
Lube	8.8 (3.1)	78.7 (27.7)	15.9 (5.6)	7.1 (2.5)	3.5 (1.24)	0.074 (0.026)	3.7 (1.3)	0.072 (0.025)	0.18 (0.064)	0.0037 (0.00127)	0.36 (0.127)
Integrated	11.1 (3.9)	85.5 (30.1)	20.2 (7.1)	7.4 (2.6)	3.7 (1.30)	0.077 (0.027)	4.8 (1.7)	0.077 (0.027)	0.19 (0.066)	0.0037 (0.00131)	0.38 (0.131)
Runoff, (2) 0	.014 (0.12)	0.056 (0.47)	0.017 (0.14)	0.011 (0.09)	0.0050 (0.04	(2)			. ,		
Ballast (3) 0	.014 (0.12)	0.072 (0.60)	0.017 (0.14)	0.011 (0.09)	0.0050 (0.04	.2)					

⁽¹⁾ Feedstock - Crude oil and/or natural gas liquids.

⁽¹⁾ The additional allocation being allowed for contaminated storm runoff flow (kg/1000 liters (1bs/1000 gallons) shall be based solely on that storm flow which passes though the treatment system. All additional storm runoff, that has been segregated from the main waste stream, shall not show a visible sheen or exceed a TOC concentration of 15 mg/l when discharged.

⁽³⁾ This is an additional allocation, based on ballast water intake (daily average) kilograms per 1000 liters (pounds per 1000 gallons).

TABLE 68

METRIC UNITS

CONVERSION TABLE

MULTIPLY (ENGLISH UNITS)

Ъу

TO OBTAIN (METRIC UNITS)

ENGLISH UNIT	ABBREVIATION	CONVERSION	ABBREVIATION	METRIC UNIT
acre	ac	0.405	ha	hectares
acre - feet	ac ft	1233.5	cu m	cubic meters
British Thermal				
Unit	BTU	. 0.252	kg cal	kilogram - calories
British Thermal			_	
Unit/pound	BTU/1b	0.555	kg cal/kg	kilogram calories/kilogram
cubic feet/minute	cfm	0.028	cu m/min	cubic meters/minute
cubic feet/second	cfs	1.7	cu m/min	cubic meters/minute
cubic feet	cu ft	0.028	cu m	cubic meters
cubic feet	cu ft	28.32	1	liters
cubic inches	cu in .	16.39	cu cm	cubic centimeters
degree Fahrenheit	F°	0.555(°F-32)*	°C	degree Centigrade
feet	ft	0.3048	m	meters
gallon'	gal	3.785	1	liters
gallon/minute	gpm	0.0631	1/sec	liters/second
horsepòwer	hp	0.7457	kw	killowatts
inches	in	2.54	cm	centimeters
inches of mercury	in Hg	0.03342	atm	atmospheres
pounds	1b	0.454	kg	kilograms
million gallons/day	mgd	3,785	cu m/day	cubic meters/day
mile	mi	1.609	km	kilometer
pound/square	•			,
inch (gauge)		$(0.06805 \text{ psig } +1)^*$	atm	atmospheres (absolute)
square feet	sq ft	0.0929	sq m	square meters
square inches	sq in	6.452	sq cm	square centimeters
tons (short)	t	0.907	kkg _	metric tons (1000 kilograms)
yard	у	0.9144	m	meters

^{*} Actual conversion, not a multiplier

SECTION XII

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SECTION XIII

BIBLIOGRAPHY

- 1. American Petroleum Institute, "Petroleum Industry Raw Waste Load Survey," December, 1972.
- 2. Annessen, R. J., and Gould, G. D., "Sour Water Processing Turns Problem Into Payout," Chemical Engineering, March 22, 1971.
- 3. Annual Refining Surveys, "Survey of Operating Refineries in the U.S.," The Oil and Gas Journal, April 1, 1973.
- 3a. Annual Refining Surveys, "Survey of Operating Refineries in the U.S.", The Oil and Gas Journal 1967.
- 4. Armstrong, T. A., "There's Profit in Processing Foul Water," The Oil and Gas Journal, pp. 96-98, June 17, 1968.
- 5. Beavon, David K., "Add-On Process Slashes Claus Tailgas Pollution," Chemical Engineering, pp. 71-73, December 13, 1971.
- 6. "The Beavon Sulfur Removal Process for the Purification of Sulfur Plant Tailgas," Ralph M. Parsons Company Publication.
- Benger, M., "Disposal of Liquid and Solid Effluents from Oil Refineries," 21st Purdue Industrial Waste Conference, pp. 759-767, 1966.
- 8. Beychok, M. R., Aqueous Wastes from Petroleum and Petrochemical Plants, John Wiley & Sons, London, 1967.
- 9. Beychok, M. R., "Waste water Treatment of Skelly Oil Company's El Dorado, Kansas Refinery," 16th Purdue Industrial Waste Conference, pp. 292-303, 1961.
- 10. Brown, K. M., "Some Treating and Pollution Control Process for Petroleum Refineries," The Second INTERPETROL Congress, Rome, Italy, June 22-27, 1971.
- 11. Brownstein, Arthur M., U.S. Petrochemicals, The Petroleum Publishing Co., Tulsa, Oklahoma, 1972.
- 12. Brunner, D. R., and Keller, D. J., "Sanitary Landfill Design and Operation," U.S. Environmental Protection Agency, Washington, D.C., 1972.

- 13. Campbell, G. C., and Scoullar, G. R., "How Shell Treats Oakville Effluent," Hydrocarbon Processing & Petroleum Refiner, 43 (5): 137-140, May, 1964.
- 14. "Chevron Waste Water Treating Process," Chevron Research Company Publication, September, 1968.
- 15. Cohen, J. M., "Demineralization of Waste waters," Advanced Waste Treatment and Water Reuse Symposium, Volume 2, February 23-24, 1971.
- 16. Conser, R. E., "The Environmental Fuels Processing Facility," SNG Symposium, Institute of Gas Technology, Chicago, Illinois, March 12-16, 1973.
- 17. Culp, R. L., and Culp, G. L., Advanced Waste water Treatment, Van Nostrand Reinhold Company, New York, 1971.
- 18. Davis, R. W., Biehl, V. A., and Smith, R. M., "Pollution Control and Waste Treatment at an Inland Refinery," 19th Purdue Industrial Waste Conference, pp. 128-138, 1964.
- 19. Daniels, E. K., Latz, J. R., Castler, L. A., "Pollution Control at Ferndale, Washington," 23rd Midyear Meeting of the American Petroleum Institute's Division of Refining, May, 1958.
- 20. Denbo, R. T., and Gowdy, F. W., "Baton Rouge Waste Water Control Program Nears End," The Oil and Gas Journal, pp. 62-65, May 29, 1972.
- 21. Diehl, D. S., Denbo, R. T., Bhatta, M. N., and Sitman, W. D., "Effluent Quality Control at a Large Refinery," Journal of the Water Pollution Control Federation, 43 (11), November, 1971.
- 22. Dorris, T. C., Patterson, D., Copeland, B. J., "Oil Refinery Effluent Treatment in Ponds," 35th Meeting of the Water Pollution Control Federation, pp. 932-939, # October 7-11, 1962.
- 23. Easthagen, J. H., Skrylov, V., and Purvis, A. L., "Development of Refinery Waste water Control at Pascagowle, Mississippi," Journal of Water Pollution Control Federation, 37 (12): 1621-1628, December, 1965.
- 24. Elkin, H. F., "Activated Sludge Process Application to Refinery Effluent Waters," Journal of Water Pollution Control Federation, 28 (9): 1122-1129, September, 1956.
- 25. Fair, G. M., Geyer, J. C., and Okum, D. A., Water and Waste water Engineering, Volume 2, John Wiley & Sons, Inc., New York, 1968.

- 26. Fluid Bed Incineration of Petroleum Refinery Wastes for the Environmental Protection Agency, Washington, D.C., March, 1971. 12050KET
- 27. Ford, D. L., and Buercklin, M. A., "The Interrelationship of Biological-Carbon Adsorption Systems for the Treatment of Refinery and Petrochemical Waste waters," 6th International Water Pollution Research Conference, Session 11, Hall C, June 18-23, 1972.
- 28. Fosberg, T. M., "Industrial Waste Water Reclamation," 74th National Meeting American Institute of Chemical Engineers, March 11-15, 1973.
- 29. Franzen, A. E., Skogan, V. G., and Grutsch, J. F., "Successful Tertiary Treatment at American," The Oil and Gas Journal, April 3, 1972.
- 30. Gillian, A. S., and Anderegg, F. C., "Biological Disposal of Refinery Wastes," 14th Purdue Industrial Waste Conference, pp. 145-154, 1959.
- 31. Gloyna, E. G., Brady, S. O., and Lyles, H., "Use of Aerated Lagoons and Ponds in Refinery and Chemical Waste Treatment," 41st Conference of the Water Pollution Control Federation, September 22-27, 1968.
- 31a. Hale, J.H., and Myers, L.H., "The Agencies Removed by Carbon Treatment of Refinery Waste waters".
 - Hart, J. A., "Air Flotation Treatment and Reuse of Refinery Waste water," 25th Annual Purdue Industrial Waste Conference, May, 1970.
- 33. Hart, J. A., "On Improving Waste water Quality," Water and Sewage Works, IW 20-26, September-October, 1970.
- 34. Hentschel, M. L., and Cox, T. L., "Effluent Water Treating at Charter International Oil Company's Houston Refinery," 74th National Meeting American Institute of Chemical Engineers, March, 1973.
- 35. Horne, W. R., and Hurse, J. E., "Biological Reduction of Phenolic Type Industrial Wastes," Southern Engineering, January, 1965.
- 35a. Ingols, R.S., "The Toxicity of Chromium," \$ percent8th Purdue Industrial Waste Conference, \$ percent pp. 86-95, 1953.
- 36. Kaup, E. C., "Design Factors in Reverse Osmosis," Chemical Engineering, April 2, 1973.
- 37. Klett, Robert J., "Treat Sour Water for Profit," Hydrocarbon Processing, October, 1972.

- 38. Klipple, R. W., "Pollution Control Built into Guayama Petrochemical Complex," Water and Sewage Works, 116 (3): IW 2-6, March, 1969.
- 39. Lankin, J. C., and Sord, L. V., "American Oil Cleans up Wastes Aerated Lagoons," Hydrocarbon Processing & Petroleum Refiner, 43 (5): 133-136, May, 1964.
- 40. "Major Oil Fields Around the World," International Petroleum Encyclopedia, 1967, Petroleum Publishing Co., Tulsa, Oklahoma.
- 41. Mohler, E. F. Jr., Clere, L. T., "Development of Extensive Water Reuse and Bio-oxidation in a Large Oil Refinery," National Conference on Complete Water Reuse, April, 1973.
- 42. McKinney, R. E., "Biological Treatment Systems for Refinery Wastes," 39th Annual Conference of the Water Pollution Control Federation, pp. 346-359, September 24-30, 1966.
- 43. McKinney, G. M., Ferrero, E. P., and Wenger, W. J., "Analysis of Crude Oils from 546 Important Oil Fields in the United States," Report of Investigations 6819, United States Department of the Interior.
- 44. McPhee, W. T., and Smith, A. R., "From Refinery Waste to Pure Water," 16th Purdue Industrial Wastes Conference, pp. 440-448, May, 1961.
- 45. McWilliams, F. G., and Schuller, R. P., "SNG, Naptha and Low-Sulfur Fuel Oils from Crude Oils Using Commercially Proven Technology American Institute of Chemical Engineers, New York, November 26-December 2, 1972.
- 46. Patterson, J. W., and Minear, R. A., Waste water Treatment Technology, for State of Illinois Institute for Environmental Quality, 2nd, January, 1973.
- 47. "P. C. Treatment Gets Industrial Trial," Environmental Science and Technology, Vol. 7, No. 3, March, 1973, pp. 200-202.
- 48. Peoples, R. F., Krishnan, P., and Simonsery, R. N., "Nonbiological Treatment of Refinery Waste water," Journal of the Water Pollution Control Federation, 44 (11): 2120-2128, November, 1972.
- 48a. "Petroleum Refining Effluent Guidelines", for the Environmental Protection Agency, Office of Water Programs.
- 49. "Petroleum Refining Industry Waste water Profile" for the Federal Water Pollution Control Association.

- 50. Porges, R., "Industrial Waste Stabilization Ponds in the United States," Journal of the Water Pollution Control Federation, 35 (4): 456-468, April, 1963.
- 51. Prather, B. V., "Effects of Aeration on Refinery Waste Water Effluents," Western Petroleum Refiners Association Proceedings of the Waste Disposal and Stream Pollution Conference, October 7-8, 1959.
- 52. "Pretreatment of Discharges to Publicly Owned Treatment Works," Federal Guidelines, E.P.A., 1973.
- 53. Process Design Manual for Carbon Adsorption, Environmental Protection Agency, Washington, D.C., October, 1971.
- 54. Process Design Manual for Suspended Solids Removal, Environmental Protection Agency, Washington, D.C., 1971.
- 55. Purcell, W. L., and Miller, R. B., "Waste Treatment of Skelly Oil Company's El Dorado, Kansas Refinery," 16th Purdue Industrial Waste Conference, pp. 292-303, 1961.
- 56. Rambow, C. A., "Industrial Waste water Reclamation," 23rd Purdue Industrial Waste Conference, pp. 1-9, May, 1968.
- 57. Reid, G. W., and Streebin, L. E., "Evaluation of Waste Waters from Petroleum and Coal Processing," Prepared for Office of Reserach and Monitoring U.S. Environmental Protection Agency, Washington, D.C., (Contract no. EPA-R2-72-001. December, 1972).
- 58. Rodriguez, D. G., "Sour Water Strippers," 74th National Meeting American Institute of Chemical Engineers, March 11-15, 1973.
- 59. Rose, B. A., "Water Conservation Reduces Load on Sohio's Waste Treatment Plant," Water and Sewage Works, 116 (9): IW 4-8, September, 1969.
- 60. Rose, W. L., and Gorringe, G. E., "Activated Sludge Plant Handles Loading Variations," The Oil and Gas Journal, pp. 62-65, October, 1972.
- 61. Sebald, J. F., "A Survey of Evaporative and Non-Evaporative Cooling Systems," 74th National Meeting American Institute of Chemical Engineers, March 11-15, 1973.
- 62. Selvidge, C. W., Conway, J. E., and Jensen, R. H., "Deep Desulfurization of Atmospheric Reduced Crudes by RCD Isomax," Japan Petroleum Institute Meeting, Tokyo, Japan, November 29, 1972.

- 62a. Short, E., and Myers, L.H., "Pilot-Plant Activated Carbon Treatment of Petroleum Refinery Waste water".
- 63. Skamser, Robert O., "The U.S. Refining Outlook to 1980," 74 National Meeting American Institute of Chemical Engineers, March 11-15, 1973.
- 64. Solid Waste Disposal Study, Technical Report Genesee County, Michigan, U.S. Department of Health, Education and Welfare, Bureau of Solid Waste Management, Cincinnati, 1969.
- 65. Standard Industrial Classification Manual, 1967, Executive Office of the President, Bureau of the Budget.
- 66. Standard Methods for Examination of Water and Waste water, American Public Health Association, American Waterworks Association, Water Pollution Control Federation, 13th Edition.
- 67. Stern, Arthur C., Air Pollution, Vol. III, Academic Press Inc., 1968.
- 68. Stramberg, J. B., "EPA Research and Development Activities with Oxygen Aeration," Technology Transfer Design Seminar for Municipal Waste water Treatment Facilities, February 29 and March 1-2, 1972.
- 69. Strong, E. R., and Hatfield, R., "Treatment of Petrochemical Wastes by Superactivated Sludge Process," Industrial and Engineering Chemistry, 46 (2): 308-315, February, 1954.
- 70. Strother, C. W., Vermilion, W. L., and Conner, A. J., "UOP Innovations in Design of Fluid Catalytic Cracking Units," Division of Refining, 37th Midyear Meeting, New York, May 8-12, 1972.
- 71. Stroud, P. W., Sorg, L. V., and Lamkin, J. C., "The First Large Scale Industrial Waste Treatment Plant on the Missouri River," 18th Purdue Industrial Waste Conference, pp. 460-475, 1963.
- 72. Thompson, C. S., Stock, J., and Mehta, P. L., "Cost and Operating Factors for Treatment of Oil Waste Water," The Oil and Gas Journal, pp. 53-56, November 20, 1972.
- 72a. U.S. Department of Health, Education and Welfare, "Public Health Service Drinking Water Standards", PHS Publication No. 956,1962.
- 73. Walker, G. J., "A Design Method for Sour Water Stream Strippers," National Petroleum Refiners Association, March 23-26, 1969.
- 74. Watkins, C. H., and Czajkowski, G. J., "Hydrodesulfurization of Atmospheric and Vacuum Gas Oils," 68th National Meeting American

Institute of Chemical Engineers, Houston, Texas, February 28-March *4, 1971.

Wigren, A. A., and Burton, F. L., "Refinery Waste water Control," Journal of the Water Pollution Control Federation, 44 (1): 117-128, January, 1972.

SECTION XIV

GLOSSARY AND ABBREVIATIONS

Glossary

Acid Oil

Straight chain and cyclic hydrocarbon with carboxyl group(s) attached.

Act

The Federal Water Pollution Act Amendments of 1972.

Aerobic

In the presence of oxygen.

Alkylates

Branched paraffin hydrocarbons.

Anaerobic

Living or active in absence of free oxygen.

uatic Life

All living forms in natural waters, including plants, fish, shellfish, and lower forms of animal life.

Aromatics

Hydrogen compounds involving a 6-carbon, benzene ring structure.

Best Available Technology Economically Achievable (BATEA)

Treatment required by July 1, 1983 for industrial discharge to surface waters as defined by section 310 (b) (2) (A) of the Act.

Best Practicable Control Technology Currently Achievable (BPCTCA)

Treatment required by July 1, 1977 for industrial discharge to surface waters as defined by section 301 (b) (1) (A) of the Act.

Best Available Demonstrated Technology (BADT)

Treatment required for new sources as defined by section 306 of the Act.

Biochemical Oxygen Demand

Oxygen used by bacteria in consuming a waste substance.

Blowdown

A discharge from a system, designed to prevent a buildup of some material, as in boiler and cooling tower to control dissolved solids.

Butadiene

Synthetic hydrocarbon having two unsaturated carbon bonds.

By-Product

Material which, if recovered, would accrue some economic benefit, but not necessarily enough to cover the cost of recovery.

Capital Costs

Financial charges which are computed as the cost of capital times the capital expenditures for pollution control. The cost of capital is based upon a weighted average of the separate costs of debt and equity.

Catalyst

A substance which can change the rate of a chemical reaction, but whis not itself involved in the reaction.



Category and Subcategory

Divisions of a particular industry which processed different traits which affect water quality and treatability.

Chemical Oxygen Demand

Oxygen consumed through chemical oxidation of a waste.

Clarification

The process of removing undissolved materials from a liquid. Specifically, removal of solids either by settling or filtration.

Coke Petroleum

Solid residue of 90 to 95 percent fixed carbon.

Cycles of Concentration

The ratio of the dissolved solids concentration of the recirculating ter to make-up water.

Depletion or Loss

The volume of water which is evaporated, embodied in product, or otherwise disposed of in such a way that it is no longer available for reuse in the plant or available for reuse by others outside the plant.

Depreciation

The cost reflecting the deterioration of a capital asset over its useful life.

Direct-Fired Heater

A heater in which heat is supplied by combustion, as distinguished from a heat exchanger where heat is supplied by a hot liquid or gas.

Emulsion

A liquid system in which one liquid is finely dispersed in another liquid in such a manner that the two will not separate through the action of gravity alone.

End-of-Pipe Treatment

eatment of overall refinery wastes, as distinguished from treatment at individual processing units.

Filtration

Removal of solid particles or liquids from other liquids or gas streams by passing the liquid or gas stream through a filter media.

Fractionator

A generally cylindrical tower in which a mixture of liquid components is vaporized and the components separated by carefully varying the temperature and sometimes pressure along the length of the tower.

Gasoline

A mixture of hydrocarbon compounds with a boiling range between 1000 and 4000 F.

Grease

A solid or semi-solid composition made up of animal fats, alkali, water, oil and various additives.

Hydrocarbon

A compound consisting of carbon and hydrogen.

Hydrogenation

The contacting of unsaturated or impure hydrocarbons with hydrogen gas at controlled temperatures and pressures for the purpose of obtaining saturated hydrocarbons and/or removing various impurities such as sulfur and nitrogen.

Industrial Waste

All wastes streams within a plant. Included are contact and non-contact waters. Not included are wastes typically considered to be sanitary wastes.

Investment Costs

The capital expenditures required to bring the treatment or control technology into operation. These include the traditional expenditures such as design; purchase of land and materials; site preparation; construction and installation; etc., plus any additional expenses required to bring the technology into operation including expenditures to establish related necessary solid waste disposal.

Isomer

A chemical compound that has the same number, and kinds of atoms as another compound, but a different structural arrangement of the atoms.

Mercaptan

An organic compound containing hydrogen, carbon, and sulfur (RSH).

Microcrystalline Wax

A non-crystalline solid hydrogen with a melting point of about 1060 to 1950 F. Also known as petrolatum.

Motor Octane Number

An expression of the antiknock value of gasoline.

Naphtha

A petroleum fraction, including parts of the boiling range of gasoline and kerosine, from which solvents are obtained.

phthenic Acids

Partially oxidized naphthalenes.

New Source

Any building, structure, facility, or installation from which there is or may be a discharge of pollutants and whose construction is commenced after the publication of the proposed regulations.

No Discharge of Pollutants

No net increase (or detectable gross concentration if the situation dictates) of any parameter designated as a pollutant to the accuracy that can be determined from the designated analytical method.

Octane

The numerical rating of a gasoline's resistance to engine knock.

Olefins

Unsaturated straight-chain hydrocarbon compounds seldom present in crude oil, but frequently in cracking processes.

eration and Maintenance

Costs required to operate and maintain pollution abatement equipment. They include labor, material, insurance, taxes, solid waste disposal, etc.

Overhead Accumulator

A tank in which the condensed vapors from the tops of the fractionators, steam strippers, or stabilizers are collected.

Paraffin Wax

A crystalline solid hydrocarbon with a melting point of 1050 to 1550 F.

Petroleum

A complex liquid mixture of hydrocarbons and small quantities of nitrogen, sulfur, and oxygen.

pН

A measure of the relative acidity or alkalinity of water. A pH of 7.0 indicates a neutral condition. A greater pH indicates alkalinity and a lower pH indicates acidity. A one unit change in pH indicates 10 fold change in acidity and alkalinity.

Phenol

Class of cyclic organic derivatives with basic formula C6HOH.

Pretreatment

Treatment proved prior to discharge to a publicly owned treatment works.

Process Effluent or Discharge

The volume of water emerging from a particular use in the plant.

Plant Effluent or Discharge After Treatment

The volume of waste water discharge from the industrial plant. In this definition, any waste treatment device is considered part of the industrial plant.

Raffinate

The portion of the oil which remains undissolved and is not removed by solvent extraction.

Raw

Untreated or unprocessed.

Reduced Crude

The thick, dark, high-boiling residue remaining after crude oil has undergone atmospheric and/or vacuum fractionation.

Secondary Treatment

Biological treatment provided beyond primary clarification.

Sludge

The settled solids from a thickener or clarifier. Generally, almost any flocculated settled mass.

Sour

Denotes the presence of sulfur compounds, such as sulfides and mercaptans, that cause bad odors.

Spent Caustic

neous solution of sodium hydroxide that has been used to remove fides, mercaptans, and organic acids from petroleum fractions.

Stabilizer

A type of fractionator used to remove dissolved gaseous hydrocarbons from liquid hydrocarbon products.

Standard Raw Waste Loads (SRWL)

Net pollution loading produced per unit of production (or raw material) by a refining process after separation of the separables (STS).

Stripper

A unit in which certain components are removed from a liquid hydrocarbon mixture by passing a gas, usually steam, through the mixture.

Supernatant

The layer floating above the surface of a layer of solids.

Surface Waters

Navigable waters. The waters of the United States, including the

sweet

Denotes the absence of odor-causing sulfur compounds, such as sulfides and mercaptans.

Topping Plant

A refinery whose processing is largely confined to oil into raw products by simple atmospheric distillation.

Total Suspended Solids (TSS)

Any solids found in waste water or in the stream which in most cases can be removed by filtration. The origin of suspended matter may be man-made wastes or natural sources such as silt from erosion.

Waste Discharged

The amount (usually expressed as weight) of some residual substance which is suspected or dissolved in the plant effluent after treatment if any.

Waste Generated

The amount (usually expressed as weight) of some residual substangenerated by a plant process or the plant as whole and which suspended or dissolved in water. This quantity is measured before treatment.

Waste Loading

Total amount of pollutant substance, generally expressed as pounds per day.

Abbreviations

- Aerated Lagoon

AS - Activated Sludge

API - American Petroleum Institute

BADT - Best Available Demonstrated Technology

BATEA - Best Available Technology Economically Achievable

BPCTCA - Best Practicable Control Technology Currently Available

bbl - Barrel

BOD - Biochemical Oxygen Demand

bpcd - Barrels per calendar day

bpsd - Barrels per stream day (operating day)

BS and W - Bottom Sediment and Water

BTX - Benzene-Toluene-Xylene mixture

COD - Chemical Oxygen Demand

cu m - cubic meter(s)

DAF - Dissolved Air Flotation

DO - Dissolved Oxygen

qpm - Gallons per minute

k - thousand(e.g., thousand cubic meters)

kg - kilogram(s)

1 - liter

1b - pound(s).

LPG - Liquified Petroleum Gas

M - Thousand (e.g., thousand barrels)

MBCD - Thousand Barrels per calendar day

MBSD - Thousand Barrels per stream day

mgd - Million gallons per day

mg/L - Milligrams per liter (parts per million)

MM - Million (e.g., million pounds)

PP - Polishing pond

psig - pounds per square inch, gauge (above 14.7 psig)

RSH - Mercaptan

sec - Second-unit of 'time

scf - Standard cubic feet of gas at 600 F and 14.7 psig

SIC - Standard Industrial Classification

SRWL - Standard Raw Waste Load

SS - Suspended Solids

STS - Susceptible to Separation

TOC - Total Organic Carbon

TSS - Total Suspended Solids

VSS - Volatile Suspended Solids